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Laparoscopic Surgery
Smith’s General Urology, 17th edition, provides in a concise format the information necessary for the understanding, diagnosis, and treatment of diseases managed by urologic surgeons. Our goal has been to keep the book current, to the point, and readable.

Medical students will find this book useful because of its concise, easy-to-follow format and organization and its breadth of information. Interns and residents, as well as practicing physicians in urology or general medicine, will find it an efficient and current reference, particularly because of its emphasis on diagnosis and treatment.

The 17th edition is a thorough revision of the book. New chapters to this edition include: “Pharmacology of the Lower Urinary Tract,” “Female Urology,” and “The Aging Male.”

The book has been reviewed and updated throughout, with emphasis on current references. The several illustrations have been further modernized and improved, including many fine anatomic drawings and the latest imaging techniques. Since the 11th edition, the following translations have been published: Chinese, French, Greek, Italian, Japanese, Korean, Portuguese, Russian, Spanish, and Turkish.

We greatly appreciate the patience and efforts of our McGraw-Hill staff, the expertise of our contributors, and the support of our readers.
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Urology deals with diseases and disorders of the male genitourinary tract and the female urinary tract. Surgical diseases of the adrenal gland are also included. These systems are illustrated in Figures 1–1 and 1–2.

**ADRENALS**

**Gross Appearance**

**A. ANATOMY**

Each kidney is capped by an adrenal gland, and both organs are enclosed within Gerota’s (perirenal) fascia. Each adrenal weighs about 5 g. The right adrenal is triangular in shape; the left is more rounded and crescentic. Each gland is composed of a cortex, chiefly influenced by the pituitary gland, and a medulla derived from chromaffin tissue.

**B. RELATIONS**

Figure 1–2 shows the relation of the adrenals to other organs. The right adrenal lies between the liver and the vena cava. The left adrenal lies close to the aorta and is covered on its lower surface by the pancreas; superiorly and laterally, it is related to the spleen.

**Histology**

The adrenal cortex is composed of 3 distinct layers: the outer zona glomerulosa, the middle zona fasciculata, and the inner zona reticularis. The medulla lies centrally and is made up of polyhedral cells containing eosinophilic granular cytoplasm. These chromaffin cells are accompanied by ganglion and small round cells.

**Blood Supply**

**A. ARTERIAL**

Each adrenal receives 3 arteries: one from the inferior phrenic artery, one from the aorta, and one from the renal artery.

**B. VENOUS**

Blood from the right adrenal is drained by a very short vein that empties into the vena cava; the left adrenal vein terminates in the left renal vein.

**KIDNEYS**

**Gross Appearance**

**A. ANATOMY**

The kidneys lie along the borders of the psoas muscles and are therefore obliquely placed. The position of the liver causes the right kidney to be lower than the left (Figures 1–2 and 1–3). The adult kidney weighs about 150 g.

The kidneys are supported by the perirenal fat (which is enclosed in the perirenal fascia), the renal vascular pedicle, abdominal muscle tone, and the general bulk of the abdominal viscera. Variations in these factors permit variations in the degree of renal mobility. The average descent on inspiration or on assuming the upright position is 4–5 cm. Lack of mobility suggests abnormal fixation (eg, perinephritis), but extreme mobility is not necessarily pathologic.

On longitudinal section (Figure 1–4), the kidney is seen to be made up of an outer cortex, a central medulla, and the internal calices and pelvis. The cortex is homogeneous in appearance. Portions of it project toward the pelvis between the papillae and fornices and are called the columns of Bertin. The medulla consists of numerous pyramids formed by the converging collecting renal tubules, which drain into the minor calices at the tip of the papillae.

**B. RELATIONS**

Figures 1–2 and 1–3 show the relations of the kidneys to adjacent organs and structures. Their intimacy with intra-peritoneal organs and the autonomic innervation they share with these organs explain, in part, some of the gastrointestinal symptoms that accompany genitourinary disease.

**Histology**

**A. NEPHRON**

The functioning unit of the kidney is the nephron, which is composed of a tubule that has both secretory and excretory functions (Figure 1–4). The secretory portion is contained largely within the cortex and consists of a renal corpuscle.
Figure 1–1. Anatomy of the male genitourinary tract. The upper and midtracts have urologic function only. The lower tract has both genital and urinary functions.
ANATOMY OF THE GENITOURINARY TRACT

and the secretory part of the renal tubule. The excretory portion of this duct lies in the medulla. The renal corpuscle is composed of the vascular glomerulus, which projects into Bowman’s capsule, which, in turn, is continuous with the epithelium of the proximal convoluted tubule. The secretory portion of the renal tubule is made up of the proximal convoluted tubule, the loop of Henle, and the distal convoluted tubule.

The excretory portion of the nephron is the collecting tubule, which is continuous with the distal end of the ascending limb of the convoluted tubule. It empties its contents through the tip (papilla) of a pyramid into a minor calyx.

B. SUPPORTING TISSUE

The renal stroma is composed of loose connective tissue and contains blood vessels, capillaries, nerves, and lymphatics.

Blood Supply (Figures 1–2, 1–4, & 1–5)

A. ARTERIAL

Usually there is one renal artery, a branch of the aorta that enters the hilum of the kidney between the pelvis, which normally lies posteriorly, and the renal vein. It may branch before it reaches the kidney, and 2 or more separate arteries may be noted. In duplication of the pelvis and ureter, it is usual for each renal segment to have its own arterial supply.

The renal artery divides into anterior and posterior branches. The posterior branch supplies the midsegment of the posterior surface. The anterior branch supplies both upper and lower poles as well as the entire anterior surface. The renal arteries are all end arteries.

The renal artery further divides into interlobar arteries, which ascend in the columns of Bertin (between

Figure 1–2. Relations of kidney, ureters, and bladder (anterior aspect).
CHAPTER 1

the pyramids) and then arch along the base of the pyramids (arcuate arteries). The renal artery then ascends as interlobular arteries. From these vessels, smaller (afferent) branches pass to the glomeruli. From the glomerular tuft, efferent arterioles pass to the tubules in the stroma.

B. VENOUS

The renal veins are paired with the arteries, but any of them will drain the entire kidney if the others are tied off.

Although the renal artery and vein are usually the sole blood vessels of the kidney, accessory renal vessels are common and may be of clinical importance if they are so placed to compress the ureter, in which case hydronephrosis may result.

Nerve Supply

The renal nerves derived from the renal plexus accompany the renal vessels throughout the renal parenchyma.

Lymphatics

The lymphatics of the kidney drain into the lumbar lymph nodes.

CALICES, RENAL PELVIS, & URETER

Gross Appearance

A. ANATOMY

1. Calices—The tips of the minor calices (8–12 in number) are indented by the projecting pyramids (Figure 1–4). These calices unite to form 2 or 3 major calices which join to form the renal pelvis.

2. Renal pelvis—The pelvis may be entirely intrarenal or partly intrarenal and partly extrarenal. Inferomedially, it tapers to form the ureter.

3. Ureter—The adult ureter is about 30 cm long, varying in direct relation to the height of the individual. It follows a rather smooth S curve. Areas of relative narrowing are found (1) at the ureteropelvic junction, (2) where the ureter crosses over the iliac vessels, and (3) where it courses through the bladder wall.

B. RELATIONS

1. Calices—The calices are intrarenal and are intimately related to the renal parenchyma.

2. Renal pelvis—If the pelvis is partly extrarenal, it lies along the lateral border of the psoas muscle and on the quadratus lumborum muscle; the renal vascular pedicle is
Figure 1–4. Anatomy and histology of the kidney and ureter. **Upper left:** Diagram of the nephron and its blood supply (Courtesy of Merck, Sharp, Dohme: Seminar. 1947;9[3].) **Upper right:** Cast of the pelvic caliceal system and the arterial supply of the kidney. **Middle:** Renal calices, pelvis, and ureter (posterior aspect). **Lower left:** Histology of the ureter. The smooth-muscle bundles are arranged in both a spiral and a longitudinal manner. **Lower right:** Longitudinal section of kidney showing calices, pelvis, ureter, and renal blood supply (posterior aspect).
Figure 1–5.  **A:** The posterior branch of the renal artery and its distribution to the central segment of the posterior surface of the kidney. **B:** Branches of the anterior division of the renal artery supplying the entire anterior surface of the kidney as well as the upper and lower poles at both surfaces. The segmental branches lead to interlobar, arcuate, and interlobular arteries. **C:** The lateral convex margin of the kidney. Brödel’s line, which is 1 cm from the convex margin, is the bloodless plane demarcated by the distribution of the posterior branch of the renal artery.
placed just anterior to it. The left renal pelvis lies at the level of the first or second lumbar vertebra; the right pelvis is a little lower.

3. Ureter—As followed from above, downward the ureters lie on the psoas muscles, pass medially to the sacroiliac joints, and then swing laterally near the ischial spines before passing medially to penetrate the base of the bladder (Figure 1–2). In females, the uterine arteries are closely related to the juxtavesical portion of the ureters. The ureters are covered by the posterior peritoneum; their lowermost portions are closely attached to it, while the juxtavesical portions are embedded in vascular retroperitoneal fat.

The vasa deferentia, as they leave the internal inguinal rings, sweep over the lateral pelvic walls anterior to the ureters (Figure 1–6). They lie medial to the latter before joining the seminal vesicle and penetrating the base of the prostate to become the ejaculatory ducts.

Histology (Figure 1–4)

The walls of the calices, pelvis, and ureters are composed of transitional cell epithelium under which lies loose connective and elastic tissue (lamina propria). External to these are a mixture of helical and longitudinal smooth muscle fibers. They are not arranged in definite layers. The outermost adventitial coat is composed of fibrous connective tissue.

Blood Supply

A. ARTERIAL

The renal calices, pelvis, and upper ureters derive their blood supply from the renal arteries; the midureter is fed by the internal spermatic (or ovarian) arteries. The lowermost portion of the ureter is served by branches from the common iliac, internal iliac (hypogastric), and vesical arteries.

B. VENOUS

The veins of the renal calices, pelvis, and ureters are paired with the arteries.

Lymphatics

The lymphatics of the upper portions of the ureters as well as those from the pelvis and calices enter the lumbar lymph nodes. The lymphatics of the midureter pass to the internal iliac (hypogastric) and common iliac lymph nodes; the lower ureteral lymphatics empty into the vesical and hypogastric lymph nodes.

BLADDER

Gross Appearance

The bladder is a hollow muscular organ that serves as a reservoir for urine. In women, its posterior wall and dome are invaginated by the uterus. The adult bladder normally has a capacity of 400–500 mL.

A. ANATOMY

When empty, the adult bladder lies behind the pubic symphysis and is largely a pelvic organ. In infants and children, it is situated higher. When it is full, it rises well above the symphysis and can readily be palpated or percussed. When overdistended, as in acute or chronic urinary retention, it may cause the lower abdomen to bulge visibly.

Extending from the dome of the bladder to the umbilicus is a fibrous cord, the median umbilical ligament, which represents the obliterated urachus. The ureters enter the bladder posteroinferiorly in an oblique manner and at these points are about 5 cm apart (Figure 1–6). The orifices, situated at the extremities of the crescent-shaped interureteric ridge that forms the proximal border of the trigone, are about 2.5 cm apart. The trigone occupies the area between the ridge and the bladder neck.

The internal sphincter, or bladder neck, is not a true circular sphincter but a thickening formed by interlaced and converging muscle fibers of the detrusor as they pass distally to become the smooth musculature of the urethra.

B. RELATIONS

In males, the bladder is related posteriorly to the seminal vesicles, vasa deferentia, ureters, and rectum (Figures 1–7 and 1–8). In females, the uterus and vagina are interposed between the bladder and rectum (Figure 1–9). The dome and posterior surfaces are covered by peritoneum; hence, in this area the bladder is closely related to the small intestine and sigmoid colon. In both males and females, the
Figure 1–7.  A: Anatomic relationship of the bladder, prostate, prostatomembranous urethra, and root of the penis.  
B: Histology of the testis. Seminiferous tubules lined by supporting basement membrane for the Sertoli and spermatogenic cells. The latter are in various stages of development.  
bladder is related to the posterior surface of the pubic symphysis, and, when distended, it is in contact with the lower abdominal wall.

**Histology (Figure 1–10)**

The mucosa of the bladder is composed of transitional epithelium. Beneath it is a well-developed submucosal layer formed largely of connective and elastic tissues. External to the submucosa is the detrusor muscle which is made up of a mixture of smooth muscle fibers arranged at random in a longitudinal, circular, and spiral manner without any layer formation or specific orientation except close to the internal meatus, where the detrusor muscle assumes 3 definite layers: inner longitudinal, middle circular, and outer longitudinal.
Figure 1–9. Anatomy and relations of the bladder, urethra, uterus and ovary, vagina, and rectum.

Figure 1–10. Left: Histology of the prostate. Epithelial glands embedded in a mixture of connective and elastic tissue and smooth muscle. Right: Histology of the bladder. The mucosa is transitional cell in type and lies on a well-developed submucosal layer of connective tissue. The detrusor muscle is composed of interlacing longitudinal, circular, and spiral smooth-muscle bundles.
ANATOMY OF THE GENITOURINARY TRACT

Blood Supply

A. ARTERIAL

The bladder is supplied with blood by the superior, middle, and inferior vesical arteries, which arise from the anterior trunk of the internal iliac (hypogastric) artery, and by smaller branches from the obturator and inferior gluteal arteries. In females, the uterine and vaginal arteries also send branches to the bladder.

B. VENOUS

Surrounding the bladder is a rich plexus of veins that ultimately empties into the internal iliac (hypogastric) veins.

Lymphatics

The lymphatics of the bladder drain into the vesical, external iliac, internal iliac (hypogastric), and common iliac lymph nodes.

PROSTATE GLAND

Gross Appearance

A. ANATOMY

The prostate is a fibromuscular and glandular organ lying just inferior to the bladder (Figures 1–6 and 1–7). The normal prostate weighs about 20 g and contains the posterior urethra, which is about 2.5 cm in length. It is supported anteriorly by the puboprostatic ligaments and inferiorly by the urogenital diaphragm (Figure 1–6). The prostate is perforated posteriorly by the ejaculatory ducts, which pass obliquely to empty through the verumontanum on the floor of the prostatic urethra just proximal to the striated external urinary sphincter (Figure 1–11). According to the classification of Lowsley, the prostate consists of 5 lobes: anterior, posterior, median, right lateral, and left lateral. According to McNeal (1972), the prostate has a peripheral zone, a central zone, and a transitional zone; an anterior segment; and a prostatic sphincteric zone (Figure 1–12). The segment of urethra that traverses the prostate gland is the prostatic urethra. It is lined by an inner longitudinal layer of muscle (continuous with a similar layer of the vesical wall). Incorporated within the prostate gland is an abundant amount of smooth musculature derived primarily from the external longitudinal bladder musculature. This musculature represents the true smooth involuntary sphincter of the posterior urethra in males.

B. RELATIONS

The prostate gland lies behind the pubic symphysis. Located closely to the posterosuperior surface are the vasa deferentia and seminal vesicles (Figure 1–7). Posteriorly, the prostate is separated from the rectum by the 2 layers of Denonvilliers’ fascia, serosal rudiments of the pouch of Douglas, which once extended to the urogenital diaphragm (Figure 1–8).

Histology (Figure 1–10)

The prostate consists of a thin fibrous capsule under which are circularly oriented smooth muscle fibers and collagenous tissue that surrounds the urethra (involuntary sphincter). Deep in this layer lies the prostatic stroma, composed of connective and elastic tissues and smooth muscle fibers in which are embedded the epithelial glands. These glands drain into the major excretory ducts (about 25 in number) which open chiefly on the floor of the urethra between the verumontanum and the vesical neck. Just beneath the transitional epithelium of the prostatic urethra lie the periurethral glands.

Blood Supply

A. ARTERIAL

The arterial supply to the prostate is derived from the inferior vesical, internal pudendal, and middle rectal (hemorrhoidal) arteries.
The veins from the prostate drain into the periprostatic plexus, which has connections with the deep dorsal vein of the penis and the internal iliac (hypogastric) veins.

Nerve Supply
The prostate gland receives a rich nerve supply from the sympathetic and parasympathetic nerve plexuses.

Lymphatics
The lymphatics from the prostate drain into the internal iliac (hypogastric), sacral, vesical, and external iliac lymph nodes.

B. SEMINAL VESICLES

Gross Appearance
The seminal vesicles lie just cephalic to the prostate under the base of the bladder (Figures 1–6 and 1–7). They are about 6 cm long and quite soft. Each vesicle joins its corresponding vas deferens to form the ejaculatory duct. The ureters lie medial to each, and the rectum is contiguous with their posterior surfaces.

Histology
The mucous membrane is pseudostratified. The submucosa consists of dense connective tissue covered by a thin layer of muscle that in turn is encapsulated by connective tissue.

Blood Supply
The blood supply is similar to that of the prostate gland.

Nerve Supply
The nerve supply is mainly from the sympathetic nerve plexus.

Lymphatics
The lymphatics of the seminal vesicles are those that serve the prostate.

SPERMATIC CORD

Gross Appearance
The 2 spermatic cords extend from the internal inguinal rings through the inguinal canals to the testicles (Figure 1–7). Each cord contains the vas deferens, the internal and external spermatic arteries, the artery of the vas, the venous pampiniform plexus (which forms the spermatic vein superiorly), lymph vessels, and nerves. All of the preceding are enclosed in investing layers of thin fascia. A few fibers of the cremaster muscle insert on the cords in the inguinal canal.

Histology
The fascia covering the cord is formed of loose connective tissue that supports arteries, veins, and lymphatics. The vas deferens is a small, thick-walled tube consisting of an internal mucosa and submucosa surrounded by 3 well-defined layers of smooth muscle encased in a covering of fibrous tissue. Above the testes, this tube is straight. Its proximal 4 cm tends to be convoluted.

Blood Supply

A. ARTERIAL
The external spermatic artery, a branch of the inferior epigastric, supplies the fascial coverings of the cord. The internal spermatic artery passes through the cord on its way to the testis. The deferential artery is close to the vas.

B. VENOUS
The veins from the testis and the coverings of the spermatic cord form the pampiniform plexus, which, at the internal inguinal ring, unites to form the spermatic vein.
**Lymphatics**
The lymphatics from the spermatic cord empty into the external iliac lymph nodes.

**EPIDIDYMIS**

**Gross Appearance**

**A. ANATOMY**
The upper portion of the epididymis (globus major) is connected to the testis by numerous efferent ducts from the testis (Figure 1–7). The epididymis consists of a markedly coiled duct that, at its lower pole (globus minor), is continuous with the vas deferens. An appendix of the epididymis is often seen on its upper pole; this is a cystic body that in some cases is pedunculated but in others is sessile.

**B. RELATIONS**
The epididymis lie posterolateral to the testis and is nearest to the testis at its upper pole. Its lower pole is connected to the testis by fibrous tissue. The vas lies posteromedial to the epididymis.

**Histology**
The epididymis is covered by serosa. The ductus epididymidis is lined by pseudostratified columnar epithelium throughout its length.

**Blood Supply**

**A. ARTERIAL**
The arterial supply to the epididymis comes from the internal spermatic artery and the artery of the vas (deferential artery).

**B. VENOUS**
The venous blood drains into the pampiniform plexus which becomes the spermatic vein.

**Lymphatics**
The lymphatics drain into the external iliac and internal iliac (hypogastric) lymph nodes.

**TESTIS**

**Gross Appearance**

**A. ANATOMY**
The average testicle measures about $4 \times 3 \times 2.5$ cm (Figure 1–7). It has a dense fascial covering called the tunica albuginea testis, which, posteriorly, is invaginated somewhat into the body of the testis to form the mediastinum testis. This fibrous mediastinum sends fibrous septa into the testis, thus separating it into about 250 lobules.

**B. RELATIONS**
The testis is closely attached posterolaterally to the epididymis, particularly at its upper and lower poles.

**Histology (Figure 1–7)**

Each lobule contains 1–4 markedly convoluted seminiferous tubules, each of which is about 60 cm long. These ducts converge at the mediastinum testis, where they connect with the efferent ducts that drain into the epididymis.

The seminiferous tubule has a basement membrane containing connective and elastic tissue. This supports the seminiferous cells which are of 2 types: (1) Sertoli (supporting) cells and (2) spermatogenic cells. The stroma between the seminiferous tubules contains connective tissue in which the interstitial Leydig cells are located.

**Blood Supply**

The blood supply to the testes is closely associated with that to the kidneys because of the common embryologic origin of the 2 organs.

**A. ARTERIAL**
The arteries to the testes (internal spermatics) arise from the aorta just below the renal arteries and course through the spermatic cords to the testes, where they anastomose with the arteries of the vasa deferentia that branch off from the internal iliac (hypogastric) artery.

**B. VENOUS**
The blood from the testis returns in the pampiniform plexus of the spermatic cord. At the internal inguinal ring, the pampiniform plexus forms the spermatic vein.

The right spermatic vein enters the vena cava just below the right renal vein; the left spermatic vein empties into the left renal vein.

**Lymphatics**
The lymphatic vessels from the testes pass to the lumbar lymph nodes, which in turn are connected to the mediastinal nodes.

**SCROTUM**

**Gross Appearance**

Beneath the corrugated skin of the scrotum lies the dartos muscle. Deep to this are the 3 fascial layers derived
from the abdominal wall at the time of testicular descent. Beneath these is the parietal layer of the tunica vaginalis.

The scrotum is divided into 2 sacs by a septum of connective tissue. The scrotum not only supports the testes but, by relaxation or contraction of its muscular layer, helps to regulate their environmental temperature.

**Histology**

The dartos muscle, under the skin of the scrotum, is unstriated. The deeper layer is made up of connective tissue.

**Blood Supply**

**A. ARTERIAL**

The arteries to the scrotum arise from the femoral, internal pudendal, and inferior epigastric arteries.

**B. VENOUS**

The veins are paired with the arteries.

**Lymphatics**

The lymphatics drain into the superficial inguinal and subinguinal lymph nodes.

**PENIS & MALE URETHRA**

**Gross Appearance**

The penis is composed of 2 corpora cavernosa and the corpus spongiosum, which contains the urethra, whose diameter is 8–9 mm. These corpora are capped distally by the glans. Each corpus is enclosed in a fascial sheath (tunica albuginea), and all are surrounded by a thick fibrous envelope known as Buck’s fascia. A covering of skin, devoid of fat, is loosely applied about these bodies. The prepuce forms a hood over the glans.

Beneath the skin of the penis (and scrotum) and extending from the base of the glans to the urogenital diaphragm is Colles’ fascia, which is continuous with Scarpa’s fascia of the lower abdominal wall (Figure 1–8).

The proximal ends of the corpora cavernosa are attached to the pelvic bones just anterior to the ischial tuberosities. Occupying a depression of their ventral surface in the midline is the corpus spongiosum, which is connected proximally to the undersurface of the urogenital diaphragm, through which emerges the membranous urethra. This portion of the corpus spongiosum is surrounded by the bulbospongious muscle. Its distal end expands to form the glans penis.

The suspensory ligament of the penis arises from the linea alba and pubic symphysis and inserts into the fascial covering of the corpora cavernosa.

**Histology**

**A. CORPORA & GLANS PENIS**

The corpora cavernosa, the corpus spongiosum, and the glans penis are composed of septa of smooth muscle and erectile tissue that enclose vascular cavities.

**B. URETHRA**

The urethral mucosa that traverses the glans penis is formed of squamous epithelium. Proximal to this, the mucosa is transitional in type. Underneath the mucosa is the submucosa which contains connective and elastic tissue and smooth muscle. In the submucosa are the numerous glands of Littre, whose ducts connect with the urethral lumen. The urethra is surrounded by the vascular corpus spongiosum and the glans penis.

**Blood Supply**

**A. ARTERIAL**

The penis and urethra are supplied by the internal pudendal arteries. Each artery divides into a deep artery of the penis (which supplies the corpora cavernosa), a dorsal artery of the penis, and the bulbourethral artery. These branches supply the corpus spongiosum, the glans penis, and the urethra.

**B. VENOUS**

The superficial dorsal vein lies external to Buck’s fascia. The deep dorsal vein is placed beneath Buck’s fascia and lies between the dorsal arteries. These veins connect with the pudendal plexus which drains into the internal pudendal vein.

**Lymphatics**

Lymphatic drainage from the skin of the penis is to the superficial inguinal and subinguinal lymph nodes. The lymphatics from the glans penis pass to the subinguinal and external iliac nodes. The lymphatics from the deep urethra drain into the internal iliac (hypogastric) and common iliac lymph nodes.

**FEMALE URETHRA**

The adult female urethra is about 4 cm long and 8 mm in diameter. It is slightly curved and lies beneath the pubic symphysis just anterior to the vagina.

The epithelial lining of the female urethra is squamous in its distal portion and pseudostratified or transitional in the remainder. The submucosa is made up of connective and elastic tissues and spongy venous spaces. Embedded in it are many periurethral glands, which are most numerous distally; the largest of these are the periurethral glands of Skene which open on the floor of the urethra just inside the meatus.
External to the submucosa is a longitudinal layer of smooth muscle continuous with the inner longitudinal layer of the bladder wall. Surrounding this is a heavy layer of circular smooth muscle fibers extending from the external vesical muscular layer. They constitute the true involuntary urethral sphincter. External to this is the circular striated (voluntary) sphincter surrounding the middle third of the urethra; this constitutes an intrinsic element in the musculature of the urethra.

The arterial supply to the female urethra is derived from the inferior vesical, vaginal, and internal pudendal arteries. Blood from the urethra drains into the internal pudendal veins.

Lymphatic drainage from the external portion of the urethra is to the inguinal and subinguinal lymph nodes. Drainage from the deep urethra is into the internal iliac (hypogastric) lymph nodes.

**Nerve Supply to the Genitourinary Organs**

See Figures 3–2 and 3–3.

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**Prostate**

**Prostate**

**Spermatic Cord**

**Testis**

**Female Urethra**
Embryology of the Genitourinary System

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At birth, the genital and urinary systems are related only in the sense that they share certain common passages. Embryologically, however, they are intimately related. Because of the complex interrelationships of the embryonic phases of the 2 systems, they are discussed here as 5 subdivisions: the nephric system, the vesicourethral unit, the gonads, the genital duct system, and the external genitalia.

NEPHRIC SYSTEM

The nephric system develops progressively as 3 distinct entities: pronephros, mesonephros, and metanephros.

Pronephros

The pronephros is the earliest nephric stage in humans, and it corresponds to the mature structure of the most primitive vertebrate. It extends from the 4th to the 14th somites and consists of 6–10 pairs of tubules. These open into a pair of primary ducts that are formed at the same level, extend caudally, and eventually reach and open into the cloaca. The pronephros is a vestigial structure that disappears completely by the 4th week of embryonic life (Figure 2–1).

Mesonephros

The mature excretory organ of the higher fish and amphibians corresponds to the embryonic mesonephros. It is the principal excretory organ during early embryonic life (4–8 weeks). It too gradually degenerates, although parts of its duct system become associated with the male reproductive organs. The mesonephric tubules develop from the intermediate mesoderm caudal to the pronephros shortly before pronephric degeneration. The mesonephric tubules differ from those of the pronephros in that they develop a cuplike outgrowth into which a knot of capillaries is pushed. This is called Bowman’s capsule, and the tuft of capillaries is called a glomerulus. In their growth, the mesonephric tubules extend toward and establish a connection with the nearby primary nephric duct as it grows caudally to join the cloaca (Figure 2–1). This primary nephric duct is now called the mesonephric duct. After establishing their connection with the nephric duct, the primordial tubules elongate and become S-shaped. As the tubules elongate, a series of secondary branchings increase their surface exposure, thereby enhancing their capacity for interchanging material with the blood in adjacent capillaries. Leaving the glomerulus, the blood is carried by one or more efferent vessels that soon break up into a rich capillary plexus closely related to the mesonephric tubules. The mesonephros, which forms early in the 4th week, reaches its maximum size by the end of the second month.

Metanephros

The metanephros, the final phase of development of the nephric system, originates from both the intermediate mesoderm and the mesonephric duct. Development begins in the 5- to 6-mm embryo with a budlike outgrowth from the mesonephric duct as it bends to join the cloaca. This ureteral bud grows cephalad and collects mesoderm from the nephrogenic cord of the intermediate mesoderm around its tip. This mesoderm with the metanephric cap moves, with the growing ureteral bud, more and more cephalad from its point of origin. During this cephalic migration, the metanephric cap becomes progressively larger, and rapid internal differentiation takes place. Meanwhile, the cephalic end of the ureteral bud expands within the growing mass of metanephrogenic tissue to form the renal pelvis (Figure 2–1). Numerous outgrowths from the renal pelvic dilatation push radially into this growing mass and form hollow ducts that branch and rebranch as they push toward the periphery. These form the primary collecting ducts of the kidney. Mesodermal cells become arranged in small vesicular masses that lie close to the blind end of the collecting ducts. Each of these vesicular masses will form a uriniferous tubule draining into the duct nearest to its point of origin.

As the kidney grows, increasing numbers of tubules are formed in its peripheral zone. These vesicular masses develop a central cavity and become S-shaped. One end of the S coalesces with the terminal portion of the collecting
tubules, resulting in a continuous canal. The proximal portion of the S develops into the distal and proximal convoluted tubules and into Henle’s loop; the distal end becomes the glomerulus and Bowman’s capsule. At this stage, the undifferentiated mesoderm and the immature glomeruli are readily visible on microscopic examination (Figure 2–2). The glomeruli are fully developed by the 36th week or when the fetus weighs 2500 g (Osathanondh and Potter, 1964a and b). The metanephros arises opposite the 28th somite (fourth lumbar segment). At term, it has ascended to the level of the first lumbar or even the twelfth thoracic vertebra. This ascent of the kidney is due not only to actual cephalic migration but also to differential growth in the caudal part of the body. During the early period of ascent (7th to 9th weeks), the kidney slides above the arterial bifurcation and rotates 90º. Its convex border is now directed laterally, not dorsally. Ascent proceeds more slowly until the kidney reaches its final position.

Certain features of these 3 phases of development must be emphasized: (1) The 3 successive units of the system develop from the intermediate mesoderm. (2) The tubules at all levels appear as independent primordia and only secondarily unite with the duct system. (3) The nephric duct is laid down as the duct of the pronephros and develops from the union of the ends of the anterior pronephric tubules. (4) This pronephric duct serves subsequently as the mesonephric duct and as such gives rise to the ureter. (5) The nephric duct reaches the cloaca by independent caudal growth. (6) The embryonic ureter is an outgrowth of the nephric duct, yet the kidney tubules differentiate from adjacent metanephric blastema.

**Molecular Mechanisms of Renal & Uretal Development**

The kidney and the collecting system originate from the interaction between the mesonephric duct (Wolffian duct) and the metanephric mesenchyme (MM). The ureteric bud (UB) forms as an epithelial outpouching from the mesonephric duct and invades the surrounding MM. Reciprocal induction between the UB and MM results in branching and elongation of the UB from the collecting system and in condensation and epithelial differentiation of MM around the branched tips of the UB. Branching of the UB occurs approximately 15 times during human renal development, generating approximately 300,000 and 1 million nephrons per kidney (Nyengaard and Bendtsen, 1992).

This process of reciprocal induction is dependent on the expression of specific factors. Glial cell-derived neurotrophic factor (GDNF) is the primary inducer of ureteric budding (Costantini and Shakya, 2006). GDNF interacts with several different proteins from the MM (eg, Wnt-1, Pax2, Eyal, Six1, Sall 1) and from the UB itself (Pax2, Lim1, Ret) resulting in outgrowth of the UB (reviewed by Shah et al, 2004). Additional specific factors are required for (1) early branching (eg, Wnt-4 and 11, fgf...
EMBRYOLOGY OF THE GENITOURINARY SYSTEM

7-10); (2) late branching and maturation (bmp2, activin); and (3) branching termination and tubule maintenance (HGF, TGF-alpha, EGFr) (reviewed by Shah et al, 2004). BMP-7, SHH and Wnt-11 produced from the branching ureteric bud induce the MM to differentiate. These factors induce the activation of Pax-2, alpha-8-integrin and Wnt-4 in the renal mesenchymal cells, resulting in condensation of the MM and the formation of pretubular aggregate and primitive renal vesicle (reviewed by Burrow 2000). With the continued induction from the UB and the autocrine activity of Wnt-4, the pretubular aggregates differentiate into comma-shaped bodies. PDGF-beta and vEGF expression are required for initiating the migration of endothelial cells into the cleft of the comma-shaped bodies to form rudimentary glomerular capillary tufts (reviewed by Burrow 2000). Wt-1 and Pod-1 may have important functions in the regulation of gene transcription necessary for the differentiation of podocytes (Ballermann 2005).

ANOMALIES OF THE NEPHRIC SYSTEM

Failure of the metanephros to ascend leads to an ectopic kidney. An ectopic kidney may be on the proper side but low (simple ectopy) or on the opposite side (crossed

Figure 2–2. Progressive stages in the differentiation of the nephrons and their linkage with the branching collecting tubules. A small lump of metanephric tissue is associated with each terminal collecting tubule. These are then arranged in vesicular masses that later differentiate into a uniferous tubule draining into the duct near which it arises. At one end, Bowman’s capsule and the glomerulus differentiate; the other end establishes communication with the nearby collecting tubules.
ectopy) with or without fusion. Failure to rotate during ascent causes a **malrotated kidney**.

Fusion of the paired metanephric masses leads to various anomalies—most commonly a **horseshoe kidney**.

The ureteral bud from the mesonephric duct may bifurcate, causing a **bifid ureter** at various levels depending on the time of the bud’s subdivision. An accessory ureteral bud may develop from the mesonephric duct, thereby forming a **duplicated ureter**, usually meeting the same metanephric mass. Rarely, each bud has a separate metanephric mass, resulting in **supernumerary kidneys**.

If the double ureteral buds are close together on the mesonephric duct, they open near each other in the bladder. In this case, the main ureteral bud, which is the first to appear and the most caudal on the mesonephric ducts, reaches the bladder first. It then starts to move upward and laterally and is followed later by the second accessory bud as it reaches the urogenital sinus. The main ureteral bud (now more cranial on the urogenital sinus) drains the lower portion of the kidney. The 2 ureteral buds reverse their relationship as they move from the mesonephric duct to the urogenital sinus. This is why double ureters always cross (Weigert-Meyer law). If the 2 ureteral buds are widely separated on the mesonephric duct, the accessory bud appears more proximal and ends in the bladder with an ectopic orifice lower than the normal one. This ectopic orifice could still be in the bladder close to its outlet, in the urethra, or even in the genital duct system (Figure 2–3). A single ureteral bud that arises higher than normal on the mesonephric duct can also end in a similar ectopic location.

Lack of development of a ureteral bud results in a **solitary kidney** and a hemitrigone.

![Figure 2–3](image-url). The development of the ureteral bud from the mesonephric duct and the relationship of both to the urogenital sinus. The ureteral bud appears at the 4th week. The mesonephric duct distal to this ureteral bud is gradually absorbed into the urogenital sinus, resulting in separate endings for the ureter and the mesonephric duct. The mesonephric tissue that is incorporated into the urogenital sinus expands and forms the trigonal tissue.
VESICOURETHRAL UNIT

The blind end of the hindgut caudal to the point of origin of the allantois expands to form the cloaca, which is separated from the outside by a thin plate of tissue (the cloacal membrane) lying in an ectodermal depression (the proctodeum) under the root of the tail. At the 4-mm stage, starting at the cephalic portion of the cloaca where the allantois and gut meet, the cloaca progressively divides into 2 compartments by the caudal growth of a crescentic fold, the urorectal fold. The 2 limbs of the fold bulge into the lumen of the cloaca from either side, eventually meeting and fusing. The division of the cloaca into a ventral portion (urogenital sinus) and a dorsal portion (rectum) is completed during the 7th week. During the development of the urorectal septum, the cloacal membrane undergoes a reverse rotation, so that the ectodermal surface is no longer directed toward the developing anterior abdominal wall but gradually is turned to face caudally and slightly posteriorly. This change facilitates the subdivision of the cloaca and is brought about mainly by development of the intraumbilical portion of the anterior abdominal wall and regression of the tail. The mesoderm that passes around the cloacal membrane to the caudal attachment of the umbilical cord proliferates and grows, forming a surface elevation, the genital tubercle. Further growth of the intraumbilical part of the abdominal wall progressively separates the umbilical cord from the genital tubercle. The division of the cloaca is completed before the cloacal membrane ruptures, and its 2 parts therefore have separate openings. The ventral part is the primitive urogenital sinus, which has the shape of an elongated cylinder and is continuous cranially with the allantois; its external opening is the urogenital ostium. The division of the cloaca and meatus is no longer directed toward the developing anterior abdominal wall but gradually is turned to face caudally and slightly posteriorly. This change facilitates the subdivision of the cloaca and is brought about mainly by development of the intraumbilical portion of the anterior abdominal wall and regression of the tail. The mesoderm that passes around the cloacal membrane to the caudal attachment of the umbilical cord proliferates and grows, forming a surface elevation, the genital tubercle. Further growth of the intraumbilical part of the abdominal wall progressively separates the umbilical cord from the genital tubercle. The division of the cloaca is completed before the cloacal membrane ruptures, and its 2 parts therefore have separate openings. The ventral part is the primitive urogenital sinus, which has the shape of an elongated cylinder and is continuous cranially with the allantois; its external opening is the urogenital ostium. The dorsal part is the rectum, and its external opening is the anus.

The urogenital sinus receives the mesonephric ducts. The caudal end of the mesonephric duct distal to the ureteral bud is progressively absorbed into the urogenital sinus. By the 7th week, the mesonephric duct and the ureteral bud have independent opening sites. This introduces an island of mesodermal tissue amid the surrounding endoderm of the urogenital sinus. As development progresses, the opening of the mesonephric duct (which will become the ejaculatory duct) migrates downward and medially. The opening of the ureteral bud (which will become the ureteral orifice) migrates upward and laterally. The absorbed mesoderm of the mesonephric duct expands with this migration to occupy the area limited by the final position of these tubes (Figure 2–3). This will later be differentiated as the trigonal structure, which is the only mesodermal inclusion in the endodermal vesicourethral unit.

The urogenital sinus can be divided into 2 main segments. The dividing line, the junction of the combined Müllerian ducts with the dorsal wall of the urogenital sinus, is an elevation called Müller’s tubercle, which is the most fixed reference point in the whole structure and which is discussed in a subsequent section. The segments are as follows:

1. The ventral and pelvic portion forms the bladder, part of the urethra in males, and the whole urethra in females. This portion receives the ureter.
2. The urethral, or phallic, portion receives the mesonephric and the fused Müllerian ducts. This will be part of the urethra in males and forms the lower fifth of the vagina and the vaginal vestibule in females.

During the third month, the ventral part of the urogenital sinus starts to expand and forms an epithelial sac whose apex tapers into an elongated, narrowed urachus. The pelvic portion remains narrow and tubular; it forms the whole urethra in females and the supramontanl portion of the prostatic urethra in males. The splanchnic mesoderm surrounding the ventral and pelvic portion of the urogenital sinus begins to differentiate into interlacing bands of smooth muscle fibers and an outer fibrous connective tissue coat. By the 12th week, the layers characteristic of the adult urethra and bladder are recognizable (Figure 2–4).

The part of the urogenital sinus caudal to the opening of the Müllerian duct forms the vaginal vestibule and contributes to the lower fifth of the vagina in females (Figure 2–5). In males, it forms the inframontatal part of the prostatic urethra and the membranous urethra. The penile urethra is formed by the fusion of the urethral folds on the ventral surface of the genital tubercle. In females, the urethral folds remain separate and form the labia minora. The glandular urethra in males is formed by canalization of the urethral plate. The bladder originally extends up to the umbilicus, where it is connected to the allantois that extends into the umbilical cord. The allantois usually is obliterated at the level of the umbilicus by the 15th week. The bladder then starts to descend by the 18th week. As it descends, its apex becomes stretched and narrowed, and it pulls on the already obliterated allantois, now called the urachus. By the 20th week, the bladder is well separated from the umbilicus, and the stretched urachus becomes the middle umbilical ligament.

PROSTATE

The prostate develops as multiple solid outgrowths of the urethral epithelium both above and below the entrance of the mesonephric duct. These simple tubular outgrowths begin to develop in 5 distinct groups at the end of the 11th week and are complete by the 16th week (112-mm stage). They branch and rebranch, ending in a complex duct system that encounters the differentiating mesenchymal cells.
around this segment of the urogenital sinus. These mesenchymal cells start to develop around the tubules by the 16th week and become denser at the periphery to form the prostatic capsule. By the 22nd week, the muscular stroma is considerably developed, and it continues to increase progressively until birth.

From the 5 groups of epithelial buds, 5 lobes are eventually formed: anterior, posterior, median, and 2 lateral lobes. Initially, these lobes are widely separated, but later they meet, with no definite septa dividing them. Tubules of each lobe do not intermingle with each other but simply lie side by side.

The anterior lobe tubules begin to develop simultaneously with those of the other lobes. Although in the early stages, the anterior lobe tubules are large and show multiple branches; gradually they contract and lose most of the branches. They continue to shrink, so that at birth they show no lumen and appear as small, solid embryonic epithelial outgrowths. In contrast, the tubules of the posterior lobe are fewer in number yet larger, with extensive branching. These tubules, as they grow, extend posterior to the developing median and lateral lobes and form the posterior aspect of the gland, which may be felt rectally.

**ANOMALIES OF THE VESICOURETHRAL UNIT**

Failure of the cloaca to subdivide is rare and results in a persistent cloaca. Incomplete subdivision is more frequent, ending with rectovesical, rectourethral, or rectovestibular fistulas (usually with imperforate anus or anal atresia).

Failure of descent or incomplete descent of the bladder leads to a urinary umbilical fistula (urachal fistula), urachal cyst, or urachal diverticulum depending on the stage and degree of maldescent.

Development of the genital primordia in an area more caudal than normal can result in formation of the corpora cavernosa just caudal to the urogenital sinus outlet, with the urethral groove on its dorsal surface. This defect results in complete or incomplete epispadias depending on its degree. A more extensive defect results in vesical extrophy. Failure of fusion of urethral folds leads to various grades of hypospadias. This defect, because of its mechanism, never extends proximal to the bulbous urethra. This is in contrast to epispadias, which usually involves the entire urethra up to the internal meatus.
GONADS

Most of the structures that make up the embryonic genital system have been taken over from other systems, and their readaptation to genital function is a secondary and relatively late phase in their development. The early differentiation of such structures is therefore independent of sexuality. Furthermore, each embryo is at first morphologically bisexual, possessing all the necessary structures for either sex. The development of one set of sex primordia and the gradual involution of the other are determined by the sex of the gonad.

The sexually undifferentiated gonad is a composite structure. Male and female potentials are represented by specific histologic elements (medulla and cortex) that have alternative roles in gonadogenesis. Normal differentiation involves the gradual predominance of one component.

The primitive sex glands make their appearance during the 5th and 6th weeks within a localized region of the thickening known as the urogenital ridge (this contains both the nephric and the genital primordia). At the 6th week, the gonad consists of a superficial germinal epithelium and an internal blastema. The blastemal mass is derived mainly from proliferative ingrowth from the superficial epithelium, which comes loose from its basement membrane.

During the 7th week, the gonad begins to assume the characteristics of a testis or ovary. Differentiation of the ovary usually occurs somewhat later than differentiation of the testis.

If the gonad develops into a testis, the gland increases in size and shortens into a more compact organ while achieving a more caudal location. Its broad attachment to the mesonephros is converted into a gonadal mesentery known as the mesorchium. The cells of the germinal epithelium grow into the underlying mesenchyme and form cordlike masses. These are radially arranged and converge toward the mesorchium, where a dense portion of the blastemal mass is also emerging as the primordium of the rete testis. A network of strands soon forms that is continuous with the testis cords. The latter also split into 3–4 daughter cords. These eventually become differentiated into the seminiferous tubules by which the spermatozoa are produced. The rete testis unites with the mesonephric components that will form the male genital ducts, as discussed in a subsequent section (Figure 2–6).

If the gonad develops into an ovary, it (like the testis) gains a mesentery (mesovarium) and settles in a more

Figure 2–5. Differentiation of the urogenital sinus and the Müllerian ducts in the female embryo. At 9 weeks, the urogenital sinus receives the fused Müllerian ducts at Müller’s tubercle (sinovaginal node), which is solidly packed with cells. As the urogenital sinus distal to Müller’s tubercle becomes wider and shallower (15 weeks), the urethra and fused Müllerian duct will have separate openings. The distal part of the urogenital sinus forms the vaginal vestibule and the lower fifth of the vagina (shaded area), and that part above Müller’s tubercle forms the urinary bladder and the entire female urethra. The fused Müllerian ducts form the uterus and the upper four-fifths of the vagina. The hymen is formed at the junction of the sinovaginal node and the urogenital sinus.
Figure 2–6. Transformation of the undifferentiated genital system into the definitive male and female systems.
caudal position. The internal blastema differentiates in the 9th week into a primary cortex beneath the germinal epithelium and a loose primary medulla. A compact cellular mass bulges from the medulla into the mesovarium and establishes the primitive rete ovarii. At 3–4 months of age, the internal cell mass becomes young ova. A new definitive cortex is formed from the germinal epithelium as well as from the blastema in the form of distinct cellular cords (Pflüger’s tubes), and a permanent medulla is formed. The cortex differentiates into ovarian follicles containing ova.

**Descent of the Gonads**

**A. TESTIS**

In addition to its early caudal migration, the testis later leaves the abdominal cavity and descends into the scrotum. By the third month of fetal life, the testis is located retroperitoneally in the false pelvis. A fibromuscular band (the gubernaculum) extends from the lower pole of the testis through the developing muscular layers of the anterior abdominal wall to terminate in the subcutaneous tissue of the scrotal swelling. The gubernaculum also has several other subsidiary strands that extend to adjacent regions. Just below the lower pole of the testis, the peritoneum herniates as a diverticulum along the anterior aspect of the gubernaculum, eventually reaching the scrotal sac through the anterior abdominal muscles (the processus vaginalis). The testis remains at the abdominal end of the inguinal canal until the seventh month. It then passes through the inguinal canal behind (but invaginating) the processus vaginalis. Normally, it reaches the scrotal sac by the end of the eighth month.

**B. OVARY**

In addition to undergoing an early internal descent, the ovary becomes attached through the gubernaculum to the tissues of the genital fold and then attaches itself to the developing urovaginal canal at its junction with the uterine (fallopian) tubes. This part of the gubernaculum between the ovary and uterus becomes the ovarian ligament; the part between the uterus and the labia majora becomes the round ligament of the uterus. These ligaments prevent extra-abdominal descent, and the ovary enters the true pelvis. It eventually lies posterior to the uterine tubes on the superior surface of the urogenital mesentery, which has descended with the ovary and now forms the broad ligament. A small processus vaginalis forms and passes toward the labial swelling, but it is usually obliterated at full term.

**GONADAL ANOMALIES**

Lack of development of the gonads is called **gonadal agenesis**. Incomplete development with arrest at a certain phase is called **hypogenesis**. **Supernumerary gonads** are rare. The commonest anomaly involves descent of the gonads, especially the testis. Retention of the testis in the abdomen or arrest of its descent at any point along its natural pathway is called **cryptorchidism**, which may be either unilateral or bilateral. If the testis does not follow the main gubernacular structure but follows one of its subsidiary strands, it will end in an abnormal position, resulting in an **ectopic testis**.

Failure of union between the rete testis and mesonephric results in a testis separate from the male genital ducts (the epididymis) and azoospermia.

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**GENITAL DUCT SYSTEM**

Alongside the indifferent gonads, there are, early in embryonic life, 2 different yet closely related ducts. One is primarily a nephric duct (Wolffian duct), yet it also serves as a genital duct if the embryo develops into a male. The other (Müllerian duct) is primarily a genital structure from the start.

Both ducts grow caudally to join the primitive urogenital sinus. The Wolffian duct (known as the pronephric duct at the 4-mm stage) joins the ventral part of the cloaca, which will be the urogenital sinus. This duct gives rise to the ureteral bud close to its caudal end. The ureteral bud grows cranially and meets metanephrogenic tissue. The part of each mesonephric duct caudal to the origin of the ureteric bud becomes absorbed into the wall of the primitive urogenital sinus, so that the mesonephric duct and ureter open independently. This is achieved at the 15-mm stage (7th week). During this period, starting at the 10-mm stage, the Müllerian ducts start to develop. They reach the urogenital sinus relatively late—at the 30-mm stage (9th week)—their partially fused blind ends producing the elevation called Müller’s tubercle. Müller’s tubercle is the most constant and reliable point of reference in the whole system.

If the gonad starts to develop into a testis (17-mm stage, 7th week), the Wolffian duct will start to differentiate into the male duct system, forming the epididymis, vas deferens, seminal vesicles, and ejaculatory ducts. At this time, the Müllerian duct proceeds toward its junction with the urogenital sinus and immediately starts to degenerate. Only its upper and lower ends persist, the former as the appendix testis and the latter as part of the prostatic utricle.

If the gonad starts to differentiate into an ovary (22-mm stage, 8th week), the Müllerian duct system forms the uterine (fallopian) tubes, uterus, and most of the vagina. The Wolffian ducts, aside from their contribution to the urogenital sinus, remain rudimentary.
MALE DUCT SYSTEM

Epididymis

Because of the proximity of the differentiating gonads and the nephric duct, some of the mesonephric tubules are retained as the efferent ductules, and their lumens become continuous with those of the rete testis. These tubules, together with the part of the mesonephric duct into which they empty, will form the epididymis. Each coiled ductule makes a conical mass known as the lobule of the epididymis. The cranial end of the mesonephric duct becomes highly convoluted, completing the formation of the epididymis. This is an example of direct inclusion of a nephric structure into the genital system. Additional mesonephric tubules, both cephalad and caudal to those that were included in the formation of the epididymis, remain as rudimentary structures, that is, the appendix of the epididymis and the paradidymis.

Vas Deferens, Seminal Vesicles, & Ejaculatory Ducts

The mesonephric duct caudal to the portion forming the epididymis forms the vas deferens. Shortly before this duct joins the urethra (urogenital sinus), a localized dilatation (ampulla) develops, and the saccular convoluted structure that will form the seminal vesicle is evaginated from its wall. The mesonephric duct between the origin of the seminal vesicle and the urethra forms the ejaculatory duct. The whole mesonephric duct now achieves its characteristic thick investment of smooth muscle, with a narrow lumen along most of its length.

Both above and below the point of entrance of the mesonephric duct into the urethra, multiple outgrowths of urethral epithelium mark the beginning of the development of the prostate. As these epithelial buds grow, they meet the developing muscular fibers around the urogenital sinus, and some of these fibers become entangled in the branching tubules of the growing prostate and become incorporated into it, forming its muscular stroma (Figure 2–4).

FEMALE DUCT SYSTEM

The Müllerian ducts, which are a paired system, are seen alongside the mesonephric duct. It is not known whether they arise directly from the mesonephric ducts or separately as an invagination of the celomic epithelium into the parenchyma lateral to the cranial extremity of the mesonephric duct, but the latter theory is favored. The Müllerian duct develops and runs lateral to the mesonephric duct. Its opening into the celomic cavity persists as the peritoneal ostium of the uterine tube (later it develops fimбриae). The other end grows caudally as a solid tip and then crosses in front of the mesonephric duct at the caudal extremity of the mesonephros. It continues its growth in a caudomedial direction until it meets and fuses with the Müllerian duct of the opposite side. The fusion is partial at first, so there is a temporary septum between the 2 lumens. This later disappears, leaving one cavity that will form the uterovaginal canal. The potential lumen of the vaginal canal is completely packed with cells. The solid tip of this cord pushes the epithelium of the urogenital sinus outward, where it becomes Müller’s tubercle (33-mm stage, 9th week). The Müllerian ducts actually fuse at the 63-mm stage (13th week), forming the sinovaginal node, which receives a limited contribution from the urogenital sinus. (This contribution forms the lower fifth of the vagina.)

The urogenital sinus distal to Müller’s tubercle, originally narrow and deep, shortens, widens, and opens to form the floor of the pudendal or vulval cleft. This results in separate openings for the vagina and urethra and also brings the vaginal orifice to its final position nearer the surface. At the same time, the vaginal segment increases appreciably in length. The vaginal vestibule is derived from the infratubercular segment of the urogenital sinus (in males, the same segment will form the inframontanal part of the prostatic urethra and the membranous urethra). The labia minora are formed from the urethral folds (in males they form the pendulous urethra). The hymen is the remnant of the Müllerian tubercle. The lower fifth of the vagina is derived from the portion of the urogenital sinus that combines with the sinovaginal node. The remainder of the vagina and the uterus are formed from the lower (fused) third of the Müllerian ducts. The uterine tubes (fallopian tubes, oviducts) are the cephalic two-thirds of the Müllerian ducts (Figure 2–6).

ANOMALIES OF THE GONADAL DUCT SYSTEM

Nonunion of the rete testis and the efferent ductules can occur and, if bilateral, causes azoospermia and sterility. Failure of the Müllerian ducts to approximate or to fuse completely can lead to various degrees of duplication in the genital ducts. Congenital absence of one or both ureterine tubes or of the uterus or vagina occurs rarely.

Arrested development of the infratubercular segment of the urogenital sinus leads to its persistence, with the urethra and vagina having a common duct to the outside (urogenital sinus).

EXTERNAL GENITALIA

During the 8th week, external sexual differentiation begins to occur. Not until 3 months, however, do the progressively developing external genitalia attain characteristics that can be recognized as distinctively male or female. During the indifferent stage of sexual development, 3
small protuberances appear on the external aspect of the cloacal membrane. In front is the genital tubercle, and on either side of the membrane are the genital swellings.

With the breakdown of the urogenital membrane (17-mm stage, 7th week), the primitive urogenital sinus achieves a separate opening on the undersurface of the genital tubercle.

**MALE EXTERNAL GENITALIA**

The urogenital sinus opening extends on the ventral aspect of the genital tubercle as the urethral groove. The primitive urogenital orifice and the urethral groove are bounded on either side by the urethral folds. The genital tubercle becomes elongated to form the phallus. The corpora cavernosa are indicated in the 7th week as paired mesenchymal columns within the shaft of the penis. By the 10th week, the urethral folds start to fuse from the urogenital sinus orifice toward the tip of the phallus. At the 14th week, the fusion is complete and results in the formation of the penile urethra. The corpus spongiosum results from the differentiation of the mesenchymal masses around the urethral groove. The glandular urethra develops as a result of canalization of an ectodermal epithelial cord that has grown through the glans. This canalization reaches and communicates with the distal end of the urogenital orifice and the urethral groove are bounded on either side of the membrane are the genital swellings. Meanwhile, the genital swellings shift caudally and, 2 months later, surrounds the glans. This forms the prepuce. Meanwhile, the genital swellings shift caudally and are recognizable as scrotal swellings. They meet and fuse, resulting in the formation of the scrotum, with 2 compartments partially separated by a median septum and a median raphe, indicating their line of fusion.

**FEMALE EXTERNAL GENITALIA**

Until the 8th week, the appearance of the female external genitalia closely resembles that of the male genitalia except that the urethral groove is shorter. The genital tubercle, which becomes bent caudally and lags in development, becomes the clitoris. As in males (though on a minor scale), mesenchymal columns differentiate into corpora cavernosa, and a coronary sulcus identifies the glans clitoridis. The most caudal part of the urogenital sinus shortens and widens, forming the vaginal vestibule. The urethral folds do not fuse but remain separate as the labia minora. The genital swellings meet in front of the anus, forming the posterior commissure, while the swellings as a whole enlarge and remain separated on either side of the vestibule and form the labia majora.

**ANOMALIES OF THE EXTERNAL GENITALIA**

Absence or duplication of the penis or clitoris is very rare. More commonly, the penis remains rudimentary or the clitoris shows hypertrophy. These anomalies may be seen alone or, more frequently, in association with pseudohermaphroditism. Concealed penis and transposition of penis and scrotum are relatively rare anomalies.

Failure or incomplete fusion of the urethral folds results in **hypospadias** (see preceding discussion). Penile development is also anomalous in cases of **epispadias** and **extrophy** (see preceding discussion).

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Gonadal Anomalies


Symptoms of Disorders of the Genitourinary Tract

Jack W. McAninch, MD, FACS

In the workup of any patient, the history is of paramount importance; this is particularly true in urology. It is necessary to discuss here only those urologic symptoms that are apt to be brought to the physician’s attention by the patient. It is important to know not only whether the disease is acute or chronic but also whether it is recurrent, since recurring symptoms may represent acute exacerbations of chronic disease.

Obtaining the history is an art that depends on the skill and methods used to elicit information. The history is only as accurate as the patient’s ability to describe the symptoms. This subjective information is important in establishing an accurate diagnosis.

SYSTEMIC MANIFESTATIONS

Symptoms of fever and weight loss should be sought. The presence of fever associated with other symptoms of urinary tract infection may be helpful in evaluating the site of the infection. Simple acute cystitis is essentially an afebrile disease. Acute pyelonephritis or prostatitis is apt to cause high temperatures (to 40°C [104°F]), often accompanied by violent chills. Infants and children who have acute pyelonephritis may have high temperatures without other localizing symptoms or signs. Such a clinical picture, therefore, invariably requires bacteriologic study of the urine.

A history of unexplained attacks of fever occurring even years before may represent otherwise asymptomatic pyelonephritis. Renal carcinoma sometimes causes fever that may reach 39°C (102.2°F) or more. The absence of fever does not by any means rule out renal infection, for it is the rule that chronic pyelonephritis does not cause fever.

Weight loss is to be expected in the advanced stages of cancer, but it may be noticed also when renal insufficiency due to obstruction or infection supervenes. In children who have “failure to thrive” (low weight and less than average height for age), chronic obstruction, urinary tract infection, or both should be suspected.

General malaise may be noted with tumors, chronic pyelonephritis, or renal failure. The presence of many of these symptoms may be compatible with human immunodeficiency virus (HIV; see Chapter 15).

LOCAL & REFERRED PAIN

Two types of pain have their origins in the genitourinary organs: local and referred. The latter is especially common.

Local pain is felt in or near the involved organ. Thus, the pain from a diseased kidney (T10–12, L1) is felt in the costovertebral angle and in the flank in the region of and below the 12th rib. Pain from an inflamed testicle is felt in the gonad itself.

Referred pain originates in a diseased organ but is felt at some distance from that organ. The ureteral colic (Figure 3–1) caused by a stone in the upper ureter may be associated with severe pain in the ipsilateral testicle; this is explained by the common innervation of these 2 structures (T11–12). A stone in the lower ureter may cause pain referred to the scrotal wall; in this instance, the testis itself is not hyperesthetic. The burning pain with voiding that accompanies acute cystitis is felt in the distal urethra in females and in the glandular urethra in males (S2–3).

Abnormalities of a urologic organ can also cause pain in any other organ (eg, gastrointestinal, gynecologic) that has a sensory nerve supply common to both (Figures 3–2 and 3–3).

Kidney Pain (Figure 3–1)

Typical renal pain is felt as a dull and constant ache in the costovertebral angle just lateral to the sacrospinalis muscle and just below the 12th rib. This pain often spreads along the subcostal area toward the umbilicus or lower abdominal quadrant. It may be expected in the renal diseases that cause sudden distention of the renal capsule. Acute pyelonephritis (with its sudden edema) and acute ureteral obstruction (with its sudden renal back pressure) both cause this typical pain. It should be pointed out, however, that many urologic renal diseases are painless because their progression is so slow that sudden capsular distention does not occur. Such diseases include cancer, chronic pyelonephritis, staghorn calculus, tuberculosis, polycystic kidney, and hydronephrosis due to chronic ureteral obstruction.
Ureteral Pain (Figure 3–1)

Ureteral pain is typically stimulated by acute obstruction (passage of a stone or a blood clot). In this instance, there is back pain from renal capsular distention combined with severe colicky pain (due to renal pelvic and ureteral muscle spasm) that radiates from the costovertebral angle down toward the lower anterior abdominal quadrant, along the course of the ureter. In men, it may also be felt in the bladder, scrotum, or testicle. In women, it may radiate into the vulva. The severity and colicky nature of this pain are caused by the hyperperistalsis and spasm of this smooth muscle organ as it attempts to rid itself of a foreign body or to overcome obstruction.

The physician may be able to judge the position of a ureteral stone by the history of pain and the site of referral. If the stone is lodged in the upper ureter, the pain radiates to the testicle, since the nerve supply of this organ is similar to that of the kidney and upper ureter (T11–12). With stones in the midportion of the ureter on the right side, the pain is referred to McBurney’s point and may therefore simulate appendicitis; on the left side, it may resemble diverticulitis or other diseases of the descending or sigmoid colon (T12, L1). As the stone approaches the bladder, inflammation and edema of the ureteral orifice ensue, and symptoms of vesical irritability such as urinary frequency and urgency may occur. It is important to realize, however, that in mild ureteral obstruction, as seen in the congenital stenoses, there is usually no pain, either renal or ureteral.

Vesical Pain

The overdistended bladder of the patient in acute urinary retention causes agonizing pain in the suprapubic area. Other than this, however, constant suprapubic pain not related to the act of urination is usually not of urologic origin.
Figure 3-2. Diagrammatic representation of autonomic nerve supply to gastrointestinal and genitourinary tracts.
Figure 3–3. Diagrammatic representation of sensory nerves of gastrointestinal and genitourinary tracts.
The patient in chronic urinary retention due to bladder neck obstruction or neurogenic bladder may experience little or no suprapubic discomfort even though the bladder reaches the level of the umbilicus.

The most common cause of bladder pain is infection; the pain is usually not felt over the bladder but is referred to the distal urethra and is related to the act of urination. Terminal dysuria may be a major complaint in severe cystitis.

**Prostatic Pain**

Direct pain from the prostate gland is not common. Occasionally, when the prostate is acutely inflamed, the patient may feel a vague discomfort or fullness in the perineal or rectal area (S2–4). Lumbosacral backache is occasionally experienced as referred pain from the prostate, but is not a common symptom of prostatitis. Inflammation of the gland may cause dysuria, frequency, and urgency.

**Testicular Pain**

Testicular pain due to trauma, infection, or torsion of the spermatic cord is very severe and is felt locally, although there may be some radiation of the discomfort along the spermatic cord into the lower abdomen. Uninfected hydrocele, spermatocele, and tumor of the testis do not commonly cause pain. A varicocele may cause a dull ache in the testicle that is increased after heavy exercise. At times, the first symptom of an early indirect inguinal hernia may be testicular pain (referred). Pain from a stone in the upper ureter has typical renal and ureteral colic and, usually, hematuria and may experience severe nausea and vomiting as well as abdominal distention. However, the urinary symptoms so far overshadow the gastrointestinal symptoms that the latter are usually ignored. Inadvertent overdistention of the renal pelvis (eg, with opaque material in order to obtain adequate retrograde urograms) may cause the patient to become nauseated, to vomit, and to complain of cramplike pain in the abdomen. This clinical experiment demonstrates the renointestinal reflex, which may lead to confusing symptoms. In the very common “silent” urologic diseases, some degree of gastrointestinal symptomatology may be present, which could mislead the clinician into seeking the diagnosis in the intraperitoneal zone.

**Epididymal Pain**

Acute infection of the epididymis is the only painful disease of this organ and is quite common. The pain begins in the scrotum, and some degree of neighborhood inflammatory reaction involves the adjacent testis as well, further aggravating the discomfort. In the early stages of epididymitis, pain may first be felt in the groin or lower abdominal quadrant. (If on the right side, it may simulate appendicitis.) This may be a referred type of pain but can be secondary to associated inflammation of the vas deferens.

**GASTROINTESTINAL SYMPTOMS OF UROLOGIC DISEASES**

Whether renal or ureteral disease is painful or not, gastrointestinal symptoms are often present. The patient with acute pyelonephritis not only has localized back pain, symptoms of vesical irritability, chills, and fever but also generalized abdominal pain and distention. A patient who is passing a stone down the ureter has typical renal and ureteral colic and, usually, hematuria and may experience severe nausea and vomiting as well as abdominal distention. However, the urinary symptoms so far overshadow the gastrointestinal symptoms that the latter are usually ignored. Inadvertent overdistention of the renal pelvis (eg, with opaque material in order to obtain adequate retrograde urograms) may cause the patient to become nauseated, to vomit, and to complain of cramplike pain in the abdomen. This clinical experiment demonstrates the renointestinal reflex, which may lead to confusing symptoms. In the very common “silent” urologic diseases, some degree of gastrointestinal symptomatology may be present, which could mislead the clinician into seeking the diagnosis in the intraperitoneal zone.

**Cause of the Mimicry**

**A. RENOINTESTINAL REFLEXES**

Renointestinal reflexes account for most of the confusion. They arise because of the common autonomic and sensory innervations of the two systems (Figures 3–2 and 3–3). Afferent stimuli from the renal capsule or musculature of the pelvis may, by reflex action, cause pylorospasm (symptoms of peptic ulcer) or other changes in tone of the smooth muscles of the enteric tract and its adnexa.

**B. ORGAN RELATIONSHIPS**

The right kidney is closely related to the hepatic flexure of the colon, the duodenum, the head of the pancreas, the common bile duct, the liver, and the gallbladder (Figure 1–3). The left kidney lies just behind the splenic flexure of the colon and is closely related to the stomach, pancreas, and spleen. Inflammations or tumors in the retroperitoneum thus may extend into or displace intraperitoneal organs, causing them to produce symptoms.

**C. PERITONEAL IRRITATION**

The anterior surfaces of the kidneys are covered by peritoneum. Renal inflammation, therefore, causes peritoneal irritation, which can lead to muscle rigidity and rebound tenderness.

The symptoms arising from chronic renal disease (eg, noninfected hydronephrosis, staghorn calculus, cancer, chronic pyelonephritis) may be entirely gastrointestinal and may simulate in every way the syndromes of peptic ulcer, gallbladder disease, or appendicitis, or other, less specific gastrointestinal complaints. If a thorough survey of the gastrointestinal tract fails to demonstrate suspected disease processes, the physician should give every consideration to study of the urinary tract.
SYMPTOMS RELATED TO THE ACT OF URINATION

Many conditions cause symptoms of “cystitis.” These include infections of the bladder, vesical inflammation due to chemical or x-radiation reactions, interstitial cystitis, prostatitis, psychoneurosis, torsion or rupture of an ovarian cyst, and foreign bodies in the bladder. Often, however, the patient with chronic cystitis notices no symptoms of vesical irritability. Irritating chemicals or soap on the urethral meatus may cause cystitis-like symptoms of dysuria, frequency, and urgency. This has been specifically noted in young girls taking frequent bubble baths.

Frequency, Nocturia, & Urgency

The normal capacity of the bladder is about 400 mL. Frequency may be caused by residual urine, which decreases the functional capacity of the organ. When the mucosa, submucosa, and even the muscularis become inflamed (eg, infection, foreign body, stones, tumor), the capacity of the bladder decreases sharply. This decrease is due to 2 factors: the pain resulting from even mild stretching of the bladder and the loss of bladder compliance resulting from inflammatory edema. When the bladder is normal, urination can be delayed if circumstances require it, but this is not so in acute cystitis. Once the diminished bladder capacity is reached, any further distention may be agonizing, and the patient may urinate involuntarily if voiding does not occur immediately. During very severe acute infections, the desire to urinate may be constant, and each voiding may produce only a few milliliters of urine. Day frequency without nocturia and acute or chronic frequency lasting only a few hours suggest nervous tension.

Diseases that cause fibrosis of the bladder are accompanied by frequency of urination. Examples of such diseases are tuberculosis, radiation cystitis, interstitial cystitis, and schistosomiasis. The presence of stones or foreign bodies causes vesical irritability, but secondary infection is almost always present.

Nocturia may be a symptom of renal disease related to a decrease in the functioning renal parenchyma with loss of concentrating power. Nocturia can occur in the absence of disease in persons who drink excessive amounts of fluid in the late evening. Coffee and alcoholic beverages, because of their specific diuretic effect, often produce nocturia if consumed just before bedtime. In older people who are ambulatory, some fluid retention may develop secondary to mild heart failure or varicose veins. With recumbency at night, this fluid is mobilized, leading to nocturia in these patients. A very low or very high urine pH can irritate the bladder and cause frequency of urination.

Dysuria

Painful urination is usually related to acute inflammation of the bladder, urethra, or prostate. At times, the pain is described as “burning” on urination and is usually located in the distal urethra in men. Women usually localize the pain to the urethra. The pain is present only with voiding and disappears soon after micturition is completed. More severe pain sometimes occurs in the bladder just at the end of voiding, suggesting that inflammation of the bladder is the likely cause. Pain also may be more marked at the beginning of or throughout the act of urination. Dysuria often is the first symptom suggesting urinary infection and is often associated with urinary frequency and urgency.

Enuresis

Strictly speaking, enuresis means bedwetting at night. It is physiologic during the first 2 or 3 years of life but becomes troublesome, particularly to parents, after that age. It may be functional or secondary to delayed neuromuscular maturation of the urethrovessical component, but it may present as a symptom of organic disease (eg, infection, distal urethral stenosis in girls, posterior urethral valves in boys, neurogenic bladder). If wetting occurs also during the daytime, however, or if there are other urinary symptoms, urologic investigation is essential. In adult life, enuresis may be replaced by nocturia for which no organic basis can be found.

Symptoms of Bladder Outlet Obstruction

A. Hesitancy

Hesitancy in initiating the urinary stream is one of the early symptoms of bladder outlet obstruction. As the degree of obstruction increases, hesitancy is prolonged and the patient often strains to force urine through the obstruction. Prostate obstruction and urethral stricture are common causes of this symptom.

B. Loss of Force and Decrease of Caliber of the Stream

Progressive loss of force and caliber of the urinary stream is noted as urethral resistance increases despite the generation of increased intravesical pressure. This can be evaluated by measuring urinary flow rates; in normal circumstances with a full bladder a maximal flow of 20 mL/s should be achieved.

C. Terminal Dribbling

Terminal dribbling becomes more and more noticeable as obstruction progresses and is a most distressing symptom.

D. Urgency

A strong, sudden desire to urinate is caused by hyperactivity and irritability of the bladder, resulting from obstruction, inflammation, or neuropathic bladder disease. In most circumstances, the patient is able to control temporarily the
sudden need to void, but loss of small amounts of urine may occur (urgency incontinence).

E. ACUTE URINARY RETENTION

Sudden inability to urinate may supervene. The patient experiences increasingly agonizing suprapubic pain associated with severe urgency and may dribble only small amounts of urine.

F. CHRONIC URINARY RETENTION

Chronic urinary retention may cause little discomfort to the patient even though there is great hesitancy in starting the stream and marked reduction of its force and caliber. Constant dribbling of urine (paradoxical incontinence) may be experienced; it may be likened to water pouring over a dam.

G. INTERRUPTION OF THE URINARY STREAM

Interruption may be abrupt and accompanied by severe pain radiating down the urethra. This type of reaction strongly suggests the complication of vesical calculus.

H. SENSE OF RESIDUAL URINE

The patient often feels that urine is still in the bladder even after urination has been completed.

I. CYSTITIS

Recurring episodes of acute cystitis suggest the presence of residual urine.

Incontinence (See also Chapter 27)

There are many reasons for incontinence. The history often gives a clue to its cause.

A. TRUE INCONTINENCE

The patient may lose urine without warning; this may be a constant or periodic symptom. The more obvious causes include previous radical prostatectomy, exstrophy of the bladder, epispadias, vesicovaginal fistula, and ectopic ureteral orifice. Injury to the urethral smooth muscle sphincters may occur during prostatectomy or childbirth. Congenital or acquired neurogenic diseases may lead to dysfunction of the bladder and incontinence.

B. STRESS INCONTINENCE

When slight weakness of the sphincteric mechanisms is present, urine may be lost in association with physical strain (eg, coughing, laughing, rising from a chair). This is common in multiparous women who have weakened muscle support of the bladder neck and urethra and in men who have undergone radical prostatectomy. Occasionally, neuropathic bladder dysfunction can cause stress incontinence. The patient stays dry while lying in bed.

C. URGE INCONTINENCE

Urgency may be so precipitate and severe that there is involuntary loss of urine. Urge incontinence not infrequently occurs with acute cystitis, particularly in women, since women seem to have relatively poor anatomic sphincters. Urge incontinence is a common symptom of an upper motor neuron lesion.

D. OVERFLOW INCONTINENCE

Paradoxical incontinence is loss of urine due to chronic urinary retention or secondary to a flaccid bladder. The intravesical pressure finally equals the urethral resistance; urine then constantly dribbles forth.

Oliguria & Anuria

Oliguria and anuria may be caused by acute renal failure (due to shock or dehydration), fluid-ion imbalance, or bilateral ureteral obstruction.

Pneumaturia

The passage of gas in the urine strongly suggests a fistula between the urinary tract and the bowel. This occurs most commonly in the bladder or urethra but may be seen also in the ureter or renal pelvis. Carcinoma of the sigmoid colon, diverticulitis with abscess formation, regional enteritis, and trauma cause most vesical fistulas. Congenital anomalies account for most urethroenteric fistulas. Certain bacteria, by the process of fermentation, may liberate gas on rare occasions.

Cloudy Urine

Patients often complain of cloudy urine, but it is most often cloudy merely because it is alkaline; this causes precipitation of phosphate. Infection can also cause urine to be cloudy and malodorous. A properly performed urinalysis will reveal the cause of cloudiness.

Chyluria

The passage of lymphatic fluid or chyle is noted by the patient as passage of milky white urine. This represents a lymphatic–urinary system fistula. Most often, the cause is obstruction of the renal lymphatics, which results in forniceal rupture and leakage. Filariasis, trauma, tuberculosis, and retroperitoneal tumors have caused the problem.

Bloody Urine

Hematuria is a danger signal that cannot be ignored. Carcinoma of the kidney or bladder, calculi, and infection are a few of the conditions in which hematuria is typically demonstrable at the time of presentation. It is important to know whether urination is painful or not, whether the
hematuria is associated with symptoms of vesical irritability, and whether blood is seen in all or only a portion of the urinary stream. The hemoglobinuria that occurs as a feature of the hemolytic syndromes may also cause the urine to be red.

A. BLOODY URINE IN RELATION TO SYMPTOMS & DISEASES

Hematuria associated with renal colic suggests a ureteral stone, although a clot from a bleeding renal tumor can cause the same type of pain.

Hematuria is not uncommonly associated with non-specific, tuberculous, or schistosomal infection of the bladder. The bleeding is often terminal (bladder neck or prostate), although it may be present throughout urination (vesical or upper tract). Stone in the bladder often causes hematuria, but infection is usually present, and there are symptoms of bladder neck obstruction, neurogenic bladder, or cystocele.

Dilated veins may develop at the bladder neck secondary to enlargement of the prostate. These may rupture when the patient strains to urinate, resulting in gross or microscopic hematuria.

Hematuria without other symptoms (silent hematuria) must be regarded as a symptom of tumor of the bladder or kidney until proved otherwise. It is usually intermittent; bleeding may not recur for months. Complacency because the bleeding stops spontaneously must be condemned. Less common causes of silent hematuria are staghorn calculus, polycystic kidneys, benign prostatic hyperplasia, solitary renal cyst, sickle cell disease, and hydronephrosis.

Painless bleeding is common with acute glomerulonephritis. Recurrent bleeding is occasionally seen in children suffering from focal glomerulitis. Joggers and people who engage in participatory sports frequently develop transient proteinuria and gross or microscopic hematuria.

B. TIME OF HEMATURIA

Learning whether the hematuria is partial (initial, terminal) or total (present throughout urination) is often of help in identifying the site of bleeding. Initial hematuria suggests an anterior urethral lesion (eg, urethritis, stricture, meatal stenosis in young boys). Terminal hematuria usually arises from the posterior urethra, bladder neck, or trigone. Among the common causes are posterior urethritis and polyps and tumors of the vesical neck. Total hematuria has its source at or above the level of the bladder (eg, stone, tumor, tuberculosis, nephritis).

OTHER OBJECTIVE MANIFESTATIONS

Urethral Discharge

Urethral discharge in men is one of the most common urologic complaints. The causative organism is usually Neiseria gonorrhoeae or Chlamydia trachomatis. The discharge is often accompanied by local burning on urination or an itching sensation in the urethra (see Chapter 15).

Skin Lesions of the External Genitalia (See Chapters 15 & 40)

An ulceration of the glans penis or its shaft may represent syphilitic chancre, chancroid, herpes simplex, or squamous cell carcinoma. Venereal warts of the penis are common.

Visible or Palpable Masses

The patient may notice a visible or palpable mass in the upper abdomen that may represent renal tumor, hydronephrosis, or polycystic kidney. Enlarged lymph nodes in the neck may contain metastatic tumor from the prostate or testis. Lumps in the groin may represent spread of tumor of the penis or lymphadenitis from chancroid, syphilis, or lymphogranuloma venereum. Painless masses in the scrotal contents are common and include hydrocele, varicocele, spermatocele, chronic epididymitis, hernia, and testicular tumor.

Edema

Edema of the legs may result from compression of the iliac veins by lymphatic metastases from prostatic cancer. Edema of the genitalia suggests filariasis, chronic ascites, or lymphatic blockage from radiotherapy for pelvic malignancies.

Bloody Ejaculation

Inflammation of the prostate or seminal vesicles can cause hematospermia.

Gynecomastia

Often idiopathic, gynecomastia is common in elderly men, particularly those taking estrogens for control of prostatic cancer. It is also seen in association with choriocarcinoma and interstitial cell and Sertoli cell tumors of the testis. Certain endocrinologic diseases, for example, Klinefelter syndrome, may also cause gynecomastia.

COMPLAINTS RELATED TO SEXUAL PROBLEMS

Many people have genitourinary complaints on a purely psychological or emotional basis. In others, organic symptoms may be increased in severity because of tension states. It is important, therefore, to seek clues that might give evidence of emotional stress.

In women, the relationship of the menses to ureteral pain or vesical complaints should be determined,
although menstruation may exacerbate both organic and functional vesical and renal difficulties.

Many patients recognize that the state of their “nerves” has a direct effect on their symptoms. They often realize that their “cystitis” develops after a tension-producing or anxiety-producing episode in their personal or occupational environment.

**Sexual Difficulties in Men**

Men may complain directly of sexual difficulty. However, they are often so ashamed of loss of sexual power that they cannot admit it even to a physician. In such cases, they may ask for “prostate treatment” and hope that the physician will understand that they have sexual complaints and that they will be treated accordingly. The main sexual symptoms include impaired quality of erection, premature loss of erection, absence of ejaculate with orgasm, premature ejaculation, and even loss of desire.

**Sexual Difficulties in Women**

Women who have the psychosomatic cystitis syndrome almost always admit to an unhappy sex life. They notice that frequency or vaginal-urethral pain often occurs on the day following the incomplete sexual act. Many of them recognize the inadequacy of their sexual experiences as one of the underlying causes of urologic complaints; too frequently, however, the physician either does not ask them pertinent questions or, if patients volunteer this information, ignores it.

**REFERENCES**


A careful history and assessment of symptoms will suggest whether a complete or limited examination is indicated, and also help direct the appropriate selection of subsequent diagnostic studies.

## EXAMINATION OF THE KIDNEYS

### Inspection

A mass that is visible in the upper abdominal area may be difficult to palpate if soft, as with hydronephrosis. Fullness in the costovertebral angle may be consistent with cancer or perinephric infection. The presence and persistence of indentations in the skin from lying on wrinkled sheets suggest edema of the skin secondary to perinephric abscess.

### Palpation

The kidneys lie rather high under the diaphragm and lower ribs and are therefore well protected from injury. Because of the position of the liver, the right kidney is lower than the left. The kidneys are difficult to palpate in men because of (1) resistance from abdominal muscle tone and (2) more fixed position than in women, moving only slightly with change of posture or respiration. The lower part of the right kidney can sometimes be felt, particularly in thin patients, but the left kidney usually cannot be felt unless it is enlarged or displaced.

The most successful method of renal palpation is carried out with the patient lying in the supine position on a hard surface (Figure 4–1). The kidney is lifted by one hand in the costovertebral angle (CVA). On deep inspiration, the kidney moves downward; the other hand is pushed firmly and deeply beneath the costal margin in an effort to trap the kidney. When successful, the anterior hand can palpate the size, shape, and consistency of the organ as it slips back into its normal position.

Alternatively, the kidney may be palpated with the examiner standing behind the seated patient. At other times, if the patient is lying on one side, the uppermost kidney drops downward and medially, making it more accessible to palpation. Perlman and Williams (1976) described an effective method of identifying renal anomalies in newborns. The fingers are placed in the costovertebral angle, with the thumb anterior and performing the palpation.

An enlarged renal mass suggests compensatory hypertrophy (if the other kidney is absent or atrophic), hydronephrosis, tumor, cyst, or polycystic disease. However, a mass in this area may also represent a retroperitoneal tumor, spleen, lesion of the bowel (eg, tumor, abscess), lesion of the gallbladder, or pancreatic cyst. Tumors may have the consistency of normal tissue or be nodular, while hydronephrosis may be either firm or soft. Polycystic kidneys are usually nodular and firm.

An acutely infected kidney is tender, but the presence of marked muscle spasm may make this difficult to elicit. In addition, this sign may not always be helpful since the normal kidney is also often tender.

Although renal pain may be diffusely felt in the back, tenderness is usually well localized just lateral to the sacrospinalis muscle and below the 12th rib (ie, CVA). Symptoms may be elicited by palpation or sharp percussion over the CVA.

### Percussion

At times, an enlarged kidney cannot be felt, particularly if it is soft as in some cases of hydronephrosis. However, such masses may be outlined by both anterior and posterior percussion and this part of the examination should not be omitted.

Percussion is of particular value in outlining an enlarging mass (progressive hemorrhage) in the flank following renal trauma, when tenderness and muscle spasm prevent palpation.

### Transillumination

Transillumination may prove quite helpful in children under age 1 year who present with a suprapubic or flank mass. A dark room is required along with a flashlight with an opaque flange protruding beyond the lens. The flashlight
is applied at right angles to the abdomen. The fiberoptic light cord, used to illuminate various optical instruments, is an excellent source of cold light. A distended bladder or cystic mass will transilluminate; a solid mass will not. Flank masses may be assessed by applying the light posteriorly.

**Differentiation of Renal & Radicular Pain**

Radicular pain is commonly felt in the costovertebral and subcostal areas. It may also spread along the course of the ureter and is the most common cause of so-called “kidney pain.” Every patient who complains of flank pain should be examined for evidence of nerve root irritation. Frequent causes are poor posture (scoliosis, kyphosis), arthritic changes in the costovertebral or costotransverse joints, impingement of a rib spur on a subcostal nerve, hypertrophy of costovertebral ligaments pressing on a nerve, and intervertebral disk disease (Smith and Raney, 1976).

Radicular pain may be noted as an aftermath of a flank incision where a rib is dislocated, causing impingement of costal nerve on the edge of a ligament. Pain experienced during the preeruptive phase of herpes zoster involving any of the segments between T11 and L2 may simulate pain of renal origin.

Radiculitis usually causes hyperesthesia of the area of skin served by the irritated peripheral nerve. This hypersensitivity can be elicited by means of the pinwheel or grasping and pinching both skin and fat of the abdomen and flanks. Pressure exerted by the thumb over the costovertebral joints reveals local tenderness at the point of emergence of the involved nerve.

**Auscultation**

Auscultation of the costovertebral areas and upper abdominal quadrants may reveal a systolic bruit, often associated with stenosis or aneurysm of the renal artery. Bruits over the femoral arteries may be found in association with Ler-iche syndrome, which may be a cause of impotence.

### EXAMINATION OF THE BLADDER

The bladder cannot be felt unless it is moderately distended. In adults, it contains at least 150 mL of urine if it can be percussed. In acute or chronic urinary retention, the bladder may reach or even rise above the umbilicus, when its outline may be seen and usually felt. In chronic retention, the bladder may be difficult to palpate due to the flabby bladder wall, in which case percussion is of value.

In male infants or young boys, palpation of a hard mass deep in the center of the pelvis is compatible with a thickened, hypertrophied bladder secondary to obstruction caused by posterior urethral valves.

A sliding inguinal hernia containing some bladder wall can be diagnosed by compression of the scrotal mass when the bladder is full, leading to additional distension.

A few instances have been reported where marked edema of the legs has developed secondary to compression of the iliac vessels by a distended bladder. Bimanual (abdominorectal or abdominovaginal) palpation may reveal the extent of a vesical tumor; to be successful, it should be done under anesthesia.

### EXAMINATION OF THE EXTERNAL MALE GENITALIA

#### PENIS

**Inspection**

If the patient has not been circumcised, the foreskin should be retracted. This may reveal tumor or balanitis as the cause of foul discharge. If retraction is not possible due to phimosis, surgical correction (dorsal slit or circumcision) is indicated.

The observation of a poor urinary stream is significant: in newborns, neurogenic bladder or the presence of posterior urethral valves should be considered while in men such a finding suggests urethral stricture or prostatic obstruction.

The scars of healed syphilis may be an important clue. An active ulcer requires bacteriologic or pathologic study (eg, syphilitic chancre, epithelioma). Superficial ulcers or...
vesicles are compatible with herpes simplex and often interpreted by the patient as a serious sexually transmitted disease (eg, syphilis). Venerable warts may be observed.

Meatal stenosis is a common cause of bloody spotting in male infants. On occasion, it may be of such degree as to cause advanced bilateral hydrenephrosis.

The position of the meatus should be noted. It may be located proximal to the tip of the glans on either the dorsal (epispadias) or the ventral surface (hypospadias). In either instance, there is apt to be abnormal curvature (chordee) of the penis in the direction of the displaced meatus.

Micropenis or macropenis may be noted during examination of the penis. In the neonate, the presence of hypospadias and bilateral undescended testes should raise the possibility of an intersex condition.

**Palpation**

Palpation of the dorsal surface of the shaft may reveal a fibrous plaque involving the tunica albuginea covering of the corpora cavernosa, typical of Peyronie’s disease. Tender areas of induration felt along the urethra may signify periurethritis secondary to urethral stricture.

**Urethral Discharge**

Urethral discharge is the most common complaint referable to the male sex organ. Gonococcal pus is usually profuse, thick, and yellow or gray-brown. Nongonorrheal discharges may be similar in appearance but are often thin, mucoid, and scant. Although gonorrhea must be ruled out as the cause of urethral discharge, a significant percentage of cases is found to be caused by chlamydiae. Patients with urethral discharge also should be examined for other sexually transmitted diseases since multiple infection is not uncommon.

Bloody discharge suggests the possibility of a foreign body in the urethra, urethral stricture, or tumor.

Urethral discharge must always be sought before the patient is asked to void.

**SCROTUM**

Angioneurotic edema and infections and inflammations of the skin of the scrotum are not common. Small sebaceous cysts are occasionally seen while malignant tumors are rare. The scrotum is bifid when microlateral or perineal hypospadias is present.

Elephantiasis of the scrotum is caused by obstruction to lymphatic drainage and endemic in the tropics due to filariasis. Genital lymphedema may also result from radical resection of the lymph nodes of the inguinal and femoral areas, in which case the skin of the penis is involved. Small hemangiomas of the skin are common and may bleed spontaneously.

Scrotal ultrasound is a helpful adjunct in evaluating the scrotal contents.

**TESTIS**

The testes should be carefully palpated with the fingers of both hands. A hard area in the testis proper must be regarded as a malignant tumor until proven otherwise. Transillumination of scrotal masses should be done routinely. With the patient in a dark room, a flashlight or fiberoptic light is placed against the scrotal sac posteriorly. A hydrocele will cause the intrascrotal mass to glow red; conversely light is not transmitted through a solid tumor. Tumors are often smooth but may be nodular and the testes may seem abnormally heavy. A testis replaced by tumor or damaged by gumma is insensitive to pressure, and the usual sickening sensation is absent. About 10% of tumors are associated with a secondary hydrocele that may require aspiration before definitive palpation can be performed.

The testis may be absent from the scrotum, and this may be transient (physiologic retractile testis) or true cryptorchidism. Palpation of the groins may reveal the presence of the organ.

The atrophic testis (postoperative orchiopexy, mumps orchitis, or torsion of the spermatic cord) may be flabby and at times hypersensitive but is usually firm and hyposensitive. Although spermatogenesis may be absent, androgen function is occasionally maintained.

**EPIDIDYMIS**

The epididymis is sometimes rather closely attached to the posterior surface of the testis, and at other times, it is quite free of it. The epididymis should be carefully palpated for size and induration, which implies infection since primary tumors are exceedingly rare.

In the acute stage of epididymitis, the testis and epididymis are indistinguishable by palpation; the testicle and epididymis may be adherent to the scrotum, which is usually quite red and exquisitely tender. With few exceptions, the infecting organism is Neisseria gonorrhoeae, Chlamydia trachomatis, or Escherichia coli.

Chronic painless induration suggests tuberculosis or schistosomiasis, although nontuberculous chronic epididymitis is also possible. Other signs of genitourinary tuberculosis include sterile pyuria, thickened seminal vesicle, nodular prostate, and “beading” of the vas deferens.

**SPERMATIC CORD & VAS DEFERENS**

A swelling in the spermatic cord may be cystic (eg, hydrocele or hernia) or solid (eg, connective tissue tumor), although the latter is rare. Lipoma in the investing fascia of the cord may simulate a hernia. Diffuse swelling and induration of the cord are seen with filarial funiculitis.

Careful palpation of the vas deferens may reveal thickening (“beading” caused by tuberculosis), or even absence of the vas. The latter finding is of importance in infertile males.
and may be associated with cystic fibrosis or ipsilateral Wolffian duct abnormality (eg, renal agenesis).

When a male patient stands, a mass of dilated veins (varicocele) may be noted behind and above the testis. The degree of dilatation decreases with recumbency and can be increased by the Valsalva maneuver. The major potential sequela of varicocele is infertility (see Chapter 42).

**TESTICULAR TUNICS & ADNEXA**

Hydroceles are usually cystic but on occasion so tense that they simulate solid tumors; transillumination confirms the diagnosis. The fluid may accumulate secondary to nonspecific acute or tuberculous epididymitis, trauma, or tumor of the testis. The latter is a distinct possibility if the hydrocele appears spontaneously between the ages of 18 and 35. It should be aspirated to permit careful palpation of underlying structures or further characterized with ultrasonography.

Hydroceles usually surround the testis completely. Cystic masses that are separate from but in the region of the upper pole of the testis are typically spermatoceles. Aspiration reveals the typical thin, milky fluid which contains sperm.

### EXAMINATION OF THE FEMALE GENITALIA

#### VAGINAL EXAMINATION

Diseases of the female genital tract may secondarily involve the urinary organs, making a thorough gynecologic examination essential, which should be performed by the male physician in the presence of a female nurse or health professional. Commonly associated conditions include urethrocystitis secondary to urethral diverticulitis or cervicitis, pyelonephritis during pregnancy, and ureteral obstruction from metastatic nodes or direct extension from cervical cancer.

**Inspection**

In newborns and children, the vaginal vestibule should be inspected for a single opening (common urogenital sinus), labial fusion, split clitoris and lack of fusion of the anterior fourchette (epispadias), or hypertrophi ed clitoris and scrotalization of the labia majora (adrenogenital syndrome).

The urinary meatus may reveal a reddened, tender, friable lesion (urethral caruncle) or a reddened, everted posterior lip often seen with senile urethritis and vaginitis. Biopsy is indicated if a malignant tumor cannot be ruled out. The diagnosis of senile vaginitis and urethritis is established by staining a smear of the vaginal epithelium with Lugol’s solution. Cells lacking glycogen (hypoestrogenism) do not take up the stain, whereas normal cells do.

Multiple painful small ulcers or blister-like lesions may be noted, probably representing herpes virus type 2 infection, which may have serious sequelae.

Smears and cultures of urethral or vaginal discharge should be made. Gonococci are relatively easy to identify; culture of chlamydia requires techniques seldom available to the physician.

The presence of sksenitis and Bartholinitis may reveal the source of persistent urethritis or cystitis. The condition of the vaginal wall should be observed. Urethrocele and cystocoele, often found with stress incontinence, may be associated with residual urine and lead to persistent infection of the bladder. A bulge in the anterior vaginal wall may represent a urethral diverticulum. The cervix should be inspected for cancer or infection. Taking biopsy specimens or making Papanicolaou smears may be indicated.

**Palpation**

At times, the urethra, base of the bladder, and lower ureters may be tender on palpation, but little can be deduced from this finding. Induration of the urethra or trigonal area, or a mass involving either, may be a clue to an existing tumor. A soft mass could be a urethral diverticulum, and pressure may cause pus to extrude from the urethra. A stone in the lower ureter may be palpable. Evidence of enlargement of the uterus (eg, pregnancy, myoma) or diseases or inflammations of the colon or adnexa may afford a clue to the cause of urinary symptoms (eg, compression of ureter by ovarian tumor, endometriosis, or diverticulitis of the sigmoid colon adherent to the bladder).

Carcinoma of the cervix may invade the base of the bladder, causing vesical irritability or hematuria; metastases to iliac lymph nodes may compress the ureters.

Rectal examination may provide further information and is the obvious route of examination in children and virgins.

### RECTAL EXAMINATION IN MALES

#### SPHINCTER & LOWER RECTUM

The estimation of sphincter tone is of importance. Laxity of the muscle suggests similar changes in the urinary sphincter and detrusor and the possibility of neurogenic disease; the same is true for a spastic anal sphincter. In addition to the digital prostatic examination, the examiner should palpate the entire lower rectum to rule out
stenosis, internal hemorrhoids, cryptitis, rectal fistulae, mucosal polyps, and rectal cancer. Testing perianal sensation is mandatory.

**PROSTATE**

A specimen of urine for routine analysis should be collected before the rectal examination. This is of importance, since prostatic massage or even palpation at times forces prostatic secretion into the posterior urethra. If this secretion contains pus, a specimen of voided urine after the rectal examination will be contaminated.

**Size**

The average prostate is about 4 cm in both length and width. As the gland enlarges, the lateral sulci become relatively deeper and the median furrow becomes obliterated. The clinical importance of prostatic hyperplasia is measured by the severity of symptoms and the amount of residual urine rather than by the size of the gland on palpation. The prostate may be of normal size and consistency on examination in a patient with acute urinary retention or severe obstructive urinary complaints.

**Consistency**

Normally, the consistency of the gland is similar to that of the contracted thenar eminence of the thumb (with the thumb completely opposed to the little finger) and is rather rubbery. It may be mushy if congested (due to lack of intercourse or chronic infection with impaired drainage), indurated (due to chronic infection with or without calculi), or stony hard (due to advanced carcinoma).

The difficulty lies in differentiating firm areas in the prostate: fibrosis from nonspecific infection, granulomatous prostatitis, nodularity from tuberculosis, or firm areas due to prostatic calculi or early cancer. Generally, nodules caused by infection are raised above the surface of the gland. At their edges, the induration gradually fades to the normal softness of surrounding tissue. Conversely, the suspicious lesion in cases of prostate cancer is usually not raised; rather, it is hard and has a sharp edge (ie, there is an abrupt change in consistency on the same plane). It tends to arise in the lateral sulcus (Figure 4–2).

Even the most experienced clinician can have trouble differentiating cancer from other conditions. The serum prostate-specific antigen (PSA) level can be helpful if elevated, and is currently the most common method of diagnosing prostate cancer (clinical stage T1c). Transrectal ultrasound-guided biopsy of the prostate can be diagnostic. Recent evidence suggests that rectal examination after radical prostatectomy is unnecessary when PSA is undetectable, since no case of locally recurrent cancer was identified in the absence of an elevated PSA.

**Mobility**

The mobility of the gland varies. Occasionally, it has great mobility while at other times, very little. With advanced carcinoma, it is fixed because of local extension through the capsule. In adults, the prostate should be routinely massaged and its secretion examined microscopically. However, prostatic massage should be avoided in the presence of an acute urethral discharge, acute prostatitis, or acute prostatocystitis; in men near the stage of complete urinary retention (because it may precipitate complete retention); or in men suffering from obvious cancer of the gland.

*Figure 4–2. Differential diagnosis of prostatic nodules.*

*A:* Inflammatory area is raised above the surface of the gland; induration decreases gradually at its periphery. 
*B:* Cancerous nodules is not raised; there is an abrupt change in consistency at its edges.
 Massage & Prostatic Smear

Copious amounts of secretion may be obtained from some prostate glands and little or none from others. The amount obtained depends to some extent on the vigor with which the massage is carried out. If no secretion is obtained, the patient should be asked to void even a few drops of urine, as these will contain adequate secretion for examination. Microscopic examination of the secretion is done under low-power magnification. Normal secretion contains numerous lecithin bodies, which are refractile like red blood cells but much smaller. Only an occasional white cell is present. A few epithelial cells and, rarely, corpora amylacea are seen. Sperm may be present, but its absence is of no significance.

The presence of large numbers or clumps of leukocytes suggests prostatitis. Stained smears are usually impractical because it is difficult to fix the material on the slide; even when fixation and staining are successful, pyogenic bacteria are usually not found. Acid-fast organisms can often be found by appropriate staining methods.

On occasion, it may be necessary to obtain cultures of prostatic secretion in order to demonstrate nonspecific organisms, tubercle bacilli, gonococci, or chlamydiae. After thorough cleansing of the glans and emptying of the bladder to mechanically cleanse the urethra, the prostate is massaged. Drops of secretion are collected in a sterile tube of appropriate culture medium.

SEMINAL VESICLES

Palpation of the seminal vesicles should be attempted. The vesicles are situated under the base of the bladder and diverge from below upward (Figure 1–8). Normal seminal vesicles are usually impalpable, but may feel cystic when overdistended. In the presence of chronic infection (particularly tuberculosis or schistosomiasis) or in association with advanced carcinoma of the prostate, they may be indurated. Stripping of the seminal vesicles should be done in association with prostatic massage, for the vesicles are usually infected when prostatitis is present. Primary tumors of the vesicles are very rare. A cystic mass may rarely be felt over the prostate or just above it. This probably represents a cyst of the Müllerian duct or the utricle; the latter is occasionally associated with severe hypospadias.

LYMPH NODES

It should be remembered that generalized lymphadenopathy usually occurs early in human immunodeficiency syndrome (see Chapter 15).

Inguinal & Subinguinal Lymph Nodes

With inflammatory lesions of the skin of the penis and scrotum or vulva, the inguinal and subinguinal lymph nodes may be involved. Such diseases include chancroid, syphilitic chancre, lymphogranuloma venereum, and on occasion, gonorrhea.

Malignant tumors (squamous cell carcinoma) involving the penis, glans, scrotal skin, or distal urethra in women metastasize to the inguinal and subinguinal nodes. Testicular tumors do not spread to these nodes unless they have invaded the scrotal skin or the patient has previously undergone orchidopexy.

Other Lymph Nodes

Tumors of the testis and prostate may involve the left supracaolic nodes (Vinchow’s or Troisier’s node). Tumors of the bladder and prostate typically metastasize to the internal iliac, external iliac, and preaortic nodes, although rarely are they so large as to be palpable. Upper abdominal masses near the midline in a young man should suggest metastases from cancer of the testis; the primary growth may be minute and hidden in the substance of what appears to be a normal testicle.

NEUROLOGIC EXAMINATION

A careful neurologic survey may uncover sensory or motor impairment that account for residual urine or incontinence. Since the bladder and sphincter are innervated by the second to the fourth sacral segments, information can be gained by testing anal sphincter tone, evaluating sensation of the perianal skin, and testing Achilles tendon and bulbocavernous reflexes. The bulbocavernous reflex is elicited by placing a finger in the rectum and squeezing the glans penis or clitoris, or by jerking on an indwelling Foley catheter. The normal reflex involves contraction of the anal sphincter and bulbocavernous muscles in response to these maneuvers.

It is wise, particularly in children, to seek a dimple over the lumbosacral area. One should palpate the sacrum to ensure it is present and normally formed. Sacral agenesis or partial development is compatible with deficits of S2–4. If findings seem abnormal, x-ray examination is indicated.

REFERENCES

Examination of the Kidneys

**External Genitalia in Males**


**External Genitalia in Females**


**Prostate**


**Neurologic Examination**


Examination of specimens of urine, blood, and genitourinary secretions or exudates commonly directs the subsequent urologic workup and frequently establishes a diagnosis. Since approximately 20% of patients who visit a primary physician’s office have a urologic problem, it is important for the physician to have a broad knowledge of the laboratory methods available to test appropriate specimens. Judicious use of such tests permits rapid, accurate, and cost-effective determination of the probable diagnosis and directs the management of patients with urologic disease.

**EXAMINATION OF URINE**

Urinalysis is one of the most important and useful urologic tests available, yet all too often the necessary details are neglected and significant information is overlooked or misinterpreted. Reasons for inadequate urinalyses include (1) improper collection, (2) failure to examine the specimen immediately, (3) incomplete examination (eg, most laboratories do not perform a microscopic analysis unless it is specifically requested by the provider), (4) inexperience of the examiner, and (5) inadequate appreciation of the significance of the findings.

The necessity of routine urinalysis as a screen in asymptomatic individuals, those admitted to hospitals, or those undergoing elective surgery continues to be debated. Numerous studies indicate that in the situations above urinalysis is not routinely necessary (Godbole and Johnstone, 2004). Patients presenting with urinary tract symptoms or signs, however, should undergo urinalysis. Studies also indicate that if macroscopic urinalysis (dip-strip) is normal, microscopic analysis is not necessary. If the patient has signs or symptoms suggestive of urologic disease, or the dip-strip is positive for protein, heme, leukocyte esterase, or nitrite, a complete urinalysis, including microscopic examination of the sediment, should be completed (Simerville, Maxted, and Pahira, 2005).

**Urine Collection**

**A. TIMING OF COLLECTION**

It is best to examine urine that has been properly obtained in the office. First-voided morning specimens are helpful for qualitative protein testing in patients with possible orthostatic proteinuria and for specific gravity assessment as a presumptive test of renal function in patients with minimal renal disease due to diabetes mellitus or sickle cell anemia or in those with suspected diabetes insipidus. Evaluation of sequential morning specimens may be required to obviate the variability often encountered. Urine specimens that are obtained immediately after the patient has eaten or that have been left standing for a few hours become alkaline and thus may contain lysed red cells, dis-integrated casts, or rapidly multiplying bacteria; therefore, a freshly voided specimen obtained a few hours after the patient has eaten and examined within 1 hour of voiding is most reliable. The patient’s state of hydration may alter the concentration of urinary constituents. Timed urine collections may be required for definitive assessment of renal function or proteinuria.

**B. METHOD OF COLLECTION**

Proper collection of the specimen is particularly important when patients have hematuria or proteinuria or are being evaluated for urinary tract infection. Examination of a urine specimen collected sequentially during voiding in several containers may help to identify the site of origin of hematuria or urinary tract infection. To gather consistent and meaningful urinalysis data, urine must be collected by a uniform method in the physician’s office or laboratory. The specimen should be obtained before a genital or rectal examination in order to prevent contamination from the introitus or expressed prostatic secretions. Urine obtained from a condom, chronic catheter, or intestinal conduit drainage bag is not a proper specimen for urinalysis.

1. **Men**—It is usually simple to collect a clean-voided midstream urine sample from men. Routine instructions may be printed on a sheet given to the patient or placed on the lavatory wall. The procedure should include (1) retraction of the foreskin (a common source of contamination of the specimen) and cleansing of the meatus with benzalkonium chloride or hexachlorophene; (2) passing the first part of the stream (15–30 mL) without collection; and (3) collecting the next or midstream portion (approximately 50–100 mL) in a sterile specimen container, which is capped immediately afterward. A portion of the specimen is prepared immediately for both macroscopic and microscopic examination, and the rest is saved in the sterile container for subsequent culture if this proves necessary.
With this midstream clean-catch method, the likelihood that the specimen will be contaminated by meatal or urethral secretions is markedly decreased, although not completely eliminated. In adult males, it is rarely necessary to collect urine by catheterization unless urinary retention is present.

2. Women—The best method for collecting a clean-voided midstream specimen from a woman is as follows: (1) the patient is placed on the examining table in the lithotomy position; (2) the vulva and urethral meatus are cleansed with benzalkonium chloride or hexachlorophene; (3) the labia are separated; and (4) the patient is instructed to initiate voiding into a container held close to the vulva. After she has passed the first 10–20 mL of urine, the next 50–100 mL is collected in a sterile container that is immediately capped. Because this technique requires considerable effort, it is acceptable to have the patient provide an initial specimen in a nonsterile container in the office lavatory. If results of urinalysis are normal, no further study is indicated; if abnormal, a urine specimen must be obtained by the more exacting technique. In either case, the specimen should be prepared for immediate examination.

If a satisfactory specimen cannot be obtained by the method described, one should not hesitate to obtain a specimen by catheterization to eliminate nonvaginal sources of abnormal urinary constituents.

3. Children—Urine for analysis, other than bacterial cultures, can be obtained from males or females by covering the cleansed urethral meatus with a plastic bag; a urine specimen for culture may require catheterization or suprapubic needle aspiration. In girls, catheterization with a small catheter attached to a centrifuge tube is appropriate, but boys should not be routinely catheterized. It is often preferable in either sex to proceed with suprapubic needle aspiration. This is easier if the patient has been previously hydrated, so that the bladder is full. Suprapubic needle aspiration is performed as follows: (1) Cleanse the suprapubic area by sponging with alcohol. (2) With a small amount of local anesthetic, raise an intradermal wheal on the midline 1–2 cm above the pubis (the bladder lies just above the pubis in young children). (3) Attach a 10-mL syringe to a 22-gauge needle. Insert the needle perpendicularly through the abdominal wheal into the bladder wall, maintaining gentle suction with the syringe so that urine will be aspirated as soon as the bladder is entered.

Macroscopic Examination

Macroscopic examination of urine often provides a clue when diagnosis is difficult.

A. Color & Appearance

Urine is often colored owing to drugs; phenazopyridine (Pyridium) will turn the urine orange; rifampin will turn it yellow-orange; nitrofurantoin will turn it brown; and l-dopa, α-methyldopa, and metronidazole will turn it redish-brown. Red urine does not always signify hematuria. A red discoloration unassociated with intact erythrocytes in the urine can result from betacyanin excretion after beet ingestion, phenolphthalein in laxatives, ingestion of vegetable dyes, concentrated urine excretion, myoglobinuria due to significant muscle trauma, or hemoglobinuria following hemolysis. In addition, *Serratia marcescens* bacteria can cause the “red diaper” syndrome. However, whenever red urine is seen, hematuria must be ruled out by microscopic analysis. Cloudy urine is commonly thought to represent pyuria, but more often the cloudiness is due to large amounts of amorphous phosphates, which disappear with the addition of acid, or urates, which dissolve with the use of alkali. The odor of urine is rarely clinically significant.

B. Specific Gravity

The specific gravity of urine (normal, 1.003–1.030) is often important for diagnostic purposes: that of patients with significant intracranial trauma may be low owing to a lack of antidiuretic hormone (vasopressin); that of patients with primary diabetes insipidus is <1.010 even after overnight dehydration; that of patients with extensive acute renal tubular damage is consistently 1.010 (similar to the specific gravity of plasma); and a low specific gravity can be an early sign of renal damage from conditions such as sickle cell anemia. Urine specific gravity is the simplest time-honored test for evaluating hydration in postoperative patients. The specific gravity of urine may affect the results of other urine tests: in dilute urine, a pregnancy test may be falsely negative; in concentrated urine, protein may be falsely positive on dip-strips yet unconfirmed on quantitative tests. The specific gravity of urine may be falsely elevated by the presence of glucose, protein, artificial plasma expanders, or intravenous contrast agents.

Studies of specific-gravity reagent strips (method based on ionic alteration of a polyelectrolyte solution) have shown the method to be rapid, reliable, and unaffected by elevated amounts of glucose or contrast medium; however, alkaline pH may falsely lower the result (0.005 per pH unit >7.0). In the routine office setting, these strips are as reliable as either the hydrometer or refractometer methods.

C. Chemical Tests

Chemically impregnated reagent strips are accurate and have simplified routine urinalysis greatly. However, they must be monitored routinely by appropriate standardized quality-control reagents. The dip-strips are reliable only when not outdated and when used with room temperature urine.

1. **pH**—The pH of urine is important in a few specific clinical situations. Patients with uric acid stones rarely have a urinary pH over 6.5 (uric acid is soluble in alkaline urine). Patients with calcium stones, nephrocalcinosis, or both may have renal tubular acidosis and will be unable to
acidify urine below pH 6.0. With urinary tract infections caused by urea-splitting organisms (most commonly Proteus species), the urinary pH tends to be over 7.0. It should be reemphasized that urine obtained within 2 hours of a large meal or left standing at room temperature for several hours tends to be alkaline. The indicator paper in most dip-strips is quite accurate; however, confirmation by a pH meter is occasionally required.

2. Protein—Dip-strips containing bromphenol blue can be used to determine the presence of >10 mg/dL protein in urine, but persistent proteinuria detected in this manner requires quantitative protein testing for confirmation. The dip-strip measures primarily albumin and is not sensitive to Bence-Jones proteins (immunoglobulins). Concentrated urine may give a false-positive result, as will urine containing numerous white blood cells (leukocytes) or vaginal secretions replete with epithelial cells. Orthostatic proteinuria can be demonstrated by detecting elevated protein levels in a urine specimen obtained after the patient has been in the upright position for several hours, whereas normal levels are found before ambulation. Prolonged fever and excessive physical exertion are also common causes of transient proteinuria.

Persistently elevated protein levels in the urine (>150 mg/24 h) may indicate significant disease. Therefore, specific quantitative protein tests, electrophoretic studies of the urine, or both may be required to determine the specific type of protein that is present.

3. Glucose—The glucose oxidase-peroxidase tests used in dip-strips are quite accurate and specific for urinary glucose. False-positive results may be obtained when patients have ingested large doses of aspirin, ascorbic acid, or cephalosporins. An occasional patient has a blood glucose level below 180 mg/dL and yet has significant glucosuria; this indicates a low renal threshold of glucose excretion. However, most patients with a positive reading have diabetes mellitus.

4. Hemoglobin—The dip-strip test for hemoglobin is not specific for erythrocytes and should be used only to screen for hematuria, with microscopic analysis of the urinary sediment used for confirmation. Free hemoglobin or myoglobin in the urine may give a positive reading; ascorbic acid in the urine can inhibit the dip-strip reaction and give a false-negative result. Note that dilute urine (<1.008) will lyse erythrocytes and thus provide a positive dip-strip reading for hemoglobin but no visible erythrocytes on microscopic analysis.

5. Bacteria and leukocytes—Test strips to determine the number of bacteria (nitrite) or leukocytes (leukocyte esterase) as predictors of bacteriuria are as accurate as microscopic sediment analysis in studies using quantitative urine cultures as the standard. The nitrite reductase test depends on the conversion of nitrate to nitrite. Many of the bacteria responsible for urinary tract infections, particularly enterobacteria, are capable of reducing nitrate to nitrite and therefore detectable by this test. When the nitrite test is positive, it suggests the presence of >100,000 organisms per mL; however, several factors can lead to false-negative results. The nitrite test is positive only for coagulase-splitting bacteria and thus when used alone is only 40–60% accurate. Urine must be in the bladder for a sufficient time before sampling for the reduction of nitrate to occur (>4 hours); therefore, this test is most likely to be positive when first-voided morning urine is tested. A false-negative test will also result if the bacteria present do not contain nitrate reductase or if dietary nitrate is absent. A false-negative nitrite study may occur in a patient taking vitamin C. The leukocyte esterase test is a widely used chemical test that depends on the presence of esterase in granulocytic leukocytes. The leukocyte esterase test is an indication of pyuria and will remain positive even after the leukocytes have degenerated. The test accurately identifies patients with 10–12 leukocytes per high-power field in a centrifuged specimen. Although this test is a good indicator of pyuria, it does not detect bacteriuria. Therefore, it is often combined with the nitrite test to detect both bacteriuria and inflammation to maximize the chances of predicting urinary tract infection. Used together, the 2 tests are as predictive as the microscopic analysis but not as accurate as a urine culture. A false-negative leukocyte esterase study can be caused by glucosuria, or by phenazopyridine hydrochloride (Pyridium), nitrofurantoin, vitamin C, or rifampin in the urine.

Microscopic Examination

To be most accurate, the microscopic sediment examination should be done personally by an experienced physician or technician. Early-morning urine is the best specimen if it can be examined within a few minutes of collection. In most cases, the sediment can be prepared as follows: (1) Centrifuge a 10-mL specimen at 2000 rpm for 5 minutes. (2) Decant the supernatant. (3) Resuspend the sediment in the remaining 1 mL of urine by tapping the tube gently against a countertop. (4) Place 1 drop of the mixture on a microscope slide, cover with a coverslip, and examine first under a low-power (10×) and then under a high-power (40×) lens. For maximal contrast of the elements in the sediment, the microscope diaphragm should be nearly closed to prevent overillumination. Significant elements (particularly bacteria) are more easily seen if the slide is stained with methylene blue, but staining is not essential. Figure 5–1 shows typical findings in the urinary sediment.

Interpretation

1. Bacteria—The significance of bacteria in the urinary sediment is discussed in the section that follows on bacteriuria.
2. Leukocytes—Just as the presence of bacteria in the sediment is not an absolute indication of infection, neither is the finding of pyuria. In the sediment from clean-voided midstream specimens from men and those obtained by suprapubic aspiration or catheterization in women, a finding of more than 5 leukocytes per high-power field is generally considered abnormal (pyuria). If the patient has symptoms of a urinary tract infection as well as pyuria and bacteriuria, one is justified in making a diagnosis of infection and initiating empiric therapy. However, in female patients with symptoms of urinary tract infection, 60% of those with pyuria will have no bacterial growth from bladder urine obtained by catheterization or suprapubic aspiration emphasizing the need for confirmation by bacterial cultures.

Renal tuberculosis can cause “sterile” acid-pyuria and should be considered in any patient with persistent pyuria and negative results on routine bacterial cultures. Specific fluorescent staining of the urinary sediment for acid-fast bacteria can be diagnostic; however, results will be positive from the sediment of spot specimens in only approximately 50% of patients with renal tuberculosis, whereas they are positive in the sediment of 24-hour specimens in 70–80% of such cases. Mycobacterium smegmatis, a commensal organism, may be present in the urine (particularly in uncircumcised men) and can give false-positive results on acid-fast stains.

Urolithiasis can also cause pyuria. In patients with persistent pyuria, the physician should consider obtaining at least a plain x-ray of the abdomen and possibly a CT urogram to determine whether urolithiasis is present. Similarly, a retained foreign body such as a self-induced bladder object or a forgotten internal ureteral stent can cause pyuria. A plain x-ray (KUB film) of the abdomen should reveal the offender.

3. Erythrocytes—The presence of even a few erythrocytes in the urine (hematuria) is abnormal and requires further investigation. Although gross hematuria is more alarming to the patient, microscopic hematuria is no less significant. Infrequent causes of hematuria include strenu-
ous exercise (long-distance running), vaginal bleeding, and inflammation of organs near or directly adjoining the urinary tract, for example, diverticulitis or appendicitis. Hematuria associated with cystitis or urethritis generally clears after treatment. Persistent hematuria in an otherwise asymptomatic patient of either sex and any age signifies disease and is an indication for further testing.

In patients with microscopic hematuria, a 3-container method for collection of urine can provide information on the site of origin of erythrocytes: (1) Give the patient 3 containers, labeled 1, 2, and 3 (or initial, mid, and final). (2) Instruct the patient to urinate and to collect the initial portion of the urine stream (10–15 mL) in the first container, the middle portion (30–40 mL) in the second, and the final portion (5–10 mL) in the third. (3) Using methods described previously, centrifuge the 3 specimens individually, prepare slides of the urinary sediment (with or without staining), and examine the slides microscopically. If erythrocytes predominate in the initial portion of the specimen, they are usually from the anterior urethra; those in the final portion are generally from the bladder neck or posterior urethra; and the presence of equal numbers of erythrocytes in all 3 containers usually indicates a source above the bladder neck (bladder, ureters, or kidneys). It is important to collect the urine before physical examination (particularly before rectal examination in men) to avoid misleading results.

The 3-container test may not be necessary in patients with gross hematuria, since the patients (men in particular) can usually tell the physician which portion of the stream contains the darkest urine (ie, the most erythrocytes). A specific dysmorphic erythrocyte configuration that can be detected with phase-contrast microscopy or by particle analyzer study of the urinary sediment and is highly indicative of active glomerular disease (Figure 5–2) can be useful. This dysmorphism is thought to be a result of extreme changes in osmolality and the high concentration of urinary chemical constituents affecting erythrocytes during passage through the kidney tubules. An automated system, iQ200, has been shown to be highly accurate for detecting, enumerating, and sizing erythrocytes in urine (Wah, Wises, and Butch, 2005).

4. Epithelial cells—Squamous epithelial cells in the urinary sediment indicate contamination of the specimen from the distal urethra in males and from the introitus in females; no other significance should be placed on them.

![Figure 5–2. Left: Dysmorphic erythrocytes in urine (arrows), viewed under light microscopy (magnification ×400). Right: Dysmorphic erythrocytes in urine (identical field), viewed under phase-contrast microscopy. (Reproduced, with permission, from Stamey TA, Kindrachuk RW: Urinary Sediment and Urinalysis: A Practical Guide for the Health Science Professional. PA, Saunders, 1985.)](image-url)
is not uncommon to find transitional epithelial cells in the normal urinary sediment; however, if they are present in large numbers or clumps and are abnormal histologically (including large nuclei, multiple nucleoli, and an increased ratio of nucleus to cytoplasm), they are indicative of a malignant process affecting the urothelium (Figure 5–3).

5. Casts—Casts are formed in the distal tubules and collecting ducts and, for the most part, are not seen in normal urinary sediment; therefore, they commonly signify intrinsic renal disease.

Although leukocyte casts have been considered suggestive of pyelonephritis, they are not an absolute indicator and should not be used as the sole criterion for diagnosis. Leukocyte casts must be distinguished from epithelial cell casts, because the latter have little significance when present in small numbers. The distinction can be made easily if a small amount of acetic acid is added under the coverslip to enhance nuclear detail. (Note that casts tend to congregate near the edges of the coverslip.) Epithelial cell or leukocyte casts in large numbers signify underlying intrinsic renal disease requiring further diagnostic workup. In renal transplant recipients, an increase in the number of epithelial cells or casts from the renal tubules may be an early indication of acute graft rejection.

Erythrocyte casts are pathognomonic of underlying glomerulitis or vasculitis. Hyaline casts probably represent a mixture of mucus and globulin congealed in the tubules; in small numbers, they are not significant. Hyaline casts are commonly seen in urine specimens taken after exercise and in concentrated or highly acidic urine specimens. Casts are rarely seen in alkaline urine and are therefore not usually present in urine specimens that have been left standing or in specimens from patients unable to acidify.

Granular casts most commonly represent disintegrated epithelial cells, leukocytes, or protein; they usually indicate intrinsic renal tubular disease.

6. Other findings—The finding of crystals in urine can be helpful in some instances, but the mere presence of crystals does not indicate disease. Crystals form in normal urine below room temperature. Cystine, leucine, tyrosine,
cholesterol, bilirubin, hematoidin, and sulfonamide crystals are abnormal findings of varying importance. Several types of crystals that may be found on microscopic examination of urinary sediment are shown in Figure 5–1.

The use of protease inhibitors for treatment of human immunodeficiency virus (HIV) has resulted in urolithiasis due to indinavir crystal formation in urine. The characteristic crystals are flat, rectangular plates, often in a fan or star-burst pattern. The presence of trichomonads or yeast cells in the stained or unstained smear of sediment from a properly obtained urine specimen establishes a diagnosis and the need for treatment.

Artifacts present in the urine sediment can be difficult to differentiate from real abnormalities. Dirt and small pieces of vegetable fiber or hair are frequently found, but the most common artifacts are starch granules from examination gloves.

**Bacteriuria**

**A. MICROSCOPIC EXAMINATION**

A presumptive diagnosis of bacterial infection may be made on the basis of results of microscopic examination of the urinary sediment. If several bacteria per high-power field are found in a urine specimen obtained by suprapubic aspiration or catheterization in a woman or in a properly obtained clean-voided midstream specimen from a man, a provisional diagnosis of bacterial infection can be made and empiric treatment started. The findings should be confirmed by bacterial culture. Finding several bacteria per high-power field are found in a urine specimen obtained by suprapubic aspiration or catheterization in a woman or in a properly obtained urine specimen establishes a diagnosis and the need for treatment.

Cultures can be confirmed by culture.

1. **Indications and interpretation**—Cultures can be used to estimate the number of bacteria in the urine (quantitative cultures), to identify the exact organism present, and to predict which drugs will be effective in treating the infection. Cultures are particularly important in patients with recurrent or persistent infections, renal insufficiency, or drug allergies.

The number of bacteria present in the urine (colony count) is influenced by the method used to collect the urine specimen, the patient’s hydration status, and whether the patient has been taking antimicrobial drugs. The concept that urinary tract infection is present only when the urine specimen contains 10^5 or more bacteria per milliliter is not an absolute rule; a lower count does not exclude the possibility of an infection, particularly in a symptomatic patient. Cultures with growth of multiple organisms usually signify contamination. The presence of a few organisms in a specimen with a low specific gravity is more significant than the same finding in a specimen with a high specific gravity, because the former is more dilute.

Identifying the drugs to which the bacteria are sensitive may or may not be necessary. *Escherichia coli*, which causes 85% of “routine” urinary tract infections, is known to be sensitive to numerous oral antimicrobial drugs. However, in patients with septicemia, renal insufficiency, diabetes mellitus, or suspected enterococcal, *Proteus*, or *Pseudomonas* infections, it is important to determine the antibiotic sensitivity of the organism and the drug concentration necessary for efficacious treatment. Monitoring antibiotic levels in blood and urine during treatment may be indicated, especially in severely ill patients and those receiving highly toxic drugs. These measurements can be done by most hospital laboratories.

2. **Rapid tests for bacteriuria**—In general, seriously ill or hospitalized patients with urinary tract infections should have cultures processed by an accredited bacteriology laboratory. However, for “routine” infections encountered in office practice, there are many satisfactory, cost-effective testing methods.

Rapid methods to screen for bacteria include growth-independent systems and growth-dependent systems. Several growth-dependent systems are available. One measures the turbidity of urine incubated in a broth medium for several hours. Positive results can be determined in as short a time period as 4 hours; however, 12 hours of growth is required before a test sample can be regarded as negative. A single non–growth dependent screening test uses the leukocyte esterase test and the nitrite test. If both tests are positive, the specificity increases to 98–99.5%, which indicates probable urinary tract infection (Young and Soper, 2001).

Reliable culture methods involve use of small strips or glass slides coated with eosin–methylene blue agar on one side and nutrient agar on the other. The strips or slides are dipped in the urine specimen and then incubated for 24 hours. Although these methods are easy to use, their disadvantages are that (1) not all bacteria will grow under these conditions, and (2) the accuracy of colony counts is debatable.

Perhaps preferable for the physician’s office (but still subject to some of the same limitations) is use of a divided plastic culture plate with blood agar on one side and deoxycholate agar on the other. A known amount of urine is inoculated onto the agar on each side of the plate, and colony counts are determined at 24–48 hours. The numbers of bacteria in 1 mL of the original urine specimen can be determined by multiplying the number of colonies by the volume (in milliliters) and dilution (if any) of the inoculum. If antibiotic sensitivity testing is also desired, an additional culture plate can be inoculated and small antibiotic-impregnated disks placed on the agar. Zones of growth inhibition seen around the disks at 12–24 hours indicate sensitivity.
3. Cultures for tuberculosis—A microscopic examination (fluorescent stain) that shows acid-fast bacilli can give a presumptive diagnosis of urinary tuberculosis. The rapidity of recovering mycobacteria in culture depends somewhat on the patient’s bacillary load. Thus, if the smear is highly positive (3–4+), cultures would become positive in 1–2 weeks. At that time, a DNA culture probe can be done for tuberculosis. It should be noted that the probe cannot distinguish between tuberculosis and patients treated with BCG; if the patient had not received BCG treatment, then \textit{M. tuberculosis} infection is likely. The total time from receipt of the specimen to presumptive diagnosis is typically about 2 weeks. Susceptibility tests, if positive for tuberculosis, would require another week.

Other Urine Tests

Many other tests of urine can be helpful in determining the presence of urologic disease.

A. UROTHELIAL CANCER TESTS

1. Urine cytology—The evaluation of voided or bladder wash (barbotage) urine for bladder urothelial cancer cells has been quite successful for higher grade (2–3) transitional cell cancers. Lower grade tumors less commonly shed abnormal cells. Cystoscopy remains the standard diagnostic test for initial diagnosis and surveillance of bladder cancer.

2. Bladder tumor antigen-TRAK test—The bladder tumor antigen test (BTA; Bard Diagnostic Sciences, Inc, Redmond, WA) is an assay for the qualitative detection of bladder tumor antigen in the urine.

3. Nuclear matrix protein 22—The nuclear matrix protein 22 test (NMP22; Matritech, Inc, Newton, MA) is an immunoassay. Normal subjects will have low levels of NMP22 in the urine, whereas patients with active transitional cell carcinoma may have high levels of urinary NMP22 (Grossman, Messing, and Soloway et al, 2005).

4. QUANTICYT System—The QUANTICYT System is a computer-based cytologic image analysis system. This system evaluates 50 randomly selected images containing 100–500 nuclei for DNA content and nuclear shape. The characteristics of these bladder cancer tests and others are compared in Table 5–1 (Konety and Getzenberg, 2001).

B. HORMONAL STUDIES

Tests for abnormalities in adrenal hormone secretion are important in the workup of patients with suspected adrenal tumors. Pheochromocytoma and neuroblastoma can be detected by measuring the excretion of vanillylmandelic acid. However, urinary levels of metanephrine, epinephrine, and norepinephrine are more sensitive indicators, particularly in cases of pheochromocytoma. Although high levels of aldosterone in urine usually indicate an aldosterone-secreting tumor, drug interference may cause false-positive or false-negative results. Other adrenocortical tumors may be detected by their production of elevated levels of urinary 17-ketosteroids.

C. STUDIES OF STONE CONSTITUENTS

Patients with recurrent urolithiasis may have an underlying abnormality of excretion of calcium, uric acid, oxalate, magnesium, or citrate. Samples of 24-hour urine collections can be tested to determine abnormally high levels of each. A few patients may have elevated cystine levels in urine. The nitroprusside test, a simple qualitative screening test for cystine, may indicate the need for quantifying cystine levels in timed urine collections. Whenever a stone is recovered, a formal stone analysis is recommended.

D. MISCELLANEOUS STUDIES

In patients with suspected fistulas of the urinary tract and bowel (eg, cancer of the colon, diverticulitis, regional ileitis), discoloration of the urine after ingestion of a poorly absorbed dye such as phenol red will confirm the diagnosis. In an equally satisfactory test for fistulas, the patient is instructed to ingest gelatin capsules filled with granulated charcoal and to submit a urine sample several days later. Examination of the centrifuged urinary sediment will reveal the typical black granules if a fistula is present.

Table 5–1. Comparison of Different Urine Tests for Bladder Transitional Cell Carcinoma.

<table>
<thead>
<tr>
<th>Test</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
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<tbody>
<tr>
<td>NMP22</td>
<td>71</td>
<td>75</td>
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<tr>
<td>BTA</td>
<td>52</td>
<td>85</td>
</tr>
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<td>67</td>
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<tr>
<td>FDP</td>
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<td>79</td>
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<tr>
<td>QUANTICYT</td>
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<td>100</td>
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<tr>
<td>Flow cytometry</td>
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<td>84</td>
</tr>
<tr>
<td>BLCA-4</td>
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<tr>
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<tr>
<td>Cytology</td>
<td>49</td>
<td>96</td>
</tr>
</tbody>
</table>

BTA, bladder tumor antigen; FDP, fibrin/fibrinogen degradation products; NMP22, nuclear matrix protein 22.

EXAMINATION OF URETHRAL DISCHARGE & VAGINAL EXUDATE

Urethral Discharge

Examination of urethral discharge in males can be particularly helpful in establishing a diagnosis. The following procedure, although exacting, provides proper specimens for determining the site of origin of bacteriuria or pyuria. Four sterile containers are labeled VB1, VB2, EPS, and VB3 (VB = voided bladder urine; EPS = expressed prostatic secretions). The patient is instructed to retract the foreskin and cleanse the meatus with benzalkonium chloride or hexachlorophene and to collect the urine specimens, capping the containers immediately afterward. The initial 10–15 mL of urine is collected in container VB1 and the subsequent 15–30 mL in container VB2. The prostate is then massaged, and secretions are collected in container EPS. The patient voids a final time, collecting the specimen in container VB3. An aliquot of each specimen is tested for nitrite and leukocyte esterase and then centrifuged, and the sediment is prepared for microscopic examination as described previously. A separate aliquot of each VB specimen and the EPS specimen are saved for subsequent culture if necessary. The presence of leukocytes or bacteria (or both) only in VB1 indicates anterior urethritis; if present in all 3 VB specimens, they may indicate cystitis or upper urinary tract infection; if present in EPS or VB3 only, they indicate a prostatic source of infection. Quantitative cultures can be similarly interpreted. Patients with positive results should be treated with appropriate antimicrobial drugs.

If the patient presents with the thick yellowish discharge typical of *Neisseria gonorrhoeae* infection, the discharge should be stained with Gram’s stain and examined for gram-negative intracellular diplococci. It is important to remember that commensal bacteria in smegma may to some extent contaminate sputum samples obtained from the urethra or from VB1 should be stained with methylene blue or Gram’s stain and examined microscopically. The presence of trichomonads, yeast cells, or bacteria in properly collected specimens indicates disease requiring treatment.

In cases of acute epididymitis, urinalysis and urine culture are often helpful in establishing the cause. Epididymitis is most commonly caused by *Chlamydia* species in young men and by *Escherichia coli* in men over 35 years of age. Culturing chlamydiae is time-consuming and expensive. Although a rapid immunofluorescence method of identifying *Chlamydia* is available, it is usually best to proceed with therapy based on the age of the patient and guided by clinical results.

The diagnosis of any sexually transmitted disease should raise the question of acquired immunodeficiency syndrome (AIDS). A recent study from the U.S. Centers for Disease Control and Prevention, in which a national sample of 12,571 males and females 15–44 years of age in 2002 were interviewed, demonstrated that one-half of this population reported that they had been tested at least once for HIV (other than through blood donation), and just over 15% had been tested within the past 12 months (Anderson et al, 2005). However, of those in this population deemed at high risk for HIV infection, one-third reported that they had never had an HIV test. This translates to an equivalent of 4.1–5.5 million, at-risk persons, aged 15–44 years, in the general population, having not been tested in the past year.

Vaginal Exudate

The underlying cause of vaginitis is often a viral, yeast, or protozoal infection or the presence of a foreign body (eg, retained tampon), and a simple physical examination may be all that is required for diagnosis.

Vaginal secretions obtained by use of a swab can be examined either stained or unstained. A drop of saline is added to a drop of specimen on a glass slide, mixed thoroughly, and covered with a coverslip. Examination under a low- or high-power lens may reveal yeast cells or trichomonads, thus suggesting appropriate therapy. Since bacteria are always present in the vagina, they generally are not significant findings in a wet smear.

RENAL FUNCTION TESTS

Urine Specific Gravity

With diminished renal function, the ability of the kidneys to concentrate urine lessens progressively until the specific gravity of urine reaches 1.006–1.010. However, the ability to dilute urine tends to be maintained until renal damage is extreme. Even in uremia, although the concentrating power of the kidneys is limited to a specific gravity of 1.010, dilution power in the specific gravity range of 1.002–1.004 may still be found. Determination of urine osmolality is undoubtedly a more meaningful measurement of renal function, but determination of specific gravity lends itself to office diagnosis.

Serum Creatinine

Creatinine, the end product of the metabolism of creatine in skeletal muscle, is normally excreted by the kidneys. Because individual daily creatinine production is constant, the serum level is a direct reflection of renal function. Serum creatinine levels remain within the normal range (0.8–1.2 mg/dL in adults; 0.4–0.8 mg/dL in young children) until approximately 50% of renal function has been lost. Unlike most other excretory products, the serum creatinine level generally is not influenced by dietary intake or hydration status.
Endogenous Creatinine Clearance

Because creatinine production is stable and creatinine is filtered through the glomerulus (although a small amount is probably secreted), its renal clearance is essentially equal to the glomerular filtration rate. The endogenous creatinine clearance test has thus become the most accurate and reliable measure of renal function available without resorting to infusion of exogenous substances such as radionuclides. Determination of creatinine clearance requires only the collection of a timed (usually 24-hour) urine specimen and a serum specimen. The resulting clearance is expressed in milliliters per minute, with 90–110 mL/min considered normal.

Because muscle mass differs among individuals, further standardization has been achieved and a corrected clearance level of 70–140 mL/min is considered normal.

Although creatinine is highly reliable as an estimate of renal function, values may be falsely low, particularly if only part of the urine is collected over the timed period or if a serum specimen is not collected concurrently.

Blood Urea Nitrogen

Urea is the primary metabolite of protein catabolism and is excreted entirely by the kidneys. The blood urea nitrogen (BUN) level is therefore related to the glomerular filtration rate. Unlike creatinine, however, BUN is influenced by dietary protein intake, hydration status, and gastrointestinal bleeding. Approximately two-thirds of renal function must be lost before a significant rise in BUN level becomes evident. For these reasons, an elevated BUN level is less specific for renal insufficiency than an elevated serum creatinine level. However, the BUN-creatinine (BUN-Cr) ratio can provide specific diagnostic information. It is normally 10:1; in dehydrated patients and those with bilateral urinary obstruction or urinary extravasation, the ratio may range from 20:1 to 40:1; patients with advanced hepatic insufficiency and overhydrated patients may exhibit a lower than normal BUN level and BUN-Cr ratio. In patients with renal insufficiency, extremely high BUN levels may develop that can be partially controlled by a decrease in dietary protein.

EXAMINATION OF BLOOD, SERUM, & PLASMA

Complete Blood Count

Normochromic normocytic anemia is often seen with chronic renal insufficiency. Chronic blood loss from microscopic hematuria is usually not sufficient to cause anemia, although gross hematuria certainly can be. A specific increase in the number of erythrocytes, as manifested by elevated hemoglobin and hematocrit levels (erythrocytosis, not polycythemia), may be indicative of a paraneoplastic syndrome associated with renal cell cancer. The leukocyte count is usually nonspecific, although marked elevations may indicate an underlying leukemia that may be the cause of urologic symptoms.

Blood Clotting Studies

Clotting studies are generally not necessary unless an insidious disorder such as von Willebrand disease, hepatic disease, or sensitivity to ingested salicylates is suspected in a patient with unexplained hematuria. The determination of prothrombin time and bleeding time (and perhaps partial thromboplastin time) is usually sufficient. A platelet count is important in patients receiving chemotherapy and those who have received extensive radiation therapy.

Electrolyte Studies

Serum sodium and potassium determinations may be indicated in patients taking diuretics or digitalis preparations and in patients who have just undergone transurethral prostatectomy. Serum calcium determinations are useful in patients with calcium urolithiasis. Elevated calcium levels are occasionally indicative of a paraneoplastic syndrome in patients with renal cell cancer. Serum albumin levels should be measured simultaneously with calcium levels to adequately assess the significance of the latter.

Prostate Cancer Markers

Prostate-specific antigen (PSA) is an extremely important prostate cancer marker. PSA is prostate-specific but not cancer specific. Serum elevation >4.0 ng/mL is correlated with prostatic cancer; however, serum levels vary with prostate volume, inflammation, and amount of cancer within the gland. PSA has become useful as a screening tool and is most useful as a marker of effective treatment (falls to zero following removal of organ-confined cancer) and early recurrence (Hernandez, Canby-Higano, and Thompson, 2005). The percentage of free PSA (ratio of unbound to total PSA) in the serum is useful for increasing the specificity of PSA for diagnosing prostate cancer. If the percentage of free PSA is <10%, approximately 60% of men will have prostate cancer, whereas if the percentage of free PSA is >25%, only 8% will have it.

Hormonal Studies

Serum parathyroid hormone studies are useful in determining the presence of a parathyroid adenoma in patients with urolithiasis and an elevated serum calcium level. Measurement of parathyroid hormone is not reliable, however, as a sole screening test for parathyroid adenoma and should not be used routinely in all patients with urolithiasis. Serum renin levels may be elevated in patients with renal hypertension, although many conditions can cause false-positive results. Studies of adrenal steroid hormones (eg, aldosterone, cortisol, epinephrine, norepinephrine) are useful in
determining adrenal function or the presence of adrenal tumors. Determinations of serum levels of the beta-subunit of hCG and of alpha-fetoprotein are indispensable in staging and in treatment follow-up for testicular tumors. One of these tumor markers is usually elevated in up to 85% of patients with nonseminomatous testicular tumors and can predict the recrudescence of tumor several months before disease is clinically evident. Serum testosterone studies can help to establish the cause of impotence or infertility.

Other Studies

The finding of elevated fasting plasma glucose levels in patients with urologic disease can establish the diagnosis of diabetes mellitus and thus indicate a possible cause of renal insufficiency, neurovesical dysfunction, impotence, or recurrent urinary tract infection. Serum uric acid levels are often elevated in patients with uric acid stones. Elevated serum complement levels may be diagnostic of underlying glomerulopathies.

LABORATORY VALUES IN ELDERLY PATIENTS

Clearly, some laboratory values change as patients age, others stay the same, and the effects of aging on some are as yet unknown. Laboratory values that do not change with increasing age include complete blood count, serum electrolytes, and hepatic function tests, among others (Table 5–2). Laboratory values that change as patients age include creatinine clearance, alkaline phosphatase, uric acid, and cholesterol (Table 5–3).

Other factors that may make laboratory interpretation more difficult include atypical disease presentation, multiple concurrent diseases, and prescription and nonprescription drug use.

REFERENCES


The field of diagnostic radiology continues to evolve, particularly in refinements to cross-sectional techniques. Imaging of the urinary tract, as a result, has become more precise, with new procedures offering a great selection of options, and new imaging algorithms being implemented. Ultrasonography, computed tomography (CT), and magnetic resonance imaging (MRI) provide higher soft-tissue contrast resolution than conventional radiography, as well as multiplanar imaging capability, resulting in significant advances in almost all areas of uroradiology. While such advances have produced new algorithms to approach the diagnostic imaging evaluation, each particular case also depends greatly on the equipment and professional talent available. In summary, ever-changing uroradiology remains indispensable in the diagnosis and treatment of patients with urologic disorders. This chapter will discuss the imaging techniques used in uroradiology, with summaries of the advantages and disadvantages of the various techniques, and will end with a brief discussion comparing imaging methods.

## RADIOGRAPHY

X-rays are electromagnetic waves with photon energies that typically fall between those of gamma rays and ultraviolet radiation. Radiography is possible because tissues differ in their ability to absorb x-rays. A radiopaque contrast medium is frequently employed to enhance soft-tissue contrast.

Although newer imaging techniques have largely replaced conventional radiography for diagnosis of many urologic problems, general radiography remains useful for some urologic disorders; therefore, the urologist should be familiar with x-ray equipment and uroradiologic techniques. The basic types of uroradiologic studies are plain (conventional) abdominal films, (also known as KUB, which stands for kidney, ureter, bladder) intravenous urograms (IVU), cystourethrogram, urethrogram, and angiograms. These studies are described separately in sections that follow.

### Basic Equipment & Techniques

1. **Radiography fluoroscopy**—Many conventional x-ray units contain both radiographic and fluoroscopic capabilities. These require a high voltage power supply, an x-ray tube, a collimating device, and an x-ray detector or film. Fluoroscopic units also use an electronic image intensifier and an image display system. Today, more radiology departments have become completely “filmless” as digital recording, displaying, and archiving of images are replacing film-based techniques.

2. **Image intensification**—Image intensifiers, coupled to video cameras, electronically augment the ordinary dim fluoroscopic image. Acquired images can be recorded and simultaneously viewed on monitors, usually in the x-ray room.

3. **Image recording**—Conventional recording of an x-ray image uses film and intensifying screens. The image intensifier and camera can be used to capture dynamic and static images. Real-time images may be recorded using photographic cine cameras, though this has been largely replaced by conventional or digital video. Conventional spot images may be acquired on x-ray film or digitally recorded, including rapid sequence images.

4. **Contrast media**—Radiographic contrast media used in uroradiology are water-soluble iodinated compounds that are radiopaque. Similar compounds are used for basic radiographic techniques and CT, though iodine concentrations will differ depending on preference and route of administration. In general, intravenous administration for CT or IVU is performed with iodine 200 mg/lb body weight in adults, and direct instillation to the collecting system or bladder is used similar media diluted to 15–45% concentration. The extracellular distribution of these agents results in improved contrast resolution and conspicuity of various structures.

Significant advances in water-soluble contrast media occurred with the introduction of low-osmolality (nonionic) organic iodine-containing compounds. When compared with conventional high-osmolality agents, these nonionic agents significantly improve patient tolerance and decrease the incidence of adverse reactions. Whether they reduce the mortality associated with the use of contrast media has not been proven. The major
Advantages & Disadvantages

Radiography produces anatomic images of almost any body part. Costs are moderate compared with cross-sectional imaging systems. Space requirements are modest, and portable equipment is available for use in hospital wards, operating rooms, and intensive care units. Because there are a great many specialists trained in radiography, its use is not confined to large medical centers. The major disadvantage of radiographic imaging is the use of ionizing radiation and relatively poor soft-tissue contrast. The evaluation of the urinary tract almost always requires opacification by iodine contrast media.

1. Plain Film of the Abdomen (Figures 6–1 through 6–3)

A plain film of the abdomen, frequently called a KUB film, is the simplest uroradiologic examination. It is generally the preliminary radiograph in extended radiologic examinations, such as intravenous urography, and is usually taken with the patient supine. It may demonstrate osseous abnormalities, abnormal calcifications, or large soft-tissue masses.

Kidney outlines usually can be seen on the plain film, so that their size, number, shape, and position can be assessed. The size of normal adult kidneys varies widely. The long diameter (the length) of the kidney is the most widely used and most convenient radiographic measurement. The average adult kidney is about 12–14 cm long. In children older than 2 years of age, the length of a normal kidney is approximately equal to the distance from the top of the first to the bottom of the fourth lumbar vertebral body. Patterns of calcification in the urinary tract (Figures 6–1 and 6–2) may help to identify specific diseases.

2. Urography (Figures 6–4 through 6–8)

The collecting structures of the kidneys, ureters, and bladder can be demonstrated radiologically with contrast media by the following methods:

Intravenous Urography

The IVU, also known as excretory urography (EU) (Figure 6–4), or intravenous pyelography (IVP), can demonstrate a wide variety of urinary tract lesions (Figures 6–4 and 6–5), is simple to perform, and is well tolerated by most patients.

Sonography, CT, and MRI have replaced urography in many cases. Nevertheless, urography is still occasionally used and is useful for demonstrating small lesions in the urinary tract (eg, papillary necrosis, medullary sponge kidney, uroepithelial tumors, pyeloureteritis cystica).

A. Patient Preparation

At one time dehydration was advocated as optimal preparation for intravenous urography (EU). This is no longer required. Furthermore, dehydration is to be avoided in infants, debilitated and elderly patients, and patients with
diabetes mellitus, renal failure, multiple myeloma, or hyperuricemia.

It is controversial whether preliminary bowel cleansing is beneficial. The choice may be made according to individual preference.

**B. STANDARD TECHNIQUE**

Following a preliminary plain film of the abdomen, additional radiographs are taken at timed intervals after the intravenous injection of iodine-containing contrast medium. Normal kidneys promptly excrete contrast agents, almost entirely by glomerular filtration.

The volume and speed of injection of the contrast medium, as well as the number and type of films taken, vary by preference, patient tolerance, and the particular clinical scenario.

**C. TECHNIQUE MODIFICATIONS**

Radiographic tomography, x-ray imaging of a selected plane in the body, permits recognition of kidney structures that otherwise are obscured on standard radiograms by extrarenal shadows, for example, those due to bone or feces (Figure 6–6). Image-intensified fluoroscopy permits study of urinary tract dynamics. “Immediate” films, which are taken immediately after the rapid (bolus) injection of contrast, typically show a dense nephrogram and permit better visualization of renal outlines. Abdominal (ureteral) compression devices temporarily obstruct the upper urinary tract during EU and improve the filling of renal collecting structures. “Delayed” films, taken hours later or on the following day, can contribute useful information. “Upright” films, taken with the patient standing or partially erect, reveal the degree of mobility and drainage of the kidneys and, if taken imme-
diately after the patient has voided ("postvoiding" film), show any residual urine in the bladder.

**Retrograde Urograms**

Retrograde urography is a minimally invasive procedure that requires cystoscopy and the placement of catheters in the ureters. A radiopaque contrast medium is introduced into the ureters or renal collecting structures through the ureteral catheters (Figures 6–7 and 6–8), and radiographs of the abdomen are taken. This study must be performed by a urologist or experienced interventional uroradiologist. Some type of local or general anesthesia should be used, and the procedure occasionally causes later morbidity or urinary tract infection.

Retrograde urograms may be necessary if excretory urogams or CT urogram (CTU) are unsatisfactory, if the patient has a history of adverse reaction to intravenous contrast media, or if other methods of imaging are unavailable or inappropriate.

**Percutaneous Urograms**

Outlining the renal collecting structures and ureters by percutaneous catheter is occasionally done when excretory or retrograde urography has failed or is contraindicated, or when there is a nephrostomy tube in place and delineation of the collecting system is desired. For antegrade studies, contrast medium is introduced either through nephrostomy tubes (nephrography) or by direct injection into the renal collecting structures via a percutaneous puncture through the patient’s back. Percutaneous retrograde urograms of the upper urinary tract are made by retrograde injection of contrast medium through the opening of a skin ureterostomy or pyelostomy (skin ureterogram, skin urogram) or through the ostium of an interposed conduit, usually a segment of small bowel (loopogram).

**3. Cystography, Voiding, Cystourethrography, & Urodynamics (Figures 6–9 through 6–12)**

Direct instillation of contrast media into the urinary bladder (cystography) is preferred over EU for more focused examination of the bladder. Contrast is usually instilled via a transurethral catheter, but when necessary can be administered via percutaneous suprapubic bladder puncture. For urodynamic studies, pressure transducers are used within the bladder lumen and rectum for dynamic measurement of intraluminal and intra-abdominal pressures, respectively. Radiographs can be taken using standard overhead
Figure 6–3. Plain films of the abdomen with abnormal radiolucencies. **Left:** Emphysematous pyelonephritis. Interstitial striated pattern of radiolucent gas throughout the entire left kidney. Similar changes were present in the right kidney. 58-year-old diabetic man with pyuria and septic shock. **Right:** Gas pyelogram. No interstitial gas, but gas fills dilated left kidney calices, pelvis, and ureter. 50-year-old diabetic woman with sepsis and left upper urinary tract infection due to gas-forming microorganisms.

Figure 6–4. Abnormal excretory urograms. **Left:** Medullary sponge kidney. Pronounced medullary tubular ectasia (arrows) of entire right kidney. Similar findings were present in upper pole pyramids of left kidney, and small medullary calculi were present in some areas of tubular ectasia in both kidneys. 34-year-old woman with repeated bouts of chills, fever, and left flank pain. **Right:** Renal tuberculosis. Irregular cavitation of lower pole pyramid (arrow). 22-year-old woman with positive urine culture for tuberculosis.
Figure 6–5. Abnormal excretory urograms. **Left:** Crossed fused ectopia. Composite of 2 films from an excretory urogram shows ectopic right kidney (R) fused to left kidney (L). Right ureter (arrows) crosses midline and enters normally into right side of bladder. Healthy 31-year-old female potential kidney donor. **Right:** Infantile polycystic kidney disease. Very large kidneys with radiopaque spoke pattern radiating out to cortex. 26 hours after administration of intravenous contrast medium. 4-month-old girl with bilateral abdominal masses.

Figure 6–6. Radiographic tomography. Tomography is used to image a plane in the body. The technique was widely used in uroradiology, often permitting demonstration of lesions otherwise hidden by overlying soft tissues or obscuring bowel shadows. However, computed tomography (CT) is rapidly replacing conventional excretory urograms, and thus tomography is declining in its use as well. **Left:** Transitional cell carcinoma. The tumor in the pelvis (arrow) is clearly shown free of obscuring gas shadows present on the nontomographic films. 56-year-old man with history of renal calculi. **Right:** Renal cell carcinoma (T). Displacement of mid-kidney collecting structures and a nephrogram defect are seen free of obscuring splenic flexure fecal shadows that were present on the nontomographic films. 44-year-old woman with fever, weight loss, anemia, and history of contralateral nephrectomy for carcinoma 15 years earlier.
Figure 6–7. Retrograde urograms and nephrostograms; lower ureters not all shown.

Upper left: Normal retrograde urogram. Intrarenal collecting structures, pelvis, and ureter are normal. Adult male with microscopic hematuria and previous technically unsatisfactory excretory urogram. Upper right: Squamous cell carcinoma. Marked irregular filling defects involving calices, pelvis, and proximal ureter, with communicating abscess cavity in upper pole (arrow). Kidney also showed squamous metaplasia and contained calculi. 51-year-old woman with 2-week history of left flank cellulitis and tenderness.

Lower left: Transitional cell carcinoma. Severe deformity with filling defects in right upper pole calices (curved arrow) and blood clots in lower calices and at ureteropelvic junction (straight arrow). 65-year-old man with gross hematuria and right flank pain.

Lower right: Fungus balls. Nephrostogram revealing 2 filling defects (arrows) in renal pelvis. Copious fungal matter aspirated through nephrostomy catheter. 65-year-old diabetic woman who had undergone left nephrectomy, with percutaneous nephrostomy catheter (white arrow) for obstruction of right kidney.
Figure 6–8. Abnormal retrograde urograms. **Upper left:** Idiopathic retroperitoneal fibrosis. Smooth narrowing of both mid ureters (arrows), with bilateral proximal ureterectasis and hydronephrosis. 51-year-old woman with no urinary tract symptoms. **Upper right:** Functional ureteral obstruction. Obstruction was due to congenitally abnormal muscle arrangements in the affected very distal ureter (curved arrow). Pronounced hydronephrosis and dilatation of ureter (U) proximal to the short segment of abnormal ureter. 13-year-old boy with repeated urinary tract infections. **Lower left:** Transitional cell carcinoma of the ureter. No contrast medium has passed beyond the large, bulky, right ureteral tumor (arrow). The ureteral widening below the tumor is distinctive and is sometimes referred to as the “champagne glass” sign (in this instance, the glass is tipped on its side). 76-year-old man with nonfunctioning right kidney. **Lower right:** Ureteral constrictions secondary to extension of carcinoma of the colon. Bilateral distal ureteral narrowings (arrows) with upper tract obstruction. Composite of separate retrograde urograms. E = unintended extravasation about tip of left ureteral catheter. 76-year-old man with cancer of the sigmoid colon.
x-rays, or during fluoroscopy. Voiding cystourethograms are radiographs of the bladder and urethra obtained during micturition. Cystography and cystourethrography are important radiologic techniques for detecting vesicoureteral reflux and may be used in the workup of patients with urinary stress incontinence. CT cystography (CT of the pelvis after the instillation of dilute contrast medium into the bladder) has been shown useful in the evaluation of traumatic bladder rupture.

4. Urethrography (Figures 6–13 through 6–15)

The urethra can be imaged radiographically by retrograde injection of radiopaque fluid or in antegrade fashion with voiding cystourethrography, or with voiding following EU. The antegrade technique is required when lesions of the posterior urethra, for example, posterior urethral valves, are suspected; the retrograde technique is more useful for examining the anterior (penile) urethra.

5. Vasography (Figure 6–16)

Vasoseminal vesiculography is most often used in the investigation of male sterility. The radiopaque contrast medium is introduced into the ductal system by direct injection into an ejaculatory duct following panendoscopy or, more commonly, by injection into the vas deferens after it has been surgically exposed through a small incision in the scrotal neck.

6. Lymphangiography (Figure 6–17)

Lymphangiography has been largely abandoned and replaced by CT and MRI.

7. Angiography

Nearly 50 years after Seldinger described techniques for percutaneous arteriography, catheter angiography maintains a role in treatment of some urologic disorders but is being replaced by CT or MRI for diagnostic examinations. Although an established imaging technique with proven value and an acceptable incidence of complications and morbidity, angiography is moderately invasive and relatively expensive.

Aortorenal & Selective Renal Arteriography (Figure 6–18)

Conventional arteriographic studies are performed almost exclusively by percutaneous needle puncture and catheterization of the common femoral arteries. Rapid sequence images are obtained during catheter injection of nonionic contrast. Aortograms at the level of the renal vessels using multihole “flush” catheters demonstrate renal vessels, including anomalies. Selective catheterization of renal arteries follows. CT and MR angiography involve peripheral injection of contrast media with breath hold rapid image acquisition through the targeted region of interest. This is usually performed after a timing bolus sequence.
Figure 6–10. Abnormal cystograms: retrograde cystograms or “cystograms” as part of excretory urogram studies. **Upper left:** Ectopic ureterocele. Giant ureterocele (straight arrows) to hydrenephrotic, nonfunctioning upper portion (curved arrow) of duplex right kidney. 9-month-old-girl with urinary tract infections. **Upper right:** Pelvic lipomatosis. Pear-shaped bladder and increased radiolucency of the pelvic soft tissues secondary to pelvic lipomatosis of severity sufficient to produce obstructive dilatation of the upper urinary tracts. Filling defects (arrows) at bladder base due to cystitis glandularis. 62-year-old man with intermittent left flank pain. **Lower left:** Rupture of the membranous urethra. Pear-shaped bladder secondary to extraperitoneal extravasation (E) and perivesical hematoma. Arrow = inflated balloon of Foley catheter. 41-year-old man with renal transplant, after a motor vehicle accident that resulted in pelvic bone fractures, separation of the sacroiliac joints, and dislocation of the left (L) but not the right hip prosthesis (patient has bilateral hip prostheses). **Lower right:** Bladder hernia. Bilateral obstructive ureterectasis (small arrows) secondary to remarkable herniation of the entire bladder (large arrow, B) into the inguinal region, 5’5”, 225 lb, 53-year-old man with panniculus reaching to mid thigh, complaining of difficulty voiding.
Figure 6–11. Abnormal cystograms: retrograde cystograms or “cystograms” as part of excretory urogram studies. Upper left: Neurogenic bladder. This neurogenic bladder has a “Christmas-tree” shape, with gross trabeculation and many diverticula. Residual myelographic contrast medium in spinal canal (straight arrow). Right vesicoureteral reflux (curved arrow). 70-year-old man with urinary incontinence. Upper right: Congenital “hourglass” bladder. Transverse concentric muscular band (arrows) separates upper and lower bladder segments, both of which contracted and emptied simultaneously and completely with voiding. 66-year-old woman with urinary stress incontinence. Lower left: Hodgkin’s disease of bladder. Global thickening of the bladder wall (arrows), more apparent on the left. 54-year-old man with generalized Hodgkin’s disease. Lower right: Papillary transitional cell bladder carcinoma. Huge (12 cm) cauliflower-like bladder mass (arrows) filling almost the entire bladder. “Cystogram” film of an excretory urogram in a 40-year-old man with recurrent bladder tumor.
Figure 6–12. Abnormal prostate and posterior urethra: cystograms and urethrograms. **Upper left:** Benign prostatic hyperplasia. Gross enlargement of prostate gland producing marked elevation (arrows) of the bladder base. The bladder shows small diverticula and slight trabeculation. Excretory urogram (cystogram) in a 65-year-old man with history of obstructive voiding symptoms. **Upper right:** Foreign body (eyeliner pencil cover) lodged in bladder and prostatic urethra, with urethrectal fistula. Radiopaque medium enters rectum and sigmoid colon (S) through fistula (arrow) from prostatic urethra. Retrograde urethrogram in a 43-year-old man. **Lower left:** Rhabdomyosarcoma of prostate. Lobulated filling defects (large arrow) encroaching on widened prostatic urethra. Voiding cystourethrogram in a 5-year-old boy with voiding difficulties. Small arrow = penile urethra. **Lower right:** Posterior urethral valves. Marked dilatation and elongation of prostatic urethra (P), with reflux into prostatic ducts (straight arrow) secondary to posterior urethral valves (curved arrow) with bilateral vesicoureteral reflux into dilated ureters (U). Voiding cystourethrogram in a 10-day-old boy.
Figure 6–13. Abnormal anterior urethras: voiding cystourethograms and retrograde urethograms. **Upper left:** Voiding cystourethrogram in a 78-year-old man with a history of urethral diverticulum of unknown etiology. 4-cm anterior urethral diverticulum (large arrow) and left vesicoureteral reflux (small arrow). **Upper right:** Urethral diverticulum in a woman. Large irregular diverticulum (arrow). Voiding cystourethrogram in a 51-year-old woman with voiding difficulties and suspected urethral stricture. **Lower left:** Ruptured urethra. Extravasation of contrast medium around the membranous urethra (arrows). Retrograde urethrogram in a 16-year-old boy in whom blunt perineal trauma was followed by bloody urethral discharge and inability to void. **Lower right:** Urethroscrotal fistula. Extravasation (E) into extraurethral tissues from fistula in bulbous urethra (arrow). Retrograde urethrogram in a 26-year-old man after end-to-end urethroplasty for stricture.
Figure 6-14. Abnormal anterior urethras: retrograde urethrograms. **Upper left:** Urethral carcinoma. Gross irregularities with filling defects involving most of penile urethra. Poorly differentiated carcinoma of anterior urethra in a 59-year-old man with obstructive voiding symptoms and inguinal adenopathy. **Upper right:** Urethral carcinoma. Filling of irregular sinus tracts and channels in a large epidermoid carcinoma of the bulbocavernous urethra (straight arrow). There are multiple thin transverse strictures of the penile urethra (curved arrow). 75-year-old man with obstructive voiding symptoms and 30-year history of urethral strictures requiring dilatations. **Lower left:** Focal urethral stricture (arrow). Middle-aged man with obstructive voiding symptoms who denied any previous urethritis. **Lower right:** Urethral strictures. Multiple strictures in the bulbocavernous urethra (lower arrow) with reflux into Cowper’s gland (middle arrow) and prostatic ducts (upper arrow). B = bladder; D = bladder diverticulum. 62-year-old man with 25-year history of urethral strictures requiring frequent dilatations.
Figure 6–15. Congenital genitourinary anomalies: voiding cystograms and retrograde urethrogram. **Upper left:** Utricle. Midline outpouching (arrow) from verumontanum between orifices of ejaculatory ducts, representing Müllerian duct cyst. **Upper right:** Gas cystogram combined with injection of utricle, oblique view. M = grossly dilated utricle (Müllerian duct cyst); straight arrows = bladder distended with air; curved arrow = coincident partial filling of left seminal vesicle and vas deferens. 34-year-old man with urgency, frequency, and suspected retrograde ejaculation. **Lower left:** Common urogenital sinus. Vagina (V) and urethra (U) join (at arrow) into a common urogenital sinus (S). Voiding cystourethrogram in a 3-week-old female pseudohermaphrodite with ambiguous genitalia and congenital adrenal hyperplasia. **Lower right:** Male pseudohermaphrodite. Bladder is distended with urine (black arrows). Retrograde urethrogram via hypospadiac meatus has fortuitously and selectively filled with contrast medium an extensive müllerian duct remnant consisting of vagina (V), cervix and cervical canal (C), and retroverted uterus (U). Residual contrast medium in hypoplastic anterior urethra (white arrow). 27-year-old man with small external genitalia, hypospadias, and perineal pain.
Figure 6–16. Vasoseminal vesiculography (vasography). Left: Normal left vasoseminal vesiculogram. V = vas deferens; S = seminal vesicle; E = ejaculatory duct. 40-year-old man with hypospermia. Right: Seminal vesiculitis. Bilateral vasogram. Mass (M) produced by the swollen, nonfilling right seminal vesicle has displaced both ejaculatory ducts (E) toward the left and indented the medial aspect of the proximal left seminal vesicle and vas deferens (V). 33-year-old man with painful ejaculations after repair of right varicocele.

Figure 6–17. Lymphangiography. Hodgkin’s disease. Extensive retroperitoneal adenopathy. The kidneys and ureters were displaced by grossly involved abdominal and pelvic lymph nodes as well. Lymphangiography has now been essentially replaced by computed tomography (CT) or magnetic resonance (MR) imaging. 52-year-old woman with stage IV Hodgkin’s disease.
CHAPTER 6

CT angiography offers higher spatial resolution than magnetic resonance angiography (MRA), but carries the risks of radiation exposure and iodinated contrast usage.

Indications for renal arteriography include suspected renal artery stenosis (renovascular hypertension), vascular malformations, tumor embolization to minimize surgical blood loss or treat bleeding tumors, and trauma. Diagnostic renal angiography to demonstrate renal vascular anatomy is uncommon today, as this information may usually be obtained noninvasively. Complications from conventional catheter angiography include bleeding at the puncture site, contrast allergy or nephrotoxicity, and renal or distal emboli.

**Inferior Venacavography & Selective Venography (Figures 6–19 and 6–20)**

The common femoral veins or less commonly the internal jugular vein are catheterized for catheter angiography of the inferior vena cava, renal, and adrenal veins. Risks of bleeding and emboli present in arterial studies are virtually eliminated. Venography is rarely used today since the

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**Figure 6–18.** Angiography: aortorenal arteriography. **Upper left:** Normal abdominal aortogram. The aortic catheter is hidden by the opacified normal aorta. Right (R) and left renal arteries and branches are well shown, as are the splenic (S) and hepatic (H) arteries arising from the celiac axis. The superior mesenteric artery is superimposed over the aortic silhouette and is not visible on this study. 28-year-old healthy female potential kidney donor. **Upper right:** Bilateral renal artery stenoses. Typical angiographic appearance and location of stenoses caused by atherosclerosis (small arrows) and fibromuscular dysplasia (large arrows). 58-year-old woman with abdominal bruits and a 16-year history of hypertension. **Lower left:** 3D coronal CT angiography image demonstrates an inferior accessory left renal artery (posterior view). **Lower right:** The left accessory renal artery origin (*) is better demonstrated rotating the model in the axial plane. 65-year-old man undergoing preoperative evaluation for laparoscopic partial nephrectomy.
**Figure 6–19.** Angiography: inferior venacavography. **Upper left:** Normal inferior vena cava (C). Unusual retrograde filling of morphologically normal renal veins (arrows) from antegrade injection into the inferior vena cava is probably due to reduced venous outflow from the kidneys with the patient in Valsalva maneuver. B = retained contrast material in the cecum from previous barium enema examination. Woman with arteriolar nephrosclerosis and renal failure. **Upper right:** Inferior vena cava obstruction. Complete block of the vena cava (curved arrow) by extension from right renal vein of tumor thrombus from a right renal carcinoma. Note cephalad blood return via the paralumbar veins (straight arrows). 60-year-old man with gross hematuria. **Lower left:** Double inferior vena cava (R, L). Persistent left supracardinal vein anomaly. 23-year-old man after orchiectomy for testicular teratocarcinoma. **Lower right:** Example of duplicated IVC on IV contrast enhanced axial CT. Normal IVC (arrow) and duplicated IVC (*).
information can be obtained at cross-sectional imaging (CT or MRI) in almost all cases. Adrenal and renal venography is performed occasionally in the setting of venous sampling to localize hormone secretion in patients with indeterminate noninvasive imaging studies.

Miscellaneous Urologic Angiography (Figure 6–21)

Although angiography has little or no value in examination of the ureter, bladder, adrenals, and prostate, angiograms of these structures may be indicated in particular clinical situations, in which case the studies are usually “tailored” to the clinical problem. In this era of multiple cross-sectional methods, these procedures are rarely used. Corpus cavernosograms are made by direct injection of suitable contrast material into the corpora cavernosa of the penis. They can be useful in examining for Peyronie’s disease, impotence, priapism, and traumatic penile lesions, but these also are not commonly performed.

SONOGRAPHY (FIGURES 6–22 THROUGH 6–27)

Basic Principles

Sound is the mechanical propagation of pressure changes, or waves, through a deformable medium. A wave frequency of 1 cycle/s (cps) is called a hertz (Hz). Sound frequencies greater than 20 kHz are beyond the range of human hearing and are called ultrasound. Medical sonography uses ultrasound to produce images. The frequencies commonly used in medical sonography are between 3.5 and 15 MHz.

Ultrasound waves for imaging are generated by transducers, devices that convert electrical energy to sound energy and vice versa. These transducers are special piezoelectric crystals that emit ultrasonic waves when they are deformed by an electrical voltage and, conversely, generate an electrical potential when struck by reflected sound waves. Thus, they act as both sonic transmitters and detectors. In general concept, medical sonography resembles naval submarine sonar. Ultrasound images are reflection images formed when part of the sound that was emitted by the transducer bounces back from tissue interfaces to the transducer. The sound reflected by stationary tissues forms the data set for anatomic gray-scale images. The sound reflected by moving structures (eg, flowing blood in a vessel) has an altered frequency due to the Doppler effect. By determining the Doppler shift, vascular flow direction and velocity can be encoded graphically (spectral Doppler) or by color (color Doppler). A more sensitive method of detecting flow, called power mode Doppler, is available on modern equipment. This technique displays the integrated power of the Doppler signal rather than the mean Doppler...
Figure 6–21. Angiography: miscellaneous urovenography. Upper left: Penoscrotal varices. Penile venography. Many tortuous veins in the penis (P) and scrotum (S). 14-year-old boy with long-standing penile and scrotal varicosities and numerous scrotal phleboliths. Upper right: Varicocele. Gonadal venography. Dilated, tortuous varicosities of the pampiniform plexus in the left scrotum. 31-year-old man with recurrence of scrotal pain after varicocele ligation. Lower left: Normal corpora cavernosogram. Injection of contrast medium into the left corpus (L), with normal (albeit slightly less) filling of the right corpus (R). 57-year-old man with impotence. Lower right: Penile fibrosis. Corpora cavernosogram. Injection of right corpus produces no filling of the proximal right corpus (arrow); there is normal filling of the left corpus (L). 33-year-old man with “crooked penis” following unsuccessful penile prosthesis operation.
frequency shift. Direction or velocity of flow is not displayed in the power mode.

Reflected sound received by the transducer is converted into electrical signals that are analyzed by computer algorithms, and rapidly converted into video images viewed directly on a real-time display. Images are rapidly updated on the display, giving an integrated cross-sectional anatomic depiction of the site studied. Individual frames may be frozen during an examination for motion-free analysis and recording, or continuous images may be recorded as digital or conventional video.

**Clinical Applications**

Ultrasound is commonly used for the evaluation of the kidney, urinary bladder, prostate, testis, and penis.

Ultrasound is useful for assessing renal size and growth. It is also helpful in triaging patients with renal failure. For example, small echogenic kidneys suggest renal parenchymal (medical) disease, whereas a dilated pelvocaliceal system indicates an obstructive, and potentially reversible, cause of renal failure.

Renal ultrasound is useful in detection and characterization of renal masses. Ultrasound provides an effective method of distinguishing benign cortical cysts from potentially malignant solid renal lesions. Since the most common renal lesion is a simple cortical cyst, ultrasound is a cost-effective method to confirm this diagnosis. Ultrasound may also be used to follow up mildly complicated cysts detected on CT, for example, hyperdense cysts or cysts with thin septations.

The differential diagnosis for echogenic renal masses includes renal stones, angiomyolipomas, renal cortical neoplasms (including carcinoma), and, less commonly, abscesses and hematomas. All echogenic renal masses should be correlated with clinical history and, if necessary, confirmed with another imaging modality or follow-up ultrasound. Thin-section CT showing fat within the renal lesion characterizes it as a benign angiomyolipoma, and no further investigation is required. Echogenic lesions smaller than 1 cm are more difficult to characterize by CT owing to partial volume averaging; in the correct clinical setting, follow-up ultrasound rather than repeat CT may be more useful.

Doppler sonography is useful for the evaluation of renal vessels, vascularity of renal masses, and complications following renal transplant. It can detect renal vein thrombosis, renal artery stenosis, and ureteral obstruction prior to the development of hydronephrosis, arteriovenous fistulas, and pseudoaneurysms. Perinephric fluid collections following renal transplantation, extracorporeal shockwave lithotripsy, or acute obstructions are reliably detected by ultrasound.

Developments in other imaging modalities have decreased the use of ultrasound in several clinical scenarios. Most patients with suspected renovascular hypertension are evaluated with CTA or MRA rather than Doppler

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**Figure 6–22.** Sonography of the kidney. **Upper:** Normal kidney. Renal cortex (C), normal renal sinus echoes (S). **Middle:** Moderate hydronephrosis and hydroureter; dilated renal pelvis (P). Dilated proximal ureter (prox ure). **Lower:** Severe hydronephrosis of the transplanted kidney, compound sagittal scans, dilated clubbed calices (C), dilated renal pelvis (P).
ultrasonography. Unenhanced helical CT is now the initial procedure of choice for the evaluation of the patient with acute flank pain and suspected urolithiasis. In addition to rapidly and sensitively detecting renal stones without the need for intravenous contrast medium, helical CT also has the potential for identifying other causes of flank pain such as appendicitis and diverticulitis. In the past, a combination of KUB and ultrasound was advocated for the evaluation of hematuria, but recent studies indicate that IVU, CT (CTU), or both are the preferred modalities to evaluate this common clinical problem.

Applications of bladder sonography include assessment of bladder volume and wall thickness, and detection of bladder calculi and tumors. The suprapubic transabdominal approach is most commonly used. The transurethral approach during cystoscopy has been recommended for tumor detection and staging.

Ultrasound examination of the testis has become an extension of the physical examination. The superficial location of the testis allows the use of a high-frequency transducer (10–15 MHz), which produces excellent spatial resolution. The addition of color Doppler sonography provides simultaneous display of morphology and blood flow. Normal low-resistance intratesticular arterial blood flow is consistently detected with power or color Doppler. Sonography is highly accurate in differentiating intratesticular from extratesticular disease, and in the detection of intratesticular pathology. Ultrasound is commonly used to evaluate acute conditions of the scrotum. It can distinguish between inflammatory processes, inguinal hernias, and acute testicular torsion. In addition, epididymitis not responding to antibiotics within 2 weeks should be investigated further with scrotal ultrasonography.

**Advantages & Disadvantages**

The main advantages of ultrasound are ease of use, high patient tolerance, noninvasiveness, lack of ionizing radia-
tion, low relative cost, and wide availability. Disadvantages include a relatively low signal-to-noise level, tissue non-specificity, limited field of view, and dependence on the operator’s skill and the patient’s habitus.

**COMPUTED TOMOGRAPHY SCANNING (FIGURES 6–28 THROUGH 6–34)**

**Basic Principles**

In CT scanning, a thin, collimated beam of x-rays is passed through the patient and captured by an array of solid-state or gas detectors.

The interconnected x-ray source and detector system are rapidly rotated in the gantry around the recumbent patient. Computers integrate the collected x-ray transmission data to reconstruct a cross-sectional image (tomogram).

Spiral (or helical) CT uses a slip-ring gantry that rotates continuously while the patient moves constantly through the collimated x-ray beam. Spiral CT technology affords the ability to image during specific phases of contrast bolus enhancement, including the ability to perform CT angiography, and allows improved image reformations. Multidector, or multislice, helical CT scanners have an array of multiple rows of detectors in a helical scanner such that multiple scan images can be acquired per gantry rotation, and as a by-product thinner sections and higher resolution achieved. Such systems are optimally paired with powerful computer workstations so that high-quality three-dimensional and multiplanar reformations can be quickly generated and analyzed.

Figure 6–24. Sonography of renal neoplasms. **Upper left:** Simple renal cyst (Cy) demonstrating sharp interfaces toward the renal parenchyma, no internal echoes, and increase through transmission. **Upper right:** Complex renal cyst (arrow) with lobulated margins and thick wall. **Lower left:** Solid tumor (T) in upper pole of left kidney with increased echogenicity relative to adjacent renal parenchyma. Pathology was oncocytoma. **Lower right:** Solid renal tumor (*) in the right kidney (K) with separate hyperechoic interpolar partially exophytic mass. The interpolar mass represented a known angiomyolipoma, while the upper pole mass represented renal cell carcinoma.
Clinical Applications

Renal CT is most commonly used in the evaluation of acute flank pain, hematuria, renal infection (search for abscess) and renal trauma, and in the characterization and staging of renal neoplasm. CT evaluation of renal anatomy and pathology generally requires intravenous injection of iodinated contrast media; noncontrast scans are needed, however, when renal or perirenal calcification, hemorrhage, or urine extravasation is suspected, since scans obtained after the administration of contrast media may mask these abnormalities. Also, pre- and postcontrast scans are required to determine whether a mass is solid or cystic.

Contrast media is usually administered as a rapid intravenous bolus for assessment of renal anatomy or measurement of aortorenal transit time. Using a bolus injection and rapid sequence scanning, renal arterial opacification is followed immediately by enhancement of the cortex. A nephrogram phase with medullary enhancement is reached within 60 seconds. Excretion of contrast material into the collecting structures can be expected within 2–3 minutes after initiation of contrast administration.

Although CT can detect ureteral tumors, the current role of CT in the evaluation of the ureters is predominantly for tumor staging and evaluation of the cause and level of obstruction. Helical CT without oral or intravenous contrast is the preferred imaging modality for patients with renal colic or suspected urolithiasis (Figure 6–33).

In the evaluation of the urinary bladder, CT is used primarily in staging bladder tumors and in diagnosing bladder rupture following trauma. Performing CT after filling the bladder with dilute contrast medium (CT cystography) improves the sensitivity of this modality for detecting tumors and bladder rupture. For prostate diseases, CT is used for detection of lymphadenopathy and to delineate prostatic abscesses. CT is used for detection of the abdominal location of suspected undescended testes, for staging of testicular tumors, and in the search for nodal or distant metastasis.

The addition of delayed CT imaging 10–15 minutes postintravenous contrast–enhanced CT shows high sensitivity and specificity in characterizing adrenal lesions. Benign adenomas, including lipid poor adenomas, show brisk contrast washout. CTA or MRA are replacing conventional angiography for diagnostic examinations.

Advantages & Disadvantages

The main advantages of CT include a wide field of view, the ability to detect subtle differences in the x-ray attenuation properties of various tissues, good spatial resolution, anatomical cross-sectional images, and operator independence. A considerable amount of diagnostic information
Figure 6–26. The use of transrectal ultrasound in the evaluation of the prostatic urethra. **Upper left:** Sonographic appearance of the prostatic urethra (U) following transurethral resection as seen on transrectal ultrasound in the sagittal plane of scanning. Urinary bladder (B). The urethra (U) is dilated to the level of the verumontanum (arrow). Peripheral zone (P), rectum (R). **Upper right:** The prostatic urethra (U) is dilated to the level to the membranous urethra (arrow). Urinary bladder (B). The cursors are placed to measure the length of the prostatic urethra. **Lower left and lower right:** Examples of testicular ultrasound. **Lower left:** The right testis (T) is normal. There is a hypo-echoic lesion within the left testis (asterisk). At surgery, it was a seminoma. **Lower right:** A large mixed solid and cystic intratesticular mass with foci of echogenic calcifications. Benign epidermoid cyst. This mass did not show concentric lamellation sometimes associated with epidermoid cysts. Ultrasound cannot always differentiate epidermoid cysts from malignant germ cell neoplasms.
Figure 6–27. Gray-scale and Doppler sonography: acute rejection in a renal transplant. **Upper left:** Gray-scale ultrasound image of transplant kidney shows poor corticomedullary differentiation. A small fluid collection is seen within the renal pelvis (arrow). Native external iliac vessels are seen as tubular hypoechoic structures (arrowheads). **Upper right and lower left:** Color Doppler images demonstrate flow within the native external iliac artery (arrowheads), the transplant renal artery (long arrow), and the interlobar arteries (short arrow). **Lower right:** Spectral Doppler analysis reveals an elevated resistive index of 0.84. These findings are compatible with, but not specific for, acute rejection. In the nonacute setting, cyclosporin toxicity or chronic rejection may also show elevated arterial resistive indices.
Figure 6–28. CT scans: adrenal glands. **Upper left:** Normal adrenal glands (arrows) have inverted arrowhead or Y shape. **Upper right:** CT densitometry. Thin section CT of incidental right adrenal mass (M) performed without intravenous contrast. Region of interest density measurement was below 10 Hounsfield Units (near zero), compatible with adrenal adenoma, confirmed by pathology. **Mid left:** Bilateral adrenal lymphoma. Enlarged adrenal glands (arrows) anterior to normal kidneys. L = liver. 53-year-old man with abdominal pain and histiocytic lymphoma of the central nervous system. **Mid right:** Left adrenal carcinoma. Large tumor (T) in left upper retroperitoneum with necrotic or cystic changes. Differential for CT included exophytic renal carcinoma. 52-year-old female with pulmonary nodules (metastases). **Lower left:** Axial CT image reveals a predominant solid mass (arrow) abutting the left kidney, with areas of cystic change. **Lower right:** Coronal oblique reformatted image from the same patient shows the mass (arrow) to be inseparable from the inferior left adrenal limb (*). Pathologically proven adrenal carcinoma.
available from CT scans depends on patterns of contrast enhancement, so a carefully tailored examination is essential. Reformatted helical image data in different planes and in 3D has made renal CT imaging, with renal angiography and urography, valuable in preoperative planning, such as for partial nephrectomy. Limitations of CT include restriction to the transaxial plane for direct imaging, tissue non-specificity, low soft-tissue contrast resolution, and the need for contrast media (both oral and intravenous). Even with careful use of contrast media, tissue contrast is sometimes unsatisfactory. Finally, radiation exposure is a consideration with multisequence CT imaging. On average, current CT urography technique exposes the patient to approximately 1.5 times the radiation dose of conventional urography. Ongoing studies evaluating reduced exposure, and modifying protocols are under way.

MAGNETIC RESONANCE IMAGING
(FIGURES 6–35 THROUGH 6–42)

Basic Principles

Clinical MRI has its basis in the nuclear properties of the hydrogen atoms in the body. Hydrogen nuclei, when considered as aggregates, sometimes referred to as “protons,” behave like tiny magnets, with net polarity (positive one direction, negative opposite) oriented along an axis at any given point in space. Ordinarily, the axes of the hydrogen nuclei in the body are randomly oriented. However, if the nuclei are placed in a strong magnetic field (like that produced in an MRI scanner), they precess and wobble like a spinning top around the lines of magnetic force.

When hydrogen nuclei in a strong magnetic field are additionally stimulated by short, pulsed radio waves of appropriate frequency, they absorb energy and invert their orientation with respect to the magnetic field. At the termination of radiofrequency pulses, the hydrogen nuclei return at various rates to their original orientation within the magnetic field, emitting energy in the form of radio waves. This phenomenon is called nuclear magnetic resonance (NMR). The emitted weak radio signals from the resonating hydrogen nuclei are received by sophisticated antenna, or coils, and transformed with various computer programs into cross-sectional images.

Different MR signal intensities reflect different hydrogen densities in body tissues, as well as differing physical, cellular, and chemical microenvironments and also flow (fluid) characteristics. The signals emitted from nuclei under MR investigation contain no innate spatial information. Spatial localization is achieved through varying the magnetic field in space (gradients), as emitted frequencies are proportional to magnetic field. Timing of precession (phase) is also controlled and varied to provide spatial localization of emitted signals.

There are biologically important nuclei other than hydrogen that are MR-sensitive, including those of phosphorus, sodium, and potassium, but these occur in lower physiologic concentrations than hydrogen. Imaging of these nuclei for tissue typing and mapping and as biologic tracers (MR spectroscopy) is undergoing intense research and development.

Figure 6–29. CT scans: kidneys. Left: Simple renal cyst. Cyst (arrow) has a CT number close to that of water. 49-year-old man with flank pain. Right: New Hydronephrosis, severe on left and mild on right. Dilated left renal pelvis and delayed left nephrogram (arrow). 40-year-old female with ovarian carcinoma and peritoneal metastases.
Figure 6–30. CT scans: kidneys. **Upper left:** Renal cell carcinoma. The left renal tumor (arrow) shows central necrosis. Note calcification in the arteriosclerotic abdominal aorta. 61-year-old man with previous right nephrectomy for renal carcinoma. **Upper right:** Recurrent renal adenocarcinoma. Massive recurrence in right renal fossa (T), with extensive invasion of posterior soft tissues and destruction of vertebral bodies (arrows). 51-year-old man after right nephrectomy for carcinoma. **Mid left:** Renal angiomyolipomas. Bilateral heterogeneous renal masses. The larger lesions all showed areas of macroscopic fat density (arrows). 35-year-old female with probable lymphangioleiomyomatosis. **Mid right:** Right renal pelvic laceration. Enhanced CT scan through the kidneys showing extravasation of radiopaque material (arrow). Hemorrhage into the psoas and back muscles has enlarged their image (asterisk). 22-year-old man with laceration of the right renal pelvis due to a stab wound. **Lower left:** Large palpable heterogeneous left renal mass, with confirmed hemorrhage with subcapsular extension. Wilms’s tumor in a 9-year-old female with acute onset fever and abdominal pain. **Lower right:** Large retroperitoneal neuroblastoma, encasing vessels.
Clinical Applications

Applications for MR in renal imaging include demonstration of congenital anomalies, diagnosis of renal vein thrombosis, and diagnosis and staging of renal cell carcinoma. MR angiography is useful in evaluating renal transplant vessels, renal vein tumor or thrombosis, and renal artery stenosis.

The use of contrast media in MRI of the kidney has broadened clinical applications. Using bolus injection of gadolinium and rapid sequence imaging, both anatomy and function of the kidney can be assessed. Gadolinium, similar to iodine contrast media, is an extracellular contrast agent primarily excreted by glomerular filtration. Compared to iodinated contrast media, gadolinium has superior renal tolerance in patients with preexisting renal failure. Recently, cases of nephrogenic systemic fibrosis have been reported in patients with renal failure who have received gadolinium. This is being actively investigated. Iodinated contrast agents used in radiography and CT increase attenuation linearly with their concentration. The effect of gadolinium on MR tissue signal intensity is more

Figure 6–31. CT scans: kidneys. Upper left: Transitional cell carcinoma. Delayed image CT urogram demonstrates an irregular nodular filling defect in the right upper collecting system (arrow). Upper right: 42-year-old woman with severe right pyelonephritis. An image through the mid pole of the right kidney reveals an enlarged kidney with marked destruction and striation of the renal parenchyma. Note the multiple low-density fluid collections. Lower left: Exophytic renal cell carcinoma. An image through the mid left kidney shows a mass which may be of renal cortical or collecting system origin. Lower right: A lower image in the same patient reveals that the mass arises from the kidney, as the contrast-containing collecting system is splayed by the mass. The renal vein is displaced anteriorly by the mass, but was free of thrombus.
complex, though in general at lower concentrations gadolinium causes an increase in signal intensity.

The use of gadolinium has extended the application of MRI to the evaluation of renal obstruction (MRU may be used when other studies are inconclusive) and the detection and characterization of renal tumors. Although MRI is capable of imaging blood vessels without contrast media, gadolinium bolus followed by rapid imaging is less susceptible to flow direction and overestimation of stenosis, which can be seen in noncontrast MRA imaging. Gadolinium-enhanced MRA is useful for assessing renal artery stenosis and for evaluating potential renal donors (Figure 6–39).

MRI is used primarily to stage bladder tumors and to differentiate between benign bladder wall hypertrophy and infiltrating malignant neoplasm. There may be a potential advantage for combined endorectal and surface coil MR staging for bladder carcinoma as well. In imaging the prostate gland, MRI is principally used to stage patients with prostate cancer. MR spectroscopy increases specificity and reduces interobserver variability in this setting. MRI of the testis is appropriate when other imaging studies are inconclusive and is applicable to the evaluation of undescended testis, trauma, epididymoorchitis, and tumors.

A modification of the MRI technique, called chemical shift imaging, can detect microscopic amounts of fat within lesions (Figure 6–36, lower images). This technique is commonly used to characterize adrenal masses. Adrenal masses containing fat are either adrenal adenomas or myelolipomas, so the CT or MRI demonstration of fat in

Figure 6–32. 3D computed tomography (CT) and CT angiography (CTA) for renal mass, performed on multidetector CT scanner. Coronal oblique reformed image with volume rendering shows a small enhancing mass (M) in the lower pole of the left kidney. The kidney has 2 renal arteries (arrows). The lower one, which serves the tumor, is the primary artery. 69-year-old male with solitary kidney and indeterminate lesion on prior CT.

Figure 6–33. Helical computed tomography (CT) without oral or intravenous contrast in a 42-year-old man with left flank pain. **Left:** CT image through the kidneys shows enlargement of the left kidney compared with the right, left pelvocaliectasis (P), and a stone in the mid pole of the left kidney (arrow). L = liver, S = spleen. **Right:** CT image through the base of the bladder shows an 8-mm stone (arrow) at the left ureterovesical junction with associated edema involving the left hemitrigone. Posterior to the ureteral stone is a 5-mm phlebolith (open arrow) within a pelvic vein.
Figure 6–34. CT scans: retroperitoneum, bladder, prostate. **Upper left:** Perirenal hematoma. Hematoma (arrow) displaces the left kidney anteriorly. G = gallbladder. 16-year-old boy with acute glomerulonephritis; low-grade fever and left flank pain following left renal biopsy. **Upper right:** Retroperitoneal metastatic seminoma. Huge retroperitoneal mass of metastatic nodes (T) destroying vertebral body (arrow), obliterating outlines of central abdominal and retroperitoneal structures, and displacing kidneys laterally and bowel anteriorly. 46-year-old man with metastatic anaplastic testicular seminoma. **Lower left:** CT scan, transitional cell carcinoma of the urinary bladder with tumor (T) extension into the bladder diverticulum. There is tumor extension into the perivesical fat (arrows). **Lower right:** CT urogram in arterial phase, enhancing 5-mm transitional cell carcinoma of the urinary bladder (arrow).
Figure 6–35. Magnetic resonance imaging (MRI) appearance of the normal kidney. Upper left: T1-weighted conventional spin-echo image showing detailed anatomy of the kidney with differentiation between higher signal intensity cortex (C) and lower signal intensity medulla (M). Left renal vein (arrow), inferior vena cava (I). Upper right: T1-weighted spin-echo image using fat saturation technique. Because the fat signal has been suppressed, the computer automatically adjusts the gray scale of the signal intensity rendering even better contrast between higher signal intensity cortex (C) and lower signal intensity medulla (M). Lower left: Coronal T2-weighted image shows multiple renal and hepatic cysts (*), as well as a partially solid and cystic left renal mass (T) in this adult female with polycystic kidney disease. Lower mid: Noncontrast T1-weighted fat saturation volume acquisition scan, used for dynamic imaging. The left renal mass is subtle on this precontrast scan (*). Lower right: Arterial phase scan obtained following gadolinium-DTPA injection of contrast medium. The left renal mass (*) shows avid enhancement of solid components. The addition of Gadolinium adds significant soft tissue contrast.
Figure 6–36. Magnetic resonance (MR) images

**Upper left and upper right:** Horseshoe kidney. Nonenhanced T1-weighted image (upper left); m, medulla; c, renal cortex. Gadolinium-DTPA-enhanced T1-weighted image (upper right). Following injection of the contrast medium, there is uniform enhancement of the renal cortex (C). The addition of the contrast enhancement shows that the part of the kidney in front of the aorta (A) and the inferior vena cava (I) is functioning renal parenchyma.

**Lower left and lower right:** Chemical shift imaging, adrenal adenoma. On in-phase gradient T1-weighted image (lower left) there is a soft tissue intensity 2.7-cm mass in the left adrenal gland (arrow). Opposed phase T1-weighted image (lower right) shows marked signal loss in the lesion consistent with intracellular lipid; and therefore, a benign adenoma. 30-year-old female with indeterminate adrenal lesion noted on prior computed tomography (CT).
Figure 6–37. Magnetic resonance imaging (MRI) appearance of renal cell carcinoma. The advantages of MRI are multiplanar imaging and the use of contrast media for better tumor characterization. **Upper left:** Sagittal T1-weighted image demonstrating a large renal cell carcinoma (T) arising from the inferior pole of the right kidney (K). Tumor extension in the posterior perirenal space (arrow). Liver (L). **Upper right:** Coronal image of a large renal cell carcinoma (T) replacing almost entire parenchyma of the left kidney (K). Superior displacement of the pancreas (arrows). Liver (L). **Lower left and lower right:** Fat saturation images before and after injection of the contrast medium. Heterogeneous tumor (lower right) in the posterior part of the right kidney shows heterogeneous enhancement following injection of gadolinium. The tumor is extending into the renal vein (arrows) and to the inferior vena cava (I).
Figure 6–38. Pre- and postcontrast computed tomography (CT) and magnetic resonance imaging (MRI) of a renal cell carcinoma adjacent to a cyst. **Upper left:** Precontrast CT image through the left kidney reveals a prominent posterior bulge (long arrow), which proved to be a cyst, and a subtle posterolateral convex contour deformity, which contains tiny calcifications (short arrow). **Upper right:** Postcontrast CT image reveals a nonenhancing cyst posteriorly (long arrow). The posterolateral contour deformity is caused by an enhancing renal cell carcinoma (short arrow) with a central low-density fluid collection. **Lower left:** Precontrast T1-weighted image shows similar intensities for the cyst, tumor, and normal renal parenchyma. **Lower right:** Postcontrast T1-weighted image. The cyst (long arrow) is nonenhancing. The margins of the enhancing renal cell carcinoma (short arrows) are seen. The central fluid collection does not enhance.
an adrenal lesion characterizes it as a benign lesion, even in the oncologic patient.

MR urography utilizes the sensitivity of MR imaging to demonstrate fluid (urine), producing urogram-like pictures without the need for contrast media. This technique is sensitive in the detection of ureterohydronephrosis and is particularly useful in patients in whom contrast material is contraindicated, such as patients with prior contrast reactions or renal failure.

Advantages & Disadvantages

Advantages of MRI include direct imaging in any plane desired (though transverse, sagittal, and coronal are most standard), choice of large or small field of view, excellent soft-tissue contrast, imaging without exposure to ionizing radiation, and (as compared to ultrasound) less operator dependence. MRI can image blood vessels and the urinary tract without contrast material. MR scanning, however, is not without drawbacks. The scanning time is relatively slow and as a result image clarity is often inferior compared with CT. Absolute contraindications to MRI include the presence of (1) intracranial aneurysm clips, unless the referring physician is certain that the clip is made of a nonferromagnetic material (such as titanium); (2) intraorbital metal fragments; and (3) any electrically, magnetically, or mechanically activated implants (including cardiac pacemakers, biostimulators, neurostimulators, cochlear implants, and hearing aids). Relative contraindications such as pregnancy should always be viewed in the light of risk versus benefit of the examination.

COMPARISON OF IMAGING METHODS (FIGURES 6–43 THROUGH 6–46)

As new imaging methods have been developed, changes have occurred in patterns of use for each type of imaging. For example, increased familiarity with and confidence in sonography and CT scanning have resulted in a decrease in the use of some long-established conventional uroradiologic studies such as EU.

Several factors are involved in these changes: (1) the increased effectiveness of newer imaging methods over older ones for some aspects of urodiagnosis; (2) the availability of equipment, trained technical personnel to operate it, and physicians to interpret the results; (3) increased awareness of the hazards of ionizing radiation; and (4) the desire to use noninvasive examinations if possible.

Because so many different types of imaging are available, each with different costs, risks, and areas of effectiveness, it may be difficult for the clinician to decide which method will yield the most information with the least cost and risk. A particular study may be critical in one diagnostic situation but useless in another. For example, sonography is an excellent noninvasive, relatively inexpensive method for differentiating simple cysts from other mass lesions in the kidney but is much less effective in imaging the adrenal glands and ureters than is CT scanning. Sonography also relies considerably on the skill of the operator. CT scanning produces excellent images and is currently the preferred imaging method for the examination of the retroperitoneum. MRI rivals CT scanning in imaging capability for some structures, for example, the kidney, but has surpassed CT in imaging the pelvis. With advances in equipment and techniques the use of MRI in urology will likely grow.
Figure 6–40. Magnetic resonance imaging (MRI) examination of the pelvis. The upper images are examples of the ability of MRI to evaluate bladder neoplasms. **Upper left:** Axial T2-weighted image. A large anterior bladder tumor (T) shows associated perivesicular fat stranding (*), raising the suspicion for extension through the muscularis. A prominent vessel (arrow) in the left obturator region exited through the obturator foramen on more inferior images. **Upper right:** T2-weighted sagittal image. The tumor (*) extends into the urachal remnant (arrow). **Lower left and lower right:** Prostate cancer. Axial T2-weighted image (**lower left**) shows dominant tumor at the right mid gland involving peripheral and transition zones with gross posterior extracapsular extension (arrow). Superimposed spectroscopy confirms depleted citrate (*) compared to the adjacent choline and creatine peaks in the region of tumor (**lower right**).
Figure 6–41. Magnetic resonance imaging (MRI) of seminal vesicles, ductus deferens, and ejaculatory ducts. **Upper left:** T2-weighted image, normal seminal vesicles, and ductus deferens, and ejaculatory ducts. The ampullae of the ductus deferens (long arrows) are normally of high signal intensity on T2-weighted images and are immediately medial to the seminal vesicles (short arrows). The seminal vesicles are also of high signal intensity and are draped over the prostate gland. **Upper right:** Seminal vesicle and ductus deferens calculi. Coronal T2-weighted images show low-signal calculi within the proximal ductus deferens and medial aspect of the seminal vesicle on the left side (arrows). The patient had a history of prostatitis, prostatic pain, and hemorrhagic ejaculate. **Lower left:** Axial T2-weighted image through the prostate. The peripheral zone is of normal high signal intensity. The normal ejaculatory ducts (arrows) are identified as 2 small foci of high signal intensity within the lower signal central zone. **Lower right:** Axial T2-weighted image through the prostate reveals a low-signal-intensity calculus (arrow) within the right ejaculatory duct (same patient as in upper right image).
Figure 6–42. Magnetic resonance imaging (MRI) appearance of scrotal contents. Upper left: Normal testis, T2-weighted image. The testicular tissue is of homogeneous high signal intensity. The tunica albuginea (arrows) demonstrates low signal intensity as does the mediastinum testis (open arrowhead). A small amount of fluid—hydrocele (H). Epididymis (E) is of low signal intensity. Upper right: Hydrocele of the right scrotum (T2-weighted image). Hydrocele (H) demonstrates high signal intensity. Testis (T). Tunica albuginea (curved black arrows). Varicocele (*). Lower left and lower right: Images of a testicular tumor. On the proton density image (lower left), the signal intensity from both testicles is similar. On the T2-weighted image (lower right), testicular tumor (*) demonstrates lower signal intensity as compared with the higher signal intensity of the normal testicular tissue (T).
Figure 6–43. Comparison of imaging methods in the evaluation of renal cell carcinoma. **Upper left:** Computed tomography (CT) scan showing a renal cyst (Cy) in the right kidney. There is bulging (arrow) in the contour of the left kidney (K), but it is difficult to discern if the lesion represents a neoplasm. **Upper right:** Angiogram showing small vascular lesions in the inferior pole of the left kidney (arrows). **Lower left and lower right:** MRI scans. **Lower left:** T1-weighted noncontrast scan. **Lower right:** T1-weighted postcontrast scan. The renal cyst (Cy) in the right kidney does not show any enhancement. The lesion (arrow) in the left kidney (K) shows marked enhancement, indicating that it is solid in nature. In this example, the contrast-enhanced MRI is superior to CT in the detection and characterization of the left renal mass.
Figure 6-44. Comparison of imaging methods: metastatic extra-adrenal familial pheochromocytoma. 10-year-old boy with hypertension and seizures precipitated by abdominal palpation. Family history of multiple extra-adrenal pheochromocytomas in the mother. **Upper left:** Excretory urogram. The right ureter is dilated and elevated (curved arrow), with the right posterior portion of the bladder displaced toward the left (straight arrow). The urographic diagnosis is possible extra-adrenal paravesical pheochromocytoma. **Upper right:** Right femoral arteriogram. Tumor stain (arrow) in right paravesical location. The angiographic diagnosis is extra-adrenal paravesical pheochromocytoma. **Lower left:** Computed tomography (CT) scan. Transverse tomogram through bladder (B) shows the tumor (T) indenting the bladder (arrows). R = rectum. **Lower right:** CT scan. Transverse tomogram through bladder. Recurrence of symptoms following removal of the right paravesical pheochromocytoma prompted another CT study, which shows recurrent tumor (arrow) in the bladder wall. Each imaging study complemented or supplemented the previous one. None, however, diagnosed the small liver metastases discovered at surgery.
**Figure 6–45.** Comparison of imaging methods. Upper left is an example of a nonenhanced computed tomography (CT), and upper right is an ultrasound study in the demonstration of renal calcifications. Fine calcifications in the medullary region indicate medullary nephrocalcinosis. They are of high density on CT (arrows) and are shown as echogenic foci (arrows) on an ultrasound scan. Lower left and lower right: Example of images of a transitional cell carcinoma and the calcified renal cyst. Retrograde urogram (lower left) shows filling defects due to tumor in the renal pelvis (black arrow) at ureteropelvic junction, and also seen are the calcifications in a lower pole mass (white arrows). Note that the infundibulum and calices of the lower pole failed to opacify in this 45-year-old woman with hematuria. On the CT scan (lower right) the cystic nature of the calcified renal mass (curved arrow) is well demonstrated and the CT scan shows better the extent of the tumor (T), which involves the entire lower pole of the kidney and extends into the dilated renal pelvis.
Figure 6–46. Comparison of imaging methods. **Upper left and upper right:** Renal sarcoma with vascular invasion. Axial CT (upper left) reveals a left renal mass (arrow). The visualized left renal vein is invaded and expanded with thrombus (*). On this sagittal ultrasound Doppler image (upper right) in the same patient, the thrombus extends across the midline from the left renal vein into the inferior vena cava (IVC) (*). **Lower left and lower right:** Demonstration of an arterial venous malformation to the seminal vesicles. On the CT scan (lower left), the right seminal vesicle (SV) is enlarged, but the nature of the enlargement (arrow) is difficult to discern. On a magnetic resonance imaging (MRI) scan (lower right), the enlarged vessels (arrows) are demonstrated as the flowing blood within the vessel lacks signal intensity in contrast to the medium signal intensity of the seminal vesicles (SV). Urinary bladder (B).
The patient and the clinician both benefit from careful consultation with the radiologist to ensure that the methods of imaging chosen are of value in diagnosis and treatment planning and do not duplicate or merely confirm established findings, with loss of time and additional expense.

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Interventional uroradiologic procedures can be divided into 2 major groups: vascular and percutaneous nonvascular. Percutaneous nonvascular interventional procedures are discussed elsewhere. The intravascular route is used, as the therapy of choice, for the embolization of arteriovenous fistulas (AVFs) or malformations, and for bleeding sites. Transcatheter embolization is used for tumor embolization, for the ablation of renal function, for the treatment of testicular vein and ovarian vein varices, and for the treatment of high-flow priapism. Balloon angioplasty and stenting of stenotic renal arteries are frequently performed endovascular techniques for the treatment of ischemic nephropathy and secondary hypertension. Renal artery aneurysms may also be treated using catheter-directed techniques such as stent grafting and selective embolization. Occasionally, fibrinolytic agents are delivered via an endovascular catheter to thrombosed renal arteries. Mechanical devices are also available for endovascular treatment of thrombosed renal vessels. This chapter will review these intravascular interventions.

**TRANSCATHETER EMBOLIZATION**

**Renal AVFs & Malformations**

Transcatheter embolization is the treatment of choice for renal AVFs, which may be congenital, spontaneous, or acquired. Iatrogenic AVFs are the type most commonly treated by transcatheter embolization. These occur as a complication of such procedures as percutaneous renal biopsy (Maleux et al, 2003), nephrostomy placement, and pyelolithotomy. Trauma or surgery can also result in AVFs. AVF occurring in the transplant kidney is successfully managed by embolization. The classical angiographic finding of spontaneous or acquired AVF is a feeding artery with an early draining vein. Ancillary findings include pseudoaneurysm and extravasation of contrast material. Congenital AVMs (AV malformations) consist of a group of multiple coiled communicating vessels that may be associated with enlarged feeding arteries and draining veins.

The modes of clinical presentation include hematuria; retroperitoneal or intraperitoneal hemorrhage; and congestive heart failure, cardiomegaly, or both. Hypertension can occur as a consequence of ischemia secondary to venous shunting of blood away from the affected area. A bruit may be heard on physical examination. Duplex Doppler ultrasound is the most useful diagnostic study, performed before angiographic intervention.

Successful intervention requires the angiographic identification, selective catheterization, and embolization of the feeding artery (Figure 7–1A, B). Using a transfemoral approach, an abdominal aortogram is performed to identify the arterial supply to the bleeding kidney. In the case of a renal transplant, an initial pelvic angiogram is performed in a steep oblique projection. The artery supplying the bleeding site is selectively catheterized. A 3F coaxial microcatheter is then used for subselective catheterization and embolization of the feeding artery. The use of a microcatheter allows accurate placement of the embolic material. Microcoils are used for the occlusion of iatrogenic AVFs because they can be deployed very precisely, thereby minimizing the loss of renal parenchyma due to resultant ischemia (Figure 7–2A–C). The procedure is usually performed without significant complications. Very rarely, inadvertent nontarget embolization or thrombosis of the renal artery can occur.

**Bleeding Sites**

Transcatheter embolization plays a key role in the management of hemorrhage in the urinary tract originating in the kidney, ureter, bladder, and pelvis (Sofocleous et al, 2005). Acute life-threatening hemorrhage can occur as a consequence of trauma, instrumentation, and tumors. Chronic intractable hemorrhage is associated with radiation cystitis, tumors, prostaticctomy, and infiltrative disorders. Hemodynamically stable patients undergo a noninvasive diagnostic study such as contrast-enhanced computed tomography (CT), before embolization.

Pelvic fractures resulting in life-threatening hemorrhage require embolization for control if resuscitation and external pelvic fixation have been ineffective. Embolization has been shown to be very effective at arresting hemorrhage. Using a transfemoral approach, the practitioner performs a nonselective pelvic arteriogram before selective catheteriza-
Figure 7–1. Transcatheter embolization of a large arteriovenous malformation (AVM) in a 64-year-old woman with hematuria. **A:** A conventional film midstream aortogram. An enlarged left renal artery is seen. There is a large serpiginous AVM arising from the lower pole renal artery branch with aneurysmal dilatation of the draining renal vein. **B:** Selective left renal digital subtraction arteriogram (DSA) after coil embolization shows cessation of flow in the AVM. Coils have been placed in the terminal portion of the lower pole artery and within the AVM. Embolization resulted in resolution of the hematuria.

Figure 7–2. Transcatheter embolization of a postbiopsy arteriovenous fistula (AVF) in a 14-year-old boy with hypertension and prior renal transplantation. A bruit was heard on examination. **A:** Pelvic arteriogram (DSA) shows an AVF arising from a lower pole branch artery. There is aneurysmal dilatation of the draining vein. **B:** DSA shows that the renal artery has been selectively catheterized, and a 3F coaxial catheter is positioned peripherally within the lower pole branch artery supplying the AVF. Several microcoils have been placed (arrow) and no flow is seen in the AVF. **C:** Completion DSA of the main renal artery shows absent flow in the AVF (arrow) with minimal devascularization of lower pole parenchyma (curved arrow).
tion and embolization of the hypogastric arteries. Because of contralateral crossover blood supply, pelvic lesions are treated by bilateral embolization. Gelfoam pledgets are frequently used. They can be deployed rapidly, are immediately effective if the patient has a normal coagulation profile, and produce temporary vascular occlusion. Gelfoam sponge is easily cut into pieces appropriate to the caliber of the vessel to be embolized. Coils may be used with, or instead of, Gelfoam. However, if used, they may hamper future access to the hypogastric artery in the event of rebleeding. Small embolic materials such as Gelfoam powder or Ivalon particles are not used to treat hemorrhage from pelvic trauma. They produce very peripheral occlusion of small vessels, thereby risking ischemia of nontarget organs. Complications specific to pelvic embolization are extremely uncommon. Nontarget embolization is rare.

**Tumors**

**A. RENAL CELL CARCINOMA**

Primary renal cell carcinoma (RCC) is treated by surgical excision. In some cases, preoperative occlusive embolization of the renal artery is used as an adjunct to surgery. Embolization reduces intraoperative hemorrhage and allows immediate ligation of the renal vein. It is used in patients with very large tumors and also in tumors supplied by many parasitized vessels. Embolization accentuates cleavage planes and therefore facilitates nephrectomy. The optimal time delay between embolization and surgery is probably 1 day. Embolization may also favorably impact patient survival (Zielinski, Szmigielski, and Petrovich, 2000). A new application is to use selective embolization of a renal tumor as an adjunctive measure prior to radiofrequency ablation of the tumor (Yamakado et al, 2006).

Palliation of nonresectable disease that causes pain and hematuria can be achieved by transcatheter embolization (Munro et al, 2003). Patients with bilateral RCC, and those with RCC in a single kidney, can undergo subselective embolization as an alternative to surgery, thereby sparing normal parenchyma. Embolization of RCC metastases to bone is performed before surgical resection to decrease intraoperative blood loss (Charzioannou et al, 2000). CT or magnetic resonance imaging may be used for tumor evaluation before and after intervention.

A transfemoral aortogram and selective arteriogram are performed to determine the blood supply to the kidney and tumor. An occlusive balloon catheter may be placed within the vessel and inflated before embolization to prevent reflux of embolic material and inadvertent nontarget embolization. However, many physicians use a simple selective catheter. Gelfoam pledgets are used for preoperative embolization (Figure 7–3A, B). Coils are not used because they can be dislodged during surgery when the kidney is manipulated. Absolute ethanol is the preferred embolic material for ablative palliative embolization of nonresectable tumor. Bone metastases are embolized by positioning a microcatheter in the vessel(s) supplying the tumor and injecting particles of polyvinyl alcohol (PVA) or other embolic agents such as embospheres until maximal obliteration of the angiographic tumor stain is achieved.

Tumor embolization is a safe procedure. Complications such as puncture-site hematomas and inadvertent nontarget embolization occur in <2% of patients. Almost all patients, however, experience postembolization syndrome (PES). PES consists of severe pain, nausea and vomiting, fever, and leukocytosis. It is probably caused by tissue necrosis that results from successful embolization. Transient ileus, transient hypertension, sepsis, and reversible renal failure have also been described. PES occurs within a few hours of the procedure and may last for several days. Its occurrence should not delay surgical intervention. Tissue swelling and tissue gas formation are seen on imaging studies. The severity of PES is related to the quantity of infarcted tissue. Analgesics and antibiotics are used for treatment. Administration of steroids and antibiotics before embolization may reduce the severity of PES.

**B. ANGIOMYOLIPOMA**

Selective embolization has proved to be an effective method of controlling hemorrhage from benign renal lesions while preserving normal parenchyma (Kothary et al, 2005). This technique has been used in the treatment of active bleeding from angiomylipoma and for the elective prevention of hemorrhage, especially if the tumors are multiple or bilateral, as in patients with tuberous sclerosis. Current guidelines suggest selectively embolizing any tumors that are larger than 4 cm in diameter. The procedure is effective in decreasing tumor size and in preventing or treating hemorrhage in 85–90% of patients. CT can clearly identify the fat component of the tumor and is therefore used for diagnosis before intervention and for follow-up.

The technique for embolization is similar to that described for RCC. Through a transfemoral approach, angiography is used to define the arterial supply to the kidney and tumor. The feeding vessels are then selectively catheterized using a coaxial microcatheter. The tumor volume is estimated, and an equal amount of absolute ethanol admixed with iodized oil is carefully injected until the flow ceases. Ethanol is easy to handle, is inexpensive, and produces permanent occlusion of the vascular bed. Iodized oil is radiopaque and therefore is useful for visualizing the flow of the embolic material during the embolization. It remains within the tumor, thereby allowing measurement of tumor response on follow-up CT. An occlusion balloon is reserved for more proximal embolization in a main renal artery when the risk of nontarget embolization is greatest.

The reported complications are similar to those seen after RCC embolization. Recurrence of tumor hemorrhage occurs in approximately 10–15% of patients and is treated by repeat embolization. A short-term tapered dose of pred-
nisone may reduce PES after the procedure (Bissler et al, 2002).

Ablation of Renal Function

Total renal infarction using transcatheter embolization may be indicated in certain circumstances: namely, abolition of urine production to assist in healing or palliation of terminal patients with urinary fistulas; prevention of excessive proteinuria; management of uncontrollable hypertension; rarely for benign obstructive uropathy in patients who are poor surgical candidates (De Baere et al, 2000; Toussi, McConnell, and Srinivasan, 2001); and for ablation of failed renal allografts causing the graft intolerance syndrome (Delgado et al, 2005). For patients with end-stage renal disease and intractable hypertension or nephrotic syndrome, the advantages of ablation must be weighed against the loss of production of vitamin D₃ and erythropoietic factors and, occasionally, the loss of the ability to eliminate some water. Total renal ablation must be achieved so that perfusion of surviving parenchyma via pericapsular branches cannot occur. The embolic agent must perfuse the entire renal substance but be safe if it passes through the kidney into the venous circulation. The technique is applicable in adults and children and in both native and allograft kidneys. Renal ablation appears to be a safe procedure, which is successful in most patients (De Baere et al, 2000).

Initially, a midstream aortogram is performed to identify the arterial supply to the kidneys. This is followed by renal artery catheterization and selective arteriography. An occlusion balloon catheter is placed in the vessel and contrast medium is injected to measure the volume of embolic agent needed to fill the vascular tree of the kidney. Absolute ethanol is the embolic agent of choice for the reasons mentioned above. An equal volume is carefully injected with the balloon inflated to prevent reflux to nontarget regions. The contents are then aspirated and an angiogram is performed to assess flow. The procedure is repeated as many times as necessary until complete cessation of blood flow is achieved.

PES of several days’ duration occurs in all patients and is managed with antibiotics and analgesics. Inadvertent embolization of the adrenal artery and other visceral vessels is a rare but serious complication that can be avoided by using the occlusion balloon technique. Systemic toxicity

Figure 7–3. Preoperative embolization of a large right renal cell carcinoma (RCC) in a 28-year-old woman. A: Digital subtraction arteriogram (DSA) of the right renal artery shows a large hypervascular mass involving most of the right kidney. There is tumor thrombus within the renal vein. B: Completion DSA of the right renal artery after Gelfoam embolization shows complete cessation of flow within the tumor and kidney. Contrast is seen within the main renal artery only.
from ethanol injection is not a clinical problem. Morbidity and mortality from the procedure is less than from surgical nephrectomy.

**Embolization of Primary Varicocele**

The subject of varicocele and male infertility is discussed in detail in Chapter 42. The basis for intervention is the presence of a varicocele on physical examination; most varicoceles are left-sided (Beddy et al, 2005). Color Doppler sonography is usually performed routinely before intervention. Several reports have suggested a positive association between improvement of seminal parameters and pregnancy. Surgery and percutaneous transvenous embolization appear to be equally effective. However, advantages of the percutaneous route include greater patient comfort, ease of bilateral treatment, and reduced recovery time.

The procedure is performed with conscious intravenous sedation and local anesthesia. The transjugular venous approach is preferred by many physicians, although the procedure can be performed transfemorally. Direct ultrasound guidance is used to access the jugular vein; a catheter is then guided fluoroscopically into the left gonadal vein. The left gonadal vein usually drains into the left renal vein, whereas the right gonadal vein usually drains directly into the inferior vena cava. The patient is placed in reverse Trendelenburg position and left gonadal venography is performed. A positive venogram result demonstrates incompetence with filling of collateral vessels. Embolization is achieved by placing coils within the gonadal vein, commencing at the region of the inguinal ligament, and continuing cranially toward the renal vein, until the gonadal vein and collateral vessels have been occluded. If a right-sided varicocele is also present, the right gonadal vein is then interrogated and embolized.

The recurrence rate of varicocele after embolization is approximately 4%. Minor complications include contrast extravasation from perforation of the vein, nontarget embolization, venospasm, and hematoma. Rarely, cardiac arrhythmia and contrast allergy can occur.

**Embolization of Ovarian Vein Varices (Pelvic Congestion Syndrome)**

Pelvic congestion syndrome is a recognized cause of chronic pelvic pain in women that has been associated with lumbo-ovarian vein varices. Dyspareunia, when present, may be a poor prognostic indicator. Symptoms are often worse with fatigue, before menstruation, and in the upright position. Occasionally, vulval and thigh varicosities are present on examination. The cause is probably multifactorial. The diagnosis of varicose pelvic veins is confirmed by duplex Doppler ultrasound that is also used for postintervention follow-up. CT venography or magnetic resonance venography are alternative noninvasive diagnostic tests.

Transvenous embolization has produced durable relief of symptoms in most patients (70%). Most responses occur within 2 weeks of treatment; however, continued improvement has been reported up to 12 months after the procedure (Kim et al, 2006). Using either a transfemoral or a transjugular approach, the ovarian veins are individually catheterized, and embolic agents such as coils or synthetic glue are deployed in the vessel at the level of the pelvic inlet (Figure 7–4A, B). Occasionally, embolization of internal iliac vein tributaries may be necessary. Reported complications are similar to those for varicocele embolization. The procedure does not appear to have a deleterious effect on fertility or the menstrual cycle.

**Treatment of High-Flow Priapism**

High-flow priapism is a fairly rare condition resulting from increased arterial flow into the lacunar spaces of the cavernous tissue. The “arteriocavernous fistula” most often results from penile or perineal trauma. In some cases the cause is unknown. Color Doppler sonography demonstrates the abnormality.

Transcatheter embolization is a minimally invasive, successful treatment for this condition (O’Sullivan et al, 2006). A nonselective transfemoral pelvic angiogram is performed to demonstrate the fistula, which originates from the pudendal artery. This procedure is followed by superselective catheterization of the injured artery using a microcatheter. The fistula is then closed by accurately deploying microcoils (Figure 7–5A, B). This technique preserves blood flow to the penis, thereby allowing normal erectile function in most cases. The use of microcoils prevents possible ischemic injury to the perineum. In the event of recurrence after embolization, repeat embolization is successful in nearly all patients. Surgery should be reserved for patients in whom embolization fails.

**RENAL ARTERY ANGIOPLASTY & STENTING**

Ischemic nephropathy due to atherosclerotic vascular disease and renal artery stenosis is a leading cause of progressive renal failure. Renal artery stenosis is the most common cause of secondary hypertension. Surgical revascularization is an established method of treatment, with reported success rates of >70%. In recent years, percutaneous transluminal angioplasty (PTA) and stent placement have become a viable alternative to surgery although there is controversy over the relative benefits of revascularization compared to pharmacologic treatment (White, 2006; Uder and Humke, 2005). Percutaneous transluminal angioplasty is the treatment of choice in the management of fibromuscular dysplasia occurring in a subset of hypertensive patients. The technique involves the use of an inflatable balloon catheter that is positioned endoluminally across the stenosis and then inflated (angioplasty).
Several diagnostic imaging modalities are used for patient selection and for postprocedure follow-up, including captopril radioisotope assay, duplex Doppler ultrasound, CT angiography, magnetic resonance angiography, and arteriography. The advantages and disadvantages of these techniques are beyond the scope of this chapter, but noncatheter angiography by CT or MR is replacing catheter angiography for diagnostic purposes.

Renal artery stenosis is described as ostial, nonostial, or branch vessel stenosis. An ostial lesion is located within 3 mm of the aortic lumen and is typical of atherosclerotic vascular disease. In fibromuscular dysplasia, nonostial and branch vessel lesions are more typically seen. The initial technical success rate of PTA varies. It may be as low as 35% for some atherosclerotic vascular disease ostial lesions, but in most series the overall rate approximates 95–100%. PTA results in stabilization or improvement of renal function in most patients with ischemic nephropathy and also leads to durable improvement or cure in most of those with hypertension. The best results after PTA have been achieved in hypertensive patients with fibromuscular dysplasia, in whom improvement or cure is achieved in approximately 90%.

Stent placement is usually the method of choice for endoluminal recanalization. Previously, stenting had been reserved for immediate failure or complication of PTA, such as elastic recoil or flow limiting intimal dissection, for residual stenoses of >30%, for a >20 mm Hg peak systolic pressure gradient after PTA, for early restenosis, and for ostial lesions that are difficult to treat by PTA alone. Renal artery stenting results in stabilization (38% of patients) or improvement (30% of patients) of renal function and in durable improvement (49%) or cure (20%) of hypertension (Leertouwer et al, 2000). PTA and stenting have also been used successfully to treat renal allograft artery stenosis. Primary patency rates vary after stenting. The average restenosis rate is approximately 17% after 6–12 months’ follow-up. However, the rate increased to 20–30% with longer follow-up. Secondary patency may be achieved in >90% of patients.

Figure 7–4. Embolization of ovarian vein varices in a 30-year-old multiparous woman with pelvic congestion syndrome. She complained of increasing pelvic pain and dyspareunia. On examination there were prominent vulval varicosities. A: Transjugular right ovarian venography demonstrates multiple large ovarian vein varices. B: Radiograph taken after coil embolization of both ovarian veins and tributaries of the internal iliac veins. The varices have been occluded. The patient’s symptoms resolved after the procedure.
Usually, the transfemoral approach is used, although a transaxillary approach may be required. Initially, a midstream aortogram is performed to identify the renal arteries, followed by a selective injection to evaluate the morphology and location of the stenosis, the diameter of the vessel, and the percentage of stenosis. In the presence of altered renal function with elevated creatinine levels, alternatives to iodinated contrast agents include gadolinium and carbon dioxide gas. Indicators of a significant stenosis include a reduction of cross-sectional diameter of at least 50%, poststenotic dilatation, collateral vessels to the affected kidney, and a transstenotic systolic pressure gradient of >20 mm Hg across the lesion. Before intervention, an oral antiplatelet agent such as clopidogrel is administered. The patient is heparinized and a vasodilating agent (eg, nitroglycerine) is given via the arterial catheter. Initially, the lesion is crossed with a guidewire. If a high-grade stenosis is present, predilatation with a small balloon may be necessary before definitive angioplasty or stent placement. An outer guiding catheter or sheath is used to facilitate contrast injection during the procedure and to improve catheter stability. Continuous fluoroscopic guidance and “vascular road-mapping” are also used to ensure precision. Frequently, a small tear in the vessel intima is seen after PTA. We prefer to use a balloon expandable stent because it can be very accurately deployed. The minimal recommended vessel diameter for stenting is 5 mm. A 10- to 20-mm-long stent is used, and approximately 1–2 mm should protrude into the aortic lumen when ostial lesions are treated (Figure 7–6A, B).

Success of the procedure is defined by <30% residual stenosis and by the resolution of a significant transstenotic pressure gradient. We frequently use a percutaneous closure device to achieve hemostasis. Antiplatelet medication is continued for 6 weeks after the procedure, and the patient is carefully followed up by clinical evaluation and repeat imaging.

The reported complication rate after PTA and stenting varies considerably but ranges from 3% to 10% in experienced operators. Complications include puncture-site hematoma; femoral artery pseudoaneurysm; contrast nephropathy; cholesterol embolization; stent malpositioning; and injury to the renal artery such as dissection, thrombosis, and rupture. Rarely, procedure-related deaths have occurred. The use of endovascular distal protection devices when stenting the carotid arteries may provide benefits for renal artery stenting. These devices are usually small filters that trap micro emboli that may be dislodged during stent placement. They limit end-organ embolization (Hagspiel et al, 2005).

**OTHER ENDOVASCULAR PROCEDURES**

**Renal Artery Aneurysms**

Renal artery aneurysms are rare and are usually not symptomatic. They carry the risk of rupture, with associated life-threatening hemorrhage. Occasionally, they are associated with renovascular hypertension. Distal embolization may occur, leading to parenchymal infarction. The indica-
tions for treatment of renal artery aneurysms include diameter >2.5 cm, interval enlargement, renovascular hypertension, pain, hematuria, intrarenal thromboemboli, and lesions in women of childbearing age. The diagnosis is made by color Doppler sonography, CT, or magnetic resonance imaging.

In the high-risk surgical patient, endovascular techniques may be well-suited to aneurysm repair, by excluding the aneurysm while preserving flow to the kidney. Depending on the location of the aneurysm, its relationship to branch vessels, and the presence or absence of a “neck,” either a stent graft or transcatheter embolization are possible treatment options (Horowitz et al, 2005; Eskandari and Resnick, 2005; Saltzberg et al, 2005). A stent graft is a metallic stent that is covered by surgical graft material. Its deployment across the aneurysm results in internal reconstruction of the vessel, with exclusion of blood flow to the aneurysm due to the impervious graft material. The cavity of the aneurysm may then thrombose.

Selective embolization of an aneurysm that is supplied by a branch vessel is performed by occluding blood flow proximally using coils. This approach results in distal parenchymal infarction and therefore can only be safely performed if there is sufficient renal functional reserve. If there is an identifiable neck separating the aneurysm from the native vessel, coils may be used to pack the aneurysm cavity, thereby resulting in thrombosis of the aneurysm while at the same time preserving distal flow. The risks of the procedure are similar to those reported above. Long-term results of endovascular repair are not yet available.

**CATHETER-DIRECTED FIBRINOLYSIS**

This procedure has been used extensively in the peripheral vasculature but with only limited success in the management of native renal artery or aortorenal bypass graft thrombosis. Small series or individual case reports have suggested a possible role in the treatment of recent renal artery occlusion before PTA and for the treatment of acute renal artery thromboembolic disease (Nakayama et al, 2006).

There is a variety of mechanical devices to remove clot in addition to or as an alternative to pharmacologic thrombolysis (Siablis et al, 2005). The diagnosis is made by non-invasive imaging such as duplex Doppler ultrasound, magnetic resonance imaging, or CT angiography and is then confirmed by angiography, at which time fibrinolytic therapy is begun. A diagnostic arteriogram is performed from a transfemoral approach, after which an infusion catheter or wire is embedded within the thrombosed segment. Tissue plasminogen activator (t-PA) is currently the most frequently used fibrinolytic agent in the United States. Several infusion protocols exist. Our preferred technique is continuous infusion of 1–2 mg t-PA/hour. Prophylactic antibiotics are administered. The patient is monitored for puncture-site and systemic bleeding in the intensive care or step-down unit throughout the infusion therapy. A repeat arteriogram is performed 12–24 hour after the initiation of therapy. When recanalization has been achieved, an underlying stenotic lesion is usually identified, at which time PTA or stenting is performed.

Complications include puncture-site and systemic hemorrhage and infection. Bleeding may be severe enough...
to require transfusion or to discontinue the infusion. The incidence of complications is related to the duration of therapy and to the dose administered.

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In contrast to retrograde instrumentation such as ureterorenoscopy, which invades the urinary tract via the natural route of the urethra under endoscopic guidance, techniques of antegrade instrumentation involve access via a percutaneous puncture. This approach must respect the intrarenal anatomy just as in open surgical nephrotomy, and imaging techniques are essential to guide the procedure.

First, and most important, a puncture route must be established that will provide straightforward access to the target and safe, bloodless instrumentation. Visualization of the puncture needle and target and precise guidance to the target require the use of imaging techniques such as ultrasound, fluoroscopy, and, in selected cases, computed tomography (CT).

Contraindications to percutaneous kidney puncture are blood clotting anomalies due to coagulopathies or pharmacologic anticoagulation. Preparation and draping of the surgical field are required as for open surgery, and the same standards of asepsis must be followed. Local anesthesia only is sufficient for puncture of the kidney and small-bore tract dilation (6–12F), for antegrade insertion of a ureteral stent or nephrostomy catheter. Lidocaine hydrochloride 1% USP, 10 mL, can be given for infiltration of the skin and tissues along the intended route of puncture down to the renal capsule. During dilation of the tract, administration of a local anesthetic in lubricant (eg, lidocaine hydrochloride jelly 2%) serves the dual purpose of anesthetization and lubrication. Dilation of nephrostomy tracts up to 30F and extraction of small renal stones can be done under local anesthesia.

Percutaneous nephrolithotomy (PNL) is still indicated for treatment of staghorn calculi and stones in caliceal diverticula, but the extent of intrarenal instrumentation for stone disintegration and extraction usually requires epidural or general anesthesia. Because puncture, tract dilation, and stone disintegration and removal are preferably performed as a one-stage procedure, the use of local anesthesia in PNL is limited.

**IMAGING & PUNCTURE TECHNIQUES**

Percutaneous puncture of the renal collecting system may be performed for diagnostic procedures (eg, antegrade pyelography, pressure/perfusion studies) or to establish access for therapeutic interventions (Table 8–1).

Both ultrasonic scanning and fluoroscopy provide visualization and guidance for a safe, accurate percutaneous puncture, but ultrasound has definite advantages:

1. No intravenous or retrograde administration of contrast dye
2. No radiation exposure
3. Continuous real-time control of puncture
4. Imaging of radiolucent, non-contrast-enhancing renal and extrarenal structures (eg, renal cyst, retroperitoneal tumor) for puncture
5. Imaging of all tissues along an intended nephrostomy tract (eg, bowel, lung)
6. Imaging in numerous planes simply by shifting, tilting, and rotating the scanning head
7. Three-dimensional information during puncture

Once the puncture needle has entered the renal collecting system, fluoroscopy is required for control and guidance of subsequent steps (eg, guidewire insertion, tract dilation, catheter insertion). In selected cases, insertion and placement of a nephrostomy catheter in a dilated renal system may be possible with ultrasonic control only. Fluoroscopy provides a two-dimensional image with complete integration of all information from the third (anterior-posterior) dimension, so that the entire length of radiopaque catheter, wires, and so on can be visualized.

For percutaneous puncture of the renal collecting system, the patient should be placed on the fluoroscopy table in the prone position. Radiolucent bolsters may be placed under the abdomen to correct for lumbar lordosis and to support the kidney. A standard puncture site is in the posterior axillary line midway between the 12th rib and the iliac crest; this site ensures that later the patient does not lie on the nephrostomy catheter in the supine position. Morbidly obese patients can be placed in a lateral decubitus position in order to minimize respiratory distress. Ultrasonic scanning is performed below the 12th rib to obtain a median longitudinal scan through the kidney. For optimal coupling of the ultrasonic beam to the skin, sterile gel (eg,
K-Y jelly) is applied to the skin at the scanning site. In the frontal view of an intravenous pyelogram, the long axis of the kidney usually follows the psoas muscle, forming about a 30° angle with the midline (Figure 8–1A). In the transverse view of a CT scan, the transverse axis of the kidney forms about a 45° angle with both a horizontal and a sagittal line (Figure 8–1B). The position and direction of the transducer should be oriented roughly to the following marks: below the 12th rib (if possible), cranial to the puncture site, with a 30° caudal-lateral rotation, and with a 45° lateral tilt of the scanning head.

Factors that may influence the choice of scanning technique and puncture site include patient size; position and rotation of the kidney; anomalies of bony structures; positions of the colon, spleen, liver, and lung relative to the kidney; and the target of puncture (upper, middle, or lower calyx; caliceal diverticulum). The scanning head can be positioned to provide the best visualization and optimum puncture site for each patient. Thus, a puncture site as high as above the 11th rib may be chosen if the lung is not visualized in the puncture route. A different puncture site must be chosen if bowel gas or the liver or spleen is visualized within the intended nephrostomy route.

The route of puncture should always aim through a pyramid into a dorsal calyx; puncture into an infundibulum may result in bleeding from segmental and interlobar vessels in the renal sinus, and direct puncture of the renal pelvis renders dilation of the nephrostomy tract and insertion of catheters and instruments difficult, with increased risk of accidental catheter dislodgment after successful entry. For large, complete staghorn calculi, when PNL is to be performed for debulking the stone volume (followed by extracorporeal shockwave lithotripsy [ESWL] for disintegrating retained caliceal stones), puncture is usually performed through a lower dorsal calyx, a position from which the lower caliceal group, the renal pelvis, and part of the upper caliceal group can be reached easily with rigid instruments. However, for staghorn stones that can be completely removed by PNL alone (without ESWL), another route (eg, middle or upper calyx puncture) may be chosen. Stones in caliceal diverticula are better approached by direct puncture of the diverticulum.

Once chosen, the target for access to the renal collecting system must be visualized ultrasonically. The cutaneous puncture site should be chosen in a virtual caudal extension of the perpendicular orientation (width) of the scanning plane. Skin and fascia are incised with a no. 11 blade. At this time, the scanning head may be shifted over the incision to measure the exact distance between the incision and the target. A 16- to 18-gauge puncture needle (Figure 8–2) may then be inserted blindly through the incision and aimed in the direction previously determined by ultrasonography. However, the needle should never be advanced blindly farther than through the abdominal fascia.

The scanning head is now placed in such a way that both the target and the puncture needle are visualized in the same scanning plane, and the needle is aligned so that its tip can be clearly seen. Vibrating the needle makes the tip more visible while the position of the scanning head is being adjusted. The needle can be safely moved back and

### Table 8–1. Indications for Percutaneous Puncture of the Renal Collecting System.

<table>
<thead>
<tr>
<th>Diagnostic indications</th>
<th>Therapeutic indications</th>
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<tbody>
<tr>
<td>Antegrade pyelography</td>
<td>Nephrostomy catheter drainage</td>
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<tr>
<td>Pressure/perfusion study (Whitaker test)</td>
<td>Antegrade ureteral stenting</td>
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<td></td>
<td>Dilation of ureteral strictures</td>
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<td></td>
<td>Percutaneous endopyeloplasty</td>
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<td></td>
<td>Perfusion chemolysis of renal stones</td>
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<td>PNL</td>
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<td>Percutaneous resection and coagulation of urothelial tumors</td>
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<td></td>
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<tr>
<td>PNL, percutaneous nephrolithotomy.</td>
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**Figure 8–1. Renal ultrasound.**

A: The longitudinal axis of the kidney forms a 30° angle with the midline. **B:** The transverse axis of the kidney forms a 45° angle with both a horizontal and a vertical line.
forth down to the renal capsule as often as necessary, but
the renal parenchyma ideally should be punctured only
once.

A needle guide can be used to direct the needle exactly
within the ultrasonic scanning plane. With some needle
guides, the angle of puncture relative to the longitudinal
axis of the scanning plane (depth) is also fixed and may be
indicated on the monitor by an electronically generated
beam. If a steeper or flatter angle of puncture is desired, the
entire scanning head and attached needle guide must be
tilted, and the choice of puncture site is therefore limited.

Another drawback of this device is that it does not allow
for independent adjustment of the puncture and scanning
direction if the needle deviates from its intended direction
after being advanced through the skin. This frequently
occurs in patients with scars from previous operations and
becomes more of a problem the farther the target is from
the cutaneous puncture site. Freehand puncture with indi-
vidual adjustment of puncture and scanning direction is
preferable in these cases.

Movement of the kidney during respiration may com-
plicate puncture if the target is small and is visible on the
monitor only during a specific respiratory phase. If the
direction of the needle and the position of the target are
aligned and both are clearly seen on the monitor, the nee-
dle is advanced through the renal capsule during the
appropriate phase of respiration (Figure 8–3). In this
phase, the kidney is usually pushed to some extent by the
puncture needle, so that visualization of needle and target
may be momentarily impaired. However, as soon as the tip
of the needle has penetrated the fibrous renal capsule, it is
seen even more clearly. If both the tip of the needle and
the target are visualized clearly at the same spot on the
scanning plane, the needle is in the desired space.

Antegrade injection of a small amount of contrast dye
for fluoroscopy outlines the renal collecting system after
successful puncture. However, if the collecting system has
not been successfully punctured at the first attempt, con-
trast dye may fill the interlobar veins, which form a basket-
like structure around the calyx, or may extravasate. In rare
cases in which contrast dye is injected into the adventitia of
the renal collecting system, extravasation may assume the
configuration of the collecting system, mimicking success-
ful puncture. Care must be taken to inject the least amount
of dye necessary so that further fluoroscopic and ultrasonic
orientation will not be hindered. A larger amount of dye
injected outside the collecting system may compress the
calyx to be entered and render puncture more difficult. If the position of the needle tip on ultrasound is close to its destination (ascertained by a small vibratory movement), the needle should be retracted a few millimeters only and readvanced at the appropriate angle and tilt. Once the collecting system is entered (Figure 8–4A), fluoroscopy alone is used to guide the subsequent steps of the procedure.

If fluoroscopy is used instead of ultrasound for guiding renal puncture, a fine-needle (20–22 gauge) puncture technique may be used. Intravenous or retrograde administration of contrast dye is needed. With retrograde injection, a ureteral balloon occlusion catheter can be inserted and blocked in the ureteropelvic junction (UPJ) to cause slight distention of the renal collecting system; this facilitates puncture of a nondilated system. First, a 16- to 18-gauge needle is inserted through the abdominal wall only, and a longer fine needle is inserted coaxially through the larger needle (Figure 8–4B). This technique improves control of the fine needle. As soon as the fine needle has entered the collecting system, the larger needle can be advanced over the fine needle, which serves as a guide. After withdrawal of the fine needle, a regular guidewire can be inserted through the large needle into the collecting system.

Urine aspirated from the collecting system should be cultured, especially if there is suspicion of a urinary tract infection.

**ANTEGRADE PYELOGRAPHY & PRESSURE/PERFUSION STUDIES**

Renal puncture is rarely indicated for diagnostic antegrade pyelography only, because less invasive radiographic techniques are available (eg, intravenous pyelography with tomograms, ultrasound, CT, magnetic resonance imaging [MRI], retrograde pyelography). However, obtaining a radiograph after antegrade injection of contrast dye should be an integral part of every percutaneous puncture for any indication. Before contrast dye is injected, urine must be aspirated to decompress an obstructed collecting system. The contrast dye should be diluted to 20–30% for better visualization of details; antegrade pyelography then provides images of the collecting system with about the same resolution of detail as retrograde pyelography.

Antegrade pyelography is also performed in conjunction with a percutaneous pressure/perfusion study (Whitaker test) to assess pyeloureteral resistance. Percutaneous urodynamic studies of the dilated upper urinary tract are indicated only in the 10–30% of cases in which noninvasive radioisotope studies (diuresis renogram) fail to differentiate an obstructed from a nonobstructed dilated system. (This is more likely in cases of ureterovesical obstruction than in pelvic-ureteral obstruction, in which diuresis renograms are reliable.)

The Whitaker test provides simultaneous measurements of intrapelvic and intravesical pressures during antegrade perfusion, with flow rates of 5, 10, 15, and 20 mL/min. Puncture of the renal collecting system is performed with a coaxial needle/catheter system with an outer 6F catheter for the renal pressure/perfusion study; thus, puncture and catheter insertion can be done as a one-step procedure. Perfusion is started with flow rates of 5–10 mL/min until steady-state equilibrium of pressure readings is reached and the entire upper urinary tract is opacified (Figure 8–5). Pressure readings may be obtained intermittently, from the perfusion catheter via a 3-way stopcock, or continuously, if a double-lumen nephrostomy catheter or 2 separate catheters for perfusion and pressure measurement are used. Continuous recordings during perfusion from a single-lumen perfusion catheter via a T connection yield erroneous pressure readings (the smaller the lumen of the nephrostomy catheter and the higher the perfusion rate, the higher the pressure reading), unless the resistance of the entire system was previously calibrated for each rate of perfusion. To obtain accurate pressure readings, the positions of the intrapelvic and intravesical pressure manometers must be adjusted to the level of the renal pelvis and bladder, respectively. At a flow rate of 10 mL/min, differential pressures (renal pelvic pressure minus bladder pressure) below 13 cm water are normal, between 14 and 22 cm water suggest mild obstruction, and above 22 cm water suggest moderate to severe obstruction. At flow rates
of 15 mL/min and 20 mL/min, upper limits of normal pressure are 18 cm and 21 cm water, respectively.

PERCUTANEOUS CATHETER PLACEMENT

Percutaneous nephrostomy catheter placement for drainage and decompression of the upper urinary tract is indicated if retrograde ureteral catheterization is not advisable (eg, in sepsis secondary to ureteral obstruction) or proves to be impossible (eg, impassable ureteral obstruction due to stone, tumor, or stricture). After percutaneous endourologic procedures, a nephrostomy catheter is usually left indwelling for a few days. To convert a nephrostomy catheter diversion into internal stent drainage, antegrade ureteral stenting through the nephrostomy tract may be attempted even in cases in which previous attempts at retrograde stenting have failed. The antegrade approach to stenting can be expected to be successful if failure of retrograde stenting was not related to mere mechanical ureteral obstruction but rather to ureteral tortuosity, false passage (ureterovaginal fistula, urinoma after open surgery), or inability to identify the orifice endoscopically (ureterointestinal anastomosis).

For diagnostic procedures such as pressure/perfusion studies (Whitaker test), a 6F catheter is sufficient. Catheters of this size can be placed in a one-step procedure of puncture if coaxial needle/catheter systems are used (Figure 8–2). For therapeutic interventions such as nephrostomy drainage or antegrade ureteral stenting, softer, larger catheters must be inserted, and puncture tract dilation is necessary before catheter insertion. For dilation of a puncture tract, a 0.035- or 0.038-in guidewire must be inserted into the collecting system, either directly through the puncture needle or through the outer catheter of a coaxial needle/catheter system. Curved-tip (J) guidewires are less likely to cause damage to the mucosa of the renal pelvis than are straight guidewires. One of the most common problems of tract dilation is kinking of the guidewire during insertion of fascial dilators; therefore, guidewires with a floppy tip and a stiff proximal section (Lunderquist wire) are preferable over floppy guidewires. If the tip of the guidewire cannot be advanced into the renal pelvis because it is trapped in a dilated calyx with a narrow infundibulum or because an obstructing stone hinders passage, the outer catheter of a coaxial needle/catheter system can be used to manipulate the guidewire into the collecting system (Figure 8–6A), or angiographic catheters with different curved-tip configurations may be inserted over the guidewire for this purpose. Once the guidewire is in the correct position (upper calyx, renal pelvis, upper ureter), radiopaque fascial dilators can be inserted under fluoroscopic control with rotating movement of the dilator during advancement. If flexible plastic fascial dilators are used, sequential insertion of dilators of increasing size (usually in 2F steps) is necessary. If stiff metal or Kevlar dilators are used, dilation from 6F to 10–12F is possible in a one-step procedure.

After tract dilation, relatively stiff nephrostomy catheters (eg, polyethylene catheters) can be introduced easily over the guidewire. However, if softer catheters (eg, silicone or polyurethane) are to be inserted, use of an introducer catheter is helpful. An introducer catheter is also helpful for antegrade ureteral stenting and for insertion of nephrostomy catheters with various self-retaining configurations of the tip (eg, pigtail). These catheters can be stretched into a straight configuration while being inserted through the introducer catheter and over a guidewire; the tip resumes its original configuration due to the memory function of the material once the guidewire is withdrawn. The introducer catheter can be inserted with the last fascial dilator in a one-step procedure if a coaxial dilator/introducer catheter system is used (Figures 8–6B and C). The use of an introducer catheter provides universal access to the renal collecting system for placement of all types of
catheters (nephrostomy catheters [Figure 8–6D], ureteral stents, balloon dilation catheters) and safety and working wires for different systems of large-bore nephrostomy tract dilation required for insertion of endoscopic instruments.

Nephrostomy catheters should be soft to avoid discomfort and irritation of the renal pelvis and should have a self-retaining mechanism or should be placed with enough slack to prevent dislodgment from the collecting system during movement of the kidney. Standard nephrostomy catheters are Malecot catheters, pigtail catheters, and loop catheters. Loop catheters have a very effective retaining mechanism; they may cause serious complications, however, if the catheter is accidentally pulled out of the kidney.

Antegrade ureteral stenting can be done through an introducer catheter using either open- or closed-tip stents. Catheters with open-tip configuration are advanced with a pusher catheter over a guidewire, which must be inserted through the introducer sheath down the ureter and into the bladder as a first step. Catheters with closed-tip configuration are advanced by pushing the indwelling wire. In either technique, a thread should be pulled through one of the proximal side holes of the catheter so that the catheter can be pulled back into the renal pelvis if it is advanced too far. The thread must be pulled out before the guidewire is withdrawn so that the pusher catheter can still hold the double-J stent in place.

An introducer catheter may also be used for insertion of a 7F balloon dilation catheter over a guidewire into the ureter to dilate ureteral strictures to 12–18F with balloon pressures of up to 15 atm. After successful dilation, an 8–10F stent is usually left indwelling for several weeks. This technique is most successful in ureteral strictures that are a complication of recent surgery for benign disorders, except ureteropelvic obstruction. Long-standing strictures or strictures due to tumor compression of the ureter, radiation damage, or ischemic ureteral necrosis after radical pelvic...

**Figure 8–6.** Small-bore tract dilation and nephrostomy catheter insertion. A: J-guidewire inserted through the needle-catheter system and advanced with assistance of the catheter into the renal pelvis. B: Insertion of a coaxial dilator/introducer catheter system over the guidewire. Stiff proximal section of the guidewire prevents extrarenal kinking. C: After the dilator has entered the collecting system, the introducer catheter is advanced over its tip. D: Pigtail nephrostomy catheter is inserted into the renal pelvis over the guidewire and through the introducer catheter.
surgery are not likely to respond favorably to balloon dilation. Long-term results of ureteral balloon dilation cannot be determined from published data, either because the periods of follow-up were too short or because balloon dilations were repeated periodically.

**PERFUSION-CHEMOLYSIS OF RENAL STONES**

Nephrostomy catheters may be used for perfusion of the renal collecting system with chemolytic agents for dissolution of renal stones. In principle, uric acid, cystine, struvite, or apatite stones are amenable to chemolysis. However, the success of ESWL and the possibility of oral chemolysis (for uric acid stones) have limited the use of percutaneous chemolysis to adjunctive treatment of residual stones after open surgery, PNL, or ESWL. Primary percutaneous chemolysis may still be indicated in patients who are poor anesthetic risks. Benefits of percutaneous chemolysis must be weighed against disadvantages and possible risks, for example, prolonged hospitalization for dissolution of large stones (cystine, struvite, or apatite stones) and possible complications of treating infection stones (sepsis, hypermagnesemia).

To limit risks, perfusion chemolysis should always be performed with a double-catheter system for irrigation and simultaneous continuous drainage. This is achieved by using either 2 separately or coaxially inserted nephrostomy catheters (Figure 8–7A) or a ureteral catheter in conjunction with a nephrostomy catheter (Figure 8–7B). To ensure effective flow around the stone, the irrigation catheter must be placed close to the stone. Lack of continuous, complete drainage of the perfusate with increased intrapelvic pressures above 30 cm water may lead to pyelotubular and pyelovenous reflux of chemolytic agents and, possibly, infected urine, resulting in hypermagnesemia (perfusion with hemiacidrin or Suby’s solution G or M) and sepsis. Irrigation should be started only in the absence of urinary tract infection or if infection is under control. Irrigation must first be tested with saline at the lowest possible height above kidney level to achieve a flow rate of 100–120 mL/h. Discomfort, pain, or leakage of perfusate may indicate inappropriate drainage of the irrigant, and patients should be instructed to interrupt the irrigation themselves in such instances.

Uric acid stones can be dissolved by sodium or potassium bicarbonate solution; cystine stones with D-penicillamine, acetylcysteine, or tromethamine-E solution; and struvite and apatite stones with Suby’s solution G or M or hemiacidrin (Renacidin; not U.S. Food and Drug Administration [FDA]-approved for renal irrigation). Patients must be monitored for developing urinary tract infection or fever, and serum creatinine, phosphorus (hemicidrin-perfusion), and magnesium levels (perfusion with hemicidrin, Suby’s solution G or M) must be obtained every other day.

**Figure 8–7.** Catheter placement for perfusion chemolysis of renal stones. **A:** Perfusion and drainage of the irrigating fluid through 2 nephrostomy catheters. **B:** Perfusion through nephrostomy catheter, and drainage of the irrigant through ureteral catheter.
The time necessary for complete stone dissolution depends on the composition and size of the stone and may vary from a few days (uric acid stones) to several weeks (cystine or struvite stones).

**ENDOSCOPIC INTRARENAL INSTRUMENTATION**

Nephroscopes are endoscopic instruments with sheaths of 15–26F that are inserted percutaneously through a nephrostomy tract. Standard rigid instruments are available in sizes 24–26F; these have telescopes with offset eyepieces (Figure 8–8, left). Rigid instruments such as graspers and ultrasound probes can be inserted through a central working channel (Figure 8–8, right). Flexible fiberoptic nephroscopes may be used as well. These have a deflecting mechanism for the tip that allows inspection of otherwise difficult-to-reach calyces. A smaller working channel allows insertion of flexible instruments such as stone baskets, wire graspers, and electrohydraulic or laser probes. However, instrumentation through flexible nephroscopes is limited by the size and flexibility of working instruments such as stone forceps, and flexible endoscopes do not offer the optical quality and durability of rigid nephroscopes.

Nephroscopy is rarely indicated for diagnostic purposes only; in most cases, it is performed for percutaneous lithotripsy and extraction of renal stones (PNL). However, ESWL has gradually replaced PNL for treatment of renal stones and is now used in >90% of cases. PNL is still indicated in cases for which ESWL is not the primary choice of treatment. Such cases include urinary obstruction not caused by the stone itself, large-volume stones, and stones that cannot be positioned within the focus of the shock wave apparatus. PNL can achieve stone-free rates of >90%. Nephroscopes also may be used for direct-vision internal incision of ureteropelvic stenosis and for palliative treatment of urothelial cancer of the upper urinary tract.

Insertion of a nephroscope into the renal collecting system requires dilation of the puncture tract to 24–30F. A safety wire should be inserted parallel to the working wire and advanced into an upper calyx or the upper ureter to guide the way back into the collecting system in case the dilator and working wire become dislodged accidentally. Insertion of an introducer catheter during small-bore tract dilation to 10–12F facilitates parallel insertion of safety and working wires. The central metal catheter of a coaxial metal dilator system (Figure 8–9, left), the central plastic catheter for insertion of sequential plastic dilators, or a balloon dilator catheter can be inserted over the working wire. Balloon dilator catheters of 9F size can dilate a nephrostomy tract to a diameter of 30F under pressure up to 10–12 atm in a one-step procedure. This may prove difficult or impossible if perirenal scar tissue from previous surgery prevents complete expansion of the balloon over its entire length. Sequential plastic dilators allow stepwise dilation of the tract under fluoroscopic control; however, on withdrawal for insertion of the next larger dilator, compression of the tract is lost intermittently and bleeding occurs into the collecting system, sometimes hindering subsequent endoscopy. Coaxial metal dilators (Figure 8–9, right) (each dilator slides over the next smaller one) allow stepwise tract dilation even in the presence of severe scarring with continuous nephrostomy tract compression for improved hemostasis.
With any dilation technique, the last step is insertion of a working sheath, which may be either the 24–26F metal working sheath of the nephroscope or a larger plastic sheath. With the balloon dilation technique, the working sheath must be introduced over a plastic dilator; with use of serial plastic or coaxial metal dilators, the working sheath slides over the last dilator. The Pathway Access Sheath is a novel device consisting of a balloon dilator with a coaxial external expandable sheath. It allows for balloon tract dilation and percutaneous access sheath placement in one simple step. Clinical studies comparing it to the standard two-step technique have demonstrated a quicker and gentler insertion for this new device. The combination of decreased axial forces and exclusion of multiple steps should theoretically result in less shearing of tissue. A 28–30F plastic working sheath is preferable to a metal nephroscope sheath in all cases in which extensive, prolonged instrumentation is anticipated (eg, staghorn stones). Larger plastic sheaths not only provide better irrigation with lower intrapelvic pressures than do continuous-flow nephroscope sheaths but also allow easier extraction of large stone fragments.

Renal Stones

In the era of ESWL, indications for PNL are limited to 4 types of stone disease:

1. Urinary obstruction not caused by the stone itself (eg, stone in a caliceal diverticulum [Figure 8–10, left and right], stone in association with ureteropelvic stenosis). These stones could be broken up by ESWL, but gravel would not pass spontaneously.

2. Large-volume stones (>3 cm, stone surface >500 mm²) (Figure 8–11, left and right) (eg, staghorn stones). These stones can be treated by several sessions of ESWL, but only about 30% of patients become stone free. However, problems associated with passing large quantities of gravel (eg, ureteral obstruction, pain, fever, sepsis) can be prevented by first percutaneously debulking the stone and then performing ESWL for endoscopically inaccessible stones.

3. Stones that cannot be positioned within the focus of the shock wave apparatus (eg, stones in kidneys with abnormal position due to anomalies of the urinary tract or skeleton, stones in transplanted kidneys, kidney stones in very obese patients that cannot be positioned into the focus of the shock wave source due to the increased distance from skin to stone, or when the weight limit of the ESWL table is exceeded).

4. PNL may be of benefit for lower pole caliceal calculi even under the 2–3-cm range. The overall stone-free rate for these stones with ESWL is only about 60%.

Large-volume staghorn stones are a much more common indication for PNL than stones that can be extracted in toto. Small stones can be extracted with a variety of rigid forceps and graspers (Figure 8–8, right). Stones may be retrieved from difficult-to-reach calyces with flexible wire baskets and graspers inserted through flexible nephrosopes. Large stones must be disintegrated using mechanical, ultrasonic, electrohydraulic, or laser energy. Strong nutcracker-type forceps (visual lithotrite, stone punch; derived from instruments for transurethral bladder stone extraction) are often employed.
venous pyelogram.

Figure 8–11. Staghorn stone requiring combined percutaneous nephrolithotomy and extracorporeal shockwave lithotripsy. Left: Plain abdominal radiograph. Right: Intravenous pyelogram.

disintegration) can be used only in a spacious renal pelvis. Hollow ultrasonic probes allow for controllable, systematic stone disintegration under continuous suction for removal of sand and smaller fragments. Electrohydraulic probes are more powerful than ultrasonic probes and may be used through flexible nephrosopes, but they do not provide continuous suction and are associated with a higher risk of scattering stone fragments into inaccessible calyces and of damaging the mucosa of the renal pelvis. However, with electrohydraulic probes and the holmium:YAG laser, disintegration of hard or large stones is faster.

For soft stones, continuous disintegration and evacuation of fragments with ultrasound probes is most time-efficient. Hard stones should be broken up into the largest possible fragments that can be extracted through the working sheath. The ureteropelvic portion of a staghorn stone should be left in place until the procedure is nearly completed, as it will act like a plug in a drain to prevent the loss of fragments into the ureter. An antegrade or retrograde positioned ureteral balloon occlusion catheter might serve the same purpose; however, the extra procedure of retrograde ureteral catheterization is rarely indicated.

Normal saline should be used as the irrigation fluid except in the case of electrohydraulic lithotripsy, in which one-sixth normal saline is more appropriate. However, even with the low-pressure system provided by a large plastic working sheath, considerable amounts of irrigation fluid may be absorbed if small veins are opened and intrarenal manipulation is prolonged. This may cause transurethral resection (TUR) syndrome with use of hypotonic fluids. Intraoperative administration of diuretics (eg, mannitol, 12.5 g) is advisable and also has proved effective in preventing intrarenal reflux. If there is suspicion of extravasation, contrast dye must be injected and a diagnostic radiograph obtained. On completion of the procedure, a plain film should be obtained and a nephrostomy catheter placed. A Foley catheter with a 5-mL balloon may be inserted through a fenestrated trocar or the plastic working sheath, which then is withdrawn and cut lengthwise for removal from the Foley catheter. Malecot catheters or straight polyethylene catheters (eg, chest tubes) may be used as well and should be secured to the skin with sutures. A final nephrostogram documents appropriate position of the catheter.

Nephrostomy catheters may be removed after 1–4 days after antegrade pyelography to check for unobstructed upper urinary tract drainage. In cases of postoperative profuse hemorrhage, the nephrostomy catheter can be occluded for 1–2 days to allow for tamponade formation. Blood clots usually dissolve later spontaneously due to urokinase activity without problems. If ESWL is to be performed, it can be done 1–4 days after the percutaneous procedure. The nephrostomy catheter should be left in place during and after ESWL to provide drainage for urine and stone gravel and to allow for a second endoscopic procedure if some of the stone fragments do not pass spontaneously after ESWL.

After removing a large bore nephrostomy catheter (24–30F), urinary discharge from the nephrostomy tract can persist for several days and be bothersome and concerning to the patient. To prevent this, the nephrostomy tube can be exchanged over a guidewire under fluoroscopic guidance for a smaller caliber catheter. Leaving this smaller tube in place for a few days will permit the tissues around the tract to expand and minimize leakage.

Some experienced endourologists have propagated percutaneous stone treatment and endopyelotomy without standard placement of a nephrostomy tube. Major advantages are marked reduction in analgesia requirements and length of hospitalization. Prerequisites are a small to moderate stone burden and no residual fragments, no more than 2 percutaneous tracts in 1 session, and no significant bleeding. Bleeding arising from the nephrostomy tract can be stopped nephroscopically by punctual electrocoagulation during withdrawal of the working sheath. However, this “so-called” tubeless percutaneous renal surgery is best performed with intraoperative antegrade placement of an internal ureteral stent to secure unobstructed urinary drainage. Expected patient discomfort from the nephrostomy catheter is traded against possible discomfort from the internal urinary stent and from cystoscopy to remove the stent later.

**Ureteropelvic Stenosis**

With the advent of PNL and ureterorenoscopy, other endosurgical techniques have been developed that are similar to procedures used in the lower urinary tract. Direct-vision internal incision of ureteropelvic stenosis (pyelolysis, endopyelotomy, endopyeloplasty) seems to be a natural outgrowth of endoscopic techniques in the upper urinary tract. Compared with the retrograde techniques of endopyelotomy (incision with a cold knife, Acucise cathe-
ter, Greenwald electrode, or laser) and the endoballoon rupture, the antegrade technique offers the advantage of an incision under direct vision. The cold-knife incision must be extended into the perirenal fat and is stented for 4–6 weeks to allow for healing, according to the principle of Davis’ intubated ureterotomy.

Success rates of antegrade endopyelotomy of up to 65–95% are reported for primary cases and up to 89% for secondary cases after failed open-surgical pyeloplasty.

The success rates for retrograde endopyelotomy with fewer patients and less follow-up than in antegrade endopyelotomy range between 73% and 90%; for the AcuCise endopyelotomy, between 76% and 81% (see Ureteropelvic Stenosis). The candidates for best endoscopic (antegrade and retrograde) outcome are those with less than grade II hydronephrosis and good renal function.

In most reports on endopyelotomy, the criteria of success differ from those of open pyeloplasty; relief of subjective symptoms is given priority over results of imaging studies such as decompression of a dilated collecting system on intravenous pyelography or renal ultrasound. Inadequate results after endopyelotomy may be related to a crossing vessel or to redundancy of the renal pelvis, which would be resected during open pyeloplasty. According to the law of Laplace, wall tension of a renal pelvis is, at the same intrapelvic pressures, higher in a more dilated collecting system with a larger diameter than in a less dilated system with a smaller diameter. A raised wall tension supposedly represents a more important pathogenetic factor for developing progressive dilatation than do elevated intrapelvic pressures due to anatomic obstruction of outflow. Secondary open pyeloplasty after failed endopyelotomy may be a more tedious operation with less satisfactory results in cases with extensive periureteral scarring due to extravasation of urine after endopyelotomy than primary open pyeloplasty.

Renal Pelvis Tumor

Another technique of endoscopic surgery in the upper urinary tract is use of electrosection, electrocoagulation, electrovaporization, and neodymium:YAG laser coagulation for treatment of urothelial tumors of the renal pelvis. However, with the limited reports of treatment of upper urinary tract urothelial cancer endoscopically, recurrence rates are yet to be compared with those of standard surgical treatment.

Ensuring a strict follow-up, percutaneous management of transitional cell carcinoma of the collecting system may be an alternative to nephroureterectomy for patients with grade I disease and for palliative treatment.

Percutaneous aspiration of cystic or solid lesions of the kidney and the adjacent retroperitoneum is usually performed for diagnostic purposes, in some cases in combination with therapeutic intentions such as drainage of fluid collections or obliteration of renal cysts (Tables 8–2 and 8–3). Because most of these lesions are radiolucent and are not enhanced with intravenously administered contrast dye, they cannot be visualized by fluoroscopy. Thus, ultrasound or CT is the imaging technique of choice to depict these lesions and guide percutaneous puncture. The technique of ultrasonically guided puncture is the same whether the target is the renal collecting system or a cystic or solid renal or extrarenal lesion. For cytologic aspiration, a fine-needle (20–22 gauge) aspiration technique is used that is comparable to fine-needle aspiration biopsy of the prostate. There is no evidence that one type of needle is preferable to the others. For aspiration and evacuation of renal cysts or

### Table 8–2. Indications for Puncture of Renal and Retroperitoneal Lesions.

<table>
<thead>
<tr>
<th>Diagnostic indications</th>
<th>Therapeutic indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid aspiration</td>
<td>Catheter drainage</td>
</tr>
<tr>
<td>Fluid chemistry</td>
<td>Urinoma, lymphocele</td>
</tr>
<tr>
<td>Bacteriology and sensitivity</td>
<td>Abscess, hematoma</td>
</tr>
<tr>
<td>Cytology</td>
<td>Fluid evacuation and injection of sclerosing agent</td>
</tr>
<tr>
<td>Radiography with percutaneously injected contrast dye</td>
<td>Simple renal cyst</td>
</tr>
<tr>
<td>Histology (core biopsy)</td>
<td></td>
</tr>
</tbody>
</table>

### Table 8–3. Differential Diagnosis of Renal and Retroperitoneal Lesions.

<table>
<thead>
<tr>
<th>Renal cystic lesion</th>
<th>Retroperitoneal fluid collection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benign cyst</td>
<td>Urinoma</td>
</tr>
<tr>
<td>Hydrocalix</td>
<td>Lymphocele</td>
</tr>
<tr>
<td>Abscess</td>
<td>Hematoma</td>
</tr>
<tr>
<td>Hematoma</td>
<td>Abscess</td>
</tr>
<tr>
<td>Cystic tumor</td>
<td>Cystic tumor</td>
</tr>
<tr>
<td>Tumor in cyst</td>
<td></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Solid renal and retroperitoneal lesions</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Benign tumor</td>
<td></td>
</tr>
<tr>
<td>Malignant primary tumor</td>
<td></td>
</tr>
<tr>
<td>Metastatic tumor</td>
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</tbody>
</table>
extrarenal fluid collections (urinoma, lymphocele), the same coaxial needle/catheter system can be used as for percutaneous puncture of the renal collecting system. A small catheter (6–10F) is placed for a few days to ensure complete drainage of fluid. When fluids of high viscosity (abscess, hematoma) are to be drained, large-bore catheters (14–20F) must be inserted, necessitating dilation of the percutaneous tract. Percutaneous renal biopsy for histologic diagnosis and classification of renal disease is performed with 14- to 16-gauge needles (eg, Franklin-Silverman, Tru-Cut) at the lower pole of the kidney.

**Renal Cysts**

Renal cysts are found in about 50% of autopsy specimens in persons over the age of 50 years and are a frequent accidental finding on ultrasound or CT studies. Only a few cases require diagnostic percutaneous puncture. Indications for diagnostic puncture of a cystic lesion are an irregular, thick wall and internal echoes on ultrasound examination; density numbers on CT higher than those of serous fluid; and hematuria. Puncture for therapeutic procedures (evacuation of fluid and instillation of a sclerosing agent) is indicated only if, due to its size or location, the cyst causes compression and urinary obstruction of a caliceal infundibulum or the ureter, or discomfort and pain.

Various tests may be performed on aspirated fluid. No one test is pathognomonic except cytologic findings of malignant cells. However, neoplasms within a cyst are exceedingly rare, and cystic degeneration of a renal neoplasm can usually be easily identified by ultrasound and CT. Benign cysts contain clear, straw-colored fluid with low fat and protein content and lactic acid dehydrogenase levels of <250 mIU/mL. Cancer is suspected if the fluid is bloody or murky and has a high content of fat, protein, and lactic acid dehydrogenase. After aspiration of 20–30% of the cystic fluid, the same amount of 60% contrast dye is injected, and diagnostic radiographs are obtained in the prone, supine, upright, decubitus, and Trendelenburg positions. If necessary, another 20–30% of the cystic fluid may be replaced by air for obtaining double-contrast radiographs.

For therapeutic obliteration of cysts, sclerosing agents such as Pantopaque or 95% ethanol can be injected after complete evacuation of the cystic fluid. A volume of 10–100 mL of 95% ethanol, approximating 10–20% of the original volume of cystic fluid, is injected into the cyst and should be drained after 30 minutes.

**Retroperitoneal Fluid Collections**

Low-viscosity retroperitoneal fluid collections (urinoma, lymphocele) are usually a complication of surgical procedures. However, urinoma may also be caused by exogenous trauma or by fornix rupture due to acute ureteral obstruction. Percutaneous techniques of catheter drainage eliminate the need for open surgical revision in most cases. Insertion of a small (6–10F) catheter (with numerous side holes) is usually sufficient. Adjunctive procedures are performed to ensure sealing of the fluid leak and obliteration of the cystic lesion. In cases of urinoma, the upper urinary tract must also be drained by a ureteral catheter or percutaneous nephrostomy catheter until drainage from the urinoma stops. Lymphoceles that develop following pelvic or retroperitoneal lymphadenectomy or renal transplantation often undergo spontaneous regression and usually do not require puncture and drainage. However, large lymphoceles developing after retroperitoneal lymphadenectomy may cause pain and even ureteral obstruction (Figure 8–12). Patients should be treated with parenteral nutrition and abdominal compression by bandaging, but if lymph drainage after percutaneous puncture and catheter placement persists for >1 week, surgical intervention with intraperitoneal marsupialization of the lymphocele and ligation or electrocoagulation of lymphatic vessels is indicated.

High-viscosity fluid collections (hematoma, abscess) usually require large-bore (14–20F) percutaneous catheters for sufficient drainage. Perirenal hematomas are most fre-

Figure 8–12. Percutaneous drainage of a lymphocele causing ureteral displacement and compression.
frequently caused by surgical or exogenous trauma and rarely develop spontaneously in the presence of a bleeding disorder or due to rupture of a renal tumor. Indications for percutaneous drainage are rare, as most small hematomas resolve spontaneously and should be followed by ultrasound or CT only. A hematoma that increases in size requires surgical intervention rather than percutaneous drainage. Secondary infection of a hematoma may be an indication for percutaneous drainage.

A perirenal abscess is mostly a complication of open surgery; hematogenic renal abscess (renal carbuncle) is less frequent. Indications for puncture and drainage should be based on CT finding of a unifocal process that can be effectively and safely drained percutaneously. Multifocal renal abscess formation is not amenable to percutaneous drainage.

**Renal & Retroperitoneal Tumors**

Percutaneous aspiration biopsy of renal and retroperitoneal tumors is indicated if less invasive radiographic studies are inconclusive and if cytologic findings may have an impact on further medical or surgical therapy (Figure 8–13). If curative treatment by open surgery seems to be feasible, aspiration biopsy is generally not indicated. If the identity of a renal lesion is questionable or if conservative, organ-sparing surgery is technically feasible, surgical excision of the lesion with intraoperative frozen sections is preferable over percutaneous aspiration biopsy. However, aspiration biopsy may be indicated to avoid radical nephrectomy of a possibly benign lesion. In multifocal or possibly metastatic lesions, cytologic evaluation can be crucial for planning surgical or medical therapy, and in these cases, aspiration biopsy is usually indicated. Interpretation of cytologic findings is limited by a 10–25% incidence of false-negative findings and the difficulty in discriminating normal renal tubular cells from low-grade renal cell cancer. As a rare complication, tumor seeding in the puncture tract has been described. The aspirate is immediately spread on glass slides. For standard Papanicolaou stains, alcohol fixation must be used.

![Figure 8–13. Percutaneous fine-needle biopsy. **Left:** Aspiration biopsy of a renal lesion. **Right:** Guidance with computed tomography scanning for fine-needle aspiration biopsy of an exophytic renal cell carcinoma.](image)
The widespread use of ultrasound has led to an increased rate of incidental diagnosis of small renal tumors. With the advent of minimally invasive techniques and improved radiodiagnostic possibilities, nonresectional renal tumor ablation has emerged and is gaining increased attention. However, cryotherapy, radiofrequency ablation (RFA), and high-intensity focused ultrasound (HIFU) are still considered experimental procedures in the clinical setting. The therapeutic strategies of local energy application aim at selective tumor destruction with minimal injury to the surrounding normal kidney parenchyma and reduced morbidity. The biological principle of cryotherapy is tissue destruction by repeated rapid freeze and thaw cycles down to temperatures below 20°C. Liquid argon and liquid nitrogen are the 2 most commonly used freezing agents. The cell destruction mechanism comprises intra- and extracellular ice crystal formation leading to intracellular dehydration and ultimately cell disruption. This is followed by a delayed cell death, which occurs during the thaw phase owing to vasoconstriction and microcirculatory failure. However, since the freezing effect decreases with increasing distance from the freezing probe, the “iceball” has to extend approximately 1 cm beyond the tumor margin to ensure complete tumor destruction. RFA involves coagulation of tumors by directly applying temperatures >50°C via needle electrodes. Since tissue carbonization at the electrode tip increases the impedance for radiofrequency transmission, tissue conductivity can be maintained by simultaneously irrigating saline through the tissue ("wet" RFA), resulting in larger RFA lesions for therapy of larger renal tumors. This can also be accomplished by multiple electrodes, which create overlapping ablation fields. The vicinity of larger caliber vessels results in dissipation of heat (“heat sink effect”) and influences the efficacy of RFA negatively. Thus, peripheral exophytic tumors seem to be controlled better than central tumors in the vicinity of larger vessels.

Indications are similar for both techniques and are presently restricted to patients with comorbidity and/or advanced age who are not suitable to surgical treatment, impaired renal function, multiple bilateral tumors as in von Hippel-Lindau disease, and renal tumors in a solitary kidney. Guidelines for cryotherapy do not recommend treatment of tumors >3 cm in size and for RFA not larger than 5 cm. Further relative contraindications for both procedures include hilar or central tumors and cystic tumors. An absolute contraindication is untreated coagulopathy.

The tumor mass can be approached either by open surgery, laparoscopy, or percutaneously using fine probes and high-resolution imaging techniques. However, the minimally invasive character of the procedure itself ideally deserves a less invasive approach than open surgery. Proponents of laparoscopy emphasize its advantage of mobilizing the tumor and providing excellent exposure, thus avoiding damage to adjacent structures. It also allows for precise confirmation of probe positioning and monitoring progress of the procedure such as development of the iceball in cryotherapy under direct vision. Percutaneous management requires MRI or CT with technical capabilities to construct three-dimensional pictures for monitoring probe placement and progress of therapy. The percutaneous approach may be performed as an outpatient procedure. Special patients requiring multiple procedures as in von Hippel-Lindau disease thus may benefit from a percutaneous treatment. In contrast to cryoablation, which has the advantage of intraoperative laparoscopic and sonographic monitoring, RFA lacks reliable real-time monitoring of the therapeutic progress. However, the introduction of real-time MRI guidance and monitoring of RFA may overcome this difficulty.

Both methods, cryotherapy and RFA, have shown promising results in carefully selected patients. These methods appear to achieve comparable oncologic control to surgical tumor resection with decreased operative morbidity. However, follow-up for cryotherapy is only as short as little over 3 years as compared to 10 and 15 years follow-up periods for surgical tumor resection. Results after RFA appear to be similarly effective, but average follow-up for RFA is even shorter. In contrast to cryotherapy, where tumor size decreases with time, tumor size after RF remains mostly constant. Tumor size after successful cryotherapy can decrease up to 75% over 3 years, and tumors may even completely disappear on MRI in some cases. This fact is important for posttreatment surveillance. A major drawback of ablative techniques is the lack of reliable histologic confirmation of complete tumor ablation. Assessment is commonly done by CT scan. After RFA, a successfully ablated lesion becomes fibrotic and nonperfused and does not show contrast enhancement as compared to a viable tumor.

Complication rates (major and minor) of cryotherapy are 1.4% and 12.2%, and of RFA 2.2% and 6%, respectively. The most commonly observed complications of cryotherapy and RFA are pain and paresthesia at the probe insertion site. Rare complications include perinephric hematoma, renal rupture, UPJ obstruction, and damage to adjacent organs. Anteriorly or centrally located tumors that abut the UPJ pose an increased risk for complications, especially colonic injuries or lesions of the renal collecting system and ureter. Bleeding complications have decreased with the use of ultrathin probes (1.5 mm diameter). Specifically for RFA, bleeding can be minimized by active coagulation of the puncture tract while removing the probe.

Ablative methods are still constantly evolving. Uncertainties exist regarding the exact amount of energy required, duration of treatment, mode of energy delivery and types of electrodes used, and render comparison of published results difficult. The adjunctive use of chemotherapeutic agents such as cyclophosphamide, 5-fluoroura-
cil, and bleomycin, or radiotherapy may have a synergistic effect on cryoablation and intensify its ablative capabilities.

Other techniques with limited animal and clinical experience that remain experimental include HIFU, microwave thermotherapy (MT), laser interstitial thermotherapy (LITT), chemoablation with or without RF and radiosurgery.

Renal Biopsy
Renal biopsy for diagnosis and classification of medical renal disease can be performed percutaneously or by open surgery. Because specimens, rather than aspirates, are needed for diagnostic histologic study, large-bore (14–16 gauge) Franklin-Silverman or Tru-Cut needles are used. Ultrasonic or fluoroscopic guidance is preferable to blind renal puncture. However, even with puncture aimed precisely at the dorsal aspect of the lower pole of the kidney, where accidental injury to large vessels is less likely, bleeding is to be expected because of the vascularity of the parenchyma and is the major complication of this procedure (about 5% of cases, with a mortality rate of 0.1%). Hematoma can usually be followed conservatively by ultrasound or CT, but transvascular embolization, open surgical revision, and even nephrectomy have been required following diagnostic renal biopsy. Therefore, open surgical biopsy rather than percutaneous biopsy is indicated in patients with solitary kidneys or uncontrolled hypertension.

URETERORENOSCOPY
Ureterorenoscopy is endoscopy of the ureter up to the renal pelvis for both diagnostic evaluation and therapeutic intervention (Table 8–4).

Ureterorenoscopes (Figure 8–14) are endoscopes for retrograde insertion into the ureter; however, they also may be used in an antegrade fashion via a percutaneously established nephrostomy tract. Technical improvements in the last decade have led to the introduction of smaller caliber, more versatile instruments. As a consequence of this progress not only has ureterorenoscopic management of urinary calculi become more amenable and safer, but also endoscopic oncologic treatment is gaining an increasing role. Rigid ureterorenoscopes are available in sizes 6.9–12.6F, and semirigid fiberoptic ureterorenoscopes and flexible ureterorenoscopes may be found in sizes 6.2–9.3F. The smallest instruments are for diagnostic procedures only. Larger instruments, with a 3–6F working channel, can accept stone baskets, wire graspers, stone forceps, biopsy forceps, and ultrasonic, electrohydraulic, or laser probes for stone disintegration. Flexible ureterorenoscopes follow the topographic anatomy of the ureter more easily and facilitate inspection of middle and lower renal calyces if a deflecting mechanism for the tip of the instrument is provided. Newer state-of-the-art flexible ureterorenoscopes have 270° deflecting tips (dual active deflection) that allow access to virtually every calyx of the collecting system. However, the use of instrumentation through flexible nephroscopes is limited by the size and flexibility of working instruments such as stone forceps, and flexible ureterorenoscopes do not offer the optical quality and durability of rigid instruments.

Insertion of a ureterorenoscope into the ureteral orifice may be facilitated by dilation of the intramural ureter, either with sequential plastic dilators of increasing size, which are slid over a guidewire, or with a balloon dilator catheter (Figure 8–15). Dilation of the ureter is often unnecessary if a small (3–5F) ureteral catheter is inserted through the working channel of the ureterorenoscope into the ureter as a guide, and the ureterorenoscope is then rotated 180° and introduced in an upside-down orientation (Figure 8–16). In this position, the ureteral catheter will spread the roof of the intramural ureter like a tent and the nose of the instrument will slide flat on the trigone into the orifice. The orifice and intramural ureter will thus be dilated only to the extent necessary for insertion of the instrument.

<table>
<thead>
<tr>
<th>Table 8–4. Indications for Ureterorenoscopy.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Diagnostic indications</strong></td>
</tr>
<tr>
<td>Lesions of ureter or renal pelvis</td>
</tr>
<tr>
<td>Hematuria from upper tract</td>
</tr>
<tr>
<td><strong>Therapeutic indications</strong></td>
</tr>
<tr>
<td>Ureteral stone treatment</td>
</tr>
<tr>
<td>Direct vision internal ureterotomy of ureteral strictures</td>
</tr>
<tr>
<td>Endoscopic resection and coagulation of ureteral tumors</td>
</tr>
</tbody>
</table>
Diagnostic Ureterorenoscopy

Indications for diagnostic ureterorenoscopy are those rare lesions of the ureter or renal pelvis whose nature cannot be determined with less invasive diagnostic procedures such as retrograde pyelography, selective urinary cytology, CT, or MRI. If a small ureterorenoscope without a working channel is used for a diagnostic procedure, a biopsy of a lesion cannot be obtained. If ureterorenoscopy is performed for evaluation of hematuria from the upper tract, the source of bleeding can rarely be identified during gross hematuria because of limited irrigation through ureterorenoscopes, resulting in poor visibility. If the urine has cleared, the chances of identifying a lesion that could not be detected on radiographic studies are small as well.

Ureteral Stones

Ureterorenoscopy is most frequently performed for treatment of ureteral stones, although >90% of ureteral stones can be treated by ESWL either in situ or after dislodgment of the ureteral stone into the renal pelvis using a ureteral catheter (push-back or flush-back procedures). For the remaining indications of ureteral stone treatment, ureterorenoscopy is used for extraction of stones, dislodgment of stones into the renal pelvis for subsequent ESWL, and intraureteral stone disintegration. Growing experience in general ureteroscopic techniques and the use of smaller caliber instruments have also rendered the procedure practicable and safe in the pediatric population, with no evidence of decreasing renal function, and/or renal growth or initiation of vesicoureteral reflux after the procedure.

For extraction of distal ureteral stones that are unresponsive to ESWL, short rigid ureterorenoscopes and alligator forceps or Dormia baskets are most helpful. Ureteroscopic clearance rates for distal ureteral stones reach 93–100%. Impacted proximal ureteral stones that did not respond to in situ ESWL and could not be dislodged with a ureteral catheter usually can be repositioned into the renal pelvis under direct vision for subsequent ESWL using a small semirigid or rigid ureterorenoscope. If the stones are too large or impacted, intraureteral lithotripsy for stone disintegration may be necessary in a few cases. Flexible ureteroscopic instrumentation in the upper ureter and the necessity to pass and withdraw the ureteroscope repeatedly to extract stone fragments bear an increased risk of injury to the ureter. Hence, in skilled hands stone clearance rates for proximal ureteral calculi range from 82% to 87%. The “ureteral access sheath” has been developed to facilitate ure-
teral reentry and to allow multiple reinsertions of the ureterorenoscope into the upper urinary tract while reducing trauma to the distal ureter. The outer sheath size is available in 12–16F. The increased efflux of irrigation fluid from the working sheath allows endoscopy to be performed with increased irrigation flow rates, thus enhancing visualization of the upper urinary tract while maintaining low (<40 cm) intrapelvic pressures. Use of a working sheath is also supposed to decrease wear on the ureteroscope and reduce operative time. However, controversy still exists regarding a possible risk of ureteral injury from placement of the working sheath resulting in ureteral stricture rates of 1.4%, which however are still in the range of modern series of flexible ureteroscopy without the access sheath.

To prevent the pushing back of stones or fragments into the renal pelvis during lithotripsy, a 3F wire basket can be used to hold the stone during disintegration or a 3F balloon catheter can be passed alongside the stone and be blocked proximally. The Dretler stone cone (Microvasive, Boston Scientific) is a new nitinol device which uses the same principle. It can be coiled proximal to the stone, preventing upward ureteral migration of stone fragments during lithotripsy and facilitating stone fragment retrieval after successful disintegration. Ultrasound probes allow safe stone disintegration under continuous suction, but they are not as effective as electrohydraulic and laser probes and can be used through rigid ureterorenoscopes only. Both electrohydraulic and laser probes can be used through rigid or flexible ureterorenoscopes. Laser probes for intraureteral lithotripsy have the smallest diameter (<1F) and do not injure the mucosa of the ureter if used under direct vision. Several pulsed lasers such as the holmium:YAG laser, pulsed-dye laser, and Alexandrite laser are available for intraureteral lithotripsy. The holmium:YAG laser is the most widely used laser for ureteroscopic lithotripsy because of its high efficiency. It can successfully disintegrate stones of any composition with stone fragmentation rates approaching 100%. Moreover, it can be used to coagulate, ablate, and incise tissues. Laser probes are flexible and as small as 200 mm in core diameter so that they can be easily passed through flexible ureterorenoscopes without hindering tip deflection and thus, maximize their durability. Electrohydraulic probes, which are available in sizes 1.6–5F, require less expensive equipment, but they bear the risk of ureteral damage if used inappropriately. The combination of flexible ureteroscopy with atrumatic ancillary tools, such as nitinol tipless baskets and thin holmium laser fibers (200 µm), permits ureteroscopic treatment of renal stones when ESWL or percutaneous treatment are not an option or have failed. A 79–90% success rate has been reported for lower pole stones treated ureteroscopically. However, success is highly dependent on stone size and multiple procedures are often required. If ureteral perforation occurs as a complication of intrareteral instrumentation, ureteral stenting using a 6–8F double-J stent for 2–6 weeks usually allows healing without late sequelae. Stents should be used for a few days even after uncomplicated ureterorenoscopy to prevent pain from urinary stasis because of edema of the intramural ureter after instrumentation. If there is ureteral perforation and a stent has been placed, the bladder should be on continuous drainage for a few days using a transurethral Foley catheter or a suprapubic cystostomy catheter to prevent urinoma formation from vesicoureteral reflux via the double-J stent.

**Ureteropelvic Stenosis**

Ureteroscopic endopyelotomy of UPJ obstruction is feasible with cold knife, electrocautery, or holmium laser. Success rates range between 73% and 90% in different series. However, subsequent repeat balloon dilation of the UPJ may be required in 10% of cases, and 3% may even require repeat incision. At least 10% of recurrences are treated by open or laparoscopic surgery. Thus, the advantages of this minimally invasive procedure such as a reduced hospitalization time and convalescence are at the cost of poorer outcomes compared with open or laparoscopic pyeloplasty. Also, the risk of complications of ureteroscopy, which range between 1% and 15%, like ureteral perforation, stricture, false passage, ureteral avulsion, bleeding, and sepsis must be weighed against the possible benefits.

In recent years Acucise endopyelotomy has gained increased attention as an alternative treatment of UPJ obstruction and other postoperative ureteral strictures in selected patients. It is simple and a truly minimally invasive method that can be performed in an ambulatory setting, but relies on fluoroscopic rather than visual ureteroscopic guidance. The device permits combining balloon dilatation and incision by electrocautery in 1 step. Overall success rates for UPJ range between 76% and 81%, and even better outcomes are reported in secondary UPJ obstruction. Prognosticators of treatment failure of UPJ are a stricture length >2 cm, <20% split renal function, the presence of either anterior or posterior crossing vessels, and massive hydronephrosis.

**Ureteral Strictures**

Cold-knife, electrocautery, or laser incision of a stenosis into the periureteral fat should be followed by stenting of the ureter for 4–6 weeks according to Davis’ principle of intubated ureterotomy. The intraoperative use of an endoscopic ultrasound probe can provide important information about the exact stricture location and its relation to adjacent structures, and thus help to direct the incision.

Best results are obtained in ureteral strictures that are a complication of surgery for benign disorders if treatment is established early. Proximal and distal ureteral strictures respond better to endoureterotomy than midureteral strictures. However, long-term results of this technique as compared with open surgical repair remain to be deter-
mended. Long-standing strictures or strictures due to external ureteral compression, radiation damage, or ischemic ureteral necrosis after surgery are not satisfactorily treated by internal ureterotomy. Acucise treatment of ureteral strictures should be reserved for short strictures (<1.5 cm), in a good functioning kidney.

**Ureteral Tumors**

Endoscopic electroresection and laser coagulation of ureteral tumors are the ureterorenoscopic variants of percutaneous endoscopic treatment of renal pelvis tumors (see the section: Endoscopic Intrarenal Instrumentation, Renal Pelvis Tumor). Conservative endoscopic treatment of urothelial tumors of the upper urinary tract is still reserved for small tumors in solitary kidneys, bilateral disease, or chronic renal failure. Ureterorenoscopy offers an adequate surveillance method in cases of conservative management allowing for simultaneous biopsy. Although rare, benign fibroepithelioma of the ureter (Figure 8–17) is sufficiently treated by ureterorenoscopic techniques. The same precautions and limitations apply for endoscopic treatment of urothelial cancer of the ureter as listed previously for percutaneous endoscopic treatment of renal pelvis tumors.

**REFERENCES**

**Percutaneous Puncture & Catheter Placement**


**Antegrade Pressure/Perfusion Studies**


**Figure 8–17.** Intravenous pyelogram revealing a fibroepithelioma in the midleft ureter (arrow) without urinary obstruction.
Percutaneous Renal Stone Treatment


Percutaneous Endoscopic Surgery


**Percutaneous Aspiration & Biopsy**


**Stone Basketing, Ureterorenoscopy**


Laparoscopy now plays a prominent role in urology. Current residents finish their training with considerable exposure to the techniques, and a plethora of courses educate physicians already in practice. The technology continues to improve. In particular, hand-assisted techniques have facilitated the adoption of laparoscopy and robotic assistance is gaining great popularity.

This chapter is an overview of urologic laparoscopy, including the physiology of laparoscopy, laparoscopic instrumentation and techniques, and a review of urologic laparoscopic procedures. The reader is referred to the laparoscopic textbooks listed in the reference list for additional information.

PHYSIOLOGY OF LAPAROSCOPY

During laparoscopy with pneumoperitoneum, the patient is exposed to unusual physiologic challenges. Although most challenges are met successfully with minor modifications of anesthetic management, the laparoscopic surgeon must be aware of them to prevent and manage complications.

Physiology: Cardiovascular

As intra-abdominal pressure increases with pneumoperitoneum, the systemic vascular resistance increases and venous return decreases. The degree of intra-abdominal pressure and the circulating blood volume determine the cardiovascular effects of pneumoperitoneum. A small increase in intra-abdominal pressure augments venous return and cardiac output. As intra-abdominal pressure rises above a certain point, the increase in resistance exceeds the increase in pressure and venous return (and therefore cardiac output) falls (Figure 9–1). This transition point occurs at a low intra-abdominal pressure in the hypovolemic state and at a greater pressure in the normovolemic and hypervolemic state. Given normovolemia, an intra-abdominal pressure of 15 mm Hg is associated with tolerable reduction of cardiac output.

The absorption of insufflated carbon dioxide \((\text{CO}_2)\) has several cardiovascular effects. The direct ones are primarily inhibitory, but \(\text{CO}_2\) also stimulates the sympathetic nervous system. If acidosis is allowed to develop, parasympathetic effects may be enhanced as well. Moderate hypercapnia produces an increase in cardiac output and blood pressure and a decrease in systemic vascular resistance, which counteract the mechanical effects of pneumoperitoneum.

Overall, an intra-abdominal pressure of 15 mm Hg and moderate hypercapnia in healthy patients produce a hyperdynamic state (increased central venous pressure, systemic vascular resistance, heart rate, and blood pressure) without significant alteration of cardiac output.

Physiological Complications: Cardiovascular

The cardiovascular complications of laparoscopy include tension pneumoperitoneum, cardiac dysrhythmias, fluid overload, and venous thrombosis.

When the intra-abdominal pressure is excessive, usually \(>40\) mm Hg, the rise in vascular resistance becomes overwhelming and "tension pneumoperitoneum" can occur. Venous return, cardiac output, and blood pressure drop precipitously. Volume status must be optimized to prevent tension pneumoperitoneum at even lower pressures. Brief periods of elevated intra-abdominal pressure during laparoscopy usually are tolerated well, but generally the pressure should be kept below 15–20 mm Hg. Whenever hemodynamic compromise due to excessive intra-abdominal pressure is suspected, immediate desufflation will quickly improve the situation.

Tachycardia and ventricular extrasystoles due to \(\text{CO}_2\) are usually benign, but fatal dysrhythmias can occur with very high arterial partial pressure of \(\text{CO}_2\) (\(\text{PaCO}_2\)). Avoidance of hypercapnia will prevent tachydysrhythmias. Since hypercapnia may also potentiate parasympathetic actions, vagal stimulation by peritoneal manipulation or distention during \(\text{CO}_2\) laparoscopy can occasionally produce bradydysrhythmias.

With decreased insensible fluid losses and urine output during laparoscopy there is a predisposition to volume overloading. The volume status of the patient should be optimized before insufflation, and then intraoperative fluid administration should be limited to appropriate replacement for blood loss plus a maintenance rate of 5 mL/kg/hr.

The increased abdominal pressure during laparoscopy restricts lower extremity venous return. Mechanical pressure forces blood out of the splanchnic circulation into the
lower extremities, where venous stasis during laparoscopy can be demonstrated with Doppler flow studies. Prophylaxis for venous thrombosis is recommended for major laparoscopic procedures.

**Physiology: Pulmonary, Acid–Base, & Insufflant-Related**

Increased intra-abdominal pressure and volume elevate the diaphragm, which reduces lung capacity and compliance. Gases with high tissue permeance are absorbed readily, and CO₂ has high tissue permeance. When instilled into the peritoneal cavity, CO₂ rapidly diffuses into the bloodstream. The amount of CO₂ absorbed from the peritoneal cavity during intraperitoneal CO₂ laparoscopy at typical pressures is equivalent to adding 5–25% to the body’s baseline production of CO₂. Clinical studies have suggested that subcutaneous emphysema, elevated intra-abdominal pressure, extraperitoneal insufflation, and increased duration of insufflation all increase the rate of CO₂ absorption.

**Physiological Complications: Pulmonary, Acid–Base, & Insufflant-Related**

The pulmonary, acid-base, and insufflant-related complications of laparoscopy include hypercapnia, intra-abdominal explosion, acidosis, extraperitoneal gas collections, and venous gas embolism.

Hypercapnia, an excess of CO₂ in the bloodstream, occurs when CO₂ production and absorption exceed its elimination. Moderate hypercapnia is stimulatory overall, but if PaCO₂ exceeds 60 mm Hg, cardiodepressive effects predominate. Cardiovascular collapse, severe acidosis, and fatal dysrhythmias can occur. Increasing ventilation rates and tidal volumes usually will elevate CO₂ elimination adequately. PaCO₂ is estimated intraoperatively by the capnographic measurement of the partial pressure of end-tidal CO₂ (P[et]CO₂), which generally is 3–5 mm Hg lower than PaCO₂ during general anesthesia. During prolonged operations or in patients with pulmonary disease the gradient may widen unpredictably, and arterial blood gases should be obtained for accurate monitoring.

Additional measures to prevent hypercapnia include reduction of intra-abdominal pressure (which both decreases the CO₂ absorption and allows more effective CO₂ elimination) and use of alternative gases for insufflation. CO₂ is the most popular gas for insufflation because of its rapid absorption (which offsets some of the hemodynamic burden imposed by the pneumoperitoneum and minimizes the effect of venous gas embolism) and its inability to support combustion. Since excessive levels of hypercapnia are dangerous, however, alternative gases for insufflation have been considered. Few have met with favor, although helium has been used to avoid hypercapnia in selected patients. Unfortunately, helium can exacerbate the clinical effects of a venous gas embolism (see below).

Laparoscopy with CO₂ insufflation causes a mild respiratory acidosis due to the absorption of CO₂. With gas insufflation pressures >20 mm Hg, a metabolic acidosis can also develop, likely related to retained acids from the decreased urine output. At typical pneumoperitoneum pressures, this usually is not a clinical problem.

Gases insufflated into the peritoneal cavity may leak into several extraperitoneal tissue planes or spaces. Subcutaneous emphysema is the most common site of extraperitoneal gas. Although generally innocuous, it may increase the risk of hypercapnia. Gas that is insufflated inadvertently into the properitoneal space or omentum might interfere with visualization during intraperitoneal laparoscopy. Pneumopericardium, pneumomediastinum, and pneumothorax can inhibit cardiac filling, limit lung excursion, or both. A CO₂ pneumothorax will usually resolve spontaneously, but thoracostomy should be performed for a large or symptomatic pneumothorax.

Venous gas embolism (VGE) is the passage of gas bubbles through the venous system into the heart and pulmonary circulation. When clinically significant, right heart
outflow is impeded, producing hypoxemia, hypercapnia, and depressed cardiac output. Clinically detectable VGE may be noted in almost 1% of laparoscopic cases when careful surveillance is used. Many VGEs during laparoscopy have been fatal. VGE is indicated by hypoxemia, evidence of pulmonary edema, increased airway pressure, hypotension, jugular venous distention, facial plethora, dysrhythmias, and a mill wheel murmur. The capnometer will register a sudden fall in P(et)CO2 if the CO2 embolus is large and an abrupt but transient increase if it is small. Swift response is required, including immediate desufflation, rapid ventilation with 100% oxygen, steep head-down tilt with the right side up, and general resuscitative maneuvers. The type of the gas comprising the embolus determines in part the outcome. Helium is less soluble in blood than CO2. This fact argues against its use for initial insufflation (Figure 9–2), but helium insufflation (to avoid hypercapnia) after the pneumoperitoneum has been safely created with CO2 is safer, since VGE rarely occurs beyond the first few minutes of insufflation.

**LAPAROSCOPIC INSTRUMENTATION & BASIC TECHNIQUES**

**Preoperative Preparation**

The considerations made in the selection of patients for laparoscopy are more stringent than for open surgery. Although any operation in a patient with obesity, previous abdominal surgery, and abnormal anatomy is more difficult than in a patient lacking these features, laparoscopy is rendered relatively more difficult by these factors than is open surgery. In addition, the physiological considerations discussed above suggest that open surgery may be favored over laparoscopy in patients with severe pulmonary disease or congestive heart failure. The patient being offered laparoscopic intervention should be fully informed of the risks and benefits, most appropriately in the context of a comparison to the spectrum of risks and benefits of open surgery for the particular procedure. It is important to inform the patient of the surgeon’s experience with the particular laparoscopic procedure, and that conversion to open surgery may be required. For transperitoneal laparoscopic surgery without intended bowel resection, patient preparation with a clear liquid diet and oral magnesium citrate on the preoperative day is adequate. For retroperitoneoscopic surgery, bowel preparation is not necessary.

A logical operating room setup is required. The primary laparoscopic cart (see below) is positioned opposite the surgeon. For an upper abdominal or retroperitoneal procedure, a secondary monitor for nursing staff or assistants placed on the opposite side of the patient is useful, but for a pelvic procedure only the primary laparoscopic tower at the patient’s feet is required. Operating rooms with booms from the ceiling that hold the primary equipment minimize clutter and reduce setup time (Figure 9–3). All equipment should be checked for function preoperatively or during setup. Adequate pressure in the CO2 tank should be verified, and a second tank should be available.

**Figure 9–2.** Arterial tracing after rapid intravenous injection of 7.5 cc/kg CO2 (top) and helium (bottom) in a dog. There is recovery within 1 minute after the CO2 injection but complete cardiovascular collapse after helium injection. (Borrowed with permission from Wolf JS Jr, Carrier S, Stoller ML: Gas embolism: helium is more lethal than carbon dioxide. J Laparoendoscopic Surg 1994;4:173.)
neal procedures into the retroperitoneum, the patient is placed in partial right flank position (45°) without flexion of the table. Subsequent rotation of the table can provide near supine or near flank position. Direct retroperitoneoscopic procedures benefit from full flank position with table flexion.

**Entry: Obtaining Pneumoperitoneum**

The 2 most common methods of insufflation to create pneumoperitoneum are the closed (Veress needle) and open (Hassan cannula) techniques. Either is acceptable as the first choice, but if the Veress needle is the surgeon’s typical first choice then the open technique should be learned as well, since sometimes the former approach is contraindicated or fails.

In a previously operated-on abdomen, the Veress needle should be placed away from prior incisions. The Veress needle has a solid spring-loaded stylet that retracts back only under pressure from firm tissue (ie, fascia) to expose the sharp cannula; once the tip is free in the intraperitoneal space the stylet springs forward and protects against visceral injury. The needle is usually placed at the chosen location of the first port. It is inserted almost perpendicular to the abdominal wall, tilting slightly away from the large midline vessels. An exception is when the needle is being placed at the umbilical location in thin individuals, in which case it needs to be angled caudal almost 45°. There are usually 2 “pops” of resistance as the fascial layer and then the peritoneal membrane are penetrated (Figure 9–4).

Once the needle is inserted, a series of maneuvers are performed before insufflation. First, attach a 10-cc syringe half-filled with saline and aspirate back. There should be no aspirated gas or liquid. Next, inject the saline in through the needle and attempt to aspirate it back. It should flow in easily and not return on aspiration. Finally, the saline in the hub of the Veress should drop rapidly into the abdomen. These maneuvers assess for the possibility that the needle tip is in a luminal structure (bowel, bladder, blood vessel), but will all be “normal” if the needle is preperitoneal—which is the most common erroneous placement (Figure 9–5). This possibility is assessed by the final test, the “opening pressure.” Gas insufflation is commenced through the needle held carefully in place. The pressure should not rise above 8 mm Hg within the first 1/2 L of gas, or if it does it should only be momentary and quickly correctable by a twist, slight withdrawal, and tilting up of the needle (which will free the needle tip from omental or mesenteric fat). If these conditions are met,
insufflation is continued. If these conditions are not met, then disconnect the insufflation tubing, allow the gas to escape, and withdraw the needle. Persistence when the pressure is too high risks extensive extraperitoneal gas collection that will interfere with surgery. We usually will make 3 attempts with the Veress needle, and if safe insufflation is not achieved, an alternative method is chosen.

The most common non-Veress entry method, the Hassan technique, is placement of a laparoscopic port through a small incision made into the intraperitoneal space under direct vision. This technique is used if the Veress needle fails or is contraindicated, but some surgeons use it in all cases. A 1.5- to 3-cm incision is made at a chosen site and deepened down to the peritoneal membrane, which is opened under direct vision. Stay sutures are placed, and the Hassan port (10/12 cannula containing a blunt obturator, with a conical adjustable sleeve, Figure 9–6) is inserted. The sutures are tied down tightly on the arms of the device, such that the conical sleeve is cinched down against the fascia. Gas is then insufflated. One modification of this technique is the use of a port with a retaining balloon that inflates intra-abdominally and is held up tight against the fascia by an adjustable retaining ring. Another modification is to make just a small nick in the fascia and insert a blunt port that dilates rather than cuts the fascia (see below). Finally, a visualizing trocar (see below) can be used.

**Entry: Port Placement**

Selection of port sites determines access to the operative field. The general scheme is to surround the site with the necessary number of instruments placed widely enough apart that they do not “sword fight” in the abdomen and with the laparoscope situated so that a good visual angle can be attained.

There are a variety of ports available, with the standard sizes being 5, 10, and 12 mm. “Needlescopic” ports (3 mm) as well as extra-wide ports (18 and 30 mm) are available for specialized use. Ports can be completely disposable, completely reusable, or “re-posable” ports, which contain both reusable and disposable components. Many disposable ports have sharp tips that are shielded to prevent viscous injury (Figure 9–7). Reusable ports most commonly have a metal trocar with a metal or plastic cannula (or

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*Figure 9–4. Insertion of the Veress needle into the peritoneal cavity.*

*Figure 9–5. Preperitoneal location of the tip of Veress needle.*

*Figure 9–6. Hassan cannula.*
both). Finally, there are the “re-posable” ports. The cannulae are reusable and the trocars, blades, and diaphragms are mixed disposable and reusable.

Trocars can be cutting or noncutting. Disposable sharp trocars are shielded. Some reusable trocars have non-shielded sharp tips. The Hassan-type ports employ blunt tips. There are ports with clear plastic tips (Figure 9–8) that allow visualization with the laparoscope as the port is being inserted. The Step system (AutoSuture Co, Norwalk, CT, Figure 9–9) employs an expandable sheath that is inserted with a Veress needle. The needle is removed and a dilating trocar is used to insert the port. This and other ports that dilate or screw into the fascia are increasing in popularity because the fascia defect is smaller and there is less risk of visceral injury than with sharp-tipped ports.

If the Veress needle has been used for abdominal entry, the next step is to place the first port. This is usually the port for the laparoscope, inserted at the site of Veress needle placement. Use a firm but controlled motion to drive the port into the abdomen. To avoid inadvertent removal of the port during the procedure, anchor to the skin with suture. Alternatively, special sheaths will hold the port firmly into the fascia. Insert the laparoscope and inspect the intraperitoneal contents. Subsequent ports are placed under visual control through the laparoscope.

**Entry: Extraperitoneal Approach**

The retroperitoneal route is well suited to many urologic procedures because the organ of interest is in the retroperitoneum. For flank retroperitoneoscopic surgery, a 2-cm incision is made below the tip of the 12th rib and taken down through the lumbodorsal fascia under direct vision. The retroperitoneal space is dilated with a finger, and then a dilating balloon (commercially available, Figure 9–10, or self-made) is used to dilate the working space. A self-retaining port is inserted at this location. Additional ports are placed at the base of the 12th rib, at the tip of the 11th rib, and in the mid axillary line above the iliac crest. For pelvic extraperitoneal surgery, the dilating balloon is placed directly into an infraumbilical incision and then slid down the inside of the rectus sheath to the pubic bone, where it is used to enlarge the preperitoneal space.

**Hand Assistance**

Hand-assisted laparoscopic surgery (HALS) entails the insertion of a hand into the laparoscopic field while main-
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The hand-assistance devices employ a compressive mechanism to affix the device to the abdomen and a locking or compressive mechanism (compressive balloon, gel, or disk) to prevent leakage of pneumoperitoneum around the intra-abdominal hand (Figure 9–11). HALS is predominantly used for transperitoneal laparoscopic procedures. The intra-abdominal hand is used for dissection, tissue identification, retraction, and control of injuries. The benefits of HALS include shorter operative times than for similar standard laparoscopic transperitoneal procedures, ease of learning by inexperienced surgeons, and enhanced ability to manage difficult surgical situations. Disadvantages of HALS include problems with the devices, such as gas leakage or interference with port placement, physical strain on the hand, interference of the hand in the operative field, and the need for a larger incision than might otherwise be required for a standard laparoscopic procedure.

Some urologic surgeons use HALS almost exclusively, and some never use it. Given the spectrum of its advantages and disadvantages, HALS is probably best used selectively. The most effective uses for HALS include cases where intact specimen removal is required, when difficult dissection is anticipated, early in a surgeon’s experience, or for large specimens. Most simple and radical nephrectomies, as well as pyeloplasties, adrenalectomies, renal cyst resections, and pelvic procedures, can be performed well by experienced surgeons without HALS.

**Laparoscopic Video Instrumentation & Cart**

The standard adult laparoscope is 10 mm in diameter, although 5-mm laparoscopes are improving in light transmission capability. For pediatric surgery, a 5-mm laparoscope typically is used, but 2-mm laparoscopes are also available. Most laparoscopes employ a 0° or 30° lens, with the latter providing more viewing angles. Some laparoscopes can have the camera chip on the tip of the instrument, which removes an interface and enhances resolution. One or two large monitors, a strong light source with cables in good condition, and the digital image converter (camera box) complete the video apparatus. Still-image and video-capture devices provide documentation. Other equipment on the laparoscopic cart includes the high-flow insufflator and additional energy sources (see below).

**Laparoscopic Instrumentation**

Laparoscopic operating instruments include ones for grasping and dissection, cutting, hemostasis, retraction, irrigation and aspiration, suturing, clipping or stapling, specimen entrapping, morcellating, and intraoperative imaging. Many are available in both disposable and reusable versions. Most standard instruments are 5 or 10 mm in diameter and 35 cm in length, but there are “needlescopic” 2-mm instruments as well as larger (15–18 mm) devices for particular uses. A thorough understanding of each instrument, especially the clip applicers and staplers, is necessary for their safe and effective use.

**Exiting the Abdomen**

After a laparoscopic procedure is completed, the operative field is inspected at 5-mm Hg pressure to allow exposure of any disrupted small venules that may have been compressed by the working pneumoperitoneum pressure. The ports are removed under vision so that any bleeding from the abdominal wall can be detected. All port sites 10 mm or larger are closed in the fascial layer with suture. The Carter-Thomason Needle-Suture Passer (Figure 9–12) (Inlet Medical Inc, Eden Prairie, MN) or similar devices simplify the often-difficult task of fascial closure. In children, even 5-mm port sites should be closed. Because the CO₂ is irritating to the abdomen and may be responsible for a considerable portion of pain after laparoscopy, attempts should be made to desufflate all gas before removing the last port. Close the port incisions with a subcuticular stitch or wound glue and apply sterile tape.

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**Figure 9–10.** Trocar-mounted preperitoneal distention balloon. (Origin Medsystems, Menlo Park, CA.)

**Figure 9–11.** Surgeon’s hand inside Omniport. (Weck Closure, Raleigh, NC.)
Postoperative Care

For urologic laparoscopic procedures that do not involve bowel resection, hospital stay can be limited to 1 or 2 days in most patients with aggressive postoperative management. Offer parenteral narcotics the afternoon and evening after the procedure. Unless contraindicated, use intravenous ketorolac around the clock (not as needed) for the first 24 hour, including a dose at skin closure in the operating room. Start a clear liquid diet postoperatively, but if nausea develops then hold off on intake for a few hours before trying again. On the morning of postoperative day 1, advance to a regular diet as tolerated and switch from parenteral to oral pain medication. Get the patient up to a chair on the day of the operation, and start ambulation in the hallway the next day.

SPECIFIC PROCEDURES

Pelvic Lymph Node Dissection

Pelvic lymph node dissection (PLND) was the first adult laparoscopic urologic procedure to gain widespread interest and acceptance. The technique was initially described in 1990, and most cases in the early series were performed before the generalized use of prostate-specific antigen testing, when PLND had positive results in up to one-thirds of patients. Due to stage migration, modern series of prostate cancer patients reveal that <10% of patients with localized disease will have nodes with a positive result. This has greatly decreased the number of stand-alone PLNDs being performed. The interest in learning and Perfecting laparoscopic PLND therefore has declined. The interest may once again increase in the future, however, as more laparoscopic radical prostatectomies and laparoscopic cystectomies are performed.

Laparoscopic PLND is indicated for assessment of pelvic nodal metastasis from prostate, bladder, or other pelvic malignancies. In prostate cancer patients, PLND is generally reserved for those with prostate-specific antigen level >20, those with high-grade disease (Gleason ≥8), those with clinical stage T3 or T4 disease, or those with suspicious nodal enlargement as seen on computed tomography scan, before definitive therapy is initiated (ie, radiation, cryotherapy, hormonal ablation, or prostatectomy). There are no contraindications specific to PLND other than those general to laparoscopy. Prior laparoscopic hernia repair can result in severe adhesions, limiting both open and laparoscopic pelvic procedures.

The patient is positioned supine, with Trendelenburg tilt, as would be used for open PLND. Laparoscopic PLND can be performed transperitoneal or extraperitoneal (Figure 9–13). Initial access is typically through an umbilical port followed by 2–4 additional ports in a diamond or fan-shaped pattern (Figure 9–14). Key land-
marks are the obliterated umbilical ligament, gonadal vessels, and internal inguinal ring (Figure 9–15), with the limits of the nodal dissection being the iliac bifurcation, the pubis, the obturator nerve, the pelvic side wall, and the medial umbilical ligament. The distal and proximal lymphatics are typically clipped to decrease lymphatic leakage. Dissection for bladder, penile, or urethral cancer usually involves a more extended field, bounded by the common iliac artery, the genitofemoral nerve, and the bladder.

Complications are similar with laparoscopic and open PLND. Specifically, injuries to the obturator nerve, iliac and epigastric vessels, and ureter have been reported but are uncommon. Other potential complications include lymphocele, bowel or bladder injury, subcutaneous emphysema, and scrotal swelling. Comparison studies of open versus laparoscopic PLND reveal a similar number of nodes retrieved. Patients treated with laparoscopic PLND enjoy an earlier time to oral intake, less postoperative pain, a shorter length of stay, and decreased recovery time. The disadvantages include prolonged operative time and higher complication rates in some series, especially early in one’s experience.

Varicocelectomy

As many as 15% of adolescent males will have clinically evident varicoceles. Most of these, however, do not need treatment. Varicocelectomy is reserved for those with asymmetrical testicular growth, infertility, or pain. The laparoscopic technique typically involves 3 ports. The initial camera port is placed at the umbilicus and additional ports are placed lateral to the rectus muscle. The gonadal veins are flattened by the pneumoperitoneum and therefore the artery appears more tubular than surrounding structures. After incising through the posterior peritoneum (Figure 9–16), gonadal veins are clipped and transected. Some series report meticulous dissection and salvage of lymphatic and arterial vessels; others have promoted transection of all vessels, including the artery. One study documents that testicular catch-up growth after artery-sparing varicocelectomy is the same as with non-artery-sparing procedures, therefore questioning the need to salvage the gonadal artery. Outcomes after laparoscopic varicocelectomy are comparable to those with open techniques. Recurrence rates are 2–15% depending on technique and follow-up. Complications include hydrocele formation.
and injury to the vas deferens as well as those generally associated with laparoscopic surgery. Unilateral laparoscopic varicocelectomy is rarely performed today due to the refinements in the subinguinal open technique.

**Orchidopexy**

Laparoscopy has been judged by many pediatric urologists to be the gold-standard diagnostic tool for nonpalpable testes. Although somewhat controversial, laparoscopic orchidopexy has been shown to be comparable to if not better than open orchidopexy in a recent multi-institutional analysis. A laparoscopic approach may be used for intersex evaluations. There are no specific contraindications for laparoscopic orchidopexy other than those for general laparoscopic procedures.

Positioning is usually supine with the legs frog-legged. A small laparoscope is inserted through a periumbilical site. The first step involves identifying the inguinal ring with the corresponding gonadal vessels and vas deferens. If the vas and vessels are seen entering the inguinal ring, the laparoscope is removed and an open inguinal exploration is performed as would be for a palpable cryptorchid testis. If the gonadal vessels are blind ending, the procedure is terminated. If an unsalvageable testis is identified, laparo-

scopic orchietomy can be performed. If a salvageable testis is identified intra-abdominally, working ports are placed and the testis is freed from its nonvital attachments (Figure 9–17). If sufficient cord length can be gained, a scrotal incision is made and the testis is pexed. If there is insufficient cord length, a 1- or 2-stage Fowler-Stephens procedure may be performed. A multi-institutional review found decreased testicular atrophy and a higher rate of postoperative scrotal testicular positioning after the 2-stage procedure. About 15 complications were noted in 310 procedures, with an overall success rate of 93%.

**Renal Cyst Decortication**

Although simple renal cysts are common, they rarely require surgical intervention. Occasionally, pain, infection,
or secondary obstruction can develop, necessitating laparoscopic renal cyst decortication. Infrequently, a cystic mass may need to be explored for diagnostic purposes. Decortication can also be performed for pain relief in patients with polycystic kidney disease who still have good renal function. There are no specific contraindications other than that for general laparoscopic techniques.

If there is suspicion of a connection to the collecting system, a retrograde ureteral catheter may be placed. A 3-port approach is often sufficient. The colon is reflected medially to expose the kidney, and Gerota’s fascia is incised to expose the cyst(s). Laparoscopic ultrasound may be used if necessary to help localize the lesions. The roof of the cyst typically appears blue before excision (Figure 9–18). The roof is excised, minimizing the removal of the surrounding renal parenchyma that can result in troublesome hemorrhage. A biopsy specimen of the base of the cyst may be taken and then coagulated with the argon beam coagulator or electrocautery. Retained perirenal fat may be fixed to the cyst base. Complications specific to renal cyst decortication include worsening of patient’s symptoms, infection, or urine leak.

**Nephrectomy**

Widespread interest in urologic laparoscopy was rekindled when the first total laparoscopic nephrectomy was performed in 1990. Laparoscopic nephrectomy has become the most common laparoscopic urologic procedure. First popularized for the removal of benign, nonfunctioning renal units, laparoscopy has now become a well-accepted technique for renal malignancies. Indications for laparoscopic nephrectomy include nonfunctioning kidneys, chronic infections, symptomatic polycystic kidney disease, and tumors. Initially, large tumor size was considered a contraindication for laparoscopic nephrectomy, but with experience tumor size has become less of an issue. The only limitation is the expertise of the laparoscopic urologist. Laparoscopic excision of a level 1 venous thrombus has been reported. Relative contraindications include large venous thrombus, adjacent organ involvement, and widespread metastatic disease.

**A. APPROACHES**

Transperitoneal or retroperitoneal approaches are used, depending on surgeon preference. Each technique has advantages and disadvantages. The transperitoneal route provides a capacious working space and allows direct visualization of familiar intraperitoneal anatomy compared to the retroperitoneal technique. Transperitoneal access may be difficult, however, in patients who have had extensive abdominal surgery. The retroperitoneal approach allows easier and more rapid access to the renal artery from the posterior aspect and avoidance of intra-abdominal organs and adhesions. The disadvantage of the retroperitoneal technique is the limited working space. Comparisons of the 2 techniques have shown no difference in operative times, cost, length of stay, or postoperative convalescence. The approach therefore is dictated by the familiarity of the surgeon and the patient’s condition.

Hand-assisted laparoscopic nephrectomy has been accepted as the technique of choice by many urologists. This technique has the advantage of using the surgeon’s hand to retract, dissect, and guide the laparoscopic tools during laparoscopic nephrectomy. A small (approximately 7–8 cm) incision is required for the hand-assistance device, which maintains pneumoperitoneum. Numerous retrospective comparison studies between pure laparoscopic and hand-assisted laparoscopic nephrectomy show similar postoperative pain, time to oral intake, and convalescence. Operative times for hand-assisted laparoscopic nephrectomy are shorter than laparoscopic nephrectomy in some series.

**B. TECHNIQUES**

Positioning for laparoscopic nephrectomy is similar to that of flank approaches for open renal surgery. If a Pfannenstiel incision is to be used for specimen extraction, it is best marked in the supine position because its location often changes when the patient is positioned and the abdomen insufflated. Sequential compression stockings are placed. After initiation of general anesthesia, a Foley catheter and oral/nasogastric tube are placed and the patient is turned into a (modified) lateral decubitus position (45°–90° with lesion side up, depending on the specific approach and surgeon preference). The lower knee is flexed and the upper leg is straight (figure of 4 position). All pressure points are adequately padded. The table may or may not require flexion depending on the specific approach and surgeon preference. The ipsilateral arm is flexed and secured with a pillow or arm supporter. The surgeon and assistant both stand toward the patient’s anterior body wall for the transperitoneal approach.
The techniques for a laparoscopic nephrectomy will be discussed in detail as a template for other common laparoscopic urologic procedures. Many variations on port placement have been used. For transperitoneal laparoscopic nephrectomy, a 4-port “L”-shaped configuration for left nephrectomy and a reversed “L”-shaped pattern for right nephrectomy is commonly used. The initial trocar is placed 2 fingerbreadths below the costal margin at the lateral edge of the rectus muscle and the abdomen is insufflated initially to 15–18 mm Hg. Intra-abdominal organs are inspected for inadvertent injury. Another port is placed in the mid-axillary line 2 fingerbreadths above the iliac crest. One to two additional ports are placed along the lateral edge of the rectus muscle including one able to accept the endoscopic stapler if one anticipates its use. Depending on the patient’s body habitus and relative position of the kidney and spleen/liver, other configurations may be chosen (Figure 9–19).

If hand assistance is chosen, port placement is altered to allow room for the hand-assistance device template. For left-sided lesions, the hand-assistance device is typically placed in the midline and may incorporate the umbilicus. For obese patients, the device may be placed paramedian, closer to the pathology. For right-sided lesions, the device may be placed in the midline or alternatively in the right lower quadrant using a muscle-splitting diagonal incision (Gibson). Placement of the hand-assistance device is dependent on the surgeon’s arm length, optimal position of the surgeon’s nondominant hand, and the patient’s body habitus. Some devices are designed for placement before insufflation, whereas others may be added later in the procedure.

For left-sided lesions, the left colon is mobilized from the splenic flexure toward the iliac vessels, leaving the anterior fascicle of Gerota’s fascia intact. The lienophrenic liga-

![Image](42x68 to 240x226)

**Figure 9–19.** One of the possible port configurations for right transperitoneal laparoscopic nephrectomy.
side of the patient. After an incision is made over Petit’s triangle, blunt dissection through the lumbodorsal fascia is performed and a plane is developed over the psoas muscle. A dissecting balloon trocar is placed into this space. There are commercially available dissecting balloon trocars for this purpose, or a “homemade” version can be constructed by attaching a glove to a catheter. The commercially available dissecting trocar has the added advantage of allowing the laparoscopist to visually inspect the dissecting process and ensure proper location. This expansion typically requires 500–1000 cc of balloon inflation. The renal artery, ureter, or both may be seen through the balloon’s wall. Pneumoretroperitoneum is then created and 2–3 additional port sites are placed according to surgeon preference (Figure 9–20). The kidney is retracted anteriorly to allow direct access to the hilum. Blunt dissection easily exposes the artery, which is clipped and transected. Anterior to the arterial stump, the vein is identified, dissected, and controlled in a similar fashion as previously described. After hilar control, the ureter is identified and transected. Further dissection is similar to that of transperitoneal techniques. Many laparoscopists remove the specimen intact when performing retroperitoneoscopic nephrectomy due to the limited working space. If morcellation is preferred, specimen-bag entrapment may require incising the peritoneum if the specimen is large. The final steps of ensuring hemostasis under low-pressure insufflation and inspecting the port sites remain the same.

The technique for hand-assisted laparoscopic nephrectomy is similar to that for transperitoneal laparoscopic nephrectomy. The hand-assistance device incision is made according to the manufacturer’s directions, and 2–3 subsequent ports are placed according to surgeon preference. Dissection is similar to the transperitoneal description above. Lateral renal attachments can be transected earlier because the hand can provide countertraction, facilitating hilar dissection. Specimen extraction is rapidly performed through the hand-assisted incision.

**C. COMPLICATIONS**

The complication rate from laparoscopic nephrectomy ranges from 8% to 17%. Open conversion is required in 1.7–4% of cases. Minor complications include ileus, mild hemorrhage, urinary tract infection, hernia, and wound infection. Major complications include pulmonary embolus; pneumothorax; injury to the duodenum, spleen, liver, or pancreas; and major hemorrhage from the aorta, IVC, or iliac, gonadal, lumbar, or renal vessels. Technical complications are more common during the first 30–50 cases of a surgeon’s experience.

**D. OUTCOMES**

Numerous studies have attempted to compare laparoscopic nephrectomy with open nephrectomy. Overall, laparoscopic nephrectomy offers shorter length of hospital stay, earlier time to oral intake, and less pain medication requirement with similar complication and cancer-control rates. Numerous studies have documented longer operative times for laparoscopic approaches; however, with surgeon experience, operative times may even be shorter than with open techniques. Studies comparing different laparoscopic techniques show similar outcomes, suggesting that no one approach is uniformly superior. The surgeon should be familiar with the various techniques so that the optimal procedure can be performed.

**Nephroureterectomy**

Open nephroureterectomy has been the gold-standard treatment for upper tract urothelial carcinoma. This procedure is performed through a long curved flank incision or 2 separate incisions, leading to significant postoperative morbidity. Laparoscopic nephroureterectomy (LNUx) incorporates the benefits of cancer control with less postoperative pain and earlier return to normal activity. Indications include urothelial carcinoma of the renal pelvis (T1–
3) and ureter (T1–2). LNUs can be employed for non-functioning kidneys due to reflux nephropathy, although control of the intramural ureter is not as critical as in cases of carcinoma. Transperitoneal, retroperitoneal, and hand-assisted techniques have been described.

Management of the distal ureter remains controversial. Initially, the distal ureter was transurethrally excised from the bladder using a resectoscope and delivered into the retroperitoneum. This procedure resulted in numerous reports of local recurrence. Other techniques to manage the distal ureter have since been described. Some perform a laparoscopic nephrectomy and proceed with a Pfannenstiel incision for open bladder cuff excision and specimen extraction. Alternatively, some authors have described transurethral mobilization and cautery of the distal ureter in hopes of sealing the ureter. Other techniques involve 1- or 2-port transvesical dissection of the distal ureter (Figure 9–21) with (2-port) or without (1-port) transvesical closure of the ureteral orifice. Others have described using an endoscopic stapler or clips to secure the bladder cuff with subsequent transurethral cautery of the ureteral orifice. Cancer-control rates appear adequate and complication rates are similar to those for laparoscopic radical nephrectomy. However, only short-term follow-up data are available.

Partial Nephrectomy

Open partial nephrectomy is performed in patients with renal lesions and risk for postoperative renal insufficiency (anatomically solitary kidney, bilateral lesions, significant preoperative renal insufficiency) or risk factors for future renal disease. Patients who are candidates for open partial nephrectomy have been treated with laparoscopic radical nephrectomy because of the decreased morbidity. Recently, reports of laparoscopic partial nephrectomy have become more common.

Transperitoneal, retroperitoneal, and hand-assisted techniques have been described. Although the overall technique is similar to that for laparoscopic radical nephrectomy, a few modifications are required. Some physicians perform preoperative ureteral catheterization to help ensure adequate collecting system closure. Laparoscopic ultrasound may be useful to delineate tumor margins and multifocality. The kidney is dissected from Gerota’s fascia and perirenal fat, except for the fatty tissue directly over the lesion. The renal artery may be occluded with a laparoscopic bulldog or, alternatively, a loop or hand can be used for localized compression. Intracorporeal cooling with ice slush has been described. Maneuvers to minimize reperfusion injury (intravenous fluids, mannitol) can be used as with open partial nephrectomy. Enucleation or wedge resection can be performed with endoscopic scissors, electrocautery (Figure 9–22), or coagulating shears. Frozen section may be used to assess tumor margins. Biopsies of the base of the resection bed are recommended by some authors. Vessels and collecting system entry sites are closed using manual intracorporeal suturing techniques or with a variety of tissue glues. The overlying parenchyma is frequently coagulated using the laparoscopic argon beam coagulator. Tissue glues and bulking agents (collagen, gelatin, etc) may be placed on the resection bed to help ensure hemostasis. Bolstering sutures can be placed to compress and reconstruct the remaining renal tissue. Once reperfused, the surgical area is inspected for hemostasis. This is a procedure in evolution, and the optimal technique is yet to be determined.

Figure 9–21. Transvesical dissection of the distal ureter performed with a Colling’s knife on a resectoscope placed through a 10-mm port inserted suprapubically into the bladder. After the nephrectomy portion of the procedure and clipping of the ureter, incision (dotted black line) is gradually made around the ureteral orifice (white arrowhead) until the distal ureter can be pulled free of the bladder.

Figure 9–22. Partial nephrectomy for small lower pole tumor (white arrow), using bipolar electrocautery to maintain hemostasis in the resection bed (black arrow) without clamping of the renal artery. The intra-abdominal hand for hand assistance is visible on the left side.
Complications of laparoscopic partial nephrectomy, including delayed hemorrhage and urinary leak, are similar to those of open partial nephrectomy. Port-site recurrences have not been documented. Long-term follow-up is needed to ascertain if cancer-control rates are equivalent to those of open partial nephrectomy.

Modifications to laparoscopic partial nephrectomy include cryoablation and radiofrequency coagulation of small peripheral renal lesions. The dissection techniques are similar to that for laparoscopic partial nephrectomy. A biopsy specimen is frequently taken to document localized renal cell carcinoma. For cryoablation, the probe is passed into the tumor and the lesion is frozen to −20°C, thawed, and then refrozen to ensure adequate tissue destruction. This process can be monitored by laparoscopic ultrasound to ensure that the “ice ball” extends 1 cm beyond the tumor margin. The probe is gently released after the lesion has thawed. Pressure, fibrin sealant, and gelatinized foam help to ensure hemostasis. The radiofrequency ablative technique is similar. A percutaneous radiofrequency probe is introduced into the lesion after adequate mobilization and biopsy. The probe delivers 50–100 W of energy and is preset to the estimated tumor volume plus a margin of normal surrounding tissue. Temperature at the tip of the probe is 100°C. After treatment, the lesion may be resected and a biopsy specimen taken from the base, or the coagulated mass can be left in situ. Postresection techniques for hemostasis are similar to those for laparoscopic partial nephrectomy. Cryoablation and radiofrequency coagulation techniques may decrease blood loss and operative times compared to partial nephrectomy. Renal arterial clamping is unnecessary, thus avoiding possible ischemia and reperfusion injuries associated with traditional partial nephrectomy. Complications are similar to those of laparoscopic partial nephrectomy. Long-term data are unavailable, and close follow-up is required.

Donor Nephrectomy

Approximately 53,000 people await kidney transplantation, but only 15,000 transplants were performed in 2001, according to the United Network for Organ Sharing data registry. Of these, <6000 kidneys came from living donors. Open donor nephrectomy has many disincentives, including pain, cosmetic concerns, and a long recuperation time. Laparoscopic techniques reduce these disincentives, and data suggest that their availability may have increased the number of living renal donors at centers that have incorporated this technique. The first laparoscopic donor nephrectomy was performed in 1995. Donors are able to return to their normal responsibilities of family and work after only 1–2 days of postoperative hospital stay. Most centers begin their series with left-sided, single-artery kidneys, but multiple arteries and right-sided donation have been documented with excellent graft survival rates. The positioning and dissection technique is similar to that of laparoscopic nephrectomy, except that the ligation of the vasculature is one of the last steps. The ureter, vein, and artery are circumferentially dissected and tagged. For the pure laparoscopic technique, a premarked low transverse incision is carried down to the peritoneum. The ureter, artery, and vein are clipped and transected in sequential order. To optimize renal vein length, especially on the right side, an endoscopic stapler that lays down 3 rows of staples without cutting can be used, in place of the more commonly used stapler that lays down 6 rows of staples and cuts in between rows 3 and 4. A peritoneotomy is made and the kidney is passed to the recipient team.

Many centers use hand-assisted laparoscopic techniques for live renal donation. Proponents of hand-assisted approaches argue that the incision should be made at the beginning of the procedure and utilized for assistance in dissection. Others use the retroperitoneal approach for laparoscopic donor nephrectomy, claiming decreased operative times. Surgeon’s experience and comfort level will dictate the laparoscopic technique. Laparoscopic donor nephrectomy produces renal units that function as well as those from open donation procedures, with similar complication rates. With widespread use of laparoscopic renal donation, it is hoped that kidney supply will match the demand for these lifesaving organs.

Pyeloplasty

Laparoscopic pyeloplasty was first introduced in 1993. Since that time, it has become a favored approach to treating ureteropelvic junction obstruction by many physicians. Transperitoneal, retroperitoneal, and hand-assisted techniques have been described. Common to all approaches is the need to be well skilled in laparoscopic suturing, a difficult intracorporeal task. Port placement is similar to that for laparoscopic transperitoneal or retroperitoneal nephrectomy. Dissection and reconfiguration of the ureteropelvic junction is similar to that of open techniques and is dependent on intraoperative findings. The Anderson-Hynes dismembered pyeloplasty, Y-V plasty, Heineke-Mikulicz reconstruction, Davis intubated ureterotomy, Hellstrom vascular relocation, and tubularized flap pyeloplasty have all been described laparoscopically. The procedure has been performed in infants, children, adults, and the elderly.

Some physicians use fibrin sealant to cover the repair with only minimal suturing. The largest laparoscopic pyeloplasty series to date with 100 cases reveals a 96% obstruction-free rate with 2-years follow-up. A comparison of open versus laparoscopic pyeloplasty revealed equivalent pain relief, activity level improvement, and relief of obstruction. These outcomes are better than other minimally invasive approaches to ureteropelvic junction obstruction, including retrograde and antegrade endopyelotomy or balloon dilation. The higher success of laparoscopic techniques compared to these other procedures is based on direct laparo-
scopic visualization of the pathologic areas, thus directing an appropriate repair. Although endopyelotomy may be successful in patients with an intrinsic obstruction, delayed failure is not uncommon in those with crossing vessels (Figure 9–23). Laparoscopic pyeloplasty is associated with decreased blood loss compared to open techniques. Postoperative pain and convalescence are similar to those in endopyelotomy; however, operative times are longer. Newer techniques with fibrin sealant and fewer sutures may decrease operative times. Robotic-assisted laparoscopic approaches may facilitate complex fine suturing. Complications are similar to that for laparoscopic nephrectomy. In addition, fistula, urine leak, and infection have been reported.

Adrenalectomy

Laparoscopic adrenalectomy is the standard approach for most adrenal lesions. Indications include aldosteronomas, pheochromocytomas, Cushing’s adenomas, incidentalomas, metastatic lesions, symptomatic myelolipomas, and feminizing/virilizing tumors. Large (>6 cm), invasive carcinomas are considered by most to be the only contraindication to laparoscopic adrenalectomy. Reports of laparoscopic adrenalectomy for up to 15-cm lesions have been reported by experienced laparoscopists. A complete endocrine workup is required preoperatively to help prevent the intraoperative hemodynamic lability associated with functional adrenal lesions. Hydration and medications may be needed 2–4 weeks preoperatively in some cases (calcium channel, alpha- or beta-blockers for pheochromocytomas, spironolactone and potassium for aldosteronomas).

Transperitoneal (anterior or lateral), retroperitoneal (posterior or lateral), hand-assisted, and transthoracic laparoscopic approaches have been reported. Bilateral synchronous adrenalectomy and partial adrenalectomy have been performed. The laparoscopic approach to the adrenal gland is similar to that previously described for transperitoneal laparoscopic nephrectomy. Instead of the “L”-shaped port configuration, however, 2–3 additional ports are placed in a subcostal location (Figure 9–24). The dissection can be compared to opening a book. For left-sided lesions, the spleen is mobilized medially while the characteristic yellow adrenal tissue is mobilized to the right. Dissection continues in a counterclockwise direction. Small lesions may be difficult to identify within the overlying adipose tissue. As one mobilizes the spleen and pancreas medially, the adrenal vein will be seen entering the renal vein. One should be cautious superomedially because the inferior phrenic vein will join the adrenal vein before entering the left renal vein and must be controlled. After one ligates and transects the adrenal vein, blunt and sharp dissection will allow mobilization of the adrenal gland from the psoas muscle and superior aspect of the kidney.

On the right side, the surgical approach is again analogous to opening a book, and dissection proceeds in a clockwise direction. The triangular ligament is incised with the posterior peritoneum, allowing medial retraction of the liver and colon. This exposes the IVC, and the adrenal is mobilized to the left. The adrenal vein is identified, ligated, and transected. Caution is necessary because an anomalous adrenal vein (10% incidence) may be identified connecting to the hepatic vein.

Complication rates from laparoscopic adrenalectomy are 8–29%. Open conversion rates range from 0% to 5%. Vascular injuries (transfusion rate 3%), visceral injury (3%), heart failure (pheochromocytoma), infections (Cushing’s syndrome), pneumothorax (<1%), and deep venous thrombosis have been reported. Subclinical Cushing’s syndrome may result in an Addisonian crisis, typically within the first 10 postoperative days. Laparoscopic adrenalectomy offers significantly less postoperative pain, earlier time to oral intake, shorter length of hospital stay, and more rapid recovery when compared to open adrenalectomy.

Retroperitoneal Lymph Node Dissection

Retroperitoneal lymph node dissection (RPLND) is indicated for patients with stage I, marker-negative high-risk
nonseminomatous germ cell tumors and for residual disease in patients with stage I–IIIB nonseminomatous germ cell tumors after chemotherapy. The open version of this procedure involves a xiphoid-to-pubis midline incision and significant morbidity related to ileus, blood loss, and postoperative pain. The laparoscopic RPLND was initiated in 1992 in an effort to reduce morbidity. The dissection template is the same as in the open approach. For left-sided lesions, the para-aortic, preaortic, and retroaortic nodes are removed. The boundaries are the renal vessels, the medial edge of the aorta, and the ureter inferiorly down to the iliac vessels. For right-sided lesions, the interaortocaval, precaval, retrocaval, and preaortic nodes are removed (Figure 9–25). The boundaries are the renal vessels, the aorta, and the ureter inferiorly down to the iliac vessels. Additionally, the para-aortic nodes between the renal hilum and the inferior mesenteric artery are removed. Laparoscopic RPLND should be approached with caution in patients with bulky nodal disease.

The patient is positioned in a modified lateral decubitus fashion. The initial port is placed near the umbilicus. Three to five additional ports are placed to optimize dissection and retraction. As with open RPLND, strict hemostasis is vital to identify the planes of dissection. The dissection is similar to the previously described transperitoneal laparoscopic nephrectomy except that on the left side, the spleen and pancreas are rotated further medially and the descending colon is mobilized lower into the pelvis. This gives excellent exposure to the retroperitoneum. On the right side, the posterior peritoneum is incised under the liver to reveal the superior margin of the IVC, and the colon and duodenum are mobilized further medially to expose the retroperitoneal area of interest. A methodical approach, using the same split-and-roll technique as in open RPLND, is performed. Nerve-sparing techniques are similar to the open approach in hopes of preserving ejaculatory function.

The largest series of laparoscopic RPLND includes 125 patients. There were 2 conversions to open surgery. Other complications encountered were hemorrhage, lymphocele (7), chylous ascites (6), injury to renal and lumbar veins, and bowel injury. Postchemotherapy dissections are more difficult, with higher morbidity and open conversion rates. Follow-up studies at an average of 3–4 years show excellent relapse-free survival, comparable to that of open techniques.

Figure 9–25. Exposure of paraspinous tissue during right-sided laparoscopic retroperitoneal lymph node dissection.

Radical Prostatectomy

Interest in laparoscopic urologic surgery has intensified with successful laparoscopic techniques for radical prostatectomy. The first laparoscopic prostatectomies, described in 1992, required a mean operative time of 9.4 hours and offered no obvious advantages. The technique was modified in Europe and has become routine in some centers, with operative times similar to those for open prostatectomy. The indications for laparoscopic radical prostatectomy are similar to the open approach in patients with localized prostate cancer. Contraindications to this procedure are the same as those for general laparoscopy.

The patient is positioned in a modified lithotomy fashion with the thighs abducted for access to the perineum and the table in steep Trendelenburg position. A periumbilical insufflation port is placed with 4–5 additional ports in a fan-shaped pattern. Typically, laparoscopic dissection begins by incising the peritoneum posterior to the bladder to expose the seminal vesicles and transect the vas deferens. Denonvillier’s fascia is incised, allowing dissection anterior to the rectum in a caudal direction toward the prostatic apex. Retropubic dissection is then performed to expose the endopelvic fascia and control the dorsal vein complex. Dissection can be approached from the bladder neck distally to the apex or from the apex proximally to the bladder neck. The lateral pedicles can be controlled with electrocautery, clips, coagulating shears, or endoscopic staplers. If nerve sparing is desired, electrocautery is limited in this area. The bladder neck is inspected and any necessary reconstruction is performed. The laparoscopic approach has the advantage of performing the anastomosis under direct magnified vision. Robotic assistance may facilitate the vesicourethral anastomosis.

Although the reported data for laparoscopic radical prostatectomy are still preliminary, oncologic results seem comparable to those in open radical prostatectomy. Three-year biochemical recurrence-free survival was 91% for T2 patients and 81% for T3 patients. A retrospective comparison of open retropubic, open perineal, and laparoscopic prostatectomy showed comparable prostate-specific antigen-free survival. Positive margin, incontinence, and sexual dysfunction rates are comparable with those of open techniques. Blood loss is less than with open techniques because the pneumoperitoneum helps reduce venous hemorrhage. Conversion to open surgery is rare after the initial
learning curve. Many centers remove the urethral catheter earlier than with open techniques. The benefits of laparoscopic radical prostatectomy as compared with open surgery remain unclear, however, because open surgical prostatectomy provides excellent results. The relative benefit of laparoscopy over open surgery in the duration and intensity of convalescence is likely much less in the case of prostatectomy than it is for upper retroperitoneal procedures such as nephrectomy and adrenalectomy. Future refinement of the technique may allow for better neurovascular bundle and urethral dissection and thus improved postoperative outcomes. Survival rates after 10–15 years will determine if laparoscopic radical prostatectomy is comparable to open techniques.

Cystectomy/Cystoprostatectomy with Urinary Diversion

Laparoscopic cystectomy/cystoprostatectomy is feasible, yet experience is limited. The first laparoscopic cystectomy was performed for pyocystis in a patient with an existing urinary diversion. In 1995, the first series of laparoscopic cystectomies for bladder cancer was reported. Initial reports included diversions performed entirely extracorporeally, but intracorporeal ileal conduit (1995), rectal sigmoid pouch (2001), and Studer ileal neobladder (2002) are now being performed.

The technique for laparoscopic cystectomy/cystoprostatectomy is similar to that of laparoscopic radical prostatectomy, with minor modifications including controlling the pedicles of the bladder with the endoscopic stapler and transecting the ureters. Bowel diversion requires division and reanastomosis as in open surgery. Endoscopic staplers and free-hand laparoscopic suturing facilitate the creation of the urinary diversion. Totally intracorporeal cystectomy with neobladder reconstruction highlights the rapid advances made in the field of laparoscopic urology; these techniques will continue to evolve.

Miscellaneous Laparoscopic Procedures

Numerous other laparoscopic procedures have been described. Interesting examples include laparoscopic ureteroneocystotomy for vesicoureteral reflux, augmentation enterocystoplasty with or without biodegradable grafts, ileal ureter, Boari flap, flank herniorrhaphy, and catheterizable cecal tubes (ACE Malone). Continued improvement in technology, surgical skills, and patient demand will likely deliver laparoscopic approaches into mainstream urology.

COMPLICATIONS

During the first decade of urologic laparoscopy, much attention was paid to its “steep learning curve.” Many large reports indicate a considerable complication rate at the outset of a center’s series, which falls markedly as experience is accrued. The surgeons in these series were usually some of the first to perform advanced laparoscopy. More recent data suggest that when the surgeons have received laparoscopic training during their residency or fellowship, this “learning curve” is less dramatic. In general, the rates of minor and major complications of laparoscopy are similar to those in open surgery. The intraoperative complications that are most feared during laparoscopy, just as in open surgery, are vascular and viscous injury. The problem in laparoscopy is that such injuries may require an emergent change in the approach (ie, conversion to hand-assistance or open surgery), whereas in open surgery the problem can be managed right there. As such, maneuvers during laparoscopic surgery must be more deliberate and careful, not because complications are easier to create but because they may be harder to manage.

FUTURE OF LAPAROSCOPY

Laparoscopy is now well established in urology. Certainly there are many applications for which laparoscopy has not yet reached its fullest potential, most notably radical prostatectomy and radical cystectomy with urinary diversion. In addition, new and evolving technologies such as surgical robots (Figure 9–26), virtual reality, and telemedicine will likely enhance not only laparoscopy but also all surgical endeavors. The challenge for the next decade is not only to continue developing procedures and techniques but also to integrate laparoscopy into urologic practice such that it is no longer viewed as a technique for technical specialists but rather a technique for disease specialists. In other words, rather than having endourologists or other minimally invasive specialists perform all of the laparoscopic procedures, those with subspecialization in oncology should perform laparoscopic oncologic procedures, those with subspecialization in reconstructive urology should...
perform reconstructive laparoscopic procedures, and so on. In this way the urologic patient will benefit most from minimally invasive surgery.

REFERENCES

**General and Instrumentation**


**Physiology**


**Specific Procedures**


The ability to manipulate the urinary tract without the need for an open surgical incision differentiates urology from other disciplines. Such intervention may be required for diagnostic or therapeutic purposes (or both). Understanding the various catheters, guidewires, stents, endoscopes, and associated instrumentation is key in helping physicians accomplish their desired tasks. Manipulation of the urinary tract should be performed in a gentle fashion; instruments need not be forced. An understanding of anatomy and alternative instrumentation should allow physicians to accomplish their tasks with finesse. The patient should understand the proposed procedure and potential complications. For example, the attempt to place a retrograde ureteral catheter to drain an infected kidney may ultimately lead to a percutaneous nephrostomy if the surgeon is unable to achieve retrograde drainage. Knowing when to stop is as important as knowing when to start.

Many procedures are performed at the bedside or in a cystoscopy suite under local anesthesia. A patient who is comfortable, informed, and assured will more likely cooperate and tolerate the procedure. A physician who is familiar with the proposed instrumentation and understands its limitations and alternatives will win the patient’s confidence.

Manipulation of the urinary tract can result in significant injury. Anticipated prolonged procedures should be covered with appropriate antibiotics directed by preoperative urine cultures and sensitivities. Generous use of a water-soluble lubricant and low-pressure irrigation decreases the likelihood of significant iatrogenic infections. Patient positioning is as important as proper choice of instrumentation. Pressure points must be identified and adequately padded, especially when the patient is placed in the dorsal lithotomy position. Additionally, the legs should be secured in their stirrups to prevent accidental injury, such as might result from a leg hitting the surgeon after an unexpected obturator reflex during endoelectric surgery.

**URETHRAL CATHETERIZATION**

Urethral catheterization is the most frequent retrograde manipulation performed on the urinary tract. Catheters are placed to drain the bladder during and after surgical procedures requiring anesthetics, to assess urinary output in critically ill patients, to collect reliable urine specimens, for urodynamic evaluation, for radiographic studies (eg, cystograms), and to assess residual urine. Such catheters can be left indwelling with a self-retaining balloon, as is done with a Foley catheter. An in-and-out procedure to drain a bladder does not require a self-retaining device. Adequate lubrication and sufficient frequency to keep the bladder at reasonable volumes are critical and must be emphasized to the patient performing self-intermittent catheterization; sterility is secondary. In contrast, when a catheter is left indwelling it is important to use sterile technique.

### Technique of Catheterization

#### A. IN MEN

The penis should be positioned pointing toward the umbilicus to decrease the acute angulation as the catheter traverses the bulbar urethra. On most occasions, the catheter passes without difficulty. When difficulties arise, a careful history relating to previous urologic manipulations is critical. Strictures are not infrequent and can occur after endourologic surgery. Urethral strictures can be found from the meatus to the bladder neck. History of a straddle injury may suggest a bulbar urethral stricture. Adequate lubrication injected into the urethra and instruction of the patient to relax his pelvic floor eases the passage beyond the striated rhabdosphincter. A large-caliber catheter of approximately 18F should be used. Narrow, stiff, small catheters have greater potential of creating false passages and possible perforation. Coudé (elbowed) tipped catheters frequently help negotiate a high bladder neck, as seen with benign prostatic hyperplasia. With self-retaining Foley catheters, complete advancement until the elbowed valve is at the meatus or until the urine returns is important. Inflating the balloon prematurely (while it is in the urethra) may result in severe pain and possible urethral rupture. This must be emphasized to ancillary nursing personnel dealing with patients who are unable to communicate effectively, because under such circumstances, urethral rupture may present only after severe infection is evident.
B. IN WOMEN

It may be difficult to identify the meatus, especially in patients with obesity or hypospadias. Lateral and outward traction on the labia and the use of the posterior bill of a vaginal speculum may be helpful. With adequate instruction and a mirror to visualize the meatus, women can learn to catheterize themselves. For repeat catheterizations, a finger inserted into the vagina can help to guide the catheter.

C. DIFFICULT PLACEMENT & REMOVAL

When a urethral catheter cannot be placed, filiforms and followers may be used. The narrow filiform leaders are stiff and can puncture the urethra if too much force is used. Thus, gentle advancement should stop when resistance is encountered, and the initial filiform should be left in place. A second and third filiform, and possibly additional ones, should be placed next to the previously placed catheters in hope that the existing catheter occupies false passages or tortuous kinks. Eventually, one of the filiforms should pass and coil into the bladder. A screw adaptor at the end of the filiform can be used to connect progressively larger followers to dilate the narrowed urethra. After adequate dilatation, an open-tipped Councill catheter can be placed over the filiform and into the bladder. If at any stage a problem or undue resistance is encountered, the procedure should be aborted and a suprapubic cystostomy should be placed to achieve adequate drainage.

Indwelling catheters should be secured to a closed gravity drainage system. Drainage tubing connected to catheters should be positioned to limit dependent curls and thereby limit airlocks that will frequently limit bladder evacuation. For long-term requirements in males, the catheter should be secured to the abdominal wall to decrease urethral traction pressure and potential stricture formation. Meatal care is needed to ensure adequate egress of urethral secretions.

Difficulty is much less common when removing indwelling urethral catheters. Here, the retention balloon is deflated prior to removal. On occasion, the balloon may not deflate. Inspection of the valve frequently reveals any problem. One may cut proximal to the valve in hopes of evacuating the balloon contents, but this is not always successful. Other options include transperineal or transabdominal balloon puncture, or injection of an organic agent such as ether through the balloon port (with a full bladder to prevent chemical cystitis) to dissolve the balloon wall. Occasionally, a narrow pediatric endoscope must be placed next to the catheter in the urethra for evaluation. An unintended retaining suture may be present after recent open surgery. These sutures can be cut. Another complication of urethral catheters is incrustation, especially when a catheter is left indwelling for a long time.

Catheter Design

Catheters differ in size, shape, type of material, number of lumens, and type of retaining mechanism (Figure 10–1). Standard sizes of external catheter diameters and most endoscopic instruments are given according to Charrière’s French scale (units of 0.33 mm = 1 French [F] or 1 Charrière [Charr]). Thus, 3F equals 1 mm in diameter and 30F equals 10 mm in diameter.

The choice of catheter size is dependent on the patient and the purpose. Large-caliber catheters are used to evacuate blood clots or other debris. Other catheters are used to stabilize grafts after open urethroplasties, for stenting after endoscopic incisions of strictures, for support of external ureteral catheters, or to assess urinary output. Triple-lumen catheters (one port for balloon inflation and deflation, and one each for inflow and outflow) have smaller lumens than 2-way catheters. Other catheter variables include balloon size and construction materials; smaller catheters have smaller balloons. Large balloons (eg, 30 mL) can be inflated well over 50 mL to decrease the likelihood of the balloon migrating into the prostatic fossa, especially after transurethral resection of the prostate. They can be used as traction devices against the bladder neck to control hemorrhage from the prostatic fossa after tranurethral resection of the prostate (TURP).

The rigidity of the catheter, the ratio between internal and external diameters, and the biocompatibility depend on the material with which the catheter is made. The standard latex catheter can result in severe reactions in patients with latex allergies, most commonly those with myelomeningoceles. Silicone varieties are good alternatives in such situations. Mucosal irritation is decreased when catheters with a low coefficient of friction are used. Hydromers are placed onto catheters to allow for transient coating, creating an interface between biologic tissues and the foreign catheter; this interface lasts for approximately 5 days. Permanent hydrogel coatings last the life of the catheter. Decreasing the coefficient of friction of these catheters brings about a decrease in mucosal irritation and better biocompatibility. Catheters with a longer lasting interface result in decreased incrustation.

URETHROSCOPY

To identify and aid in treating urethral pathology, endoscopic inspection via a urethroscope with a 0° lens is helpful. Stricture disease can be identified or confirmed after radiographic studies. Strictures are characterized by circumferential narrowing. Sequential dilation of urethral strictures by inserting catheters of increasing size exerts shear and tear forces to the mucosa and is likely to produce extended scarring. Thus, stricture recurrence is common if periodic urethral dilation is terminated. Balloon dilation of a stricture with 7–9F balloon dilators (which can be passed
over guidewires and inflated up to 30F with pressures of up to 15 atm) does not exert shear force, but the long-term results are poor. Limited circumferential strictures can be incised under direct vision with an endoscopic cold knife. The incision is usually made at the 12-o’clock position, adequate to allow passage of the urethroscope. The bladder then can be evacuated and adequate irrigation used if further incision results in hemorrhage. It is difficult to identify the true extent and depth of a stricture solely by vision because scarring can involve deeper tissues. Here, urethral ultrasonography is an adjunct.

Urethral diverticulum can be identified with urethroscopy. A catheter can be placed through the neck of the diverticulum to help confirm its location during definitive open surgical repair. Urethroscopy can be used to direct injection of dye into rare retained müllerian duct cysts, to identify and extract foreign bodies or rare calculi, and to access biopsy-suspicious lesions. Urethroscopy allows endoscopic treatment of urethral condylomata.

**Figure 10–1.** Urethral catheters, metal stylet, catheter, and guidewire techniques for catheter insertion.

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**CYSTOSCOPY**

Endoscopic inspection of the lower urinary tract requires irrigation, illumination (fiberoptics), and optics. The optics and illumination are offset by the irrigating and working port. To optimize a complete examination, the rigid endoscope should be rotated, and 0°, 30°, 70°, and 120° lenses may be required. Suprapubic pressure facilitates inspection of the bladder dome, which frequently has an air bubble. A systematic approach is required when evaluating the urethra, prostate, bladder walls, dome and neck, and ureteral orifices (including location, number, shape, and character of efflux). The bladder should be evaluated at different levels of filling. It is only after full distention of the bladder that characteristic glomerulations and ecchymoses are seen in interstitial cystitis. Rectal examination with the endoscope in place is informative, especially in assessing prostate size and length of prostatic urethra. Similarly concurrent vaginal examination in women can be useful in evaluation of cystoceles.
Choice of irrigant during endoscopic manipulation is important. There are conductive and nonconductive irrigants. Conductive irrigants, which include saline and lactated Ringer’s solution, would be inappropriate during traditional endoelectric surgery because the electrical charge would be diffused by the irrigant. Nonconductive irrigants include water and glycine. Water has a theoretic advantage of increasing visibility, and because it is hypotonic, it can lyse tumor cells. If the potential exists for increased intravascular absorption, iso-osmotic or other nonhemolyzing agents are preferred to hypotonic solutions.

Rigid endoscopy results in discomfort, which can be minimized with 1% lidocaine per urethra as a local anesthetic. Flexible endoscopes decrease patient discomfort and allow for instrumentation in the supine rather than the dorsal lithotomy position. They are now used routinely in an office setting for hematuria/tumor surveillance and double-J stent retrieval. Videoendoscopy with flexible scopes allows patients to visualize normal and abnormal anatomy and thus helps them understand their pathology. Videoendoscopy reduces fluid contact to the urologist and can help reduce cervical neck disease. However, there are disadvantages. Flexible scopes have smaller irrigating ports, and they do not have a working sheath. As a result, changing lenses, assessing residual urine, and repeat evacuation of irrigant cannot be completed without entirely removing the endoscope. Rigid endoscopy allows for a greater variety of instrumentation, better optics, and increased durability.

Instrumentation similar to that used to evaluate the urethra and bladder can be used to inspect continent urinary reservoirs or conventional ileal loops. A Robinson or a Foley catheter placed prior to the endoscope gives the operator a visual landmark and an exit port for irrigation to keep the procedure at a low pressure. Alternatively, the Foley balloon can be inflated and the catheter plugged to transiently expand the intestinal segment in hopes of helping to identify landmarks or pathologic lesions. Endoscopic inspection allows for identification of calculi, foreign bodies and mucous plugs, and also has the potential for intubation of ureterointestinal anastomoses.

**URETERAL CATHETERIZATION**

Ureteral catheterization is required in performing retrograde pyelography, collecting urine for cytologic examination or cultures, and performing brush biopsies (Figure 10–2). Other procedures (Figure 10–3) that require ureteral catheterization include draining an obstructed kidney due to either intrinsic or extrinsic compressions and placement of an internal double-J stent. Finding the ureteral orifice can be difficult. Long-term indwelling Foley catheters, infection, history of ureteral reimplantation, or renal transplantation can hinder identification of the ureteral orifice. One must first try to identify the interureteric ridge and then look for a jet of urinary efflux. Varying bladder volumes and use of intravenous methylene blue may be helpful. However, it may take up to 5–20 minutes for intravenous agents to be excreted out of the ureteral orifice. Once the orifice is identified, catheters usually are placed uneventfully. However, in the setting of benign prostatic hyperplasia with J-hooking of the distal ureter, previous retroperitoneal surgery, reimplantation of the ureter, decreased lower extremity mobility or other skeletal abnormalities, or edema or kinking secondary to long-standing impacted ureteral calculi, catheterization procedures can be difficult or impossible. An Albiron bridge may help direct catheters and guidewires.

There are many configurations of catheter tips (Figure 10–4). Acorn or cone-tipped catheters are excellent for routine retrograde pyelography. Care should be taken to eliminate air in the catheter before injection to avoid confusing air with a filling defect. Fluoroscopy helps determine the appropriate volume of radiocontrast material to decrease the likelihood of pyelolymphatic or pyelovenous reflux or fornical rupture. The average collecting system holds 7–9 mL of contrast material. If performed under local anesthesia, overdistention is recognized by severe ipsilateral flank pain. With low-pressure injections, there is no systemic absorption of contrast material. A coudé-tipped catheter allows for dramatic mobility of the tip of the catheter by merely twisting it; there is no need for exaggerated motion of the endoscope. This is helpful in orifices that are difficult to identify because of edema or tumor infiltration.

To bypass severe angulations, passage of a guidewire must first be attempted. Straight guidewires can be made floppy if they have removable cores, and frequently this
allows easy passage. At times, hydrophilic coudé-tipped torque guidewires are helpful. If the orifice can be engaged with the tip of the guidewire but the guidewire cannot be advanced, the tip of the endoscope should be pivoted toward the contralateral orifice while the guidewire is advanced through the endoscope just enough to keep the guidewire engaged in the orifice. The guidewire should then be advanced against the back wall of the bladder, effectively changing the vector force so that the wire can be advanced through a severe J deformity (Figure 10–5). With the guidewire advanced, an exchange catheter can be advanced over the guidewire for injection of contrast material, to be exchanged later for another guidewire or an open-ended catheter. A coudé-tipped guidewire or floppy-tipped guidewire (with a removable core guide) can be advanced through such exchange catheters to facilitate bypassing stones or severe kinks. A push-pull maneuver (pulling the exchange catheter while pushing the guide-wire) frequently straightens the ureter as a result of resistance from the exchange catheter, allowing advancement of the guidewire. To increase resistance, an occlusion balloon ureteral catheter can be inflated and with gentle traction can help straighten a kinked or tortuous ureter. Additional helpful maneuvers include deep exhalation, thus elevating the diaphragm, external cephalad pressure by an assistant, or Trendelenburg patient positioning.

Double-J catheters are used to facilitate internal drainage due to obstruction from ureteral angulation and internal or external ureteral compression; they are also used to help decrease the likelihood of sepsis or obstruction in the presence of steinstrasse after extracorporeal shock wave lithotripsy. Double-J stents increase the internal lumen of the ureter. This increase may be used to one’s advantage in the setting of a narrow ureter. Placing a double-J catheter and postponing the ureteroscopy for a few days significantly decreases the difficulty of the subsequent procedure. Double-J stents disrupt normal ureteral peristalsis. These stents can be placed over a guidewire or with a closed leading end. With proper placement of the proximal end into the renal pelvis, the J should project in the lateral position when seen on fluoroscopy or x-ray. Projecting in an anterior-posterior position suggests a proximal ureteral location. Proximal J stent placement can be confirmed by renal ultrasonography during placement in pregnant patients. If it is too long, the distal end in the bladder can result in severe irritative voiding symptoms; if too short, it is more likely to migrate proximally beyond the ureteral orifice into the ureter. In the latter situations, drainage cannot be ensured, and the stent must be extracted with a ureteroscope or snared with a ureteral stone basket.

Patients must be informed that internalized stents have been placed. Frequently, they will not feel the stent. When the stent is left in place for long periods, the likelihood of incrustations, poor drainage, and difficult extraction is increased. It is unclear whether double-J stents facilitate drainage because of drainage around the catheter or via the numerous side holes communicating with the internal lumen. New helically ridged double-J ureteral stents likely enhance ureteral stone passage through unidirectional ratchet-like motion over the external ridges during respiratory and body wall motion. Other complications include distal migration into the bladder, distal migration beyond the bladder neck (resulting in total urinary incontinence), and flank pain during micturition secondary to reflux. The catheter can be removed with cold cup forceps via a flexible or rigid cystoscope or by pulling a string that has been attached to the distal end of the catheter and left exiting through the meatus. Although double-J catheters have potential complications, they can help ensure internal urinary drainage.

Balloon dilators can be used to ease passage of ureteroscopes (rigid or flexible; see Chapter 8) and extract intact large calculi. Balloons are routinely passed over a guide-
Wire. Woven balloons have a tight, unfolded outer surface that shortens in longitudinal length when inflated. In contrast, nonwoven varieties are folded and may be difficult to pass after initial inflation and deflation; however, they do not shorten in length with inflation. Balloons inflated alongside distal ureteral calculi can result in balloon perforation or extrusion of the calculus outside the ureteral lumen. Balloon inflation is best achieved with ratcheted or torqued syringe aids directed with pressure gauges. Ureteral access sheaths, frequently made with a hydrophilic coating, can be placed over a guidewire. They dilate the ureter without the need for a ureteral balloon and simplify multiple passages up the ureter.

Retrograde endopyelotomy is an alternative to laparoscopic and open surgical repair, and percutaneous antegrade approaches. After documentation of the exact location of the ureteropelvic junction obstruction under fluoroscopic control, a 150-cm superstiff Lunderquist guidewire is advanced into the renal pelvis. The endoscope is removed and the retrograde endopyelotomy device (Acucise) is advanced over the guidewire under fluoroscopic control. After the incising wire is directed laterally, the balloon is inflated during cautery. Successful results are seen in approximately 80% of patients. An internal endopyelotomy double-J stent, 14F at the proximal end, straddling the ureteropelvic junction, and tapered to 6–8F as it enters and coils in the bladder, or a routine 7F double-J ureteral stent is placed over the stiff guidewire and left in place for 6 weeks. Placing a standard double-J catheter before this procedure dilates the ureter and eases passage of the Acucise and the endopyelotomy double-J catheter.

A large selection of endoscopic baskets is available to entrap and remove material including calculi, sloughed papillae, bulky tumors, fungal bezoars, and foreign bodies. Baskets are designed with and without filiform leaders and can be advanced on their own or, more commonly, through the working ports of flexible and rigid ureteroscopes. Round wire baskets can be torqued to help entrap the target material. A few (2–3) wired baskets are used for large material, while numerous (4–6) wired baskets are
used for small or numerous objects. Flat wired baskets can engage stones effectively. If twisted, however, the wire can fold and transform into a knifelike edge. Once the basket is engaged, one should ensure that the endothelium is not entrapped. Gentle traction helps extract these foreign materials. Engaged baskets may be difficult to disengage. Occasionally, one must cut the handle and place a ureteroscope alongside the basket to facilitate stone and basket extraction. Nitinol baskets have rounded tops and decrease potential endothelial trauma.

Transurethral injections can be performed through a variety of endoscopes. Newer injectables include fibrin glue, Botox, and bulking agents.

**TRANSURETHRAL SURGERY**

Resectoscopes are endoscopes with sheaths of 10–30F (Figure 10–6) designed for transurethral surgery; they allow urologists to excise, fulgurate, or vaporize tissue from the lower genitourinary tract. Applying an alternating current at high frequencies decreases muscular contractions and allows for cutting and coagulation properties. A pure sine wave is optimal for cutting, whereas dampened oscillating waveforms are best for coagulation. It is possible to combine the 2 waves to allow for simultaneous cutting and coagulation. A ground plate, as an indifferent electrode, usually applied over the hip, is required. The cutting current results in rapid vaporization of tissue, allowing the cutting loop to move easily through tissue with minimal resistance, and separates the chip, enabling easy flow into the bladder. Rapid succession of cutting sweeps allows rapid surgical excision. In contrast, a coagulation current results in less rapid vaporization and thus decreased separation of tissue from the cutting current. If a traditional resectoscope does not cut tissue, the operator should check for a broken resecting loop, a broken or disconnected cable or generator, or a conductive irrigant (as with saline) that diffuses the current. Newer plasmakinetic or bipolar resectoscopes send current from one edge of the endoscopic loop to the other. A high electrical current is generated locally at the loop and effectively vaporizes tissue on contact. Due to the bipolar design, conductive irrigants are used with these bipolar resectoscopes. Resection can also be performed with lasers in a similar fashion.

Before transurethral surgery, the urethra should be calibrated with sounds to ensure ease in placing the resecto-
Urethral sounds and probes come in numerous varieties (Figure 10–7). An Otis urethrotome can be used to incise the urethra at the 12-o’clock position, thereby decreasing the potential for stricture disease in narrow urethras. Generous use of a water-soluble lubricant is indicated. Before placement of the resectoscope, the loop should be inspected for defects and for appropriate engagement to ensure complete retraction into the sheath, thereby allowing resected tissue to flow easily into the bladder. The endoscope can be placed under direct vision, especially if the patient has not recently been cystoscoped. Alternatively, a Timberlake obturator allows for blind placement of the resectoscope sheath. Most endoscopes require the operator to intermittently remove the working element to allow evacuation of bladder contents. Other endoscopes have an additional channel for continuous operation. An alternative is a percutaneous suprapubic drainage catheter, which allows for continuous flow. Orientation with identification of landmarks, such as the verumontanum and the ureteral orifices, before resection dramatically decreases the potential for complications. Bladder lesions are best resected with minimal bladder distention to decrease the likelihood of perforation. A Bugbee electrode can be used for point coagulation of bleeding points or pathologic lesions. A rollerball can be used to coagulate large areas. Transurethral resection can be used to resect an obstructing prostate gland, to drain prostatic abscesses, or to unroof the ejaculatory duct in select infertility patients.

TURP is a time-tested procedure for resecting prostatic tissue and decreasing symptoms of urinary obstruction. In experienced hands, this can be done with minimal complications. New, alternative procedures are being investigated, especially with patients who are poor anesthetic risks, whose life expectancy is limited, or who are averse to TURP. Those with small glands or with bladder neck contractures have been treated by transurethral incision of the prostate from a point just distal to the ureteral orifices to the verumontanum. Transcystoscopic urethroplasty, also known as prostatic balloon dilatation, dilates the prostatic urethra under visual and fluoroscopic control. Intracysthral coils can be placed in high-risk patients to avoid permanent catheter drainage. Thermotherapy treatment delivers temperatures of 41°C to 44°C for 60 minutes.
Obstructing median lobes are unsuited for such newer techniques. Interstitial laser ablation of the prostate gland is another method for the management of prostatic enlargement. Long-term studies are required to compare this technique with TURP.

There are various cutting techniques for resecting an obstructing prostate gland during TURP. All require good vision, a comfortable operator, identification of the surgical capsule, and established goals that are met before starting additional stages of the procedure. Pulsatile arterial bleeders should be coagulated first and venous hemorrhage next. Occasionally, arterial bleeders cannot be coagulated without additional resection of tissue. An Ellik bulb or piston syringe should be available to evacuate resected tissue. At the conclusion of the procedure, one should ensure adequate resection and hemostasis, and perform an inspection for forgotten chips of tissue and possible injuries. A Foley catheter should be placed into the bladder and irrigated to confirm unobstructed flow and adequate hemostasis. If an undermined trigone is suspected, a coudé-tipped catheter, a finger in the rectum, or a stylet inserted into the catheter can help in proper placement. The Foley balloon should be inflated 20 mL + 1 mL for each gram of resected tissue.

Video cameras can be attached to the optical eyepiece while transurethral surgery is performed. Use of the camera can reduce the risk of cervical disc disease and distance the surgeon from blood products. It is an excellent resource to improve the teaching of endoscopic surgery.

Acute complications include intra- or extraperitoneal rupture of the bladder, rectal perforation, incontinence, incision of a ureteral orifice with possible reflux or stricture, hemorrhage, gas explosion (especially during resection of a bladder lesion at the dome in the presence of accumulated gas), epididymitis, sepsis, and transurethral resection syndrome. The transurethral resection syndrome is characterized by delusional hyponatremia resulting in possible confusion, congestive heart failure, or pulmonary edema. It is secondary to a large amount of fluid being absorbed, usually through a perforation of a low-pressure system such as the venous sinuses. If perforations are noted, especially into a sinus, the height of the irrigating solution should be lowered, hemostasis achieved, and the procedure brought to a rapid conclusion. Other complications include impotence (with excessive coagulation) and urethral stricture disease. After an adequate transurethral resection of the prostate, retrograde ejaculation almost always occurs.

LOWER TRACT CALCULI

Most bladder calculi originating from the upper tract pass spontaneously through the urethra. In contrast, bladder calculi resulting from outlet obstruction may require endoscopic extraction. Many of these calculi can be washed out or extracted with the aid of various forceps or a resectoscope loop. Calculi too large to pass through an endoscopic sheath first require fragmentation. Visual lithotripters with crushing jaws or a punch-type mechanism are effective. Introduction of these bulky devices is potentially dangerous. A distended bladder facilitates effective engagement of the stone without injuring the bladder wall. Twisting the lithotrite before crushing ensures that the bladder wall is free from the instrument.

Other methods available to fragment bladder calculi include ultrasonic, electrohydraulic, laser, and pneumatic lithotripters. Ultrasonic lithotripters use vibratory energy delivered via a rigid metal transducer. An offset endoscopic lens is required. Gentle pressure of the transducer against the stone facilitates fragmentation; excessive pressure can erode or perforate the bladder wall. A hollow core with suction extracts the fragments. Electrohydraulic lithotripsy generates a spark-gap, resulting in a shock wave. It is delivered at the end of a flexible catheter and can be applied as single or repetitive shocks. Fragmentation can be performed with normal saline. A rheostat can adjust the power output. A high setting can result in the stone caroming to various locations in the bladder; lower settings result in suboptimal fragmentation. To optimize fragmentation, the tip of the lithotrite should be a few millimeters away from the stone. To protect the endoscopic optics, the endoscope should be kept at a distance. Shock waves fragment brittle material, such as the stone or a lens. Biologic tissues are elastic and are unharmed as long as the spark-gap does not touch them. Air-driven, jack-hammer-like devices (rigid and flexible) can be used for stone fragmentation. The photothermal mechanism of holmium lasers is effective in fragmenting very large bladder stones. Uric acid calculi produce small amounts of cyanide gas when fragmented with holmium lasers; no clinical sequelae have been documented. Pneumatic lithotripters are effective with minimal tissue trauma. They use reusable probes and are powered by compressed gas.

ADVANCED INSTRUMENTATION

Lasers

Lasers (light amplification by stimulated emission of radiation) have been used through flexible and rigid endoscopes. Carbon dioxide and argon lasers result in tissue penetration that is inadequate for the needs of urology. Neodymium:YAG lasers give adequate tissue coagulation and are useful for various lesions. The holmium:YAG system is excellent for stone fragmentation and tissue ablation and is now the most popular system in use. Disadvantages are the lack of adequate tissue for histopathologic evaluation and the initial cost of machinery.

Ultrasonography

Ultrasound has found increased application to the lower genitourinary tract. It results in minimal discomfort; gives a
three-dimensional appreciation of the shape, size, and volume of organs and disease; and can provide direct intervention. Various transducers are available; high-megahertz transducers are required for superficial structures (for example, scrotal structures) to assess testicular disease (including tumors and torsion), while low-megahertz transducers are reserved for deep structures (for example, guiding percutaneous access for kidneys and bladders). Intervening tissue can significantly reduce image quality.

Transrectal ultrasound is valuable in evaluating the prostate to determine size and confirm digital information on the presence and stage of a suspected malignant tumor. Because of the low incidence of detecting malignancies (1.6–7%), mass screening programs are not cost effective. Direct needle biopsies, with automatic biopsy mechanisms, are quick, well tolerated, and result in reliable tissue cores and less pain than traditional needles (such as Tru-Cut) directed under digital palpation. Percutaneous drainage tubes, radioactive seed implants, and temperature coils used for cryosurgery of the prostate can be placed safely under transrectal ultrasonic guidance. Transrectal ultrasonography can yield unreliable images that often are misinterpreted by the novice. Pitfalls include faulty instrument settings, poor coupling caused by feces or gas, and unrecognized artifacts resulting from reverberation, deflection, shadowing, or enhancement.

Suprapubic ultrasonography can help to assess prostate anatomy, especially size and intravesical extension. It can help to evaluate the bladder for residual urine and for calculi that are questionable on plain abdominal radiographs. (Changing the patient’s position can shift the position of a calculus.) Distal ureteral stones can be identified, especially when visualized through a full bladder used as an acoustic window. Double-J stents, incrustations, diverticula, and large malignant lesions can be identified. The procedure also can direct placement of suprapubic cystostomy drainage catheters.

Additional applications include endocavitary, color, Doppler, and dynamic ultrasonography. Endocavitary ultrasound, which includes transvaginal, transurethral (Figure 10–8), and transcystoscopic techniques, can delineate vaginal, urethral, and bladder disease. Endoureteral ultrasound can help in the identification of crossing vessels, preferably before an endopyelotomy. Color and Doppler ultrasound can assess blood flow as related to erectile dysfunction. Dynamic ultrasound can supplement urodynamic findings. Ultrasound applied to the lower genitourinary tract causes minimal discomfort and provides valuable information.

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Cytology, Biopsy Histology

Endoscopy

Lithotripsy
Because of their damaging effect on renal function, obstruction and stasis of urinary flow are among the most important urologic disorders. Either leads eventually to hydronephrosis, a peculiar type of atrophy of the kidney that may terminate in renal insufficiency or, if unilateral, complete destruction of the organ. Furthermore, obstruction leads to infection, which causes additional damage to the organs involved.

**Classification**

Obstruction may be classified according to cause (congenital or acquired), duration (acute or chronic), degree (partial or complete), and level (upper or lower urinary tract).

**Etiology**

Congenital anomalies, more common in the urinary tract than in any other organ system, are generally obstructive. In adult life, many types of acquired obstructions can occur.

**A. CONGENITAL**

The common sites of congenital narrowing are the external meatus in boys (meatal stenosis) or just inside the external urinary meatus in girls, the distal urethra (stenosis), posterior urethral valves, ectopic ureters, ureteroceles, and the ureterovesical and ureteropelvic junctions. Another congenital cause of urinary stasis is damage to sacral roots 2–4 as seen in spina bifida and myelomeningocele. Vesicoureteral reflux causes both vesical and renal stasis (see Chapter 12).

**B. ACQUIRED**

Acquired obstructions are numerous and may be primary in the urinary tract or secondary to retroperitoneal lesions that invade or compress the urinary passages. Among the common causes are (1) urethral stricture secondary to infection or injury; (2) benign prostatic hyperplasia or cancer of the prostate; (3) vesical tumor involving the bladder neck or one or both ureteral orifices; (4) local extension of cancer of the prostate or cervix into the base of the bladder, occluding the ureters; (5) compression of the ureters at the pelvic brim by metastatic nodes from cancer of the prostate or cervix; (6) ureteral stone; (7) retroperitoneal fibrosis or malignant tumor; and (8) pregnancy.

Neurogenic dysfunction affects principally the bladder. The upper tracts are damaged secondarily by ureterovesical obstruction or reflux and, often, complicating infection. Severe constipation, especially in children, can cause bilateral hydronephrosis from compression of the lower ureters.

Elongation and kinking of the ureter secondary to vesicoureteral reflux commonly lead to ureteropelvic obstruction and hydronephrosis. Unless a voiding cystourethrogram is obtained in children with this lesion, the primary cause may be missed and improper treatment given.

**Pathogenesis & Pathology**

Obstruction and neuropathic vesical dysfunction have the same effects on the urinary tract. These changes can best be understood by considering (1) the effects on the lower tract (distal to the bladder neck) of severe external urinary meatal stricture and (2) the effects on the midtract (bladder) and upper tract (ureter and kidney) of benign prostatic hyperplasia.

**A. LOWER TRACT (EG, URETHRAL STRICTURE)**

Hydrostatic pressure proximal to the obstruction causes dilation of the urethra. The wall of the urethra may become thin, and a diverticulum may form. If the urine becomes infected, urinary extravasation may occur, and periurethral abscess can result. The prostatic ducts may become widely dilated.

**B. MIDTRACT (EG, PROSTATIC HYPERPLASIA)**

In the earlier stages (compensatory phase), the muscle wall of the bladder becomes hypertrophied and thickened. With decompensation, it becomes less contractile and, therefore, weakened.

1. **Stage of compensation**—To balance the increasing outlet resistance, the bladder musculature hypertrophies. Its thickness may double or triple. Complete emptying of the bladder is thus made possible.

   Hypertrophied muscle may be seen endoscopically. With secondary infection, the effects of infection are often superimposed. There may be edema of the submucosa, which may be infiltrated with plasma cells, lymphocytes, and polymorphonuclear cells. At cystoscopy, surgery, or
autopsy, the following evidence of this compensation may be visible (Figure 11–1):

**a. Trabeculation of the bladder wall**—The wall of the distended bladder is normally quite smooth. With hypertrophy, individual muscle bundles become taut and give a coarsely interwoven appearance to the mucosal surface. The trigonal muscle and the interureteric ridge, which normally are only slightly raised above the surrounding tissues, respond to obstruction by hypertrophy of their smooth musculature. The ridge then becomes prominent. This trigonal hypertrophy causes increased resistance to urine flow in the intravesical ureteral segments owing to accentuated downward pull on them. It is this mechanism that causes relative functional obstruction of the ureterovesical junctions, leading to back pressure on the kidney and hydronephrosis. The obstruction increases in the presence of significant residual urine, which further stretches the ureterotr生殖的 complex. (A urethral catheter relieves the obstruction somewhat by eliminating the trigonal stretch. Definitive prostatectomy leads to permanent release of stretch and gradual softening of trigonal hypertrophy with relief of the obstruction.)

**b. Cellules**—Normal intravesical pressure is about 30 cm of water at the beginning of micturition. Pressures 2–4 times as great may be reached by the trabeculated (hype rtrophied) bladder in its attempt to force urine past the obstruction. This pressure tends to push mucosa between the superficial muscle bundles, causing the formation of small pockets, or cellules (Figure 11–1).

**c. Diverticula**—If cellules force their way entirely through the musculature of the bladder wall, they become saccules, then actual diverticula, which may be embedded in perivesical fat or covered by peritoneum, depending on their location. Diverticula have no muscle wall and are therefore unable to expel their contents into the bladder efficiently even after the primary obstruction has been removed. When secondary infection occurs, it is difficult to eradicate; surgical removal of the diverticula may be required. If a diverticulum pushes through the bladder wall on the anterior surface of the ureter, the ureterovesical junction will become incompetent (see Chapter 12).

**d. Mucosa**—In the presence of acute infection, the mucosa may be reddened and edematous. This may lead to temporary vesicoureteral reflux in the presence of a “borderline” junction. The chronically inflamed membrane may be thinned and pale. In the absence of infection, the mucosa appears normal.

**2. Stage of decompensation**—The compensatory power of the bladder musculature varies greatly. One patient with prostatic enlargement may have only mild symptoms of prostatism but a large obstructing gland that can be palpated rectally and observed cystoscopically; another may suffer acute retention and yet have a gland of normal size

*Figure 11–1. Changes in the bladder developing from obstruction. Upper left: Normal bladder and prostate. Upper right: Obstructing prostate causing trabeculation, cellule formation, and hypertrophy of the interureteric ridge. Bottom: Marked trabeculation (hypertrophy) of the vesical musculature; diverticulum displacing left ureter.*
on rectal palpation and what appears to be only a mild obstruction cystoscopically.

In the face of progressive outlet obstruction, possibly aggravated by prostatic infection with edema or by congestion from lack of intercourse, decompensation of the detrusor may occur, resulting in the presence of residual urine after voiding. The amount may range up to 500 mL or more.

C. UPPER TRACT

1. Ureter—In the early stages of obstruction, intravesical pressure is normal while the bladder fills and is increased only during voiding. The pressure is not transmitted to the ureters and renal pelves because of the competence of the ureterovesical “valves.” (A true valve is not present; the ureterotrigonal unit, by virtue of its intrinsic structure, resists the retrograde flow of urine.) However, owing to trigonal hypertrophy (see the section Trabeculation of the bladder wall) and to the resultant increase in resistance to urine flow across the terminal ureter, there is progressive back pressure on the ureter and kidney, resulting in ureteral dilatation and hydronephrosis. Later, with the phase of decompensation accompanied by residual urine, there is an added stretch effect on the already hypertrophied trigone that increases appreciably the resistance to flow at the lower end of the ureter and induces further hydroureretonephrosis. With decompensation of the ureterotrigonal complex, the valve-like action may be lost, vesicoureteral reflux occurs, and the increased intravesical pressure is transmitted directly to the renal pelvis, aggravating the degree of hydroureretonephrosis.

Secondary to the back pressure resulting from reflux or from obstruction by the hypertrophied and stretched trigone or by a ureteral stone, the ureteral musculature thickens in its attempt to push the urine downward by increased peristaltic activity (stage of compensation). This causes elongation and some tortuosity of the ureter (Figure 11–2). At times, this change becomes marked, and bands of fibrous tissue develop. On contraction, the bands further angulate the ureter, causing secondary ureteral obstruction. Under these circumstances, removal of the obstruction below may not prevent the kidney from undergoing progressive obstruction due to the secondary ureteral obstruction.

Finally, because of increasing pressure, the ureteral wall becomes attenuated and therefore loses its contractile power (stage of decompensation). Dilatation may be so extreme that the ureter resembles a loop of bowel (Figures 11–3 and 12–8, upper right).

2. Kidney—The pressure within the renal pelvis is normally close to zero. When this pressure increases because of obstruction or reflux, the pelvis and calyces dilate. The degree of hydronephrosis that develops depends on the duration, degree, and site of the obstruction (Figure 11–4). The higher the obstruction, the greater the effect on the kidney. If the renal pelvis is entirely intrarenal and the obstruction is at the ureteropelvic junction, all the pressure will be exerted on the parenchyma. If the renal pelvis is extrarenal, only part of the pressure produced by a ureteropelvic stenosis is exerted on the parenchyma; this is because the extrarenal renal pelvis is embedded in fat and dilates more readily, thus “decompressing” the calyces (Figure 11–2).

In the earlier stages, the pelvic musculature undergoes compensatory hypertrophy in its effort to force urine past the obstruction. Later, however, the muscle becomes stretched and atonic (and decompensated).

The progression of hydronephrotic atrophy is as follows:

1. The earliest changes in the development of hydronephrosis are seen in the calyces. The end of a normal calyx is concave because of the papilla that projects into it; with increase in intrapelvic pressure, the fornices become blunt and rounded. With persistence of increased intrapelvic pressure, the papilla becomes flattened, then convex (clubbed) as a result of compression enhanced by ischemic atrophy (Figure 11–5). The parenchyma between the calyces is affected to a lesser extent. The changes in the renal parenchyma are due to (a) compression atrophy from increase in intrapelvic pressure (more accentuated with intrarenal pelves) and (b) ischemic atrophy from hemodynamic changes, mainly manifested in arcuate vessels that run at the base of the pyramids parallel to the kidney outline and are more vulnerable to compression between the renal capsule and the centrally increasing intrapelvic pressure.

2. This spotty atrophy is caused by the nature of the blood supply of the kidney. The arterioles are “end arteries”; therefore, ischemia is most marked in the areas farthest from the interlobular arteries. As the back pressure increases, hydronephrosis progresses, with the cells nearest the main arteries exhibiting the greatest resistance.

3. This increased pressure is transmitted up the tubules. The tubules become dilated, and their cells atrophy from ischemia. It should be pointed out that a few instances of dilated renal pelves and calyces are not due to the presence of obstruction. Rarely, the renal cavities are congenitally capacious and thus simulate hydronephrosis. More commonly, hydronephrosis may occur in childhood owing to the back pressure associated with vesicoureteral reflux. If the valvular incompetence resolves (and this is common), some degree of the hydronephrotic changes may persist. These persisting changes may cause the physician to suspect the presence of obstruction, which may lead to unnecessary surgery. An isotope renogram or the Whitaker test can be performed to determine whether organic obstruction is present.

4. Only in unilateral hydronephrosis are the advanced stages of hydronephrotic atrophy seen. Eventually the kidney is completely destroyed and appears as a
Figure 11–2. Mechanisms and results of obstruction. **Upper left:** Early stage. Elongation and dilatation of ureter due to mild obstruction. **Upper center:** Later stage. Further dilatation and elongation with kinking of the ureter; fibrous bands cause further kinking. **Upper right:** Intrarenal pelvis. Obstruction transmits all back pressure to parenchyma. **Lower:** Extrarenal pelvis, when obstructed, allows some of the increased pressure to be dissipated by the pelvis.
Figure 11–3. Pathogenesis of bilateral hydronephrosis. Progressive changes in bladder, ureters, and kidneys from obstruction of an enlarged prostate: thickening of bladder wall, dilatation and elongation of ureters, and hydronephrosis.
thin-walled sac filled with clear fluid (water and electrolytes) or pus (Figure 11–6).

If obstruction is unilateral, the increased intrarenal pressure causes some suppression of renal function on that side. The closer the intrapelvic pressure approaches the glomerular filtration pressure (6–12 mm Hg), the less urine can be secreted. Glomerular filtration rate and renal plasma flow are reduced, concentrating power is gradually lost, and the urea-creatinine concentration ratio of urine from the hydronephrotic kidney is lower than that of urine from the normal kidney.

Hydronephrotic atrophy is an unusual type of pathologic change. Other secretory organs (e.g., the submaxillary gland) cease secreting when their ducts are obstructed. This causes primary (disuse) atrophy. The completely obstructed kidney, however, continues to secrete urine. (If this were not so, hydronephrosis could not occur, since it depends on increased intrarenal pressure.) As urine is excreted into the renal pelvis, fluid and, particularly, soluble substances are reabsorbed, through either the tubules or the lymphatics. This has been demonstrated by injecting phenolsulfonphthalein (PSP) into the obstructed renal pelvis. It disappears (is reabsorbed) in a few hours and is excreted by the other kidney. If the intrapelvic pressure in the hydronephrotic kidney rapidly increases to a level approaching filtration pressure (resulting in cessation of filtration), a safety mechanism is activated that produces a break in the surface lining of the collecting structure at the weakest point—the fornices. This leads to escape and extravasation of urine from the renal pelvis into the parenchymal interstitium (pyelointerstitial backflow). The extravasated fluid is absorbed by the renal lymphatics, and the pressure in the renal pelvis drops, allowing further filtration of urine. This explains the process by which the markedly hydronephrotic kidney continues to function. Further evidence of the occurrence of extravasation and reabsorption is that the markedly hydronephrotic kidney does not contain urine in the true sense; only water and a few salts are present.

Functional impairment in unilateral hydronephrosis, as measured by excretory urograms or renal scans, is greater and increases faster than that seen in bilateral hydronephrotic kidneys showing comparable damage on urography. As unilateral hydronephrosis progresses, the normal kidney undergoes compensatory hypertrophy (particularly in children) of its nephrons (renal counterbalance), thereby assuming the function of the diseased kidney in order to maintain normal total renal function. For this reason, successful anatomic repair of the ureteral obstruction of such a kidney may fail to improve its powers of waste elimination.

If both kidneys are equally hydronephrotic, a strong stimulus is continually being exerted on both to maintain maximum function. This is also true of a hydronephrotic solitary kidney. Consequently, the return of function in
Figure 11–6. Pathogenesis of unilateral hydronephrosis. Progressive changes in ureter and kidney secondary to obstructing calculus (arrowheads). As the right kidney undergoes gradual destruction, the left kidney gradually enlarges (compensatory hypertrophy).
Physiologic Explanation of Symptoms of Bladder Neck Obstruction

The following hypothesis has been proposed to explain the syndrome known as “prostatism,” which occurs with progressive vesical obstruction:

The bladder, like the heart, is a hollow muscular organ that receives fluid and forcefully expels it. And, like the heart, it reacts to an increasing work load by going through the successive phases of compensation and finally decompensation.

Normally, contraction of the detrusor muscle and the trigone pulls the bladder neck open and forms a funnel through which the urine is expelled. The intravesical pressure generated in this instance varies between 20 and 40 cm of water; this force further widens the bladder neck.

With bladder neck obstruction, hypertrophy of the vesical musculature develops, allowing the intravesical voiding pressure to rise to 50–100 cm or more of water in order to overcome the increased outlet resistance. Despite this, the encroaching prostate appears to interfere with the mechanisms that ordinarily open the internal orifice. Also, the contraction phase may not last long enough for all of the urine to be expelled; “exhaustion” of the muscle occurs prematurely. The refractory phase then sets in, and the detrusor is temporarily unable to respond to further stimuli. A few minutes later, voiding may be initiated again and completed.

A. Compensatory Phase

1. Stage of irritability—In the earliest stages of obstruction of the bladder neck, the vesical musculature begins to hypertrophy. The force and size of the urinary stream remain normal because the balance is maintained between the expelling power of the bladder and urethral resistance. During this phase, however, the bladder appears to be hypersensitive. As the bladder is distended, the need to void is felt. In patients with a normal bladder, these early urges can be inhibited, and the bladder relaxes and distends to receive more urine. However, in patients with a hypertrophied detrusor, the contraction of the detrusor is so strong that it virtually goes into spasm, producing the symptoms of an irritable bladder. The earliest symptoms of bladder neck obstruction, therefore, are urgency (even to the point of incontinence) and frequency, both day and night.

2. Stage of compensation—As the obstruction increases, further hypertrophy of the muscle fibers of the bladder occurs, and the power to empty the bladder completely is thereby maintained. During this period, in addition to urgency and frequency, the patient notices hesitancy in initiating urination while the bladder develops contractions strong enough to overcome resistance at the bladder neck. The obstruction causes some loss in the force and size of the urinary stream and the stream becomes slower as vesical emptying nears completion (exhaustion of the detrusor as it nears the end of the contraction phase).

B. Decompensatory Phase

If vesical tone becomes impaired or if urethral resistance exceeds detrusor power, some degree of decompensation (imbalance) occurs. The contraction phase of the vesical muscle becomes too short to completely expel the contents of the bladder, and some urine remains in the bladder (residual urine).

1. Acute decompensation—The tone of the compensated vesical muscle can be temporarily emissary by rapid filling of the bladder (high fluid intake) or by overstretching of the detrusor (postponement of urination though the urge is felt). This may cause increased difficulty of urination, with marked hesitancy and the need for straining to initiate urination; a very weak and small stream; and termination of the stream before the bladder completely empties (residual urine). Acute and sudden complete urinary retention may also occur.

2. Chronic decompensation—As the degree of obstruction increases, a progressive imbalance between the power of the bladder musculature and urethral resistance develops. Therefore, it becomes increasingly difficult to expel all the urine during the contraction phase of the detrusor. The symptoms of obstruction become more marked. The amount of residual urine gradually increases, and this diminishes the functional capacity of the bladder. Progressive frequency of urination is noted. On occasion, as the bladder decompensates, it becomes overdistended and attenuated. It may contain 1000–3000 mL of urine. It loses its power of contraction, and overflow (paradoxic) incontinence results.

Clinical Findings

A. Symptoms

1. Lower and mid tract (urethra and bladder)—Symptoms of obstruction of the lower and mid tract are typified by the symptoms of urethral stricture, benign
prostatic hyperplasia, neurogenic bladder, and tumor of the bladder involving the vesical neck. The principal symptoms are hesitancy in starting urination, lessened force and size of the stream, and terminal dribbling; hematuria, which may be partial, initially, with stricture or total with prostatic obstruction or vesical tumor; and burning on urination, cloudy urine (due to complicating infection), and occasionally acute urinary retention.

2. Upper tract (ureter and kidney)—Symptoms of obstruction of the upper tract are typified by the symptoms of ureteral stricture or ureteral or renal stone. The principal complaints are pain in the flank radiating along the course of the ureter, gross total hematuria (from stone), gastrointestinal symptoms, chills, fever, burning on urination, and cloudy urine with onset of infection, which is the common sequel to obstruction or vesicoureteral reflux. Nausea, vomiting, loss of weight and strength, and pallor are due to uremia secondary to bilateral hydronephrosis. A history of vesicoureteral reflux in childhood may be significant. Obstruction of the upper tract may be silent even when uremia supervenes.

B. Signs

1. Lower and midtract—Palpation of the urethra may reveal induration about a stricture. Rectal examination may show atony of the anal sphincter (damage to the sacral nerve roots) or benign or malignant enlargement of the prostate. Vesical distention may be found.

Although observation of the force and caliber of the urinary stream affords a rough estimate of maximum flow rate, the rate can be measured accurately with a urine flowmeter or, even more simply, by the following technique: Have the patient begin to void. When observed maximum flow has been reached, interpose a container to collect the urine and simultaneously start a stopwatch. After exactly 5 seconds, remove the container. The flow rate in milliliters per second can easily be calculated. The normal urine flow rate is 20–25 mL/s in males and 25–30 mL/s in females. Any flow rate under 15 mL/s should be regarded with suspicion. A flow rate under 10 mL/s is indicative of obstruction or weak detrusor function. Flow rates associated with anatomic neurogenic (neuropathic) bladder (diminished detrusor power), or with urethral stricture or prostatic obstruction (increased urethral resistance) may be as low as 3–5 mL/s. A cystometrogram can differentiate between these 2 causes of impaired flow rate. After definitive treatment of the cause, the flow rate should return to normal.

In the presence of a vesical diverticulum or vesicoureteral reflux, although detrusor power is normal, the urinary stream may be impaired because of the diffusion of intravesical pressure into the diverticulum and vesicoureteral junction as well as the urethra. Excision of the diverticulum or repair of the vesicoureteral junctions leads to efficient expulsion of urine via the urethra.

2. Upper tract—An enlarged kidney may be discovered by palpation or percussion. Renal tenderness may be elicited if infection is present. Cancer of the cervix may be noted; it may invade the base of the bladder and occlude one or both ureteral orifices, or its metastases to the iliac lymph nodes may compress the ureters. A large pelvic mass (tumor, pregnancy) can displace and compress the ureters. Children with advanced urinary tract obstruction (usually due to posterior urethral valves) may develop ascites. Rupture of the renal fornices allows leakage of urine retroperitoneally; with rupture of the bladder, urine may pass into the peritoneal cavity through a tear in the peritoneum.

C. Laboratory Findings

Anemia may be found secondary to chronic infection or in advanced bilateral hydronephrosis (stage of uremia). Leukocytosis is to be expected in the acute stage of infection. Little, if any, elevation of the white blood count accompanies the chronic stage.

Large amounts of protein are usually not found in the obstructive uropathies. Casts are not common from hydronephrotic kidneys. Microscopic hematuria may indicate renal or vesical infection, tumor, or stone. Pus cells and bacteria may or may not be present. In the presence of significant bilateral hydronephrosis, urine flow through the renal tubules is slowed. Thus, urea is significantly reabsorbed but creatinine is not. Blood chemistry therefore reveals a urea-creatinine ratio well above the normal 10:1.

D. X-Ray Findings (Figure 11–7)

A plain film of the abdomen may show enlargement of renal shadows, calcific bodies suggesting ureteral or renal stone, or tumor metastases to the bones of the spine or pelvis. Metastases in the spine may be the cause of spinal cord damage (neuropathic bladder); if they are osteoblastic, they are almost certainly from cancer of the prostate.

Excretory urograms reveal almost the entire story unless renal function is severely impaired. They are more informative when obstruction is present because the radiopaque material is retained. These urograms demonstrate the degree of dilatation of the pelves, calyces, and ureters. The point of ureteral stenosis is revealed. Segmental dilatation of the lower end of a ureter implies the possibility of vesicoureteral reflux (Figure 11–7), which can be revealed by cystography. The cystogram may show trabeculation as an irregularity of the vesical outline and may show diverticula. Vesical tumors, nonopaque stones, and large intravesical prostatic lobes may cause radiolucent shadows. A film taken immediately after voiding will show residual urine. Few tests that are as simple and inexpensive give the physician so much information.

Retrograde cystography shows changes of the bladder wall caused by distal obstruction (trabeculation, diverticula) or demonstrates the obstructive lesion itself (enlarged prostate, posterior urethral valves, cancer of the bladder). If
Figure 11–7. Changes in bladder, ureters, and kidneys caused by obstruction. Upper left: Cystogram showing benign prostatic enlargement and multiple diverticula. Arrows point to femoral hernia that probably developed as a result of straining to urinate. Upper right: Pregnancy. Significant dilatation and elongation of upper right ureter due to compression at the pelvic line. Left side normal. Lower left: Excretory urogram, 70 minutes after injection. Advanced right hydronephrosis secondary to ureteropelvic obstruction. Mild ureteropelvic obstruction on left. Lower right: Stone in left ureter (at arrow) with mild hydronephrosis.
the ureterovesical valves are incompetent, ureteropyelograms are obtained by reflux. Retrograde urograms may show better detail than the excretory type, but care must be taken not to overdistend the passages with too much opaque fluid; small hydronephroses can be made to look quite large. The degree of ureteral or ureterovesical obstruction can be judged by the degree of delay of drainage of the radiopaque fluid instilled. Computed tomography scanning and sonography can also help determine the extent of dilatation and parenchymal atrophy.

E. ISOTOPE SCANNING

In the presence of obstruction, the radioisotope renogram may show depression of both the vascular and secretory phases and a rising rather than a falling excretory phase due to retention of the radiopaque urine in the renal pelvis. The $^{131}$I activity recorded on the gamma camera will show that the isotope is poorly taken up, slowly transported through the parenchyma, and accumulated in the renal pelvis.

F. INSTRUMENTAL EXAMINATION

Exploration of the urethra with a catheter or other instrument is a valuable diagnostic measure. Passage may be blocked by a stricture or tumor. Spasm of the external sphincter may make passage difficult. Passage of the catheter immediately after voiding allows estimation of the amount of residual urine in the bladder. Residual urine is common in bladder neck obstruction (enlarged prostate), cystocele, and neurogenic (neuropathic) bladder. Residual urine is usually absent with urethral stricture, even though the urinary stream may be markedly impaired.

Measurement of vesical tone by means of cystometry is helpful in diagnosing neurogenic bladder and in differentiating between bladder neck obstruction and vesical atony. Inspection of the urethra and bladder by means of cystoscopy and panendoscopy may reveal the primary obstructive agent. Catheters may be passed to the renal pelvis and urine specimens obtained. The function of each kidney may be measured and retrograde ureteropyelograms can be obtained.

G. INTERVENTIONAL URORADIOLOGY

If there is doubt about the presence of true obstruction, either the Whitaker test or an isotope renogram can be done. However, Whitaker and Buxton-Thomas (1984) have shown that neither test is without error.

Differential Diagnosis

A thorough examination usually leaves no doubt about the diagnosis. The differential diagnosis under these circumstances is rarely difficult. If seemingly simple infection does not respond to medical therapy or if infection recurs, obstruction, a foreign body, or vesicoureteral reflux is the probable cause, and complete study of the urinary tract is indicated.

Complications

Stagnation of urine leads to infection, which then may spread throughout the entire urinary system. Once established, infection is difficult and at times impossible to eradicate even after the obstruction has been relieved.

Often the invading organisms are urea-splitting (Proteus, staphylococci), which causes the urine to become alkaline. Calcium salts precipitate and form bladder or kidney stones more easily in alkaline urine. If both kidneys are affected, the result may be renal insufficiency. Secondary infection increases renal damage.

Pyonephrosis is the end stage of a severely infected and obstructed kidney. The kidney is functionless and filled with thick pus. At times, a plain film of the abdomen may show an air urogram caused by gas liberated by infecting organisms.

Treatment

A. RELIEF OF OBSTRUCTION

Treatment of the main causes of obstruction and stasis (benign prostatic hyperplasia, cancer of the prostate, neurogenic bladder, ureteral stone, posterior urethral valves, and ureteral stenosis) is described in detail elsewhere in this book.

1. Lower tract obstruction (distal to the bladder)—With patients in whom secondary renal or ureterovesical damage (reflux in the latter) is minimal or nonexistent, correction of the obstruction is sufficient. If significant reflux is demonstrated and does not subside spontaneously after relief of obstruction, surgical repair may be needed. Repair becomes imperative if there is considerable hydro-nephrosis in addition to reflux. Preliminary drainage of the bladder by an indwelling catheter or other means of diversion (eg, loop ureterostomy) is indicated in order to preserve and improve renal function. If, after a few months of drainage, reflux persists, the incompetent ureterovesical junction should be surgically repaired.

2. Upper tract obstruction (above the bladder)—If tortuous, kinked, dilated, or atonic ureters have developed secondary to lower tract obstruction (so that they are themselves obstructive), vesical drainage will not protect the kidneys from further damage; the urine proximal to the obstruction must be diverted by nephrostomy or ureterostomy. The kidneys then may regain some function. Over a period of many months, the ureter may become less tortuous and less dilated; its obstructive areas may open up. If radiopaque material instilled into the nephrostomy tube passes readily to the bladder, it may be possible to remove the nephrostomy tube. If obstruction or reflux
persists, surgical repair is indicated. Permanent urinary diversion (e.g., ureteroileal conduit) may be necessary.

If one kidney has been irreversibly damaged, as measured by kidney function tests, urography, sonography, computed tomography scan, or scintigraphy, nephrectomy may be necessary.

**B. ERADICATION OF INFECTION**

Once the obstruction is removed, every effort should be made to eradicate infection. If the infection has been severe and prolonged, antibiotics may fail to sterilize the urinary tract.

**Prognosis**

No simple statement can be made about the prognosis in this group of patients. The outcome depends on the cause, site, degree, and duration of the obstruction. The prognosis is also definitely influenced by complicating infection, particularly if the infection has been present for a long time.

If renal function is fair to good, if the obstruction or other causes of stasis can be corrected, and if complicating infection can therefore be eradicated, the prognosis is generally excellent.

**REFERENCES**


Under normal circumstances, the ureterovesical junction allows urine to enter the bladder but prevents urine from regurgitating into the ureter, particularly at the time of voiding. In this way, the kidney is protected from high pressure in the bladder and from contamination by infected vesical urine. When this valve is incompetent, the chance for development of urinary infection is significantly enhanced, and pyelonephritis is inevitable. In significant cases especially in children, pyelonephritis—acute, chronic, or healed—is secondary to vesicoureteral reflux.

**ANATOMY OF THE URETEROVESICAL JUNCTION**

An understanding of the causes of vesicoureteral reflux requires knowledge of the anatomy of the ureterovesical valve. Anatomic studies performed by Hutch (1972) and by Tanagho and Pugh (1963) (Figure 12–1) are incorporated into the following discussion.

**Mesodermal Components**

The mesodermal component, which arises from the Wolffian duct, is made up of 2 parts that are innervated by the sympathetic nervous system:

**A. THE URETER & THE SUPERFICIAL TRIGONE**

The smooth musculature of the renal calyces, pelvis, and extravesical ureter is composed of helically oriented fibers that allow for peristaltic activity. As these fibers approach the vesical wall, they are reoriented into the longitudinal plane. The ureter passes obliquely through the vesical wall; the intravesical ureteral segment is thus composed of longitudinal muscle fibers only and therefore cannot undergo peristalsis. As these smooth-muscle fibers approach the ureteral orifice, those that form the roof of the ureter swing to either side to join those that form its floor. They then spread out and join equivalent muscle bundles from the other ureter and also continue caudally, thus forming the superficial trigone. The trigone passes over the neck of the bladder, ending at the verumontanum in the male and just inside the external urethral meatus in the female. Thus, the ureterotrigonal complex is one structure. Above the ureteral orifice, it is tubular; below that point, it is flat.

**B. WALDEYER’S SHEATH & THE DEEP TRIGONE**

Beginning at a point about 2–3 cm above the bladder, an external layer of longitudinal smooth muscle surrounds the ureter. This muscular sheath passes through the vesical wall, to which it is connected by a few detrusor fibers. As it enters the vesical lumen, its roof fibers diverge to join its floor fibers, which then spread out, joining muscle bundles from the contralateral ureter and forming the deep trigone, which ends at the bladder neck.

**Endodermal Component**

The vesical detrusor muscle bundles are intertwined and run in various directions. As they converge on the internal orifice of the bladder, however, they tend to become oriented into 3 layers:

**A. INTERNAL LONGITUDINAL LAYER**

The internal longitudinal layer continues into the urethra submucosally and ends just inside the external meatus in the female and at the caudal end of the prostate in the male.

**B. MIDDLE CIRCULAR LAYER**

The middle circular layer is thickest anteriorly and stops at the vesical neck.

**C. OUTER LONGITUDINAL LAYER**

The muscle bundles of the outer longitudinal layer take a circular and spiral course about the external surface of the female urethra and are incorporated within the peripheral prostatic tissue in the male. They constitute the true vesicourethral sphincter.

The vesical detrusor muscle is innervated by the parasympathetic nerves (S₂–S₄).

**PHYSIOLOGY OF THE URETEROVESICAL JUNCTION**

Although many investigators had suspected that normal trigonal tone tended to occlude the intravesical ureter, it remained for Tanagho et al (1965) to prove it. Using non-refluxing dogs, they demonstrated the following:

1. Interruption of the continuity of the trigone resulted in reflux. An incision was made in the trigone 3
mm below the ureteral orifice, resulting in an upward and lateral migration of the ureteral orifice with shortening of the intravesical ureter. Reflux was demonstrable. After the incision healed, reflux ceased.

(2) Unilateral lumbar sympathectomy resulted in paralysis of the ipsilateral trigone. This led to lateral and superior migration of the ureteral orifice and reflux.

(3) Electrical stimulation of the trigone caused the ureteral orifice to move caudally, thus lengthening the intravesical ureter. This maneuver caused a marked rise in resistance to flow through the ureterovesical junction. Ureteral efflux of urine ceased. Intravenous injection of epinephrine caused the same reaction. On the other hand, isoproterenol caused the degree of occlusion to drop below normal. If the trigone was incised, however, electrical stimulation of the trigone or the administration of epinephrine failed to increase ureteral occlusive pressure.

(4) During gradual filling of the bladder, intravesical pressure increased only slightly, whereas pressure within the intravesical ureter rose progressively—owing, apparently, to increasing trigonal stretch. A few seconds before the expected sharp rise in intravesical pressure generated for voiding, the closure pressure in the intravesical ureter rose sharply and was maintained for 20 seconds after detrusor contraction had ceased. This experiment demonstrated that ureterovesical competence is independent of detrusor action and is governed by the tone of the trigone, which contracts vigorously just before voiding, thus helping to open and funnel the vesical neck. At the same time, significant pull is placed on the intravesical ureter, so that it is occluded during the period when intravesical pressure is high. During the voiding phase, there is naturally no efflux of ureteral urine.

One may liken this function to the phenomenon of the Chinese thimble: The harder the finger (trigone) pulls, the tighter the thimble (intravesical ureter) becomes. Conversely, a deficient pull may lead to incomplete closure of the ureterovesical junction.

It was concluded from these experiments that normal ureterotrigonal tone prevents vesicoureteral reflux. Electrical or pharmacologic stimulation of the trigone caused increased occlusive pressure in the intravesical ureter and increased resistance to flow down the ureter, whereas incision or paralysis of the trigone led to reflux. The theory that ureterovesical competence was maintained by intravesical pressure compressing the intravesical ureter against its backing of detrusor muscle was thereby disproved.

Biopsy of the trigone (and the intravesical ureter) in patients with primary reflux revealed marked deficiency in the development of its smooth muscle (Figure 12–2). Electrical stimulation of such a trigone caused only a minor contraction of the ureterotrigonal complex. This work led to the conclusion that the common cause of reflux, particularly in children, is congenital attenuation of the ureterotrigonal musculature.

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**VESICOURETERAL REFUX**

**CAUSES**

The major cause of vesicoureteral reflux is attenuation of the trigone and its contiguous intravesical ureteral musculature. Any condition that shortens the intravesical ureter
VESICOURETERAL REFLUX

may also lead to reflux, but this is less common. Familial vesicoureteral reflux has been observed by a number of authors. It appears to be a genetic trait.

Congenital Causes

A. TRIGONAL WEAKNESS (PRIMARY REFLUX)

Trigonal weakness is by far the most common cause of ureteral reflux. It is most often seen in young children, more common in girls than boys. Reflux in adults—usually women—probably represents the same congenital defect. Weakness of one side of the trigone leads to a decrease in the occlusive pressure in the ipsilateral intravesical ureter. Diffuse ureterotrigonal weakness causes bilateral reflux.

It is postulated that ureteral trigonal weakness is related to the development of the ureteral bud on the mesonephric duct. It is known that the ureter acquires its musculature from its cranial end caudally, so that if a segment is muscularily deficient, it is deficient in its most caudal part. It is also postulated that if the ureter is too close to the urogenital sinus on the mesonephric duct, it will join the latter relatively early in embryonic life, before acquiring adequate mesenchymal tissue around itself to be differentiated later into proper trigonal musculature as well as lower ureter. This embryologic hypothesis explains all the known features of refluxing ureters: their muscular weakness, their lateral placement on the bladder base with a very short submucosal segment, and their usual association with weak ureteral musculature and gaping ureteral orifices (which, in severe cases, ensures a golf-hole endoscopic appearance at their junction with the bladder wall). It also explains why, in duplicated systems, if there is only one refluxing unit, it is the upper orifice (which originated closer to the urogenital sinus on the mesonephric duct and thus has the least muscular development).

In the normal state, the intravesical ureterotrigonal muscle tone exerts a downward pull, whereas the extravesical ureter tends to pull cephalad (Figure 12–3). If trigonal development is deficient, not only is its occlusive power diminished but the ureteral orifice tends to migrate upward toward the ureteral hiatus. The degree of this retraction relates to the degree of incompetence of the

Figure 12–2. Histology of the trigone in primary reflux. 

Figure 12–3. A: Small ureterocele developing in a duplicated system (where it always involves a lower ureteral orifice). B: Expansion of submucosal segment leads to lifting and angulation of ipsilateral lower pole ureteral orifice. Duplicated system ureteroceles are rarely so small. (Diagrammatic representation.) (Reproduced, with permission, from Tanagho, EA: Ureteroceles: Embryogenesis, pathogenesis and management. J Cont Educ Urol [Feb] 1979;18:13.)
junction (Figure 12–4). If the ureteral orifice lies over the ureteral hiatus in the bladder wall (so-called golf-hole orifice), it is completely incompetent. The degree of incompetence is judged by the findings on excretory urography and cystography and the cystoscopic appearance of the ureteral orifices.

B. FAMILIAL REFLUX

There appears to be genetic predisposition for reflux. The reported prevalence of vesicoureteral reflux (VUR) among siblings of index patients with reflux has ranged from 4.7% to 51%, which is significantly higher than the incidence of reflux in the general population (1%) (Ataei et al, 2004). In addition, the incidence of reflux varies among nationalities and races. The exact form of genetic transmission has yet to be delineated.

C. URETERAL ABNORMALITIES

1. Complete ureteral duplication (Figure 12–5)—
The intravesical portion of the ureter to the upper renal segment is usually of normal length, whereas that of the ureter to the lower pole is abnormally short; this orifice is commonly incompetent. However, Stephens (1957) demonstrated that the musculature of the superiorly placed orifice is attenuated, which further contributes to its weakness.

2. Ectopic ureteral orifice—Single ureter or one of a pair may open well down on the trigone, at the vesical neck, or in the urethra. In this instance, vesicoureteral reflux is the rule. This observation makes it clear that the length of the intravesical ureter is not the sole factor in reflux. Such intravesical ureteral segments are usually devoid of smooth muscle. Thus, they have no occlusive force.

Figure 12–4. Histology of the various grades of submucosal muscular weakness of the ureteral orifice. (See also Figure 12–9.) A: Normal. Minimal deficiency. (Cone orifice.) B: More marked muscular weakness. (Stadium orifice.) C: Marked muscular deficiency. (Horseshoe orifice.) D: Extreme muscular deficiency. Only a few muscle fibers can be seen; the rest is collagen tissue.
3. Ureterocele—A ureterocele involving a single ureter rarely allows reflux, but this lesion usually involves the ureter that drains the upper pole of a duplicated kidney. Because the ureteral orifice is obstructed, the intramural ureter becomes dilated. This increases the diameter of the ureteral hiatus, further shortening the intravesical segment of the other ureter, which therefore may become incompetent. Resection of the ureterocele usually causes its ureter to reflux freely as well.

**Voiding Dysfunction**

Abnormal voiding habits have been associated with reflux. Toilet-trained children, in particular girls, may alter their bladder function by inhibiting their urge to void. This can result in abnormally high voiding pressure, bladder overactivity, and poor bladder compliance. These changes in bladder dynamics can either induce the development of primary reflux or prevent its resolution (Greenfield and Wan, 2000). In addition, alterations in the bowel function (eg, constipation) can cause further deterioration in bladder function and consequently the development or persistence of primary reflux (Bower, Yip, and Yeung, 2005).

**Vesical Trabeculation**

Occasionally, a heavily trabeculated bladder may be associated with reflux. The causes include spastic neurogenic bladder and severe obstruction distal to the bladder. These lesions, however, are associated with trigonal hypertrophy as well; the resultant extra pull on the ureterotrigonal muscle tends to protect the junction from incompetence. In a few such cases, however, the vesical mucosa may protrude through the ureteral hiatus just above the ureter to form a diverticulum, or saccule (Figure 12–6). The resulting dilatation of the hiatus shortens the intravesical segment; reflux may then occur.

**Edema of the Vesical Wall Secondary to Cystitis**

As noted previously, valves vary in their degrees of incompetence. A “borderline” junction may not allow reflux when the urine is sterile, but valvular function may be impaired when cystitis causes associated edema involving the trigone and intravesical ureter. In addition, the abnormally high voiding pressure may lead to reflux, in which case secondary pyelonephritis may ensue. After cure of the infection, cystography again reveals no reflux. It is believed that a completely normal junction will not decompensate even under these circumstances.

It has been shown that pyelonephritis of pregnancy is associated with vesicoureteral reflux. Many patients give a history of urinary tract infections during childhood. The implication is that they “outgrew” reflux at puberty, but if bacteriuria becomes established during pregnancy, their
“borderline” valves may become incompetent. This condition may be aggravated by the hormones of pregnancy, which may contribute to a further loss of tone of the ureterotrigonal complex. After delivery, reflux is usually no longer demonstrable (Hutch and Amar, 1972).

Eagle-Barrett (Prune Belly) Syndrome

The Eagle-Barrett syndrome is a relatively rare condition in which there is failure of normal development of the abdominal muscles and the smooth muscle of the ureters and bladder. Bilateral cryptorchidism is the rule. At times, talipes equinovarus and hip dislocation are also noted. Because the smooth muscle of the ureterotrigonal complex is deficient, reflux is to be expected; advanced hydroureteronephrosis is therefore found.

Iatrogenic Causes

Certain operative procedures may lead to either temporary or permanent ureteral regurgitation.

A. PROSTATECTOMY

With any type of prostatectomy, the continuity of the superficial trigone is interrupted at the vesical neck. If the proximal trigone moves upward, temporary reflux may occur. This mechanism may account for the high fever (and even bacteremia) that is sometimes observed when the catheter is finally removed. Fortunately, in 2–3 weeks the trigone again becomes anchored and reflux ceases.

Preexisting trigonal hypertrophy (due to prostatic obstruction) helps to compensate for the effect of trigonal interruption; thus, reflux may never occur.

B. WEDGE RESECTION OF THE POSTERIOR VESICAL NECK

Wedge resection of the posterior vesical neck, often ill-advised when performed in conjunction with plastic revision of the vesical neck for supposed vesical neck stenosis or dysfunction, may also upset trigonal continuity and allow reflux.

C. URETERAL MEATOTOMY

Extensive ureteral meatotomy may be followed by reflux. Fortunately, however, limited incision of the roof of the intravesical ureter divides few muscle fibers, since the fibers have left the roof to join muscle fibers on the floor. Wide resection for treatment of vesical cancer is often followed by ureteral reflux.

D. RESECTION OF URETEROCELE

If the ureteral hiatus is widely dilated, this procedure is often followed by reflux.

Contracted Bladder

A bladder that is contracted secondary to interstitial cystitis, tuberculosis, radiotherapy, carcinoma, or schistosomiasis may be associated with ureteral reflux.

COMPLICATIONS

Vesicoureteral reflux damages the kidney through one or both of 2 mechanisms: (1) pyelonephritis and (2) hydroureteronephrosis.

Pyelonephritis

Vesicoureteral reflux is one of the common contributing factors leading to the development of cystitis, particularly in females. When reflux is present, bacteria reach the kidney and the urinary tract cannot empty itself completely, so infection is perpetuated. Pyelonephritis is discussed in more detail in Chapter 13.
Hydroureteronephrosis
(See also Chapter 11)
Dilation of the ureter, renal pelvis, and calyces is usually observed in association with reflux (Figure 12–7), sometimes to an extreme degree (Figure 12–8). In males, because they have a relatively long segment of sterile urethra, such changes are often seen in the absence of infection. Sterile reflux is less damaging than infected reflux. There are 3 reasons for the dilatation:

1) Increased work load: The ureter is meant to transport the urine secreted by the kidney to the bladder only once. In the presence of reflux, variable amounts of urine go back and forth, and the work load may be doubled, quadrupled, or increased 10-fold or even more. Eventually, the ureter is not able to transport the increased volume of urine, and stasis and dilatation result.

![Figure 12–7. Excretory urogram with changes that imply right vesicoureteral reflux. Upper left: Excretory urogram showing normal right urogram and a ureter that is mildly dilated and remains full through its entire length. The ureteral change implies reflux. Upper right: Cystogram demonstrates the reflux. Note, now, the degree of dilatation of the ureter, pelvis, and calyces. Lower left: Excretory urogram shows bilateral hydroureteronephrosis with pyelonephritic scarring. These findings imply the presence of reflux. Lower right: Voiding cystourethrogram. Free reflux bilaterally.]
Figure 12–8. Cystograms revealing vesicoureteral reflux. Upper left: Saccule at right ureterovesical junction. Upper right: Meningomyelocele. Reflux with severe bilateral hydroureteronephrosis; serum creatinine, 0.6 mg/dL; phenolsulfonphthalein excretion, 5% in 1 hour. Lower left: Post-prostatectomy patient with reflux on left and bilateral saccules. Lower right: Ten-year-old boy with meningomyelocele. Bladder has been emptied. Impairment of drainage at ureterovesical junctions is demonstrated. (Courtesy of Hutch JA, Amar AD: Vesicoureteral Reflux and Pyelonephritis. Appleton-Century-Crofts, 1972.)
(2) **High hydrostatic pressure:** The ureter is protected from the high pressures of the urinary bladder by a competent ureterovesical junction. If there is free reflux, the high intravesical pressure is directly transmitted to the ureteral and pelvic walls, which results in marked stretching and dilation.

(3) **Weak ureteral musculature:** In reflux, the ureteral wall is invariably deficient in musculature to some degree. The more severe the reflux, the more apparent the muscular deficiency. Some cases show more massive dilatation than others. The properly muscularized ureter is better able to resist and compensate for overwork and hydrostatic pressure than the muscularily deficient ureter. The latter tends to undergo further dilatation once it is subjected to any increased intraluminal pressure.

Whether sterile reflux is harmful is the subject of controversy. My colleagues and I believe there is conclusive evidence that severe sterile reflux can lead to parenchymal damage. Pyelointerstitial backflow or pyelotubular backflow under the high pressures of reflux (not infrequently seen during cystographic studies) leads to extravasation of urine in the interstitium of the kidney. The presence of urine in any interstitium will result in a marked inflammatory response with cellular infiltration, resulting finally in fibrosis and scarring. On a long-term basis, this can lead to parenchymal changes indistinguishable from pyelonephritic scarring caused by inflammation due to bacterial infection. This damage may be termed reflux nephropathy. If severe, it will produce parenchymal damage serious enough to lead to end-stage kidney disease.

Ransley’s studies (1976) indicate that intrarenal reflux is more likely to occur in the presence of flat, concave, or compound papillae, because their collecting ducts tend to open with an increase in intrapelvic pressure and reflux. Papillae prone to reflux are more commonly seen in the polar segments of the kidney. Normal papillae might also permit intrarenal reflux if flattened as a result of the changes due to reflux.

Intravesical pressure is transmitted through the incompetent ureteral orifice. This back pressure is quite high at the time of voiding. Furthermore, the ureteropelvic and ureterovesical junctions are less distensible than the rest of the ureter. Either junction may have trouble passing the normal amount of secreted urine plus the refluxed urine; functional obstruction may result. A common cause of ureteropelvic and ureterovesical “obstruction” is vesicoureteral reflux. Such changes indicate the need for cystography.

**INCIDENCE**

Vesicoureteral reflux occurs in 25–40% (Fanos and Cataldi, 2004) of children with urinary tract infection but in only 8% of adults with bacteriuria. This discrepancy is explained by the fact that girls usually have pyelonephritis, whereas women usually have cystitis only. Bacteriuria does not always imply pyelonephritis.

The fairly competent (borderline) valve refluxes only during an acute attack of cystitis. Since cystography is performed in such cases only after the infection has been eradicated, the incidence of reflux found on cystography is abnormally low. On the other hand, reflux is demonstrable in 85% of patients whose excretory urograms reveal significant changes typical of healed pyelonephritis.

When infection associated with reflux occurs during the first few weeks of life, many patients are septic and uremic. Most are boys with posterior urethral valves. After 1 year of age, the female-male ratio of children with infection and reflux is approximately 3–4:1.

**CLINICAL FINDINGS**

A history compatible with acute pyelonephritis implies the presence of vesicoureteral reflux. This is most commonly seen in females, particularly young girls. Persistence of recurrent “cystitis” may suggest the possibility of reflux. Such patients often have asymptomatic low-grade pyelonephritis.

**Symptoms Related to Reflux**

**A. SYMPTOMATIC PYELONEPHRITIS**

The usual symptoms in adults are chills and high fever, renal pain, nausea and vomiting, and symptoms of cystitis. In children, only fever, vague abdominal pains, and sometimes diarrhea are apt to occur.

**B. ASYMPTOMATIC PYELONEPHRITIS**

The patient may have no symptoms whatsoever. The incidental findings of pyuria and bacteriuria may be the only clues. This fact points up the need for a screening urinalysis in all children.

**C. SYMPTOMS OF CYSTITIS ONLY**

In cases of symptoms of cystitis only, bacteriuria is resistant to antimicrobial drugs, or infection quickly recurs following treatment. These patients may have reflux with asymptomatic chronic pyelonephritis.

**D. RENAL PAIN ON VOIDING**

Surprisingly, renal pain on voiding is a rare complaint in patients with vesicoureteral reflux.

**E. UREMIA**

The last stage of bilateral reflux is uremia due to destruction of the renal parenchyma by hydronephrosis or pyelonephritis (or both). The patient often adjusts to renal insufficiency and may appear quite healthy. Many renal transplants are performed in patients whose kidneys have deteriorated secondarily to reflux and accompanying infection. Early diagnosis, based on careful urinalysis, would
have led to the proper diagnosis in childhood. Progressive pyelonephritis is, with few exceptions, preventable.

**F. HYPERTENSION**

In the later stages of atrophic pyelonephritis, a significant incidence of hypertension is observed.

**Symptoms Related to the Underlying Disease**

The clinical picture is often dominated by the signs and symptoms of the primary disease.

**A. URINARY TRACT OBSTRUCTION**

Young girls may have hesitancy in initiating the urinary stream and an impaired or intermittent stream secondary to spasm of the periurethral striated muscle (see Distal Urethral Stenosis in Chapter 41). In males, the urinary stream may be slow as a result of posterior urethral valves (infants) or prostatic enlargement (men over age 50).

**B. SPINAL CORD DISEASE**

The patient may have a serious neurogenic disease such as paraplegia, quadriplegia, multiple sclerosis, or meningomyelocele. Symptoms may be limited to those of neurogenic bladder: incontinence of urine, urinary retention or large residual volume, and vesical urgency.

**Physical Findings**

During an attack of acute pyelonephritis, renal tenderness may be noted. Its absence, however, does not rule out chronic renal infection. Palpation and percussion of the suprapubic area may reveal a distended bladder secondary to obstruction or neurogenic disease. The finding of a hard midline mass deep in the pelvis in a male infant is apt to represent a markedly thickened bladder caused by posterior urethral valves. Examination may reveal a neurologic deficit compatible with a paretic bladder.

**Laboratory Findings**

The most common complication of reflux, particularly in females, is infection. Bacteriuria without pyuria is not uncommon. In males, the urine may be sterile because of the long, sterile urethra.

The serum creatinine may be elevated in the advanced stage of renal damage, but it may be normal even when the degree of reflux and hydronephrosis is marked (Figure 12–8, upper right).

**X-Ray Findings**

The plain film may reveal evidence of spina bifida, meningomyelocele, or absence of the sacrum and thus point to a neurologic deficit. Even in vesicoureteral reflux, excretory urograms may be normal, but usually one or more of the following clues to the presence of reflux is noted (Figure 12–7): (1) a persistently dilated lower ureter, (2) areas of dilatation in the ureter, (3) ureter visualized throughout its entire length, (4) presence of hydronephrosis with a narrow juxtavesical ureteral segment, or (5) changes of healed pyelonephritis (caliceal clubbing with narrowed infundibula or cortical thinning). A normal intravenous urogram does not rule out reflux.

The presence of ureteral duplication suggests the possibility of reflux into the lower pole of the kidney. In this case, hydronephrosis or changes compatible with pyelonephritic scarring may be seen. Abnormality of the upper segment of a duplicated system can be caused by the presence of an ectopic ureteral orifice with reflux or by obstruction secondary to a ureterocele.

Reflux is diagnosed by demonstration of its existence with one of the following techniques: simple or delayed cystography, voiding cystourethrography, or voiding cinel fluoroscopy. Radionuclide scanning can be used: 1 mCi of $^{99m}$Tc is instilled into the bladder along with sterile saline solution, and the gamma camera will reveal ureteral reflux.

Reflux can be demonstrated by a technique using indigotindisulfonate sodium (indigo carmine), a blue dye. The bladder is filled with sterile water containing 5 mL of indigo carmine per 100 mL, after which the patient voids and the bladder is thoroughly flushed out with sterile water. The ureteral orifices are then viewed cystoscopically for blue-tinged efflux. This technique has the advantage that no ionizing radiation is used, and its efficiency is equal to that of voiding cystourethrography. In general, reflux demonstrable only with voiding implies a more competent valve than does reflux that occurs at low pressures. As has been pointed out, failure to demonstrate reflux on one study does not rule out intermittent reflux.

The voiding phase of the cystogram may reveal changes compatible with distal urethral stenosis with secondary spasm of the voluntary periurethral muscles in girls (Figure 39–1) or changes diagnostic of posterior urethral valves in young boys.

**Instrumental Examination**

**A. URETHRAL CALIBRATION**

In females, urethral calibration using bougies à boule should be done. Distal urethral stenosis is almost routinely found in young girls who have urinary infection. Dilatation of the ring of stenosis is an important step in improving the hydrodynamics of voiding: lowering intravesical voiding pressure and eliminating the presence of residual vesical urine (see Chapter 39). Less commonly, urethral stenosis is discovered in women and should be treated.
B. CYSTOSCOPY

Most young girls with reflux have smooth-walled or only slightly trabeculated bladders. Chronic cystitis, ureteral duplication, or ureterocele may be evident. An orifice may be ectopic and be found at the bladder neck or even in the urethra. As the bladder is filled, a small diverticulum may form on the roof of the ureteral orifice (Figure 12–6). These findings imply the possibility of reflux. The major contribution of cystoscopy is to allow study of the morphologic characteristics of the ureteral orifice and its position in relation to the vesical neck (Figure 12–9). However, cystoscopy should not be performed as a part of the workup for reflux. Rather, it can be performed prior to surgical correction to help define the anatomic and rule out other bladder and ureteral anomalies.

1. Morphology—The orifice of a normal ureter has the appearance of a volcanic cone. That of a slightly weaker valve looks like a football stadium; an even weaker one has the appearance of a horseshoe with its open end pointing toward the vesical neck. The completely incompetent junction has a golf-hole orifice that lies over the ureteral hiatus.

2. Position—By and large, the more defective the appearance of the ureteral orifice, the farther from the vesical neck it lies. The degree of lateralization of the orifice reflects the degree of ureterotrigonal deficiency.

DIFFERENTIAL DIAGNOSIS

Functional (nonocclusive) vesicoureteral obstruction may cause changes similar to those suggesting the presence of reflux on excretory urography. Multiple cystograms fail to show reflux. Tanagho, Smith, and Guthrie (1970) showed that this congenital obstruction is due to an abundance of circularly oriented smooth-muscle fibers in the ureteral musculature at this point. Its action is sphincteric.

Significant obstruction distal to the vesical neck leads to hypertrophy of both the detrusor and trigonal muscles. The latter exert an exaggerated pull on the intravesical ureter and thus cause functional obstruction (Tanagho and Meyers, 1965). Hydroptcronephrosis is therefore to be expected; vesicoureteral reflux is uncommon.

Other lesions that may cause hydronephrosis without reflux include low ureteral stone, occlusion of the ureter by cervical or prostatic cancer, urinary tract tuberculosis, and schistosomiasis.

TREATMENT

It is impossible to give a concise and definitive discourse on the treatment of vesicoureteral reflux because of the many factors involved and because there is no unanimity of opinion among urologists on this subject. In general, probably more than half of the cases of primary reflux that occur in children can be controlled by nonsurgical means; the rest require some form of operative procedure. Adults with reflux usually require vesicoureteroplasty.

Medical Treatment

A. INDICATIONS

In the majority of the cases, children with primary reflux are initially treated medically, since there is a chance of spontaneous resolution. Positive predictors for reflux resolution include: unilateral reflux, the lower grades of reflux, the earlier age of presentation, and male gender. A boy with posterior urethral valves may cease to have reflux once these valves are destroyed.

In a woman who occasionally develops acute pyelonephritis following intercourse but whose urine quickly clears on antimicrobial therapy, reflux will probably be controlled if she takes steps to prevent vesical infections. This is particularly true if reflux cannot be demonstrated cystographically when her urine is sterile. The maintenance of sterile urine allows her “borderline” valve to remain competent.

B. METHODS OF TREATMENT

Dilation of the ring of distal urethral stenosis in young girls or of posterior urethral valves in boys usually gives excellent results, reducing intravesical voiding pressure and abolishing vesical residual urine and reflux.

Urinary infection should be definitively treated with antimicrobial drugs, after which chronic suppressive therapy should be continued until the reflux has resolved or surgically corrected.
Children with reflux often have voiding dysfunction due to thin-walled bladders and do not perceive the normal urge to void when the bladder is full. Further detrusor tone is lost with overfilling, increasing the likelihood of residual urine. Such children should “void by the clock” every 3–4 hours whether they have the urge or not. Vesical residual urine may then be minimized.

C. EVALUATION OF SUCCESS OF MEDICAL TREATMENT

Cystograms should be repeated every 12–18 months. Excretory urography or nuclear renal scan should be performed periodically to be sure that renal deterioration does not occur. About half of children with reflux are cured by medical treatment.

Surgical Treatment

A. INDICATIONS

Reflux caused by the following abnormalities will not disappear spontaneously: (1) ectopic ureteral orifice, (2) ureteral duplication, (3) ureteroceles associated with ureteral duplication and reflux into the uninvolved ureter, (4) golf-hole ureteral orifice, and (5) low-pressure reflux with significant hydronephrosis.

Surgery is indicated (1) if it is not possible to keep the urine sterile and reflux persists; (2) if acute pyelonephritis recurs despite a strict medical regimen and chronic suppressive antimicrobial therapy; or (3) if increased renal damage is demonstrated by serial excretory urograms or nuclear scan.

B. TYPES OF SURGICAL TREATMENT

In cases of markedly impaired kidney function and massively dilated ureters, preliminary urinary diversion may be required to improve renal function and to allow dilated ureters to regain tone, after which definitive relief of obstruction (eg, posterior urethral valves) and ureterovesicoplasty can be performed at the optimum time. Some patients with irreversible lesions causing reflux (eg, meningomyelocele) or badly damaged and atonic ureters may require permanent diversion of the urine (ie, ureteroileocutaneous anastomosis).

1. Temporary urinary diversion—If refluxed urine drains freely into the bladder, cystostomy (or an indwelling urethral catheter in girls) may prove helpful. If the ureters are dilated and kinked, a low redundant loop can be brought to the skin. The ureter is opened at this point and urine collected into an ileostomy bag. Later, the loop and the section of ureter distal to it can be resected and the ureter proximal to the loop reimplanted into the bladder. Nephrostomy may be necessary if there is no ureteral redundancy.

2. Permanent urinary diversion—If it is felt that successful ureterovesicoplasty cannot be accomplished, a Bricker type of diversion is indicated. If renal function is poor and the ureters are widely dilated and atonic, ureteroileocutaneous diversion may be the procedure of choice.

3. Other surgical procedures—

a. If reflux is unilateral, with the affected kidney badly damaged and the other kidney normal, nephrectomy is indicated.

b. If one renal pole of a duplicated system is essentially functionless, heminephrectomy with removal of its entire ureter should be done. If there is moderate hydronephrosis of one renal pole with duplication, an alternative is anastomosis of the dilated ureter or pelvis to the normal ureter or pelvis. The remainder of the dilated refluxing ureter should be removed.

c. In unilateral reflux, anastomosis of the lower end of the refluxing ureter into the side of its normal mate (transureteroureterostomy) has a few proponents.

4. Definitive repair of ureterovesical junction (ureterovesicoplasty)—

a. Principles of repair (Tanagho, 1970)—

1. Resect the lower 2–3 cm of the ureter in which the muscle is underdeveloped.

2. Free up enough extravesical ureter so that an intravesical segment 2.5 cm long can be formed.

3. Place the intravesical ureter in a submucosal position.

4. Suture the wall of the new ureteral orifice to the cut edge of the trigonal muscle.

b. Types of operation—The following procedures satisfy the preceding principles and have been successful in a high percentage of cases: suprahiatal repair, increasing the length of intravesical ureter above the level of the ureteral hiatus (Paquin, 1959; Politano and Leadbetter, 1958); infrahiatal repair, the advancement procedures of Hutch (1963) and Glenn and Anderson (1967); combined supravaginal and infrahiatal repair, which is the most attractive; and transtrigonal repair (Cohen, 1975).

If the ureters are unduly tortuous, the redundant portions must be resected. If they are widely dilated, the lower ends must be tailored to a more normal size.

c. Results of ureterovesicoplasty—About 93% of patients no longer show reflux after ureterovesicoplasty. About 3% develop ureterovesical stenosis that requires reoperation. At least 75% have and maintain sterile urine without antimicrobial drugs 3–6 months after surgery. Many patients in whom bacteriuria persists have cystitis only. This has been demonstrated by the finding that renal urine specimens collected by ureteral catheters are sterile. Febrile attacks cease. Considering that only the most severe and advanced cases are submitted to surgical repair, these are impressive results, and they exceed by far the cure rates reported when only antimicrobial drugs are used (10–15%). This operation is rightly considered one of the most significant accomplishments of modern urology.
**C. Subureteric Transurethral Injection (STING)**

More recently, endoscopic treatment of reflux has become popular treatment for reflux. A biocompatible material such as Deflux (a mixture of dextranomer microspheres and non-animal-stabilized hyaluronic acid) can be injected into the intramural ureter. The bulking agent allows for the coaptation of the ureteral orifice and intramural ureter (Aaronson 2005) in order to prevent reflux. The success rate for the STING procedure varies from 70% to 90%, depending upon the grade of the reflux treated. While it does not approximate the cure rate from open surgery, Technical improvements and risk factors for failure are gradually being identified that are aimed at improving performance.

**PROGNOSIS**

In patients with reflux who are judged to have fairly competent valves, conservative therapy as outlined previously is highly successful in the cure of the reflux and therefore of infection.

Patients with very incompetent ureterovesical valves subjected to surgical repair also have an excellent prognosis. A few children, however, have such badly damaged urinary tracts when finally submitted to diagnostic procedures that little help other than permanent urinary diversion can be offered.

**REFERENCES**

**General**


Bacterial Infections of the Genitourinary Tract

Hiep T. Nguyen, MD

Urinary tract infection (UTI) is a term that is applied to a variety of clinical conditions ranging from the asymptomatic presence of bacteria in the urine to severe infection of the kidney with resultant sepsis. UTI is one of the more common medical problems. It is estimated that 150 million patients are diagnosed with a UTI yearly, resulting in at least $6 billion in health care expenditures (Stamm and Norrby, 2001). UTIs are at times difficult to diagnose; some cases respond to a short course of a specific antibiotic, while others require a longer course of a broad-spectrum antibiotic. Accurate diagnosis and treatment of a UTI is essential to limit its associated morbidity and mortality and avoid prolonged or unnecessary use of antibiotics. Advances in our understanding of the pathogenesis of UTI, the development of new diagnostic tests, and the introduction of new antimicrobial agents have allowed physicians to appropriately tailor specific treatment for each patient.

EPIDEMIOLOGY

The epidemiology of UTI grouped by age and sex is shown in Table 13–1. In newborns up to 1 year of age, bacteriuria is present in 2.7% of boys and 0.7% in girls (Wettergren, Jodal, and Jonasson, 1985). The incidence of UTI in uncircumcised males is higher than in circumcised males (1.12% compared to 0.11%) during the first 6 months of life (Wiswell and Roscelli, 1986). In children between 1 and 5 years of age, the incidence of bacteriuria in girls increases to 4.5%, while it decreases in boys to 0.5% (Randolph and Greenfield, 1964). Most UTIs in children younger than 5 years are associated with congenital abnormalities of the urinary tract, such as vesicoureteral reflux or obstruction. The incidence of bacteriuria remains relatively constant in children 6–15 years of age. However, the UTIs in these children are more likely to be associated with functional abnormalities of the urinary tract, such as dysfunctional voiding. During adolescence, the incidence of UTI significantly increases (to 20%) in young women, while remaining constant in young men (Sanford, 1975).

Approximately 7 million cases of acute cystitis are diagnosed yearly in young women (Shappert, 1999); this likely is an underestimate of the true incidence of UTI because at least 50% of all UTIs do not come to medical attention. The major risk factors for women 16–35 years of age are related to sexual intercourse and diaphragm use. Later in life, the incidence of UTI increases significantly for both males and females. For women between 36 and 65 years of age, gynecologic surgery and bladder prolapse appear to be important risk factors. In men of the same age group, prostatic hypertrophy/obstruction, catheterization, and surgery are relevant risk factors. For patients older than 65 years, the incidence of UTI continues to increase in both sexes. Incontinence and chronic use of urinary catheters are important risk factors in these patients. In those younger than 1 year and those older than 65 years, the morbidity and mortality of UTI are the greatest (Shortliffe and McCue, 2002).

Based on data from the Urologic Diseases in North America Project, the overall lifetime prevalence of UTI was estimated to be 14,000 per 100,000 men (Griebling, 2005a) and 53,000 per 100,000 women (Griebling, 2005b). Overall medical expenditures for the treatment of UTIs in the United States were estimated to be $1 billion for men (Griebling, 2005a) and $2.5 billion for women (Griebling, 2005b). The increased costs in treatment of UTIs for women is primarily due to an increase in the trend toward using fluoroquinolones as a first-line therapy for UTI. UTIs occurred in 2.4–2.8% of children. In this patient population, UTIs resulted in more than 1.1 million physician visits annually, accounting for 0.7% of the doctor visits (Freedman, 2005).

PATHOGENESIS

Bacterial Entry

Understanding of the mode of bacterial entry, host susceptibility factors, and bacterial pathogenic factors is essential to tailoring appropriate treatment for the diverse clinical manifestations of UTI. There are 4 possible modes of bacterial entry into the genitourinary tract. It is generally accepted that periurethral bacteria ascending into the urinary tract causes most UTI. Most cases of pyelonephritis are caused by the ascent of bacteria from the bladder, through the ureter and into the renal parenchyma. Conse-
quently, the short nature of the female urethra combined with its close proximity to the vaginal vestibule and rectum likely predisposes women to more frequent UTIs than men (Nicolle et al., 1982).

Other modes of bacterial entry are uncommon causes of UTI. Hematogenous spread can occur in immunocompromised patients and in neonates. *Staphylococcus aureus*, *Candida* species, and *Mycobacterium tuberculosis* are common pathogens that travel through the blood to infect the urinary tract. Lymphophagocytic spread through the rectal, colonic, and periuterine lymphatics has been postulated as a cause for UTI; however, currently there is little scientific support to suggest that dissemination of bacteria through lymphatic channels plays a role in the pathogenesis of UTI. Direct extension of bacteria from adjacent organs into the urinary tract can occur in patients with intraperitoneal abscesses or vesicovestibular or vesicovaginal fistulas. Relapsing infection from an inadequately treated focus in the prostate or kidney may seed other parts of the urinary tracts.

**Host Defenses**

Host factors have an essential role in the pathogenesis of UTI. Unobstructed urinary flow with the subsequent washout of ascending bacteria is essential in preventing UTI. In addition, the urine itself has specific characteristics (its osmolality, urea concentration, organic acid concentration, and pH) that inhibit bacterial growth and colonization (Sobel, 1997). It also contains factors that inhibit bacterial adherence, such as Tamm-Horsfall glycoprotein (THG; Duncan, 1988; Pak et al., 2001; Wagenlehner et al., 2005). It has been observed that the severity of the bacterialuria and the degree of inflammatory changes in the urinary tract were much greater in THG-deficit mice, suggesting that THG helps eliminate bacterial infection from the urinary tract and acts as a general host-defense factor against UTI (Raffi et al., 2005). Urinary retention, stasis, or reflux of urine into the upper urinary tract can promote bacterial growth and subsequent infection. Consequently, any anatomic or functional abnormalities of the urinary tract that impede urinary flow can increase the host’s susceptibility to UTI. These abnormalities include obstructive conditions at any level of the urinary tract, neurologic diseases affecting the function of the lower urinary tract, diabetes, and pregnancy. Similarly, the presence of foreign bodies (such as stones, catheters, and stents) allows the bacteria to hide from these host defenses.

The epithelium lining the urinary tract not only provides a physical barrier to infection but also have the capacity to recognize bacteria in order to innate host defenses. The urothelial cells express toll-like receptors (TLRs) that upon engagement by specific bacterial components lead to production of inflammatory mediators (Chowdhury, 2004). In response to the presence of bacteria, cells lining the urinary tract secrete chemoattractants such as interleukin-8 to recruit neutrophils to the area and limit tissue invasion (Frendeus et al., 2001). Specific serum and urinary antibodies are produced by the kidney to enhance bacterial opsonization and phagocytosis and to inhibit bacterial adherence. The protective role of both cellular and humoral-mediated immunity in preventing UTIs remains unclear; deficiency in B-cell or T-cell function has not been associated with the increased frequency of UTI or altered the course of the infection (Schaeffer, 2001; Svanborg Eden et al., 1988). However, it should be noted that the same host defense mechanisms that help to prevent/limit the infection (such as the inflammatory responses) can lead to cell and tissue damage. In the kidneys, cell damage and subsequent development of scarring may lead to pathological conditions such as hypertension, preclampsia during pregnancy and renal dysfunction and failure (Jahnukainen, Chen, and Celsi, 2005).

Many studies have demonstrated that there is selectivity in bacterial adherence to cells lining the urinary tract, and the degree of adherence correlates with colonization and infection. Women with recurrent UTIs have higher adherence of bacteria to their mucosal cells in vitro compared to women who never had an infection (Navas et al., 1994). The increased adherence may be due to having more binding sites for bacterial adhesins on their mucosal cells. Alternatively, these patients may not secrete soluble compounds, which normally compete for the same receptors that bind bacterial adhesins. Blood group antigens may constitute one group of these soluble compounds that inhibit bacterial adherence (Lomberg et al., 1986). These findings would suggest a genetic predisposition for UTI.

Other important host factors include the normal flora of the periurethral area or the prostate and the presence of
vesicoureteral reflux. In women, the normal flora of the periurethral area composed of organisms such as lactobacillus provides a defense against the colonization of uropathogenic bacteria (Osset et al, 2001). Alterations in the periurethral environment (such as changes in the pH or estrogen levels or the use of antibiotics) can damage the periurethral flora, allowing uropathogens to colonize and subsequently to infect the urinary tract (Schaefler et al, 1999). In men, the prostate secretes fluid containing zinc, which has potent antimicrobial activity (Fair, Couch, and Wehner, 1976). Finally, in children, the presence of vesicoureteral reflux does not increase their susceptibility to UTI but does allow bacteria to be inoculated into the upper tract and the infection to progress.

Aging is associated with an increased susceptibility to UTI, in part because of the increased incidence of obstructive uropathy in men (Matsumoto, 2001; Nicolle, 2002) and alteration in the vaginal and periurethral flora from menopause in women (Foxman et al, 2001). Other causes include soiling of the perineum from fecal incontinence, neuromuscular diseases, increased instrumentation, and bladder catheterization (Ronald, 2002).

**Bacterial Pathogenic Factors**

Not all bacteria are capable of adhering to and infecting the urinary tract. Of the many strains of *Escherichia coli*, the uropathogens belong to a limited number of O, K, and H serogroups. They have increased adherence properties to uroepithelial cells (Blanco et al, 1996; Hovance and Gorzynski, 1980; Orskov et al, 1982), resistance to the bactericidal activity of human serum (Bjorksten and Kaijser, 1978), production of hemolysin (Hughes et al, 1983; Koronakis and Hughes, 1996), and the increased expression of K capsular antigen (Whitfield and Roberts, 1999). The ability of *E. coli* to adhere to epithelial cells is mediated by ligands located on the tips of the bacterial fimbiae (pili). The ligands bind to glycolipids or glycoprotein receptors on the surface membrane of uroepithelial cells. The pili are classified by their ability to cause hemagglutination and the type of sugar that can block this process. P pili, which can agglutinate human blood, bind to glycolipid receptors on uroepithelial cells, erythrocytes (P-blood group antigens), and renal tubular cells (Svenson et al, 1983). Type 1 pili, which can agglutinate guinea pig blood, bind to mannoside residues on uroepithelial cells (Ofek et al, 2000). P pili are observed in over 90% of the *E. coli* strains causing pyelonephritis but <20% of the strains causing lower UTIs (Kallenius et al, 1981; Roberts et al, 1997). In contrast, type 1 pili may help bacteria to adhere to bladder mucosa (Connell et al, 1996; Martinez et al, 2000). Most uropathogenic *E. coli* have both types of pili. Once attachment to the uroepithelial cells occurs, other bacterial pathogenic factors become important. Most uropathogenic *E. coli* strains produce hemolysin, which initiates tissue invasion and makes iron available for the infecting pathogens (Hughes et al, 1983; Koronakis and Hughes, 1996). The presence of K antigen on the invading bacteria protects them from phagocytosis by neutrophils (Bortolussi et al, 1979; Evans et al, 1981). These factors allow the infecting pathogens to escape the various host defenses (Svanborg et al, 1996).

Recently, it has been observed that many bacteria such as *E. coli* have the ability to invade into the host cells, acting as opportunistic intracellular pathogens (Bower, Eto, and Mulvey, 2005). Cytotoxic necrotizing factor, Afa/Dr adhesions and type 1 pili have been shown to promote invasion into the host cells. The intracellular bacteria mature into biofilms, creating pod-like bulges on the urothelial surface (Anderson et al, 2003). The pods contain bacteria encased in a polysaccharide-rich matrix surrounded by a protective shell of uroplakin. The ability of the uropathogenic bacteria to transiently invade, survive, and multiply within the host cells and to create biofilms on genitourinary tract tissues may provide a mechanism for the persistence and recurrence of UTIs.

**CAUSATIVE PATHOGENS**

Most UTIs are caused by a single bacterial species. At least 80% of the uncomplicated cystitis and pyelonephritis are due to *E. coli*, with most of pathogenic strains belonging to the O serogroups (Orskov et al, 1982). Other less common uropathogens include *Klebsiella*, *Proteus*, and *Enterobacter* spp. and enterococci. In hospital-acquired UTIs, a wider variety of causative organisms is found, including *Pseudomonas* and *Staphylococcus* spp. (Wagenlehner and Naber, 2000); UTIs caused by *S. aureus* often result from hematogenous dissemination. Group B beta-hemolytic streptococci can cause UTIs in pregnant women (Wood and Dillon, 1981). *S. saprophyticus*, once often thought of as urinary contaminants, can cause uncomplicated UTIs in young women (Hovelius and Mardh, 1984). In children, the causative bacterial spectrum is slightly different from that of adults, with *Klebsiella* and *Enterobacter* spp. being more common causes of UTI (Jeena et al, 1996; Ronald, 2002; Schlager, 2001). Anaerobic bacteria, lactobacilli, corynebacteria, streptococci (not including enterococci) and *S. epidermidis* are found in normal periurethral flora. They do not commonly cause UTIs in healthy individuals and are considered common urinary contaminants.

**DIAGNOSIS**

The diagnosis of UTI is sometimes difficult to establish and relies on urinalysis and urine culture. Occasionally, localization studies may be required to identify the source of the infection. Most often, the urine is often obtained from a voided specimen. In children who are not toilet-trained, a urine collection device, such as a bag, is placed over the genitalia, and the urine is cultured from the
bagged specimen. These 2 methods of urine collection are easy to obtain, but potential contamination from the vagina and perirectal area may occur. There is a high false-positive rate, especially from bagged specimens (Al-Orifi et al, 2000). Suprapubic aspiration avoids potential contamination; however, due to its invasiveness, it is rarely used except in children and selected patients. Urine obtained from a urinary catheter is less invasive than a suprapubic aspiration and is less likely to be contaminated than that from a voided specimen. If a patient has an indwelling catheter, a urine specimen should be obtained from the collection port on the catheter.

Urinalysis

The urine can be immediately evaluated for leukocyte esterase, a compound produced by the breakdown of white blood cells (WBCs) in the urine. Urinary nitrite is produced by reduction of dietary nitrates by many gram-negative bacteria. Esterase and nitrite can be detected by a urine dipstick and are more reliable when the bacterial count is >100,000 colony-forming units (CFU) per milliliter. Microscopic examination of the urine for WBCs and bacteria is performed after centrifugation. When bacteria counts are >100,000 CFU/mL, bacteria can be detected microscopically (Jenkins, Fenn, and Matsen, 1986). More than 3 WBCs per high-power field suggests a possible infection. The sensitivity and specificity of these tests are shown in Table 13–2. The urinary nitrite test is highly specific but not sensitive, whereas the other 3 tests have a sensitivity and specificity approximately 80%. A combination of these tests may help to identify those patients in whom urine culture will be positive. Conversely, when esterase, nitrite, blood, and protein is absent in urine, <2% of the urine samples will be positive by culture, providing a >98% negative predictive value and a sensitivity of 98% (Patel et al, 2005).

Urine Culture

The gold standard for identification of UTI is the quantitative culture of urine for specific bacteria. The urine should be collected in a sterile container and cultured immediately after collection. When this is not possible, the urine can be stored in the refrigerator for up to 24 hours. The sample is then diluted and spread on culture plates. Each bacterium will form a single colony on the plates. The number of colonies is counted and adjusted per milliliter of urine (CFU/mL). Defining the CFU/mL that represents clinically significant infection can be difficult. It is dependent on the method of collection, the sex of the patient, and the type of bacteria isolated (Table 13–3). Traditionally, >100,000 CFU/mL is used to exclude contamination. However, studies have clearly demonstrated that clinically significant UTI can occur with <100,000 CFU/mL bacteria in the urine (Stamm et al, 1982).

Localization Studies

Occasionally, it is necessary to localize the site of infection. For upper urinary tract localization (Lorentz, 1979), the bladder is irrigated with sterile water and a ureteral catheter is placed into each ureter. A specimen is collected from the renal pelvis. Culture of this specimen will indicate whether infection in the upper urinary tract is present. In men, infection in the lower urinary tract can be differentiated (Figure 13–1) (Meares and Stamey, 1968). A specimen is collected at the beginning of the void and represents possible infection in the urethra. A midstream specimen is next collected and represents possible infection in the bladder. The prostate is then massaged and the patient is asked to

Table 13–2. Sensitivity and Specificity of Urinalysis.

<table>
<thead>
<tr>
<th>Tests</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Esterase</td>
<td>83 (67–94)</td>
<td>78 (64–92)</td>
</tr>
<tr>
<td>Nitrite</td>
<td>53 (15–82)</td>
<td>98 (90–100)</td>
</tr>
<tr>
<td>E or N</td>
<td>93 (90–100)</td>
<td>72 (58–91)</td>
</tr>
<tr>
<td>White blood cells</td>
<td>73 (32–100)</td>
<td>81 (45–98)</td>
</tr>
<tr>
<td>Bacteria</td>
<td>81 (16–99)</td>
<td>83 (11–100)</td>
</tr>
<tr>
<td>Any above</td>
<td>99.8 (99–100)</td>
<td>70 (60–92)</td>
</tr>
</tbody>
</table>

Table 13–3. Probability of UTI Based on Urine Culture.

<table>
<thead>
<tr>
<th>Collection</th>
<th>CFU</th>
<th>Probability of Infection (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Suprapubic</td>
<td>Gram neg. any</td>
<td>&gt;99</td>
</tr>
<tr>
<td></td>
<td>Gram pos. &gt;1000</td>
<td></td>
</tr>
<tr>
<td>Catheterization</td>
<td>&gt;10^5</td>
<td>95</td>
</tr>
<tr>
<td></td>
<td>10^4–5</td>
<td>Likely</td>
</tr>
<tr>
<td></td>
<td>10^3–4</td>
<td>Repeat</td>
</tr>
<tr>
<td></td>
<td>&lt;10^3</td>
<td>Unlikely</td>
</tr>
<tr>
<td>Clean catch</td>
<td>Male: &gt;10^6</td>
<td>Likely</td>
</tr>
<tr>
<td></td>
<td>Female: 3 specimens: &gt;10^5</td>
<td>95</td>
</tr>
<tr>
<td></td>
<td>2 specimens: &gt;10^5</td>
<td>90</td>
</tr>
<tr>
<td></td>
<td>1 specimen: &gt;10^5</td>
<td>80</td>
</tr>
<tr>
<td></td>
<td>5 x 10^4–10^5</td>
<td>Repeat</td>
</tr>
<tr>
<td></td>
<td>1–5 x 10^4 sympt.</td>
<td>Repeat</td>
</tr>
<tr>
<td></td>
<td>1–5 x 10^4 nonsympt.</td>
<td>Unlikely</td>
</tr>
<tr>
<td></td>
<td>&lt;10^5</td>
<td>Unlikely</td>
</tr>
</tbody>
</table>

CFU, colony-forming unit; gram neg., gram-negative; gram pos., gram-positive; nonsympt., nonsymptomatic; sympt., symptomatic.
void again. This specimen represents possible infection of the prostate.

**ANTIBIOTICS**

Treatment with antimicrobial agents has minimized the morbidity and mortality associated with UTIs. The goal in treatment is to eradicate the infection by selecting the appropriate antibiotics that would target specific bacterial susceptibility. However, choosing the appropriate antimicrobial agents is often difficult. Many antibiotics are available, and the lowest effective dose and length of therapy are not well defined. Many conventions for the treatment of UTI are arbitrary. The general principles for selecting the appropriate antibiotics include consideration of the infecting pathogen (antibiotic susceptibility, single-organism versus poly-organism infection, pathogen versus normal flora, community versus hospital-acquired infection); the patient (allergies, underlying diseases, age, previous antibiotic therapy, other medications currently taken, outpatient versus inpatient status, pregnancy); and the site of infection (kidney versus bladder versus prostate). Because most antimicrobials are cleared from the body by the liver or the kidney, certain antimicrobial agents need to be adjusted in the presence of liver or renal diseases (Table 13–4). Table 13–5 lists the common uropathogens and the recommended oral and intravenous antimicrobial agents for treatment. Table 13–6 lists the common sites of UTI, the recommended treatment, and the duration of therapy. In patients with recurrent UTIs or those who are at risk for UTI (such as children with vesicoureteral reflux), prophylactic antibiotics may be used. Table 13–7 lists common prophylactic regimens.

**Trimethoprim-Sulfamethoxazole**

Trimethoprim-sulfamethoxazole (TMP-SMX) is commonly used to treat many UTIs, except those caused by Enterococcus and Pseudomonas spp. It interferes with the bacterial metabolism of folate. TMP-SMX is highly effective and relatively inexpensive. Adverse reactions occur in 6–8% of patients using this medication; they include hypersensitivity reactions, rashes, gastrointestinal upset, leukopenia, thrombocytopenia, and photosensitivity. TMP-SMX should not be used in patients who have a folic acid deficiency state, glucose-6-phosphate dehydrogenase deficiency, or AIDS, or in pregnant patients. It is the most frequently prescribed antibiotic for uncomplicated UTI (Huang and Stafford, 2002). Recently, the use of TMP-SMX has declined due to the increased incidence of bacterial resistance (Brown, Freeman, and Foxman, 2002) and physicians’ preference for other newer antibiotics (Huang and Stafford, 2002).

**Fluoroquinolones**

Fluoroquinolones have a broad spectrum of activity, especially against gram-negative bacteria. Although they have
### Table 13–5. Recommended Antimicrobial Agents for Common Genitourinary Pathogens.

<table>
<thead>
<tr>
<th>Bacteria</th>
<th>Oral Therapy</th>
<th>Parenteral Therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gram-positive cocci</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Staphylococcus aureus</em></td>
<td>Nafcillin, nitrofurantoin, ciprofloxacin</td>
<td>Nafcillin, vancomycin</td>
</tr>
<tr>
<td><em>Staphylococcus epidermidis</em></td>
<td>Ampicillin, nitrofurantoin, ciprofloxacin</td>
<td>Ampicillin, penicillin G</td>
</tr>
<tr>
<td><em>Staphylococcus saprophyticus</em></td>
<td>Ampicillin, nitrofurantoin, ciprofloxacin</td>
<td>Ampicillin, penicillin G</td>
</tr>
<tr>
<td><em>Streptococcus</em>, group D</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>S. faecalis</em> (enterococci)</td>
<td>Ampicillin, nitrofurantoin</td>
<td>Ampicillin plus gentamicin</td>
</tr>
<tr>
<td><em>S. bovis</em></td>
<td>Penicillin G, ampicillin</td>
<td>Ampicillin, vancomycin</td>
</tr>
<tr>
<td><em>Streptococcus</em>, group B</td>
<td>Ampicillin, cephalosporin</td>
<td>Ampicillin, cephalosporin</td>
</tr>
<tr>
<td><strong>Gram-negative cocci</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Neisseria gonorrhoeae</em></td>
<td>Ciprofloxacin plus doxycycline</td>
<td>Ceftriaxone</td>
</tr>
<tr>
<td><strong>Gram-negative rods</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Escherichia coli</em></td>
<td>TMP-SMX, ciprofloxacin, nitrofurantoin</td>
<td>Gentamicin</td>
</tr>
<tr>
<td><em>Enterobacter</em> spp.</td>
<td>TMP-SMX, ciprofloxacin, nitrofurantoin</td>
<td>Gentamicin plus piperacillin</td>
</tr>
<tr>
<td><em>Gardnerella vaginalis</em></td>
<td>Metronidazole, ampicillin</td>
<td>Metronidazole</td>
</tr>
<tr>
<td><em>Klebsiella</em> spp.</td>
<td>TMP-SMX, ciprofloxacin</td>
<td>Gentamicin plus cephalosporin</td>
</tr>
<tr>
<td><em>Proteus</em> spp.</td>
<td>Ampicillin, TMP-SMX, ciprofloxacin</td>
<td>Ampicillin, gentamicin</td>
</tr>
<tr>
<td><em>Pseudomonas aeruginosa</em></td>
<td>Carbenicillin, tetracycline, ciprofloxacin</td>
<td>Gentamicin plus piperacillin</td>
</tr>
<tr>
<td><em>Serratia</em> spp.</td>
<td>TMP-SMX, carbenicillin</td>
<td>TMP-SMX, amikacin</td>
</tr>
<tr>
<td><strong>Other pathogens</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chlamydiae</td>
<td>Tetracycline, erythromycin</td>
<td>Tetracycline, erythromycin</td>
</tr>
<tr>
<td>Mycoplasmas, ureaplasmas</td>
<td>Tetracycline, erythromycin</td>
<td>Tetracycline, erythromycin</td>
</tr>
<tr>
<td>Obligate anaerobes</td>
<td>Metronidazole, clindamycin</td>
<td>Metronidazole, clindamycin</td>
</tr>
</tbody>
</table>

TMP-SMX, trimethoprim plus sulfamethoxazole.

### Table 13–6. Recommended Antimicrobial Agents and Duration of Therapy Based on the Type of UTI.

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Pathogen</th>
<th>Choice of Antibiotics</th>
<th>Duration of Therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cystitis</td>
<td><em>E. coli</em></td>
<td>1st: TMP-SMX</td>
<td>1–3 days</td>
</tr>
<tr>
<td></td>
<td><em>Klebsiella</em></td>
<td>2nd: Fluoroquinolone</td>
<td></td>
</tr>
<tr>
<td></td>
<td><em>Proteus</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pyelonephritis</td>
<td><em>E. coli</em></td>
<td>1st: Fluoroquinolone</td>
<td>7–10 days</td>
</tr>
<tr>
<td></td>
<td><em>Proteus</em></td>
<td>2nd: 2nd generation cephalosporin</td>
<td></td>
</tr>
<tr>
<td></td>
<td><em>Klebsiella</em></td>
<td>3rd: Aminopenicillin/BLI</td>
<td></td>
</tr>
<tr>
<td></td>
<td><em>Enterobacteria</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Complicated UTI</td>
<td><em>E. coli</em></td>
<td>1st: Fluoroquinolone</td>
<td>3–5 days</td>
</tr>
<tr>
<td></td>
<td><em>Enterococci</em></td>
<td>2nd: Aminopenicillin/BLI</td>
<td>after afebrile</td>
</tr>
<tr>
<td></td>
<td><em>Pseudomonas</em></td>
<td>3rd: 3rd generation cephalosporin</td>
<td></td>
</tr>
<tr>
<td></td>
<td><em>Staphylococci</em></td>
<td>Aminoglycosides</td>
<td></td>
</tr>
<tr>
<td>Prostatitis</td>
<td><em>E. coli</em></td>
<td>1st: Fluoroquinolone</td>
<td>Acute: 2 weeks</td>
</tr>
<tr>
<td></td>
<td><em>Enterobacteria</em></td>
<td>2nd: 2nd generation cephalosporin</td>
<td>Chronic:</td>
</tr>
<tr>
<td></td>
<td><em>Pseudomonas</em></td>
<td>3rd: 3rd generation cephalosporin</td>
<td>4–6 weeks</td>
</tr>
<tr>
<td></td>
<td><em>Enterococci</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Epididymitis</td>
<td><em>E. coli</em></td>
<td>1st: Fluoroquinolone</td>
<td>2 weeks</td>
</tr>
<tr>
<td></td>
<td><em>Enterobacteria</em></td>
<td>2nd: 2nd generation cephalosporin</td>
<td></td>
</tr>
<tr>
<td></td>
<td><em>Chlamydia</em></td>
<td>1st: Doxycycline</td>
<td></td>
</tr>
<tr>
<td></td>
<td><em>Ureaplasma</em></td>
<td>2nd: Macrolide</td>
<td></td>
</tr>
</tbody>
</table>

BLI, beta-lactamase inhibitor; TMP-SMX, trimethoprim plus sulfamethoxazole.

(Adapted from table 2 from Wagenlehner FM, Naber KG. Hospital-acquired urinary tract infections. J Hosp Infect 2000;46:171.)
Nitrofurantoin, 50 or 100 mg daily  
Nitrofurantoin macrocrystals, 100 mg daily  
TMP-SMX, 40/200 mg daily  
Cephalaxin, 250 mg daily  
Ciprofloxacin, 250 mg daily  
Trimethoprim, 100 mg daily

Table 13–7. Prophylactic Antibiotic Regimens.

adequate activity against Staphylococci species, fluoroquinolones do not have good activity against Streptococci species and anaerobic bacteria. They interfere with the bacterial DNA gyrase, preventing bacterial replication. Although they are highly effective in the treatment of UTI, fluoroquinolones are relatively expensive. Adverse reactions are infrequent and include mild gastrointestinal effects, dizziness, and lightheadedness. Fluoroquinolones should not be used in patients who are pregnant and should be used judiciously in children because of potential damage to developing cartilage. Due to their broad spectrum of activity, fluoroquinolones have gained popularity in the empiric treatment of both uncomplicated and complicated UTIs (Schaeffer, 2002).

Nitrofurantoin

Nitrofurantoin has good activity against most gram-negative bacteria (except for Pseudomonas and Proteus spp.), Staphylococci, and Enterococci species. It inhibits bacterial enzymes and DNA activity. Nitrofurantoin is highly effective in the treatment of UTI and is relatively inexpensive. Adverse reactions are relatively common and include gastrointestinal upset, peripheral polyneuropathy, and hepatotoxicity. Long-term use may result in pulmonary hypersensitivity reaction and interstitial changes. With increasing awareness of this antibiotic and its activity against common uropathogens, nitrofurantoin usage in the treatment of uncomplicated UTIs has increased from 14% to 30% in the past 5 years (Huang and Stafford, 2002).

Aminoglycosides

Aminoglycosides are commonly used in the treatment of complicated UTI. They are highly effective against most gram-negative bacteria. When combined with ampicillin, they are effective against enterococci. They inhibit bacterial DNA and RNA synthesis. The principal adverse effects of aminoglycosides are nephrotoxicity and ototoxicity. Aminoglycosides are primarily used in patients with complicated UTIs who require intravenous antibiotics (Santucci and Krieger, 2000). Aminoglycosides can be given as a single daily dosing; this regimen is directed toward obtaining higher peak and lower trough levels in order to achieve more effective microbial killing while reducing toxicity (Carapetis et al, 2001).

Cephalosporins

Cephalosporins have good activity against most uropathogens (Garcia-Rodriguez and Munoz Bellido, 2000). First-generation cephalosporins have good activity against gram-positive bacteria, E. coli, and Proteus and Klebsiella spp. Second-generation cephalosporins have increased activity against anaerobes and Haemophilus influenzae. Third-generation cephalosporins have broader coverage against gram-negative bacteria but less against gram-positive bacteria. The cephalosporins inhibit bacterial cell wall synthesis. Adverse reactions include hypersensitivity and gastrointestinal upset. Oral cephalosporins have been used effectively in the empiric treatment of uncomplicated UTIs (Lawrenson and Logie, 2001); in children with febrile UTI/pyelonephritis, oral third-generation cephalosporins such as cefixime have been shown to be safe and effective (Hoberman et al, 1999).

Penicillins

First-generation penicillins are ineffective against most uropathogens and are not commonly used in the treatment of UTI. However, the aminopenicillins (amoxicillin and ampicillin) have good activity against Enterococci, Staphylococci, E. coli, and Proteus mirabilis. However, gram-negative bacteria can quickly develop resistance to many aminopenicillins. The addition of beta-lactamase inhibitors such as clavulanic acid makes the aminopenicillins more active against the gram-negative bacteria. Although penicillins and aminopenicillins are inexpensive, the addition of the beta-lactamase inhibitors makes them more expensive. Adverse reactions include hypersensitivity (which can be immediate or delayed), gastrointestinal upset, and diarrhea. In general, penicillins are not commonly used in the treatment of UTI unless they are combined with beta-lactamase inhibitors (Sotto et al, 2001).

Antibiotic Resistance

Drug resistance among uropathogens has increased steadily during the past several years (Miller and Tang, 2004) and has much geographical variability. Local hospital antibiograms, which quantifies drug resistance seen at the hospital microbiology laboratory during a particular year, can provide information regarding local antibiotic resistance among bacteria for a specific locale. Evaluating these antibiograms together, some important trends in drug resistance can be seen (Kahlmeter, 2003). Among uropathogens particularly E. coli, resistance to ampicillin (18–54%), trimethoprim (9–27%), and sulfamethoxazole (16–49%) were high. Resistance to nitrofurantoin and fluoroquinolones were generally lower (<3%). However, with more extensive usage, resistance to these drugs is increasing.
KIDNEY INFECTION

Acute Pyelonephritis

Acute pyelonephritis is defined as inflammation of the kidney and renal pelvis, and its diagnosis is usually made clinically.

A. PRESENTATION AND FINDINGS

Patients with acute pyelonephritis present with chills, fever, and costovertebral angle tenderness. They often have accompanying lower-tract symptoms such as dysuria, frequency, and urgency. Sepsis may occur, with 20–30% of all systemic sepsis resulting from a urine infection. Urinalysis commonly demonstrates the presence of WBCs and red blood cells in the urine. Leukocytosis, increased erythrocyte sedimentation, and elevated levels of C-reactive protein are commonly seen on blood analysis. Bacteria are cultured from the urine when the culture is obtained before antibiotic treatment is instituted. *E. coli* is the most common causative organism, accounting for 80% of the cases. *Klebsiella*, *Proteus*, *Enterobacter*, *Pseudomonas*, *Serratia*, and *Citrobacter* spp. account for the remaining cases. Of the gram-positive bacteria, *Streptococcus faecalis* and *S. aureus* can be important causes of pyelonephritis. In reproductive-age women, sexual activity, patient and family history of UTI are associated with an increased risk of developing pyelonephritis. Diabetes and urinary incontinence also independently increase this risk (Scholes et al, 2005).

B. RADIOGRAPHIC IMAGING

Contrast-enhanced computed tomography (CT) scans can accurately demonstrate findings, confirming the diagnosis of pyelonephritis (Dacher et al, 1993). Acute bacterial infection causes constriction of peripheral arterioles and reduces perfusion of the affected renal segments. Perfusion defects, which can be segmental, multifocal, or diffuse, are seen as areas of reduced signal density (Figure 13–2). Renal enlargement, attenuated parenchyma, and a compressed collecting system are other characteristic findings on CT scan. However, CT scan is not necessary unless the diagnosis is unclear or the patient is not responding to therapy. Radionuclide study with 99mTc-dimercaptosuccinic acid is equally sensitive in detecting the perfusion defects of pyelonephritis (Levtchenko et al, 2001). In patients with acute pyelonephritis, renal ultrasonography is important to rule out concurrent urinary tract obstruction but cannot reliably detect inflammation or infection of the kidney.

C. MANAGEMENT

The management of acute pyelonephritis depends on the severity of the infection (Ghiro et al, 2002; Nickel, 2001). In patients who have toxicity because of associated septicemia, hospitalization is warranted. Approximately 10–30% of all adult patients with acute pyelonephritis require hospitalization, with incidence of 11.7 per 10,000 for women and 2.4 per 10,000 for men (Brown, Ki, and Foxman, 2005). Empiric therapy with intravenous ampicillin and aminoglycosides is effective against a broad range of uropathogens, including enterococci and *Pseudomonas* species. Alternatively, amoxicillin with clavulanic acid or a third-generation cephalosporin can be used. In a recent study of community-acquired UTIs in children hospitalized in a tertiary center (Marcus et al, 2005), it was noted that 40% of the culture-proved UTIs were caused by non-*E. coli* pathogens. Non-*E. coli* infections were more commonly found in males who had renal abnormalities and who had received antibiotic therapy in the prior month. Non-*E. coli* uropathogens were often resistant to cephalosporins and aminoglycosides. About 19% of the patients were initially treated with inappropriate empiric intravenous antibiotics. Fever from acute pyelonephritis may persist for several days despite appropriate therapy. Parenteral therapy should be maintained until the patient defervesces. If bacteremia is present, parenteral therapy should be continued for an additional 7–10 days and then the patient should be switched to oral treatment for 10–14 days. In patients who are not severely ill, outpatient treatment with oral antibiotics is appropriate. For adults, treatment with fluoroquinolones or TMP-SMX is well tolerated and effective. Therapy should continue for 10–14 days. Some patients in whom acute pyelonephritis develops will require follow-up radiologic examination such as voiding cystourethrogram or cystoscopy.

**Emphysematous Pyelonephritis**

Emphysematous pyelonephritis is a necrotizing infection characterized by the presence of gas within the renal parenchyma or perinephric tissue. About 80–90% of patients with emphysematous pyelonephritis have diabetes; the rest of the cases are associated with urinary tract obstruction from calculi or papillary necrosis (Shokeir et al, 1997; Tseng et al, 2005).
A. PRESENTATION AND FINDINGS

Patients with emphysematous pyelonephritis present with fever, flank pain, and vomiting that fails initial management with parenteral antibiotics (Tang et al, 2001). Pneumaturia may be present. Bacteria most frequently cultured from the urine include *E. coli*, *Klebsiella pneumoniae*, and *Enterobacter cloacae*.

B. RADIOGRAPHIC IMAGING

The diagnosis of emphysematous pyelonephritis is made after radiographic examination. Gas overlying the affected kidney may be seen on a plain abdominal radiograph (KUB). CT scan is much more sensitive in detecting the presence of gas in the renal parenchyma than renal ultrasonography.

C. MANAGEMENT

In the management of emphysematous pyelonephritis, prompt control of blood glucose and relief of urinary obstruction is essential, in addition to fluid resuscitation and parenteral antibiotics. The mortality rate is 11–54% (Michaeli et al, 1984). Poor prognostic factors include high serum creatinine level, low platelet count, and the presence of renal/perirenal fluid in association with a bubbly/loculated gas pattern or gas in the collecting system (Wan et al, 1998). In combination with medical treatment, percutaneous drainage appears to be helpful in accelerating resolution of the infection and minimizing the morbidity and mortality of the infection (Chen et al, 1997). Nephrectomy may be required if there is no function in the affected kidney. About 3–4 weeks of parenteral antibiotic therapy is usually required.

Chronic Pyelonephritis

Chronic pyelonephritis results from repeated renal infection, which leads to scarring, atrophy of the kidney, and subsequent renal insufficiency. The diagnosis is made by radiologic or pathologic examination rather than from clinical presentation.
A. PRESENTATION AND FINDINGS

Many individuals with chronic pyelonephritis have no symptoms, but they may have a history of frequent UTIs. In children, there is a strong correlation between renal scarring and recurrent UTIs (Wennerstrom et al, 2000). The developing kidney appears to be very susceptible to damage, and this susceptibility appears to be age dependent. Renal scarring induced by UTIs is rarely seen in adult kidneys. Because patients with chronic pyelonephritis often are asymptomatic, the diagnosis is made incidentally when radiologic investigation is initiated to evaluate for the complications associated with renal insufficiency, such as hypertension, visual impairments, headaches, fatigue, and polyuria. In these patients, urinalysis may show leukocytes or proteinuria but is likely to be normal. Serum creatinine levels reflect the severity of the renal impairment. Urine cultures are only positive when there is an active infection.

B. RADIOGRAPHIC IMAGING

Intravenous pyelogram or CT scan can readily demonstrate a small and atrophic kidney on the affected side. Focal coarse renal scarring with clubbing of the underlying calyx is characteristic. Ultrasonography similarly can demonstrate these findings. DMSA is the best imaging modality to look for renal scarring (Figure 13–3A–B; Stoller and Kogan, 1986). Areas of scarring can be seen as photopenic areas.

C. MANAGEMENT

The management of chronic pyelonephritis is somewhat limited because renal damage incurred by chronic pyelonephritis is not reversible. Eliminating recurrent UTIs and identifying and correcting any underlying anatomic or functional urinary problems such as obstruction or urolithiasis can prevent further renal damage. In children, evaluation for vesicoureteral reflux with a voiding cystourethrogram is important to eliminate a risk factor for recurrent UTIs and renal scarring. Rarely, removal of the affected kidney may be necessary due to hypertension or having a large stone burden in a nonfunctioning kidney.

Renal Abscesses

Renal abscesses result from a severe infection that leads to liquefaction of renal tissue; this area is subsequently sequestered, forming an abscess. They can rupture out into the perinephric space, forming perinephric abscesses. When the abscesses extend beyond the Gerota’s fascia, paranephric abscesses develop. Historically, most renal/perinephric abscesses result from hematogenous spread of staphylococci, in particular from infected skin lesions. Patients with diabetes, those undergoing hemodialysis, or intravenous drug abusers were at high risk for developing renal abscesses. With the development of effective antibiotics and better management of diseases such as diabetes and renal failure, renal/perinephric abscesses due to gram-positive bacteria are less prevalent; those caused by E. coli or Proteus species are becoming more common (Merimsky and Feldman, 1981; Thorley, Jones, and Sanford, 1974). Abscesses that form in the renal cortex are likely to arise

Figure 13–3. Chronic pyelonephritis. Multiple parenchymal defects (white and black arrows) are seen on DMSA scan (A), suggestive of scarring from recurrent infection. Voiding cystourethrogram (B) revealed high-grade reflux in this patient.
from hematogenous spread, whereas those in the corticomedullary junction are caused from gram-negative bacteria in conjunction with some other underlying urinary tract abnormalities, such as stones or obstruction.

A. PRESENTATION AND FINDINGS

The most common presenting symptoms in patients with renal/perinephric abscesses include fever, flank or abdominal pain, chills, and dysuria. Many of the symptoms have lasted for more than 2 weeks. A flank mass may be palpated in some patients. Urinalysis usually demonstrates WBCs; however, it may be normal in approximately 25% of the cases (Thorley, Jones, and Sanford, 1974). Urine cultures only identify the causative organisms in about one-thirds of cases and blood cultures in only about half of cases (Edelstein and McCabe, 1988).

B. RADIOGRAPHIC IMAGING

Renal abscesses can be accurately detected using ultrasonography or CT scans. There is a wide range of ultrasonographic findings ranging from an anechoic mass within or displacing the kidney to an echogenic fluid collection that tends to blend with the normally echogenic fat within Gerota's fascia (Corriere and Sandler, 1982). With high sensitivity, CT scans can demonstrate an enlarged kidney with focal areas of hypoattenuation early on during the course of the infection. Once the inflammatory wall forms around the fluid collection, the abscess appears as a mass with a rim of contrast enhancement, the “ring” sign (Figure 13–4). CT scans may also demonstrate thickening of Gerota’s fascia, stranding of the perinephric fat, or obliteration of the surrounding soft-tissue planes (Dalla Palma, Pozzi-Mucelli, and Ene, 1999). Intravenous pyelogram and kidneys, ureter,
and bladder tests are less sensitive tests in detecting renal/periurethral abscesses, with results being normal in about 20% of the cases (Thorley, Jones, and Sanford, 1974).

C. MANAGEMENT

The appropriate management of renal abscess first must include appropriate antibiotic therapy. Because it is often very difficult to identify the correct causative organisms from the urine or blood, empiric therapy with broad-spectrum antibiotics (ampicillin or vancomycin in combination with an aminoglycoside or third-generation cephalosporin) is usually recommended. If the patient does not respond within 48 hours of treatment, percutaneous drainage under CT or ultrasound guidance is indicated (Siegel, Smith, and Moldwin, 1996). The drained fluid should be cultured for the causative organisms. If the abscess still does not resolve, then open surgical drainage or nephrectomy may be necessary. Follow-up imaging is needed to confirm resolution of the abscesses. These patients will also require evaluation for underlying urinary tract abnormalities such as stone or obstruction after the infection has resolved.

Xanthogranulomatous Pyelonephritis

Xanthogranulomatous pyelonephritis (XGP) is a form of chronic bacterial infection of the kidney. The affected kidney is almost always hydronephrotic and obstructed. In most cases, XGP occurs unilaterally. Severe inflammation and necrosis obliterate the kidney parenchyma. Characteristically, foamy lipid-laden histiocytes (xanthoma cells) are present and may be mistaken for renal clear cell carcinoma (Iskandar, Prahlow, and White, 1993; Lorentzen and Nielsen, 1980).

A. PRESENTATION AND FINDINGS

Patients with XGP commonly present with flank pain, fever, chills, and persistent bacteriuria. A history of urolithiasis is present in about 35% of the patients (Malek and Elder, 1978). On physical examination, a flank mass can often be palpated. Urinalysis commonly demonstrates WBCs and protein. Serum blood analysis reveals anemia and may show hepatic dysfunction in approximately 50% of the patients (Malek and Elder, 1978). Because XGP primarily occurs unilaterally, azotemia or renal failure is not often seen (Goodman et al, 1979). *E. coli* or *Proteus* species are commonly cultured from the urine. However, one-third of patients with XGP have no growth in their urine, most likely because they have recently received antibiotic therapy. Approximately 10% of the patients with XGP have mixed organisms or anaerobic bacteria identified in their urine. Culture of the affected renal tissue can reliably identify the causative organism.

B. RADIOGRAPHIC IMAGING

CT scan is the most reliable method in imaging patients suspected of having XGP. It usually demonstrates a large heterogeneous, reniform mass. The renal parenchyma is often marked with multiple water-density lesions, representing dilated calyces or abscesses (Figure 13–5A–B; Goldman et al, 1984). On contrast-enhanced images, these lesions will have a prominent blush peripherally, while the central areas, which are filled with pus and debris, do not enhance. An area of central calcification surrounded by a contracted pelvis may also be seen (Eastham, Ahlering, and Skinner, 1994). The inflammatory process may be seen extending to the perinephric fat, the retroperitoneum, and adjacent organs such as the psoas muscle, spleen, colon, or the great vessels. Because of the association of urolithiasis and XGP, renal calculi may be seen (Parsons, 1993). Renal ultrasonography can also be used in performing imaging on patients with XGP (Tiu et al, 2001). It usually reveals an enlarged kidney with a large central echogenic area and anechoic parenchyma. However, ultrasonography does not provide comparable anatomic details to those obtained from CT scan. It is not uncommon for XGP to be misdiagnosed as a renal tumor because of their similar appearances on radiologic imaging (Zorzos et al, 2002).

C. MANAGEMENT

The management of XGP is dependent on accurate diagnosis. In some cases, XGP is misdiagnosed as a renal tumor. A nephrectomy is performed and a diagnosis is made pathologically. In those in whom a diagnosis of XGP is suspected, kidney-sparing surgery such as a partial nephrectomy is indicated. However, when the infection is extensive, a nephrectomy with excision of all involved tissue is warranted. There are reported cases of treating XGP with antibiotic therapy alone (Brown, Dodson, and Weintrub, 1996) or in combination with percutaneous drainage; however, these treatments are not likely to be curative in most patients and may lead to complications such as renal cutaneous fistula.

Pyonephrosis

Pyonephrosis refers to bacterial infection of a hydronephrotic, obstructed kidney, which leads to suppurative destruction of the renal parenchyma and potential loss of renal function. Because of the extent of the infection and the presence of urinary obstruction, sepsis may rapidly ensue, requiring rapid diagnosis and management.

A. PRESENTATION AND FINDINGS

Patients with pyonephrosis are usually very ill, with high fever, chills, and flank pain. Lower-tract symptoms are not usually present. Bacteriuria and pyuria may not be present when there is complete obstruction of the affected kidney.

B. RADIOGRAPHIC IMAGING

Imaging with renal ultrasonography can be performed to rapidly diagnose pyonephrosis. Ultrasonographic findings include persistent echoes in the inferior portion of the col-
Figure 13–5. Xanthogranulomatous pyelonephritis. CT scan (A) demonstrates a large heterogeneous left kidney, with dilated calyces and areas filled with lipid-laden macrophages. Xanthogranulomatous pyelonephritis is often associated with the presence of renal stones. Pathology specimen (B) better demonstrated the pockets of intraparenchymal abscesses and deposition of macrophages (arrows). PNS, percutaneous nephrostomy.
lecting system, fluid-debris level with dependent echoes that shift with positional changes (Figure 13–6), strong echoes with acoustic shadowing from air in the collecting system, and weak echoes throughout a dilated collecting system. Renal or ureteral calculi may also be identified on ultrasonography.

C. MANAGEMENT

Management of pyonephrosis includes immediate institution of antibiotic therapy and drainage of the infected collecting system. Broad-spectrum antimicrobials are indicated to prevent sepsis while the causative organism is being identified; antibiotics should be started before manipulation of the urinary tract. Performing drainage of the obstruction through the lower urinary tract (such as using a ureteral stent) should be reserved for patients who are not septic. Extensive manipulation may rapidly induce sepsis and toxemia. In the ill patient, drainage of the collecting system with a percutaneous nephrostomy tube is preferable. Once the infection is treated, additional imaging evaluation is required to identify the cause of the urinary obstruction, such as urolithiasis or ureteropelvic junction obstruction.

BLADDER INFECTION

Acute Cystitis

Acute cystitis refers to urinary infection of the lower urinary tract, principally the bladder. Acute cystitis more commonly affects women than men. The primary mode of infection is ascending from the periurethral/vaginal and fecal flora. The diagnosis is made clinically. In children, the distinction between upper and lower UTI is important. In general, those in whom acute cystitis developed do not usually require any extensive radiologic investigation (such as a voiding cystourethrogram), but those in whom pyelonephritis developed do (American Academy of Pediatrics, 1999).

Figure 13–6. Pyonephrosis. US demonstrates fluid-debris level (white arrow) within the dilated renal pelvis.
A. PRESENTATION AND FINDINGS

Patients with acute cystitis present with irritative voiding symptoms such as dysuria, frequency, and urgency. Low back and suprapubic pain, hematuria, and cloudy/foul-smelling urine are also common symptoms. Fever and systemic symptoms are rare. Typically, urinalysis demonstrates WBCs in the urine, and hematuria may be present. Urine culture is required to confirm the diagnosis and identify the causative organism. However, when the clinical picture and urinalysis are highly suggestive of the diagnosis of acute cystitis, urine culture may not be needed. *E. coli* causes most of the acute cystitis. Other gram-negative (*Klebsiella* and *Proteus* spp.) and gram-positive (*S. saprophyticus* and enterococci) bacteria are uncommon pathogens (Gupta et al, 1999). Diabetes and lifetime history of UTI are risk factors for acute cystitis. Of interest, the use of oral or vaginal estrogen was not protective in postmenopausal women with recurrent acute cystitis (Jackson et al, 2004).

B. RADIOGRAPHIC IMAGING

In uncomplicated infection of the bladder, radiologic evaluation is often not necessary.

C. MANAGEMENT

Management for acute cystitis consists of a short course of oral antibiotics. TMP-SMX, nitrofurantoin, and fluoroquinolones have excellent activity against most pathogens that cause cystitis. TMP-SMX and nitrofurantoin are less expensive and thus are recommended for the treatment of uncomplicated cystitis (Huang and Stafford, 2002). However, it is estimated that resistance to TMP-SMX by *E. coli* isolates causing uncomplicated acute cystitis is approximately 20%, compared to <2% to nitrofurantoin (Gupta, Scholes, and Stamm, 1999). In adults and children, the duration of treatment is usually limited to 3–5 days (Abramsson et al, 2002; Naber, 1999). Longer therapy is not indicated. Single-dose therapy for the treatment of recurrent cystitis/UTI appears to be less effective (Philbrick, 1986); however, fluoroquinolones with long half-lives (floxacin, pefloxacin, and rufloxacin) may be suitable for single-dose therapy (Naber, 1999). Resistance to penicillins and aminopenicillins is high and thus they are not recommended for treatment.

Recurrent Cystitis/UTI

A. PRESENTATION AND FINDINGS

Recurrent cystitis/UTI is caused either by bacterial persistence or reinfection with another organism. Identification of the cause of the recurrent infection is important, because the management of bacterial persistence and reinfection are distinct. If bacterial persistence is the cause of recurrent UTI, the removal of the infected source is often curative, whereas preventative therapy is effective in treating reinfection.

B. RADIOGRAPHIC IMAGING

When bacterial persistence is the suspected cause, radiologic imaging is indicated. Ultrasonography can be obtained to provide a screening evaluation of the genitourinary tract. More detailed assessment with intravenous pyelogram, cystoscopy, and CT scans may occasionally be necessary. In patients who have frequent, recurrent UTI, bacterial localization studies and more extensive radiologic evaluation (such as retrograde pyelograms) is warranted. When bacterial reinfection is the suspected cause of recurrent cystitis, the patient should be carefully evaluated for evidence of vesicovaginal or vesicoenteric fistula. Otherwise, radiologic examination is often not necessary in these patients.

C. MANAGEMENT

Management of recurrent cystitis, again, depends on its cause. Surgical removal of the infected source (such as urinary calculi) is needed to treat bacterial persistence. Similarly, fistulas need to be repaired surgically to prevent bacterial reinfection. In most cases of bacterial reinfection, medical management with prophylactic antibiotics is indicated. Low-dose continuous prophylactic antibiotic has been shown to reduce the recurrences of UTI by 95% compared to placebo or historical controls (Mangiarotti, Pizzini, and Fanos, 2000; Nicolle and Ronald, 1987). Alternatively, intermittent self-start antibiotic therapy can be used in treating recurrent cystitis in some women. Motivated patients self-identify episodes of infection on the basis of their symptoms and treat themselves with a single dose of antibiotics such as TMP-SMX. This regimen has been shown to be effective and economical in selected patients (Pfau and Sacks, 1993; Raz et al, 1991). When the recurrent cystitis/UTI is related to sexual activity, frequent emptying of the bladder and a single dose of antibiotic taken after sexual intercourse can significantly reduce the incidence of recurrent infection (Pfau and Sacks, 1994). Alternatives to antibiotic therapy in the treatment of recurrent cystitis/UTI include intravaginal estriol (Raz and Stamm, 1993), lactobacillus vaginal suppositories (Reid and Burton, 2002), and cranberry juice taken orally (Lowe and Fagelman, 2001).

Malacoplakia

Malacoplakia is an uncommon inflammatory disease of the bladder that can also affect other parts of the urinary tract, including the ureters and kidneys (Stanton and Maxted, 1981). In the bladder, it manifests as plaques or nodules made of large histiocytes (von Hansemann cells) with laminar inclusion bodies (Michaelis-Gutmann bodies) (McClure, Cameron, and Garrett, 1981).
**A. PRESENTATION AND FINDINGS**

Malacoplakia more commonly affects women than men (Stanton and Maxted, 1981) and is associated with a history of UTI. Patients with malacoplakia often have chronic illness or are immunosuppressed. In patients with malacoplakia of the bladder, irritative voiding symptoms (urgency and frequency) and hematuria are common (Curran, 1987). When the disease affects the ureter or kidney, the patient may present with fever, flank pain, or flank mass. When it affects both kidneys, signs or symptoms of azotemia or renal failure may be present (Dobyan, Truong, and Eknayan, 1993). Treatment with fluoroquinolone has significantly decreased the mortality rate associated with renal malacoplakia (Tam et al, 2003).

**B. RADIOLOGIC IMAGING**

Radiologic imaging with ultrasonography or CT may demonstrate a mass in the bladder and evidence of obstruction if the disease extends to the ureter (Vas et al, 1985). When the disease involves the kidney, focal or diffuse, hypodense, parenchymal masses may be seen on CT imaging (Frederic, D’Hondt, and Potvliege, 1981). It is often difficult to distinguish malacoplakia from malignancy (transitional cell or renal cell carcinoma) with radiologic imaging. The diagnosis is often established after biopsy.

**C. MANAGEMENT**

Management of malacoplakia primarily consists of antibiotic therapy, in particular those that produce high intracellular levels. Consequently, TMP-SMX and fluoroquinolones are recommended in the treatment of malacoplakia. Bethaneol and ascorbic acid, which enhance phagolysosomal activity, may have some benefits (Stanton and Maxted, 1981; Trujillo-Santos et al, 1999). In patients with malacoplakia limited to the lower urinary tract, antibiotic therapy alone is usually sufficient. However, when malacoplakia involves the ureter or kidney, surgical excision may be needed in addition to the antibiotic therapy (Dasgupta et al, 1999; Long and Althausen, 1989). The prognosis is poor and the mortality rate is high in patients who have bilateral renal involvement, regardless of treatment.

**PROSTATE INFECTION**

**Acute Bacterial Prostatitis**

Acute bacterial prostatitis refers to inflammation of the prostate associated with a UTI. It is thought that infection results from ascending urethral infection or reflux of infected urine from the bladder into the prostatic ducts. In response to bacterial invasion, leukocytes (polymorphonuclear leukocytes, lymphocytes, plasma cells, and macrophages) are seen within and surrounding the acini of the prostate. Edema and hyperemia of the prostatic stroma frequently develop. With prolonged infection, variable degree of necrosis and abscess formation can occur.

**A. PRESENTATION AND FINDINGS**

Acute bacterial prostatitis is uncommon in prepubertal boys but frequent affects adult men. It is the most common urologic diagnosis in men younger than 50 years (Collins et al, 1998). Patients with acute bacterial prostatitis usually present with an abrupt onset of constitutional (fever, chills, malaise, arthralgia, myalgia, lower back/rectal/perineal pain) and urinary symptoms (frequency, urgency, dysuria). They may also present with urinary retention due to swelling of the prostate. Digital rectal examination reveals tender, enlarged glands that are irregular and warm. Urinalysis usually demonstrates WBCs and occasionally hematuria. Serum blood analysis typically demonstrates leukocytosis. Prostate-specific antigen levels are often elevated. The diagnosis of prostatitis is made with microscopic examination and culture of the prostatic expressate and culture of urine obtained before and after prostate massage. In patients with acute prostatitis, fluid from the prostate massage often contains leukocytes with fat-laden macrophages. However, at the onset of acute prostatitis, prostatic massage is usually not suggested because the prostate is quite tender and the massage may lead to bacteremia. Similarly, urethral catheterization should be avoided. Culture of urine and prostate expressate usually identifies a single organism, but occasionally, polymicrobial infection may occur. E. coli is the most common causative organism in patients with acute prostatitis. Other gram-negative bacteria (Proteus, Klebsiella, Enterobacter, Pseudomonas, and Serratia spp.) and enterococci are less frequent pathogens. Anaerobic and other gram-positive bacteria are rarely a cause of acute prostatitis (RO Roberts et al, 1997).

**B. RADIOLOGIC IMAGING**

Radiologic imaging is rarely indicated in patients with acute prostatitis. Bladder ultrasonography may be useful in determining the amount of residual urine. Transrectal ultrasonography is only indicated in patients who do not respond to conventional therapy.

**C. MANAGEMENT**

Treatment with antibiotics is essential in the management of acute prostatitis. Empiric therapy directed against gram-negative bacteria and enterococci should be instituted immediately, while awaiting the culture results. Trimethoprim and fluoroquinolones have high drug penetration into prostatic tissue and are recommended for 4–6 weeks (Wagenlehner et al, 2005). The long duration of antibiotic treatment is to allow complete sterilization of the prostatic tissue to prevent complications such as chronic prostatitis and abscess formation (Childs, 1992; Nickel, 2000). Patients who have sepsis, are immunocompromised or in
acute urinary retention, or have significant medical comorbidities would benefit from hospitalization and treatment with parenteral antibiotics. Ampicillin and an aminoglycoside provide effective therapy against both gram-negative bacteria and enterococci. Patients with urinary retention secondary to acute prostatitis should be managed with a suprapubic catheter because transurethral catheterization or instrumentation is contraindicated.

Chronic Bacterial Prostatitis

In contrast to the acute form, chronic bacterial prostatitis has a more insidious onset, characterized by relapsing, recurrent UTI caused by the persistence of pathogen in the prostatic fluid despite antibiotic therapy.

A. PRESENTATION AND FINDINGS

Most patients with chronic bacterial prostatitis typically present with dysuria, urgency, frequency, nocturia, and low back/perineal pain. These patients usually are afebrile and not uncommonly have a history of recurrent or relapsing UTI, urethritis, or epididymitis caused by the same organism (Nickel and Moon, 2005). Others are asymptomatic, but the diagnosis is made after investigation for bacteriuria. In patients with chronic bacterial prostatitis, digital rectal examination of the prostate is often normal; occasionally, tenderness, firmness, or prostatic calculi may be found on examination. Urinalysis demonstrates a variable degree of WBCs and bacteria in the urine, depending on the extent of the disease. Serum blood analysis normally does not show any evidence of leukocytosis. Prostate-specific antigen levels may be elevated. Diagnosis is made after identification of bacteria from prostate expressate or urine specimen after a prostatic massage, using the 4-cup test (Table 13–8). The causative organisms are similar to those of acute bacterial prostatitis. It is currently believed that other gram-positive bacteria, *Mycoplasma, Ureaplasma*, and *Chlamydia* spp. are not causative pathogens in chronic bacterial prostatitis.

B. RADIOLOGIC IMAGING

Radiologic imaging is rarely indicated in patients with chronic prostatitis. Transrectal ultrasonography is only indicated if a prostatic abscess is suspected.

C. MANAGEMENT

Antibiotic therapy is similar to that for acute bacterial prostatitis (Bjerklund Johansen et al, 1998). Interestingly, the presence of leukocytes or bacteria in the urine and prostatic massage does not predict antibiotic response in patients with chronic prostatitis (Nickel et al, 2001). In patients with chronic bacterial prostatitis, the duration of antibiotic therapy may be 3–4 months. Using fluoroquinolones, some patients may respond after 4–6 weeks of treatment. The addition of an alpha blocker to antibiotic therapy has been shown to reduce symptom recurrences (Barbalias, Nikiforidis, and Liatsikos, 1998). Despite maximal therapy, cure is not often achieved due to poor penetration of antibiotic into prostatic tissue and relative isolation of the bacterial foci within the prostate. When recurrent episodes of infection occur despite antibiotic therapy, suppressive antibiotic (TMP-SMX 1 single-strength tablet daily, nitrofurantoin 100 mg daily, or ciprofloxacin 250 mg daily) may be used (Mears, 1987). Transurethral resection of the prostate has been used to treat patients with refractory disease; however, the success rate has been variable and this approach is not generally recommended (Barnes, Hadley, and O’Donoghue, 1982).

Table 13–8. Technique of Localization Cultures (4-Cup Test) for the Diagnosis of Prostatitis.

<table>
<thead>
<tr>
<th>Preparation:</th>
<th>- Require that the patient have a full bladder</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Retract foreskin of uncircumcised men</td>
<td></td>
</tr>
<tr>
<td>- Clean glans with soap/water or providone-iodine</td>
<td></td>
</tr>
<tr>
<td>Collection:</td>
<td>- Collect first 10 mL of voided urine (VB1)</td>
</tr>
<tr>
<td>- Discard the next 100 mL</td>
<td></td>
</tr>
<tr>
<td>- Collect the next 10 mL of voided urine (VB2)</td>
<td></td>
</tr>
<tr>
<td>- Massage prostate and collect prostate expressate (EPS)</td>
<td></td>
</tr>
<tr>
<td>- Collect first 10 mL of voided urine after massage (VB3)</td>
<td></td>
</tr>
<tr>
<td>- Immediately culture and microscopically examine all specimens</td>
<td></td>
</tr>
<tr>
<td>Interpretation:</td>
<td>- All specimens &lt;10^3 CFU/mL → not bacterial prostatitis</td>
</tr>
<tr>
<td>- VB3 or EPS &gt;10× CFU of VB1 → chronic bacterial prostatitis</td>
<td></td>
</tr>
<tr>
<td>- VB1 &gt; other specimens → urethritis or specimen contamination</td>
<td></td>
</tr>
<tr>
<td>Caution:</td>
<td>- Sensitivity of the test may not be high (Lipsky, 1999)</td>
</tr>
<tr>
<td>Alternative:</td>
<td>- Voided specimen before and after prostate massage (Nickel, 1997)</td>
</tr>
</tbody>
</table>

CFU, colony-forming unit.

Granulomatous Prostatitis

Granulomatous prostatitis is an uncommon form of prostatitis. It can result from bacterial, viral, or fungal infection, the use of bacillus Calmette-Guérin therapy (Rischmann et al, 2000), malacoplakia, or systemic granulomatous diseases affecting the prostate. Two-third of the cases have no specific cause. There are 2 distinct forms of nonspecific granulomatous prostatitis: noneosinophilic and eosinophilic. The former represents an abnormal tissue response to extravasated prostatic fluid (O’Dea, Hunting, and Greene, 1977).
The latter is a more severe, allergic response of the prostate to some unknown antigen.

**A. PRESENTATION AND FINDINGS**

Patients with granulomatous prostatitis often present acutely, with fever, chills, and obstructive/irritative voiding symptoms. Some may present with urinary retention. Patients with eosinophilic granulomatous prostatitis are severely ill and have high fevers. Digital rectal examination in patients with granulomatous prostatitis demonstrates a hard, indurated, and fixed prostate, which is difficult to distinguish from prostate carcinoma. Urinalysis and culture do not show any evidence of bacterial infection. Serum blood analysis typically demonstrates leukocytosis; marked eosinophilia is often seen in patients with eosinophilic granulomatous prostatitis. The diagnosis is made after biopsy of the prostate.

**B. MANAGEMENT**

Some patients respond to antibiotic therapy, corticosteroids, and temporary bladder drainage. Those with eosinophilic granulomatous prostatitis dramatically respond to corticosteroids (Ohkawa, Yamaguchi, and Kobayashi, 2001). Transurethral resection of the prostate may be required in patients who do not respond to treatment and have significant outlet obstruction.

**Prostate Abscess**

Most cases of prostatic abscess result from complications of acute bacterial prostatitis that were inadequately or inappropriately treated. Prostatic abscesses are often seen in patients with diabetes; those receiving chronic dialysis; or patients who are immunocompromised, undergoing urethral instrumentation, or who have chronic indwelling catheters.

**A. PRESENTATION AND FINDINGS**

Patients with prostatic abscess present with similar symptoms to those with acute bacterial prostatitis. Typically, these patients were treated for acute bacterial prostatitis previously and had a good initial response to treatment with antibiotics. However, their symptoms recurred during treatment, suggesting development of prostatic abscesses. On digital rectal examination, the prostate is usually tender and swollen. Fluuctuance is only seen in 16% of patients with prostatic abscess (Weinberger et al, 1988).

**B. RADIOLOGIC IMAGING**

Retrograde urethrogram is only indicated in patients with recurrent infection and obstructive voiding symptoms. Most patients with uncomplicated urethritis do not require any radiologic imaging.

**C. MANAGEMENT**

Pathogen-directed antibiotic therapy is required. In patients with gonococcal urethritis, ceftriaxone (250 mg intramuscularly) or fluoroquinolones (ciprofloxacin 250 mg) (David, Wildman, and Rajamanoharan, 2000) or norfloxacin (800 mg) may be used. For patients with nongonococcal urethritis, treatment is with tetracycline or erythromycin (500 mg 4 times daily) or doxycycline (100 mg twice daily) for 7–14 days (O’Mahony, 1999). However, the most essential component of treatment is prevention. Sexual partners of the affected patients should be treated, and protective sexual practices (such as using condoms) are recommended.

**URETHRITIS**

**Types of Urethritis**

Infection/inflammation of the urethra can be categorized into those types caused by *Neisseria gonorrhoeae* and by other organisms (*Chlamydia trachomatis*, *Ureaplasma urealyticum*, *Trichomonas vaginalis*, and herpes simplex virus) (Dixon, Pearson, and Clutterbuck, 2002). Most cases are acquired during sexual intercourse.

**A. PRESENTATION AND FINDINGS**

Patients with urethritis may present with urethral discharge and dysuria. The amount of discharge may vary significantly, from profuse to scant amounts. Obstructive voiding symptoms are primarily present in patients with recurrent infection, in whom urethral strictures subsequently develop. It is important to note that approximately 40% of patients with gonococcal urethritis are asymptomatic (John and Donald, 1978). The diagnosis is made from examination and culture of the urethra. It is important to obtain the specimen from within the urethra, rather than from just the discharge. Approximately 30% of men infected with *N. gonorrhoeae* will have concomitant infection with *C. trachomatis*.

**B. RADIOLOGIC IMAGING**

Retrograde urethrogram is only indicated in patients with recurrent infection and obstructive voiding symptoms. Most patients with uncomplicated urethritis do not require any radiologic imaging.

**C. MANAGEMENT**

Antibiotic therapy in conjunction with drainage of the abscess is required. Transrectal ultrasonography or CT scan can be used to direct transrectal drainage of the abscess (Barozzi et al, 1998). Transurethral resection and drainage may be required if transrectal drainage is inadequate. When properly diagnosed and treated, most cases of prostatic abscess resolve without significant sequelae (Weinberger et al, 1988).

**EPIDIDYMITIS**

**Causes of Epididymitis**

Infection and inflammation of the epididymis most often result from an ascending infection from the lower urinary
tract. Most cases of epididymitis in men younger than 35 years are due to sexually transmitted organisms (N. gonorrhoeae and C. trachomatis); those in children and older men are due to urinary pathogens such as E. coli. In homosexual men who practice anal intercourse, E. coli and other coliform bacteria are common causative organisms. The infection in the epididymis may spread to involve the testis.

A. PRESENTATION AND FINDINGS

Patients with epididymitis present with severe scrotal pain that may radiate to the groin or flank. Scrotal enlargement due to the inflammation of the epididymis/testis or a reactive hydrocele may develop rapidly. Other symptoms of urethritis, cystitis, or prostatitis may be present before or concurrent with the onset of scrotal pain. On physical examination, an enlarged and red scrotum is present, and it is often difficult to distinguish the epididymis from the testis during the acute infection. A thickened spermatic cord can occasionally be palpated. Urinalysis typically demonstrates WBCs and bacteria in the urine or urethral discharge. Serum blood analysis often reveals leukocytosis.

B. RADIOLOGIC IMAGING

Frequently, it is difficult to distinguish epididymitis from acute testicular torsion based on the history and physical examination alone (Petrack and Hafeez, 1992). Scrotal Doppler ultrasonography or radionuclide scanning can be used to confirm the diagnosis (Paltiel et al, 1998). The presence of blood flow in the testis on Doppler ultrasonography or uptake of the tracers into the center of the testis on radionuclide scanning rules out torsion. On scrotal ultrasonography, patients with epididymitis commonly have an enlarged epididymis with increased blood flow. A reactive hydrocele or testicular involvement may also be seen. Prepubertal children who are diagnosed with epididymitis will require radiologic investigation for urinary tract anomalies such as reflux or ureteral ectopia (Likit-
nukul et al, 1987). Postpubertal children who are diagnosed with epididymitis should be educated about sexually transmitted diseases and safe sexual practices.

C. Management

Oral antibiotic treatment is directed against specific causative organisms, as mentioned in the previous sections on urethritis and UTI. In addition, bed rest, scrotal elevation, and the use of nonsteroidal anti-inflammatory agents are helpful in reducing the duration of the symptoms. In patients with epididymitis caused by sexually transmitted organisms, treatment of their sexual partners is recommended to prevent reinfection. For patients with sepsis or severe infection, hospitalization and parenteral antibiotic therapy may be needed. Open drainage is indicated in cases in which an abscess develops. Occasionally, patients with chronic, relapsing epididymitis and scrotal pain may require epididymectomy for relief of their symptoms.

SPECIAL CIRCUMSTANCES

UTI Related to Pregnancy

With pregnancy, there are anatomic and physiologic changes to the urinary tract due to compression by the gravid uterus and alterations in the hormonal milieu. Renal length increases approximately by 1 cm during normal pregnancy as a result of increased vascular and interstitial volume (Waltzer, 1981). The glomerular filtration rate increases by 30–50%, most likely secondary to the increase in cardiac output (Waltzer, 1981). Typically, there is significant ureteral dilation with resultant urinary stasis during the second and third trimesters of gestation. This hydroureter is attributed to the smooth muscle-relaxing effects of progesterone and the mechanical compression of the ureters by the uterus at the level of the pelvic brim (Waltzer, 1981). The bladder is also affected, both physiologically and physiologically. The enlarged uterus displaces the bladder superiorly and anteriorly. The bladder becomes hyperemic, and its capacity is increased, most likely due to the effects of progesterone (Waltzer, 1981).

Because of these changes in the urinary tract during normal pregnancy, bacteriuria is a clinically relevant finding in pregnant women. It is estimated that the prevalence of bacteriuria is 4–6% (Sweet, 1977), which is not significantly different from that in nonpregnant women of comparable age. Interestingly, approximately 30% of those who have bacteriuria on screening evaluation later have pyelonephritis, compared to only 1–2% in those who do not have bacteriuria (Sweet, 1977). Treatment of bacteriuria decreases the incidence of pyelonephritis during pregnancy to approximately 3% (Christensen, 2000; Sweet, 1977).

Overall, the incidence of acute bacterial pyelonephritis is 1–4% in pregnant women (Gilstrap, Cunningham, and Whalley, 1981; Wing, 1998). About 60–70% of the episodes of pyelonephritis occur during the second and third trimesters of pregnancy, when urinary stasis is the greatest. In 10–20%, recurrent episodes of pyelonephritis develop before delivery (Gilstrap, Cunningham, and Whalley, 1981). Significant maternal risk factors include diabetes and history of UTI. When left untreated, pyelonephritis during pregnancy is associated with a high rate of infant prematurity and its associated perinatal mortality (Locksmith and Duff, 2001; McGregor and French, 1998; Schieve et al, 1994). It remains unclear whether treated pyelonephritis during pregnancy has any effects on the developing fetus (Gilstrap and Ramin, 2001).

Consequently, it is recommended that women be screened for bacteriuria during pregnancy to prevent the development of pyelonephritis. A voided urine specimen should be obtained at the first prenatal visit and at 16 weeks of gestation (Stenqvist et al, 1989). For asymptomatic individuals, significant bacteriuria is defined as 2 voided urine cultures with >10^5 CFU/mL of a single organism. For symptomatic pregnant women, >10^3 CFU/mL is considered to be significant (Rubin, Beam, and Stamm, 1992). Pregnant women who are found to have bacteriuria should be treated with penicillins, oral cephalosporins (Christensen, 2000; Wing et al, 1999), or fosfomycin trometamol (Minassian et al, 1998). Table 13–9 lists the antibiotics and their effects on pregnancy. However, amoxicillin is not recommended because of the rate of bacterial resistance (Hart et al, 2001). A 3-day course is suggested, although single-dose therapy may be effective in some patients (Tincello and Richmond, 1998). Repeat urine culture to document eradication of bacteriuria is necessary in all patients. Patients with acute bacterial pyelonephritis should be treated with parenteral cephalosporins,

Table 13–9. Antibiotics and Their Effects on Pregnancy.

<table>
<thead>
<tr>
<th>Drugs</th>
<th>Side Effects on the Developing Fetus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfonamides</td>
<td>Kernicterus</td>
</tr>
<tr>
<td>Trimethoprim</td>
<td>Interferes with neural tube development</td>
</tr>
<tr>
<td>Tetracyclines</td>
<td>Dysplasia and discoloration of teeth and bones</td>
</tr>
<tr>
<td>Nitrofurantoin</td>
<td>Hemolysis and G6PD deficiency</td>
</tr>
<tr>
<td>Aminoglycosides</td>
<td>Nerve damage</td>
</tr>
<tr>
<td>Fluoroquinolones</td>
<td>Interferes with cartilage formation</td>
</tr>
<tr>
<td>Penicillins</td>
<td>Safe</td>
</tr>
<tr>
<td>Cephalosporins</td>
<td>Safe</td>
</tr>
<tr>
<td>β-Lactamase inhibitors</td>
<td>Safe</td>
</tr>
<tr>
<td>Monobactams</td>
<td>Safe</td>
</tr>
<tr>
<td>Fosfomycin trometamol</td>
<td>Safe</td>
</tr>
</tbody>
</table>
penicillins with beta-lactamase inhibitors, or monolactams (Rubin, Beam, and Stamm, 1992). Periodic surveillance urine culture is recommended because many of these women will have recurrent episodes of pyelonephritis.

**UTI in Patients with Human Immunodeficiency Virus or Acquired Immunodeficiency Syndrome**

Human immunodeficiency virus (HIV) alters the normal host defense against bacterial infection. When the CD4 lymphocyte count falls to <200 per mm³, the risk of bacterial and opportunistic UTI increases dramatically (Evans et al, 1995; Hoepelman et al, 1992). In addition, antiretroviral medications used to treat HIV (eg, zidovudine) can further suppress normal immune response and increase the risk of UTI in these patients.

**A. UTI/CYSTITIS**

Hoepelman et al (1992) obtained urine cultures from HIV-positive men prospectively and when they had symptoms suggestive of a UTI. They observed that positive urine cultures were identified in 30% of HIV-infected men with CD4 <200 per mm³ and in 11% with CD4 = 200–500 per mm³, while none with CD4 >500 per mm³ had evidence of a urine infection. Gugino et al (1998) similarly observed that the incidence of bacteriuria in asymptomatic HIV-infected women was the same as that in uninfected women. Causative organisms include common uropathogens such as *E. coli* and *Klebsiella* and *Enterococcus* spp. Urinary infection with *S. aureus* and *Pseudomonas aeruginosa* is more common in HIV-infected patients (Schonwald, Begovac, and Skerk, 1999). Because of the common prophylactic use of TMP-SMX to prevent *Pneumocystis carinii* pneumonia in AIDS patients, the incidence of UTI in this group is decreased. However, when a UTI develops in these patients, the infecting organism is typically resistant to TMP-SMX (van Dooyeweert et al, 1996).

**B. PROSTATITIS**

In HIV patients, the incidence of bacterial prostatitis is approximately 3% and is 14% in patients with AIDS, compared to 1–2% in noninfected men of similar age (Leport et al, 1989). Causative organisms include common prostatitis pathogens such as *E. coli* and *Proteus* spp. and other less common organisms such as *Salmonella typhi*, *S. aureus*, *P. aeruginosa*, and *N. gonorrhoeae* (Staiman and Lowe, 1995). Prolonged treatment (4–6 weeks) with fluoroquinolones may be necessary because of a high risk of reinfection and lower immunity status in these patients. Prostatic abscess is more common in patients with AIDS compared to that in the general population (Staiman and Lowe, 1995; Trauzzi et al, 1994). Causative organisms include *E. coli* and other gram-negative bacteria or opportunistic fungus or mycobacterial infection (Lee, Dinneen, and Ahmad, 2001). Effective drainage and prolonged antimicrobial or antifungal therapy are needed.

**C. EPIDIDYMITIS AND URETHRITIS**

In HIV-infected men, epididymitis may be caused by *N. gonorrhoeae* and *C. trachomatis*. However, infection by coliform bacteria such as *E. coli* is more common, especially in patients having unprotected anal intercourse (Berger, Kessler, and Holmes, 1987). In HIV-infected patients with suppressive or antibiotic-resistant epididymitis, infection with fungi or mycobacteria should be considered (Desmond et al, 1993). In HIV-infected men who present with urethritis, treatment for both *Chlamydia* and *N. gonorrhoeae* is indicated even when gonococcus is only isolated from culture. Due to increased viral shedding with genital infections, it is recommended that HIV-infected patients abstain from sexual intercourse until 7 days after treatment is completed.

**D. INFECTION BY UNCOMMON ORGANISMS**

Urinary infection with *Mycobacterium* species can develop in HIV-infected patients. The kidneys are first infected and the infection spreads to the lower urinary tract. In patients with AIDS, it is estimated that 6–23% have renal tuberculosis (Marques et al, 1996). *M. tuberculosis* is the most common pathogen, with *M. avium* and *M. intracellulare* being less common (Sepkowitz et al, 1995). In HIV-infected patients who present with irritative/obstructive voiding symptoms but have no evidence of bacterial infection on culture, infection of the lower urinary tract by *Mycobacterium* species should be considered. Treatment with at least 2 antituberculosis agents is needed for 6–9 months.

**UTIs in Patients with Diabetes Mellitus**

UTIs are more common and tend to have more complicated course in patients with diabetes mellitus. There is a 2- to 5-fold increase in the incidence of acute pyelonephritis in diabetics compared to nondiabetics. Complications such as emphysematous pyelonephritis, renal and perirenal abscesses are more frequently seen in the diabetic patients (Williams and Schaeffer, 2004). Interestingly, the mortality and risk of hospitalization for UTI were not increased in patients with diabetes; however, the length of hospitalization may be prolonged (Nicolle, 2005). Defects in the local urinary cytokine secretions and an increased adherence of the microorganisms to the uroepithelial cells are potential mechanisms that may contribute to the increased prevalence of both asymptomatic and symptomatic bacteriuria in these patients (Hoepelman, 2003). Asymptomatic bacteriuria occurs in diabetic women more commonly than in nondiabetics. It is associated with an increased risk of UTI among patients with type 2 diabetes. However, treatment of asymptomatic bacteriuria with antimicrobial ther-
apy has not been shown to reduce symptomatic UTIs, pyelonephritis, or hospitalization for UTI (Ooi, Frazee, Gardner, 2004).

Though resistant bacteria are more frequently found in diabetic patients with UTI, empiric treatment with antibiotic therapy for a diabetic patient with complicated UTIs is similar to that of the nondiabetic patient. One important exception is that staphylococcal infection is not uncommon in the diabetic patients and can lead to urinary tract sepsis. This should be considered especially when a diabetic patient presents with a renal carbuncle. Oral outpatient therapy is not recommended for the diabetic patient with a complicated UTI. Treatment with TMP-SMX should be avoided if possible, because it can potentiate the hypoglycemic effects of the oral hypoglycemic drugs. Fluoroquinolones are safe and effective (ie, low resistance) in the treatment of diabetic patients with complicated UTIs (Williams and Schaeffer, 2004).

REFERENCES


Lee LK, Dinneen MD, Ahmad S: The urologist and the patient infected with human immunodeficiency virus or with acquired immunodeficiency syndrome. BJU Int 2001;88:500.


Specific infections are those caused by specific organisms, each of which causes a clinically unique disease that leads to specific pathologic tissue reactions. See also Chapter 15.

TUBERCULOSIS

Tubercle bacilli may invade one or more (or even all) of the organs of the genitourinary tract and cause a chronic granulomatous infection that shows the same characteristics as tuberculosis in other organs. Urinary tuberculosis is a disease of young adults (60% of patients are between the ages of 20 and 40) and is a little more common in males than in females.

Etiology

The infecting organism is *Mycobacterium tuberculosis*, which reaches the genitourinary organs by the hematogenous route from the lungs. The primary site is often not symptomatic or apparent.

The kidney and possibly the prostate are the primary sites of tuberculous infection in the genitourinary tract. All other genitourinary organs become involved by either ascent (prostate to bladder) or descent (kidney to bladder, prostate to epididymis). The testis may become involved by direct extension from epididymal infection.

Pathogenesis (Figure 14–1)

A. KIDNEY & URETER

When a shower of tubercle bacilli hits the renal cortex, the organisms may be destroyed by normal tissue resistance. Evidence of this is commonly seen in autopsies of persons who have died of tuberculosis; only scars are found in the kidneys. However, if enough bacteria of sufficient virulence become lodged in the kidney and are not overcome, a clinical infection is established.

Tuberculosis of the kidney progresses slowly; it may take 15–20 years to destroy a kidney in a patient who has good resistance to the infection. As a rule, therefore, there is no renal pain and little or no clinical disturbance of any type until the lesion has involved the calyces or the pelvis, at which time pus and organisms may be discharged into the urine. It is only at this stage that symptoms (of cystitis) are manifested. The infection then proceeds to the pelvic mucosa and the ureter, particularly its upper and vesical ends. This may lead to stricture and obstruction (hydronephrosis).

As the disease progresses, a caseous breakdown of tissue occurs until the entire kidney is replaced by cheesy material. Calcium may be laid down in the reparative process. The ureter undergoes fibrosis and tends to be shortened and therefore straightened. This change leads to a “golf-hole” (gaping) ureteral orifice, typical of an incompetent valve.

B. BLADDER

Vesical irritability develops as an early clinical manifestation of the disease as the bladder is bathed by infected material. Tubercles form later, usually in the region of the involved ureteral orifice, and finally coalesce and ulcerate. These ulcers may bleed. With severe involvement, the bladder becomes fibrosed and contracted; this leads to marked frequency. Ureteral reflux or stenosis and, therefore, hydronephrosis may develop. If contralateral renal involvement occurs later, it is probably a separate hematogenous infection.

C. PROSTATE & SEMINAL VESICLES

The passage of infected urine through the prostatic urethra ultimately leads to invasion of the prostate and one or both seminal vesicles. There is no local pain.

On occasion, the primary hematogenous lesion in the genitourinary tract is in the prostate. Prostatic infection can ascend to the bladder and descend to the epididymis.

D. EPIDIDYMIS & TESTIS

Tuberculosis of the prostate can extend along the vas or through the perivasal lymphatics and affect the epididymis. Because this is a slow process, there is usually no pain. If the epididymal infection is extensive and an abscess forms, it may rupture through the scrotal skin, thus establishing a permanent sinus, or it may extend into the testicle.
Figure 14–1. Pathogenesis of tuberculosis of the urinary tract.
Pathology

A. Kidney & Ureter

The gross appearance of the kidney with moderately advanced tuberculous infection is often normal on its outer surface, although the kidney is usually surrounded by marked perinephritis. Usually, however, there is a soft, yellowish localized bulge. On section, the involved area is seen to be filled with cheesy material (caseation). Widespread destruction of parenchyma is evident. In otherwise normal tissue, small abscesses may be seen. The walls of the pelvis, calyces, and ureter may be thickened, and ulceration appears frequently in the region of the calyces at the point at which the abscess drains. Ureteral stenosis may be complete, causing “autonephrectomy.” Such a kidney is fibrosed and functionless. Under these circumstances, the bladder urine may be normal and symptoms absent.

Tubercle foci appear close to the glomeruli. These are an aggregation of histiocytic cells possessing a vesicular nucleus and a clear cell body that can fuse with neighboring cells to form a small mass called an epithelioid reticulum. At the periphery of this reticulum are large cells with multiple nuclei (giant cells). This pathologic reaction, which can be seen macroscopically, is the basic lesion in tuberculosis. It can heal by fibrosis or coalesce and reach the surface and ulcerate, forming an ulcerocavernous lesion. Tubercles might undergo a central degeneration and caseate, creating a tuberculous abscess cavity that can reach the collecting system and break through. In the process, progressive parenchymal destruction occurs. Depending on the virulence of the organism and the resistance of the patient, tuberculosis is a combination of caseation and cavitiation and healing by fibrosis and scarring.

Microscopically, the caseous material is seen as an amorphous mass. The surrounding parenchyma shows fibrosis with tissue destruction, small round cell and plasma cell infiltration, and epithelial and giant cells typical of tuberculosis. Acid-fast stains will usually demonstrate the organisms in the tissue. Similar changes can be demonstrated in the wall of the pelvis and ureter.

In both the kidney and ureter, calcification is common. It may be macroscopic or microscopic. Such a finding is strongly suggestive of tuberculosis but, of course, is also observed in bladder infection. Secondary renal stones occur in 10% of patients.

In the most advanced stage of renal tuberculosis, the parenchyma may be completely replaced by caseous substance or fibrous tissue. Perinephric abscess may develop, but this is rare.

B. Bladder

In the early stages, the mucosa may be inflamed, but this is not a specific change. The bladder is quite resistant to actual invasion. Later, tubercles form and can be easily seen endoscopically as white or yellow raised nodules surrounded by a halo of hyperemia. With mural fibrosis and severe vesical contracture, reflux may occur.

Microscopically, the nodules are typical tubercles. These break down to form deep, ragged ulcers. At this stage the bladder is quite irritable. With healing, fibrosis develops that involves the muscle wall.

C. Prostate & Seminal Vesicles

Grossly, the exterior surface of these organs may show nodules and areas of induration from fibrosis. Areas of necrosis are common. In rare cases, healing may end in calcification. Large calcifications in the prostate should suggest tuberculous involvement.

D. Spermatic Cord, Epididymis, & Testis

The vas deferens is often grossly involved; fusiform swellings represent tubercles that in chronic cases are characteristically described as beaded. The epididymis is enlarged and quite firm. It is usually separate from the testis, although occasionally it may adhere to it. Microscopically, the changes typical of tuberculosis are seen. Tubular degeneration may be marked. The testis is usually not involved except by direct extension of an abscess in the epididymis.

E. Female Genital Tract

Infections are usually carried by the bloodstream; rarely, they are the result of sexual contact with an infected male. The incidence of associated urinary and genital infection in females ranges from 1% to 10%. The uterine tubes may be affected. Other presentations include endarteritis, localized adnexal masses (usually bilateral), and tuberculous cervicitis, but granulomatous lesions of the vaginal canal and vulva are rare.

Clinical Findings

Tuberculosis of the genitourinary tract should be considered in the presence of any of the following situations: (1) chronic cystitis that refuses to respond to adequate therapy, (2) the finding of sterile pyuria, (3) gross or microscopic hematuria, (4) a nontender, enlarged epididymis with a beaded or thickened vas, (5) a chronic draining scrotal sinus, or (6) induration or nodulation of the prostate and thickening of one or both seminal vesicles (especially in a young man). A history of present or past tuberculosis elsewhere in the body should cause the physician to suspect tuberculosis in the genitourinary tract when signs or symptoms are present.

The diagnosis rests on the demonstration of tubercle bacilli in the urine by culture or positive polymerase chain reaction (PCR). The extent of the infection is determined by (1) the palpable findings in the epididymides, vasa deferentia, prostate, and seminal vesicles; (2) the renal and ureteral lesions as revealed by imaging; (3) involvement of the bladder as seen through the cystoscope; (4) the degree
of renal damage as measured by loss of function; and (5) the presence of tubercle bacilli in one or both kidneys.

A. SYMPTOMS

There is no classic clinical picture of renal tuberculosis. Most symptoms of this disease, even in the most advanced stage, are vesical in origin (cystitis). Vague generalized malaise, fatigue, low-grade but persistent fever, and night sweats are some of the nonspecific complaints. Even vesical irritability may be absent, in which case only proper collection and examination of the urine will afford the clue. Active tuberculosis elsewhere in the body may be noted when the bladder becomes full. Irritability may become extreme. If ulceration occurs, suprapubic pain is occasionally found and is of either renal or vesical origin. At times, particularly in a late stage of the disease, the vesical irritability may become extreme. If ulceration occurs, suprapubic pain may be noted when the bladder becomes full.

1. Kidney and ureter—Because of the slow progression of the disease, the affected kidney is usually completely asymptomatic. On occasion, however, there may be a dull ache in the flank. The passage of a blood clot, secondary calculi, or a mass of debris may cause renal and ureteral colic. Rarely, the presenting symptom may be a painless mass in the abdomen.

2. Bladder—The earliest symptoms of renal tuberculosis may arise from secondary vesical involvement. These include burning, frequency, and nocturia. Hematuria is occasionally found and is of either renal or vesical origin. At times, particularly in a late stage of the disease, the vesical irritability may become extreme. If ulceration occurs, suprapubic pain may be noted when the bladder becomes full.

3. Genital tract—Tuberculosis of the prostate and seminal vesicles usually causes no symptoms. The first clue to the presence of tuberculous infection of these organs is the onset of a tuberculous epididymitis.

Tuberculosis of the epididymis usually presents as a painless or only mildly painful swelling. An abscess may drain spontaneously through the scrotal wall. A chronic draining sinus should be regarded as tuberculous until proved otherwise. In rare cases, the onset is quite acute and may simulate an acute nonspecific epididymitis.

B. SIGNS

Evidence of extragenital tuberculosis may be found (lungs, bone, lymph nodes, tonsils, intestines).

1. Kidney—There is usually no enlargement or tenderness of the involved kidney.

2. External genitalia—A thickened, nontender, or only slightly tender epididymis may be discovered. The vas deferens often is thickened and beaded. A chronic draining sinus through the scrotal skin is almost pathognomonic of tuberculous epididymitis. In the more advanced stages, the epididymis cannot be differentiated from the testis on palpation. This may mean that the testis has been directly invaded by the epididymal abscess.

Hydrocele occasionally accompanies tuberculous epididymitis. The idiopathic hydrocele should be tapped so that underlying pathologic changes, if present, can be evaluated (epididymitis, testicular tumor). Involvement of the penis and urethra is rare.

3. Prostate and seminal vesicles—These organs may be normal to palpation. Ordinarily, however, the tuberculous prostate shows areas of induration, even nodulation. The involved seminal vesicle is usually indurated, enlarged, and fixed. If epididymitis is present, the ipsilateral seminal vesicle usually shows changes as well.

C. LABORATORY FINDINGS

Proper urinalysis affords the most important clue to the diagnosis of genitourinary tuberculosis.

(1) Persistent pyuria without organisms on culture means tuberculosis until proved otherwise. Acid-fast stains done on the concentrated sediment from a 24-hour specimen are positive in at least 60% of cases. However, this must be corroborated by a positive culture.

If clinical response to adequate treatment of bacterial infection fails and pyuria persists, tuberculosis must be ruled out by bacteriologic and imaging.

(2) Cultures for tubercle bacilli from the first morning urine are positive in a very high percentage of cases of tuberculous infection. If positive, sensitivity tests should be ordered. In the face of strong presumptive evidence of tuberculosis, negative cultures should be repeated. Three to five first morning voided specimens are ideal. It can also be infected with tubercle bacilli, or it may become hydronephrotic from fibrosis of the bladder wall (ureterovesical stenosis) or vesicoureteral reflux.

If tuberculosis is suspected, the tuberculin test should be performed. A positive test, particularly in an adult, is hardly diagnostic, but a negative test in an otherwise healthy patient speaks against a diagnosis of tuberculosis.

D. X-RAY FINDINGS (FIGURE 14–2)

A plain film of the abdomen may show enlargement of one kidney or obliteration of the renal and psoas shadows due to perinephric abscess. Punctate calcification in the renal parenchyma may be due to tuberculosis. Renal stones are found in 10% of cases. Calcification of the ureter may be noted, but this is rare (Figure 6–1).

Excretory urograms can be diagnostic if the lesion is moderately advanced. The typical changes include (1) a “moth-eaten” appearance of the involved ulcerated calyces, (2) obliteration of one or more calyces, (3) dilatation of the calyces due to ureteral stenosis from fibrosis, (4) abscess cavities that connect with calyces, (5) single or multiple ureteral strictures, with secondary dilatation, with shortening and therefore straightening of the ureter, and (6) absence of function of the kidney due to complete ureteral occlusion and renal destruction (autonephrectomy). Ultrasound and computed tomography (CT) also show the calcifications, renal contractions and scars, ureteral and calyceal strictures suggestive of genitourinary tuberculosis. Ultrasound has the
advantage of low cost and low invasiveness. Contrast CT is highly sensitive for calcifications and the characteristic anatomic changes.

E. INSTRUMENTAL EXAMINATION

Thorough cystoscopic study is indicated even when the offending organism has been found in the urine and excretory urograms show the typical renal lesion. This study clearly demonstrates the extent of the disease. Cystoscopy may reveal the typical tubercles or ulcers of tuberculosis. Biopsy can be done if necessary. Severe contracture of the bladder may be noted. A cystogram may reveal ureteral reflux.

Differential Diagnosis

Chronic nonspecific cystitis or pyelonephritis may mimic tuberculosis perfectly, especially since 15–20% of cases of tuberculosis are secondarily invaded by pyogenic organ-
isms. If nonspecific infections do not respond to adequate therapy, a search for tubercle bacilli should be made. Painless epididymitis points to tuberculosis. Cystoscopic demonstration of tubercles and ulceration of the bladder wall means tuberculosis. Urograms are usually definitive.

Acute or chronic nonspecific epididymitis may be confused with tuberculosis, since the onset of tuberculosis is occasionally quite painful. It is rare to have palpatory changes in the seminal vesicles with nonspecific epididymitis, but these are almost routine findings in tuberculosis of the epididymis. The presence of tubercle bacilli on a culture of the urine is diagnostic. On occasion, only the pathologist can make the diagnosis by microscopic study of the surgically removed epididymis.

Multiple small renal stones or nephrocalcinosis seen by x-ray may suggest the type of calcification seen in the tuberculous kidney. In renal tuberculosis, the calcium is in the parenchyma, although secondary stones are occasionally seen.

Necrotizing papillitis, which may involve all of the calyces of one or both kidneys or, rarely, a solitary calyx, shows caliceal lesions (including calcifications) that simulate those of tuberculosis. Careful bacteriologic studies fail to demonstrate tubercle bacilli.

Medullary sponge kidneys may show small calcifications just distal to the calyces. The calyces are sharp, however, and no other stigmas of tuberculosis can be demonstrated.

In disseminated coccidioidomycosis, renal involvement may occur. The renal lesion resembles that of tuberculosis. Coccidioidal epididymitis may be confused with tuberculous involvement.

Urinary bilharziasis is a great mimic of tuberculosis. Both present with symptoms of cystitis and often hematuria. Vesical contraction, seen in both diseases, may lead to extreme frequency. Schistosomiasis must be suspected in endemic areas; the typical ova are found in the urine. Cystoscopic and urographic findings are definitive for making the differential diagnosis.

**Complications**

**A. RENAL TUBERCULOSIS**

Perinephric abscess may cause an enlarging mass in the flank. A plain film of the abdomen shows obliteration of the renal and psoas shadows. Sonograms and CT scans may be more helpful. Renal stones may develop if secondary nonspecific infection is present. Uremia is the end stage if both kidneys are involved.

**B. URETERAL TUBERCULOSIS**

Scarring with stricture formation is one of the typical lesions of tuberculosis and most commonly affects the juxtavesical portion of the ureter. This may cause progressive hydronephrosis. Complete ureteral obstruction may cause complete nonfunction of the kidney (autonephrectomy).

**C. VESICAL TUBERCULOSIS**

When severely damaged, the bladder wall becomes fibrosed and contracted. Stenosis of the ureters or reflux occurs, causing hydronephrotic atrophy.

**D. GENITAL TUBERCULOSIS**

The ducts of the involved epididymis become occluded. If this is bilateral, sterility results. Abscess of the epididymis may rupture into the testis, through the scrotal wall, or both, in which case the spermatic tubules may slough out.

**Treatment**

Genitourinary tuberculosis is extrapulmonary tuberculosis. The primary treatment is medical therapy. Surgical excision of an infected organ, when indicated, is merely an adjunct to overall therapy.

**A. RENAL TUBERCULOSIS**

A strict medical regimen should be instituted. A combination of drugs is usually desirable. The following drugs are effective in combination: (1) isoniazid (INH), 200–300 mg orally once daily; (2) rifampin (RMP), 600 mg orally once daily; (3) ethambutol (EMB), 25 mg/kg daily for 2 months, then 15 mg/kg orally once daily; (4) streptomycin, 1 g intramuscularly once daily; and (5) pyrazinamide, 1.5–2 g orally once daily. It is preferable to begin treatment with a combination of isoniazid, rifampin, and ethambutol. The European Association of Urology guidelines recommends 2 or 3 months of intensive triple drug therapy (INH, RMP, and EMB) daily followed by 3 months of continuation therapy with INH and RMP two or three times per week. If resistance to one of these drugs develops, one of the others listed should be chosen as a replacement.

The following drugs are usually considered only in cases of resistance to first-line drugs and when expert medical personnel are available to treat toxic side effects, should they occur: aminosalicylic acid (PAS), capreomycin, cycloserine, ethionamide, pyrazinamide, viomycin. **Pyrazinamide may cause serious liver damage.**

**B. VESICAL TUBERCULOSIS**

Tuberculosis of the bladder is always secondary to renal or prostatic tuberculosis; it tends to heal promptly when definitive treatment for the “primary” genitourinary infection is given. Vesical ulcers that fail to respond to this regimen may require transurethral electrocoagulation. Vesical instillations of 0.2% monochloroacetic acid (Clorapactin) may also stimulate healing.

Should extreme contracture of the bladder develop, it may be necessary to divert the urine from the bladder or perform augmentation cystoplasty after subtotal cystectomy (ileocystoplasty, ileoceccystoplasty, sigmoidocystoplasty) to increase bladder capacity.
C. TUBERCULOSIS OF THE EPIDIDYMIS

This condition never produces an isolated lesion; the prostate is always involved and usually the kidney as well. Only rarely does the epididymal infection break through into the testis. Treatment is medical. If after months of treatment an abscess or a draining sinus exists, epididymectomy is indicated.

D. TUBERCULOSIS OF THE PROSTATE & SEMINAL VESICLES

Although a few urologists advocate removal of the entire prostate and the vesicles when they become involved by tuberculosis, the majority opinion is that only medical therapy is indicated. Control can be checked by culture of the semen for tubercle bacilli.

E. GENERAL MEASURES FOR ALL TYPES

Optimal nutrition is no less important in treating tuberculosis of the genitourinary tract than in the treatment of tuberculosis elsewhere. Anticholinergic medications may help with bladder irritability.

F. TREATMENT OF OTHER COMPLICATIONS

Perinephric abscess usually occurs when the kidney is destroyed, but this is rare. The abscess must be drained, and nephrectomy should be done either then or later to prevent development of a chronic draining sinus. Prolonged antimicrobial therapy is indicated. If ureteral stricture develops on the involved side, ureteral dilatations offer a better than 50% chance of cure. The severely involved bladder may cause incompetence of the ureterovesical junction on the uninvolved side. Ureteroneocystostomy cannot be done in such a bladder; some form of urinary diversion may be required. For this reason, serial imaging and assessments of renal function are necessary even when the treatment is medical.

AMICROBIC (ABACTERIAL) CYSTITIS

Amicrobic cystitis is a rare disease of abrupt onset with a marked local vesical reaction. Although it acts like an infectious disease, search for the usual urinary bacterial pathogens is negative. It affects adult men and occasionally children, usually boys.

Etiology

The patient usually gives a history of recent sexual exposure. Mycoplasmas and chlamydiae have been isolated or suspected as etiologic agents. An adenovirus has been isolated from the urine in children suffering from acute hemorrhagic cystitis.

Pathogenesis & Pathology

Whatever the source and identity of the invader, the disease is primarily manifested as an acute inflammation of the bladder. Vesical irritability is severe and often associated with terminal hematuria. The mucosa is red and edematous, and superficial ulceration is occasionally seen. A thin membrane of fibrin often lies on the wall. Similar changes may be noted in the posterior urethra. The renal parenchyma is not involved, although the pelvic and ureteral mucosa may show mild inflammatory changes. Some dilatation of the lower ureters is apt to develop. This may be due to an inflammatory reaction about the ureteral orifices, for these changes regress after successful treatment.

Microscopically, there is nothing specific about the reaction. The mucosa and submucosa are infiltrated with neutrophils, plasma cells, and eosinophils. Submucosal hemorrhages are common; superficial ulceration of the mucosa may be noted.

Clinical Findings

A. SYMPTOMS

All symptoms are local. Urethral discharge, which is usually clear and mucoid but may be purulent, may be the initial symptom in men. Symptoms of acute cystitis come on abruptly. Urgency, frequency, and burning may be severe. Terminal hematuria is not uncommon. Suprapubic discomfort or even pain may be noted; it is most apt to be present as the bladder fills and is relieved somewhat by voiding. There is no fever or malaise.

B. SIGNS

Some suprapubic tenderness may be found. Urethral discharge may be profuse or scanty and may be purulent or thin and mucoid. The prostate is usually normal to palpation. Massage is contraindicated during the acute stage of urinary tract infection. When massage is done later, infection is usually not present.

C. LABORATORY FINDINGS

Some leukocytosis may develop. The urine is grossly purulent and may contain blood as well. Stained smears reveal an absence of bacteria. Routine cultures are uniformly negative. In a few cases, mycoplasmas and TRIC agent (Chlamydia trachomatis) have been identified, but the significance of this is not yet clear. Search for tubercle bacilli is not successful.

Urethral discharge reveals no bacteria. Renal function is not impaired.

D. X-RAY FINDINGS

Excretory urograms may demonstrate some dilatation of the lower ureters, but these changes regress completely when the disease is cured. The bladder shadow is small because of its markedly diminished capacity. Cystograms may reveal reflux.
E. INSTRUMENTAL EXAMINATION

Cystoscopy is not indicated in acute inflammation of the bladder. It has been done, however, when the diagnosis was obscure and tuberculosis suspected. In such cases it reveals redness and edema of the mucosa. Superficial ulceration may be noted. Bladder capacity is markedly diminished. Biopsy of the wall shows nonspecific changes.

Differential Diagnosis

Tuberculosis causes symptoms of cystitis, which usually come on gradually and become severe only in the stage of ulceration. A painless, nontender enlargement of an epididymis suggests tuberculosis. Although both tuberculosis and amicrobic cystitis produce pus without bacteria, thorough laboratory study demonstrates tubercle bacilli only in the former. On cystoscopy, the tuberculous bladder may be studded with tubercles. The ulcers in this disease are deep and of a chronic type. The changes in amicrobic cystitis are more acute; ulceration, if present, is superficial. Excretory urograms in tuberculosis may show "moth-eaten" calyces typical of infection with acid-fast organisms.

Nonspecific (pyogenic) cystitis may mimic amicrobic cystitis perfectly, but pathogenic organisms are easily found on a smear stained with methylene blue or on culture. Cystitis secondary to chronic nonspecific prostatitis occasionally produces pus without bacteria. The findings on rectal examination, the pus in the prostatic secretion, and the response to antibiotics point to the proper diagnosis.

Vesical neoplasm may ulcerate, become infected, and bleed; hence it may mimic amicrobic cystitis. Bacteriuria, however, is found. In case of doubt, cystoscopy is indicated.

Interstitial cystitis may be accompanied by severe symptoms of vesical irritability. However, it usually affects women and urinalysis is entirely negative except for a few red cells. Cystoscopy should be diagnostic.

Complications

Amicrobic cystitis is usually self-limited. Rarely, secondary contracture of the bladder develops. Under these circumstances, vesicoureteral reflux may be noted.

Treatment

A. SPECIFIC MEASURES

One of the tetracyclines or chloramphenicol, 1 g/day orally in divided doses for 3–4 days, is said to be curative in 75% of cases. Streptomycin, 1–2 g/day intramuscularly for 3–4 days, may be tried. Neoarsphenamine is also effective and appears to be the drug of choice, but arsenicals are hard to find. The first dose is 0.3 g intravenously; subsequent dosage is 0.45 g intravenously every 3–5 days for a total of 3–4 injections.

Penicillin and the sulfonamides are without effect. In the cases reported in children, cure occurred spontaneously.

B. GENERAL MEASURES

Bladder sedatives are usually of little help if symptoms are severe. Analgesics or narcotics may prove necessary to combat pain. Hot sitz baths may relieve spasm.

The instillation of a 0.1% solution of sodium oxychlo-rosene (Clorpactin WCS-90) has been recommended.

Prognosis

The prognosis is excellent.

C. CANDIDIASIS

Candida albicans is a yeast-like fungus that is a normal inhabitant of the respiratory and gastrointestinal tracts and the vagina. The intensive use of potent modern antibiotics is apt to disturb the normal balance between normal and abnormal organisms, thus allowing fungi such as Candida to overwhelm an otherwise healthy organ. The bladder and, to a lesser extent, the kidneys have proved vulnerable; candidemia has been observed. Anogenital candidiasis is discussed in Chapter 42.

The patient may present with vesical irritability or symptoms and signs of pyelonephritis. Fungus balls may be passed spontaneously. The diagnosis is made by observing mycelial or yeast forms of the fungus microscopically in a properly collected urine specimen. The diagnosis may be confirmed by culture. Excretory urograms may show caliceal defects and ureteral obstruction (fungus masses).

Vesical candidiasis usually responds to alkalinization of the urine with sodium bicarbonate. A urinary pH of 7.5 is desired; the dose is regulated by the patient, who checks the urine with indicator paper. Should this fail, amphotericin B should be instilled via catheter three times a day. One dissolves 50 mg of the drug in 1 L of sterile water.

If there is renal involvement, irrigations of the renal pelvis with a similar concentration of amphotericin B are efficacious. In the presence of systemic manifestations or candidemia, flucytosine (Ancobon) is the drug of choice. The dose is 100 mg/kg/day orally in divided doses given for 1 week. In the face of serious involvement, 600 mg is given intravenously on the first day followed by a shift to the oral form of the drug. Nifuroxat, a nitrofuran antibiotic, is superior to flucytosine. The recommended dose is 400 mg three times daily for 1 week. The dose must be modified in the face of renal impairment. The drug is more active in acid urine. Graybill et al (1983) reported good results with ketoconazole. The dose is 200–400 mg/day for 2–3 weeks or more depending on the effect as reflected by serial cultures. Its toxicity is relatively low. Amphotericin B (Fungizone) has the disadvantages of requiring parenteral administration and being highly nephrotoxic. It is given intravenously in a dosage of 1–5
mg/day in divided doses dissolved in 5% dextrose. The concentration of the solution should be 0.1 mg/mL.

**ACTINOMYCOSIS**

Actinomycosis is a chronic granulomatous disease in which fibrosis tends to become marked and spontaneous fistulas are the rule. On rare occasions, the disease involves the kidney, bladder, or testis by hematogenous invasion from a primary site of infection. The skin of the penis or scrotum may become involved through a local abrasion. The bladder may also become diseased by direct extension from the appendix, bowel, or oviduct.

**Etiology**

*Actinomyces israelii* is the causative organism.

**Clinical Findings**

There is nothing specifically pathognomonic about the symptoms or signs in actinomycosis. Pelvic involvement can be confused with malignancy. The microscopic demonstration of the organisms, which are visible as yellow bodies called “sulfur granules,” makes the diagnosis. If persistently sought, these may be found in the discharge from sinuses or in the urine. Definitive diagnosis is established by culture.

Urographically, the lesion in the kidney may resemble tuberculosis (eroded calyces) or tumor (space-occupying lesion).

**Treatment**

Penicillin G is the drug of choice. The dosage is 10–20 million U/day parenterally for 4–6 weeks, followed by penicillin V orally for a prolonged period. If secondary infection is suspected, a sulfonamide is added; streptomycin is also efficacious. Broad-spectrum antibiotics are indicated only if the organism is resistant to penicillin. Surgical drainage of the abscess or, better, removal of the involved organ is usually indicated.

**Prognosis**

Removal of the involved organ (eg, kidney or testis) may be promptly curative. Drainage of a granulomatous abscess may cause the development of a chronic draining sinus. Chemotherapy is helpful.

**SCHISTOSOMIASIS (BILHARZIASIS)**

Schistosomiasis, caused by a blood fluke, is a disease of warm climates. In its 3 forms, it affects about 350 million people. *Schistosoma mansoni* is widely distributed in Africa, South and Central America, Pakistan, and India; *Schistosoma japonicum* is found in the Far East; and *Schistosoma haematobium* (Bilharzia haematobia) is limited to Africa (especially along its northern coast), Saudi Arabia, Israel, Jordan, Lebanon, and Syria.

Schistosomiasis is on the increase in endemic areas because of the construction of modern irrigation systems that provide favorable conditions for the intermediate host, a freshwater snail. This disease principally affects the urogenital system, especially the bladder, ureters, seminal vesicles, and, to a lesser extent, the male urethra, and prostate gland. Because of emigration of people from endemic areas, the disease is being seen with increasing frequency in both Europe and the United States. Infection with *S. mansoni* and *S. japonicum* mainly involves the colon.

**Etiology**

Humans are infected when they come in contact with larva-infested water in canals, ditches, or irrigation fields during swimming, bathing, or farming procedures. Fork-tailed larvae, the cercariae, lose their tails as they penetrate deep under the skin. They are then termed schistosomules. They cause allergic skin reactions that are more intense in people infected for the first time. These schistosomules enter the general circulation through the lymphatics and the peripheral veins and reach the lungs. If the infection is massive, they may cause pneumonia. They pass through the pulmonary circulation, to the left side of the heart, and to the general circulation. The worms that reach the vesicoprostatic plexus of veins survive and mature, whereas those that go to other areas die.

**Pathogenesis**

The adult *S. haematobium* worm, a digenetic trematode, lives in the prostatovesical plexus of veins. The male is about 10 × 1 mm in size, is folded upon itself, and carries the long, slim 20 × 0.25 mm female in its “schist,” or gynecophoric canal. In the smallest peripheral venules, the female leaves the male and partially penetrates the venule to lay her eggs in the subepithelial layer of the affected viscus, usually in the form of clusters that form tubercles. The ova are seen only rarely within the venules; they are almost always in the subepithelial or interstitial tissues. The female returns to the male, which carries her to other areas to repeat the process.

The living ova, by a process of histolysis and helped by contraction of the detrusor muscle, penetrate the overlying urothelium, pass into the cavity of the bladder, and are extruded with the urine. If these ova reach fresh water, they hatch, and the contained larvae—ciliated miracidia—find a specific freshwater snail that they penetrate. There they form sporocysts that ultimately form the cercariae, which leave the snail hosts and pass into fresh water to repeat their life cycle in the human host.
Pathology

The fresh ova excite little tissue reaction when they leave the human host promptly through the urothelium. The contents of the ova trapped in the tissues and the death of the organisms cause a severe local reaction, with infiltration of round cells, monocytes, eosinophils, and giant cells that form tubercles, nodules, and polyps. These are later replaced by fibrous tissue that causes contraction of different parts of the bladder and strictures of the ureter. Fibrosis and massive deposits of eggs in subepithelial tissues interfere with the blood supply of the area and cause chronic bilharzial ulcerations. Epithelial metaplasia is common, and squamous cell carcinoma is a frequent sequel. Secondary infection of the urinary tract is a common complication and is difficult to overcome. The trapped dead ova become impregnated with calcium salts and form sheets of subepithelial calcified layers in the ureter, bladder, and seminal vesicles.

Clinical Findings

A. SYMPTOMS

Penetration of the skin by the cercariae causes allergic reactions, with cutaneous hyperemia and itching that are more intense in people infected for the first time. During the stage of generalization or invasion, the patient complains of symptoms such as malaise, fatigue and lassitude, low-grade fever, excessive sweating, headache, and backache. When the ova are laid in the bladder wall and begin to be extruded, the patient complains of terminal, slightly painful hematuria that is occasionally profuse. This may remain the only complaint for a long time until complications set in, when vesical symptoms become exaggerated and progressive. Increasing frequency, suprapubic and back pain, urethralgia, profuse hematuria, pyuria, and necroturia are likely to occur, with secondary infection, ulceration, or malignancy. Renal pain may be due to ureteral stricture, vesicoureteral reflux, or secondary stones obstructing the ureter. Fever, rigor, toxemia, and uremia are manifestations of renal involvement.

B. SIGNS

In early uncomplicated cases, there are essentially no clinical findings. Later, a fibroed, pitted, bilharzial glans penis, a urethral stricture or fistula, or a perineal fibrous mass may be found. A suprapubic bladder mass or a renal swelling may be felt abdominally. Rectal examination may reveal a fibroed prostate, an enlarged seminal vesicle, or a thickened bladder base.

C. LABORATORY FINDINGS

Urinalysis usually reveals the terminal-spined dead or living ova, blood and pus cells, and bacteria. Malignant squamous cells may be seen. The hemogram usually shows leukocytosis with eosinophilia and hypochromic normocytic anemia. Serum creatinine and blood urea nitrogen measurements may demonstrate some degree of renal impairment.

A variety of immunologic methods have been used to confirm the diagnosis of schistosomiasis. Positive immunologic tests indicate previous exposure but not whether schistosomiasis is currently present. The cercariae, schistosomules, adult worms, and eggs are all potentially antigenic. Adult worms, however, acquire host antigen on their integument that circumvents the immunologic forces of the host. Antibody production may be manifested as hypergammaglobulinemia.

D. X-RAY FINDINGS

A plain film of the abdomen may show areas of grayness in the flank (enlarged hydronephrotic kidney) or in the bladder area (large tumor). Opacifications (stones) may be noted in the kidney, ureter, or bladder. Linear calcification may be seen in the ureteral and bladder walls (Figure 14–3). Punctate calcification of the ureter (ureteritis calcinosa) and a honeycombed calcification of the seminal vesicle may be obvious (Figure 14–3).

Excretory urograms may show either normal or diminished renal function and varying degrees of dilatation of the upper urinary tracts (Figure 14–4). These changes include hydronephrosis, dilated and tortuous ureters, ureteral strictures, or a small contracted bladder having a capacity of only a few milliliters. Gross irregular defects of the bladder wall may represent cancer (Figure 14–4). Abdominal and pelvic CT is replacing excretory urography as the initial imaging of choice in many centers.

Retrograde urethrography may reveal a bilharzial urethral stricture. Cystograms often reveal vesicoureteral reflux, particularly if the bladder is contracted.

E. INSTRUMENTAL EXAMINATION

Cystoscopy may show fresh conglomerate, grayish tubercles surrounded by a halo of hyperemia, old calcified yellowish tubercles, sandy patches of mucous membrane, and a lusterless ground-glass mucosa that lacks the normal vascular pattern. Other obvious lesions include bilharzial polyps, chronic ulcers on the dome that bleed when the bladder is deflated (weeping ulcers), vesical stones, malignant lesions, stenosed or patulous ureteric orifices, and a distorted, asymmetric trigone. All are signs of schistosomal infestation.

Differential Diagnosis

Bilharzial cystitis is unmistakable in endemic areas. The presence of schistosomal ova in the urine, together with radiographic and cystoscopic findings, usually confirms the diagnosis. Nonspecific cystitis usually responds to medical treatment unless there is a complicating factor. Tuberculous cystitis may mimic bilharzial cystitis; the detection of tubercle bacilli, together with the radiographic picture, is confirmatory, but tuberculosis may occur in a bilharzial bladder.
Vesical calculi and malignancy should be diagnosed by thorough urologic examination, although both conditions are common in association with bilharzial bladder. Complications of schistosomiasis are the result of fibrosis, which may be extreme and causes contraction of the bladder neck as well as of the bladder itself. It also causes strictures of the urethra and ureter that are usually bilateral. Vesicoureteral reflux is a frequent sequela. Secondary persistent infection and stone formation usually complicate the picture still further. Squamous cell tumors of the bladder are common. They are seen as early as the second or third decade of life and are much more common in men than in women.

**Treatment**

**A. Medical Measures**

Praziquantel, metrifonate, and oxamniquine are the drugs of choice in treating schistosomiasis. These drugs do not have the serious side effects associated with the older drugs (eg, antimonials).
Praziquantel is unique in that it is effective against all human schistosome species. It is given orally and is effective in adults and children. Patients in the hepatosplenic stage of advanced schistosomiasis tolerate the drug well. The recommended dosage for all forms of schistosomiasis is 20 mg/kg three times in 1 day only.

Metrifonate is also a highly effective oral drug. It is the drug of choice for treatment of *S. haematobium* infections but is not effective against *S. mansoni* or *S. japonicum*. For treatment of *S. haematobium* infections, the dosage is 7.5–10 mg/kg (maximum 600 mg) once and then repeated twice at 2-week intervals.

Oxamniquine is a highly effective oral drug and is the drug of choice for treatment of *S. mansoni* infections. It is safe and effective in advanced disease. It is not effective in *S. haematobium* or *S. japonicum* infections. The dosage is 12–15 mg/kg given once; for children under 30 kg, 20 mg/kg is given in 2 divided doses.

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**Figure 14–4.** Schistosomiasis. **Upper left:** Excretory urogram showing markedly contracted bladder. Lower right ureter dilated probably secondary to vesicoureteral reflux. **Right:** Excretory urogram at 2 hours showing a fairly normal right kidney. The upper ureter is distorted. Arrows point to calcified wall. The lower ureter is quite abnormal. The calyces and pelvis of the left kidney are dilated, but the kidney shows atrophy secondary to nonspecific infection. The upper ureter is dilated and displaced by elongation due to obstruction. Arrows show calcification. Linear calcification can be seen in the periphery of the lower half of the bladder wall (arrows). **Lower left:** Nodular squamous cell carcinoma of the bladder. Dilated left lower ureter probably secondary to obstruction by tumor. Nonvisualization of the right ureter caused by complete occlusion.
in 1 day, with an interval of 2–8 hours between doses. Cure rates are 70–95%.

4 Niridazole, a nitrothiazole derivative, is effective in treating *S. mansoni* and *S. haematobium* infections. It may be tried against *S. japonicum* infections. It is given orally and should be administered only under close medical supervision. The dosage is 25 mg/kg (maximum, 1.5 g) daily in 2 divided doses for 7 days. Side effects may include nausea, vomiting, anorexia, headache, T-wave depression, and temporary suppression of spermatogenesis.

5 Antimonial drugs are no longer used in the treatment of schistosomiasis if praziquantel, oxamniquine, or metrifonate is available. The antimonials (eg, sodium antimony dimercaptosuccinate [stibocaptate], stibophen, tartar emetic) are much more toxic, and a longer course of therapy is needed. Tartar emetic is nonetheless occasionally needed as a third alternative drug in the treatment of *S. japonicum* infection.

B. GENERAL MEASURES

Antibiotics or urinary antiseptics are needed to overcome or control secondary infection. Supportive treatment in the form of iron, vitamins, and a high-calorie diet is indicated in selected cases.

C. COMPLICATIONS

Treatment of the complications of schistosomiasis of the genitourinary tract makes demands on the skill of the physician. Juxtavesical ureteral strictures require resection of the stenotic segment with ureteroneocystostomy. If the ureter is not long enough to reimplant, a tube of bladder may be fashioned, turned cephalad, and anastomosed to the ureter. Vesicoureteral reflux requires a suitable surgical repair. A contracted bladder neck may need transurethral anterior commissurotomy or a suprapubic Y-V plasty.

A chronic “weeping” bilharzial bladder ulcer necessitates partial cystectomy. The contracted bladder is treated by enterocystoplasty (placing a segment of bowel as a patch on the bladder). This procedure, which significantly increases vesical capacity, is remarkably effective in lessening the severity of symptoms associated with contracted bladder. Preoperative vesicoureteral reflux may disappear.

The most dreaded complication, squamous cell carcinoma, requires total cystectomy with urinary diversion if the lesion is deemed operable. Unfortunately, late diagnosis is common.

Prognosis

With energetic treatment, mild and early cases of schistosomiasis are not likely to result in severe damage to the urinary tract. On the other hand, massive repeated infections undermine the function of the urinary tract to such an extent that patients are disabled and become chronic invalids whose life spans are shortened by one or two decades.

In many endemic areas, attempts have been made to control the disease by mass treatment of patients, proper education, mechanization of agriculture, and various methods of eradication or control of the snail population. All these efforts have failed to be fully effective.

FILARIASIS

Filaria is endemic in the countries bordering the Mediterranean, in south China and Japan, the West Indies, and the South Pacific islands, particularly Samoa. Limited infection, as seen in American soldiers during World War II, gives an entirely different clinical picture from that seen in the frequent reinfections usually encountered among the native population.

Etiology

*Wuchereria bancrofti* is a threadlike nematode about 0.5 cm or more in length that lives in the human lymphatics. In the lymphatics, the female gives off microfilariae, which are found in the peripheral blood, particularly at night. The intermediate host (usually a mosquito) bites an infected person and becomes infested with microfilariae, which develop into larvae. These in turn transferred to another human, in whom they reach maturity. Mating occurs, and microfilariae are again produced. *Brugia malayi*, a nematode that causes filariasis in Southeast Asia and adjacent Pacific islands, acts in a similar fashion.

Pathogenesis & Pathology

The adult nematode in the human host invades and obstructs the lymphatics; this leads to lymphangitis and lymphadenitis. In long-standing cases, the lymphatic vessels become thickened and fibrous; there is a marked reticuloendothelial reaction.

Clinical Findings

A. SYMPTOMS

In mild cases (few exposures), the patient suffers recurrent lymphadenitis and lymphangitis with fever and malaise. Not infrequently, inflammation of the epididymis, testis, scrotum, and spermatic cord occurs. These structures then become edematous, boggy, and at times, tender. Hydrocele is common. In advanced cases (many exposures), obstruction of major lymph channels may cause chyluria and elephantiasis.

B. SIGNS

Varying degrees of painless elephantiasis of the scrotum and extremities develop as obstruction to lymphatics progresses. Lymphadenopathy is common.
C. LABORATORY FINDINGS

Chylous urine may look normal if minimal amounts of fat are present, but in an advanced case or following a fatty meal, it is milky. On standing, the urine forms layers: the top layer is fatty, the middle layer is pinkish, and the lower layer is clear. In the presence of chyluria, large amounts of protein are to be expected. Hypoproteinemia is found, and the albumin-globulin ratio is reversed. Both white blood cells (leukocytes) and red blood cells (erythrocytes) are found.

Marked eosinophilia is the rule in the early stages. Microfilariae may be demonstrated in the blood, which should be drawn at night. The adult worm may be found by biopsy. When filariae cannot be found, an indirect hemagglutination titer of 1/128 and a bentonite flocculation titer of 1/5 in combination are considered diagnostic.

D. CYSTOSCOPY

Following a fatty meal, endoscopy to observe the efflux of milky urine from the ureteral orifices may differentiate between unilateral and bilateral cases.

E. X-RAY FINDINGS

Retrograde urography and lymphangiography may reveal the renolymphatic connections in patients with chyluria.

Prevention

In endemic areas, mosquito abatement programs must be intensively pursued.

Treatment

A. SPECIFIC MEASURES

Diethylcarbamazine (Hetrazan) is the drug of choice, but it is toxic. The dose is 2 mg/kg orally three times daily for 12 days. This drug kills the microfilariae but not the adult worms. Several courses of the drug may be necessary. Antibiotics may be necessary to control secondary infection.

B. GENERAL MEASURES

Prompt removal of recently infected patients from the endemic area almost always results in regression of the symptoms and signs in early cases.

C. SURGICAL MEASURES

Elephantiasis of the external genitalia may require surgical excision.

D. TREATMENT OF CHYLURIA

Mild cases require no therapy. Spontaneous cure occurs in 50% of cases. If nutrition is impaired, the lymphatic channels may be sealed off by irrigating the renal pelvis with 2% silver nitrate solution. Should this fail, renal decapsula-
Treatment
Nephrectomy is generally the treatment of choice for renal hydatid disease. Aspiration of the cyst is unwise; leakage or rupture may occur. Retroperitoneal cysts are best treated by marsupialization and curettage.

Prognosis
Echinococcosis of the kidney usually has a good prognosis. The problem presented by perivesical cysts is more troublesome. After surgical intervention, drainage may be prolonged. It must be remembered, too, that involvement of other organs, especially the liver, is usually present.

REFERENCES

Tuberculosis

Amicrobic (Abacterial) Cystitis

Genitourinary Candidiasis

Actinomycosis

Figure 14–5. Hydatid disease, right kidney. Plain film showing 2 calcified hydatid cysts.

**Schistosomiasis (Bilharziasis)**

**Filariasis**

**Onchocerciasis**

**Echinococcosis (Hydatid Disease)**
The usual approach to sexually transmitted diseases (STDs) considers the causative agents, emphasizing different classes, genera, species, and microbiological characteristics. This fits with most medical school curricula since the causative agents span the full spectrum of medical microbiology (viruses, bacteria, protozoa, ectoparasites, and so on). This classical approach may prove difficult in clinical practice, where many different types of agents must be considered in the differential diagnosis of an individual patient.

This chapter takes a very selective and practical approach. Because patients present with symptoms and signs that may be caused by pathogens from different microbiological classes, we will emphasize diagnosis and treatment of clinical syndromes in contrast to traditional teaching (Table 15–1). This is a large subject with much active research and a huge literature. We stress the most important conditions encountered in urology: urethritis, epididymitis, genital ulcers, genital warts, plus a brief consideration of human immunodeficiency virus (HIV) infection.

URETHRITIS & CERVICITIS

Urethritis in Men

Urethritis, or urethral inflammation, is often caused by infection. Characteristically, patients complain of urethral discharge and dysuria. On examination the discharge may be purulent or mucopurulent. Asymptomatic infections are common (McCormack and Rein, 1990; Krieger, 1996; Centers for Disease Control and Prevention, 1998). The most important pathogens are bacteria, Neisseria gonorrhoeae, and Chlamydia trachomatis.

Testing is recommended to document a specific disease because both of these infections are reportable to health departments, and because specific diagnosis may improve compliance and partner notification (Centers for Disease Control and Prevention, 1998; Centers for Disease Control and Prevention, 2006). The traditional diagnostic algorithm includes microscopic examination of the Gram-stained urethral smear for gram-negative intracellular diplococci and culture for N. gonorrhoeae. New nucleic acid amplification tests have proved accurate for detection of N. gonorrhoeae and C. trachomatis in first-void urine in high-risk populations (Mahony et al, 2001). If diagnostic testing is unavailable, patients should be treated empirically for both infections (Centers for Disease Control and Prevention, 1998; Centers for Disease Control and Prevention, 2006).


A. ETIOLOGY

Gonorrhea is diagnosed when N. gonorrhoeae is detected by Gram stain, culture, or nucleic acid amplification testing. Nongonococcal urethritis (NGU) is diagnosed when gram-negative intracellular organisms cannot be diagnosed on microscopic examination. C. trachomatis, the most common infectious cause of NGU, is responsible for 23–55% of cases in reported series, but the proportion of cases is substantially lower in urological practice. The prevalence of chlamydial infection differs by age group, with a lower prevalence among older men. In addition, the proportion of NGU caused by C. trachomatis has been declining. Documentation of chlamydial NGU is important because this diagnosis supports partner referral, evaluation, and treatment (Centers for Disease Control and Prevention, 1998).

The etiology of most cases of nonchlamydial NGU is unknown. The genital mycoplasmas, Ureaplasma urealyticum and perhaps Mycoplasma genitalium or M. hominis, are implicated in 20–30% of cases in some series (Krieger, 1996; Horner et al, 2001; Totten et al, 2001; Stamm et al, 2007). Specific diagnostic tests for these organisms are not indicated routinely. Trichomonas vaginalis, a protozoan parasite, and herpes simplex virus (HSV) may also cause NGU (Joyner et al, 2000; Madeb et al, 2000). Testing and treatment for these organisms should be considered in situations where NGU is unresponsive to treatment (McCormack and Rein, 1990; Centers for Disease Control and
B. DOCUMENT URETHRITIS

It is important to document the presence of urethritis because some patients have symptoms in the absence of inflammation. Urethritis may be documented by the presence of any of the following clinical signs: mucopurulent urethral discharge on physical examination, ≥5 leukocytes per oil immersion microscopic field of the Gram-stained urethral secretions, a positive leukocyte esterase test on first-void urine, or ≥10 leukocytes per high-power microscopic field of the first-void urine (Krieger, 1996; Centers for Disease Control and Prevention, 2006). The Gram stain is the preferred diagnostic test for documenting urethritis and for evaluating presence or absence of gonococcal infection because it is rapid, highly sensitive, and specific.

If none of the criteria for urethritis are met, then treatment should be deferred. The patient should be tested for both \( N. \) gonorrhoeae and \( C. \) trachomatis and followed up closely in the event of a positive test result.

Empiric treatment of symptoms without documenting the presence of urethritis is recommended only if the patient is at high risk for infection and is unlikely to return for follow-up. Empiric treatment should be appropriate for both gonococcal and chlamydial infection. Sex partners should be referred for appropriate evaluation and treatment.

C. TREATMENT OF GONOCOCCAL INFECTIONS

There are an estimated 600,000 new gonococcal infections per year in the United States. In men, most infections cause symptoms that cause the patient to seek treatment soon enough to prevent serious sequelae. However, this may not be soon enough to prevent transmission of infection to sex partners. In contrast, many gonococcal (and also chlamydial) infections in women do not cause recognizable symptoms until the patient presents with complications, such as pelvic inflammatory disease. Symptomatic and asymptomatic pelvic inflammatory disease both result in tubal scarring, increased rates of ectopic pregnancy, and infertility.

Dual therapy is recommended for both gonococcal and chlamydial infection because patients are often coinfected with both pathogens (Krieger, 1996; Centers for Disease Control and Prevention, 1998; Centers for Disease Control and Prevention, 2006). Quinolone-resistant \( N. \) gonorrhoeae have been reported from many geographic areas, and such infections are becoming widespread in parts of Asia (Rahman et al, 2001; Tompkins and Zenilman, 2001; Trees et al, 2001).

Increasing antimicrobial resistance resulted in substantial changes in the gonorrhea treatment guidelines (Centers for Disease Control and Prevention, 2007). Fluoroquinolones (i.e., ciprofloxacin, ofloxacin, or levofloxacin) were the most frequently used drugs for treating gonorrhea because of their high efficacy, ready availability, and convenience as a single-dose, oral therapy. Unfortunately, this practice resulted in increasing fluoroquinolone resistance in \( N. \) gonorrhoeae. Since 2000, quinolones could no longer be recommended for treating patients who acquired their infections in Asia, the Pacific Islands, or Hawaii. Progressive increases in resistance led to extension of these recommendations to patients in California in 2002, and to treatment of gonorrhea in men who have sex with men elsewhere in the United States in 2004. Recent increases in the prevalence of fluoroquinolone-resistant \( N. \) gonorrhoeae throughout the United States led to the conclusion that fluoroquinolones can no longer be recommended for treating gonococcal infections anywhere in the United States. Consequently, only one class of drugs, the cephalosporins, is still recommended and available for the treatment of gonorrhea (Centers for Disease Control and Prevention, 2007). Of the recommended cephalosporins, only cefixime
SEXUALLY TRANSMITTED DISEASES

is available in an oral formulation. However, this drug is not currently available in the United States. Spectinomycin 2 g in a single dose is considered an effective alternative regime. But this drug is also not available in the United States. This means that there is no available oral treatment recommended for gonorrhea in the United States.

Table 15–2 summarizes recommended treatment regimens for uncomplicated gonococcal infections, where the recommended treatments reliably cure ≥97% of infections (Centers for Disease Control and Prevention, 1998; Centers for Disease Control and Prevention, 2007). Pharyngeal infections are more difficult to treat, and few regimens reliably cure >90% of infections. Patients who cannot tolerate cephalosporins should be treated with spectinomycin (2 g as a single intramuscular dose). However, this regimen is only 52% effective for pharyngeal infections.

Routine test-of-cure cultures are no longer recommended for patients treated with the recommended regimens. Such patients should refer their sex partners for evaluation and treatment. However, patients should be reevaluated if their symptoms persist after therapy. Any gonococci that persist should be evaluated for antimicrobial susceptibility. Infections identified after treatment are usually reinfections rather than treatment failures. Persistent inflammation may be caused by *C. trachomatis* or other organisms.

A few patients have complications such as disseminated gonococcal infection, perihepatitis, meningitis, or endocarditis. These infections result from gonococcal bacteremia. Disseminated gonococcal infection often causes petechial or pustular skin lesions, asymmetrical arthralgias, tenosynovitis, or septic arthritis. Occasionally patients have perihepatitis, and rare patients have endocarditis or meningitis. *N. gonorrhoeae* strains that cause disseminated infection tend to cause minimal genital tract inflammation. The recommended treatment is ceftriaxone (1 g intramuscularly or intravenously every 24 hours for disseminated infection or 1 g intravenously every 12 hours for meningitis or endocarditis).

**D. TREATMENT OF NONGONOCOCCAL URETHRITIS (NGU)**

Treatment should be initiated as soon as possible after diagnosis (Table 15–2). Single-dose regimens are preferred because these treatments offer the advantages of improved compliance and directly observed therapy (Centers for Disease Control and Prevention, 1998; Centers for Disease Control and Prevention, 2006). The recommended treatments employ either azithromycin or doxycycline. Alternative choices for patients who are allergic or cannot tolerate these drugs include a 7-day course of either erythromycin or ofloxacin. Routine follow-up and repeat testing are no longer recommended for patients taking the recommended regimens. However, patients should return for reevaluation if symptoms persist or recur after completion of treatment. The presence of symptoms alone without documentation of signs or laboratory findings of inflammation is not sufficient for retreatment. Patients should refer their sex partners for appropriate evaluation and treatment.

**E. TREATMENT OF RECURRENT AND PERSISTENT URETHRITIS**

Objective signs of urethritis should be documented before prescribing a repeat course of empirical therapy (Krieger, 1996; Centers for Disease Control and Prevention, 2006). Men with persistent or recurrent urethritis should be retreated with the initial regimen if they did not comply with treatment or if they were reexposed to an untreated sex partner. Other patients should have a wet mount and urethral culture for *T. vaginalis*. For patients who were
compliant with the initial regimen and who were not reexposed, the regimen in Table 15–2 should be used. This provides treatment for both *T. vaginalis* and the genital mycoplasmas.

**Mucopurulent Cervicitis in Women**

Mucopurulent cervicitis holds many parallels to urethritis in men (Centers for Disease Control and Prevention, 1998; Mead, 1990; Rein, 1990). Characteristically, patients have a purulent or mucopurulent endocervical exudate visible in the endocervical canal or on an endocervical swab sample. Easily induced endocervical bleeding is also common, as is an increased number of polymorphonuclear cells on the Gram-stained endocervical secretions. Patients may present with abnormal vaginal discharge or abnormal vaginal bleeding, for example, after intercourse, but many are asymptomatic.

As is the case with urethritis in men, *N. gonorrhoeae* and *C. trachomatis* are the most important infectious causes of mucopurulent cervicitis. However, neither pathogen may be identified in many women. Treatment should be guided by the results of testing for gonococcal and chlamydial infection, unless the patient is considered unlikely to return for follow-up. In such cases, empirical therapy should be given for both *C. trachomatis* and *N. gonorrhoeae*.

**EPIDIDYMITIS**

Epididymitis is caused by sexually transmitted pathogens or by organisms causing urinary tract infection (Krieger, 1996; Centers for Disease Control and Prevention, 2006; Krieger, 1990). Among sexually active men <35 years old, most cases of epididymitis are caused by sexually transmitted pathogens, particularly *C. trachomatis* and *N. gonorrhoeae*. Epididymitis may also be caused by *E. coli* among men who are the insertive partners during anal intercourse. Sexually transmitted epididymitis is usually associated with urethritis, which is often asymptomatic. Patients with uncomplicated sexually transmitted epididymitis do not require thorough evaluation for anatomic abnormalities.

Most cases of epididymitis in men older than 35 years are associated with urinary tract infection. The most common pathogens are gram-negative enteric bacteria. Epididymitis associated with urinary infection is more common among men who have anatomic abnormalities or those who have recently had urinary tract instrumentation. Therefore, evaluation of genitourinary tract anatomy is indicated for men with epididymitis associated with urinary tract infection.

Epididymitis is typically associated with unilateral hemiscrotal pain and tenderness. An inflammatory hydrocele and palpable swelling of the epididymis are characteristic. Diagnostic recommendations include a Gram-stained smear for evaluation of urethritis and for presumptive identification of gonococcal infection, diagnostic testing for *N. gonorrhoeae* and *C. trachomatis*, urine Gram stain and culture, syphilis serology and HIV testing (if sexually transmitted epididymitis is likely).

**Treatment**

Outpatient management is appropriate for most patients with epididymitis. Hospitalization should be considered when severe pain suggests other possible diagnoses, such as testicular torsion, testicular infarction, or testicular abscess; when patients are febrile; or when noncompliance with medication regimens is likely (Krieger, 1996; Centers for Disease Control and Prevention, 2006; Krieger, 1990). Empiric antimicrobial regimens are summarized in Table 15–3. Adjunctive measures include bed rest, scrotal elevation, and analgesics until fever and local inflammation subsides.

Routine follow-up is recommended. Failure to respond within 3 days requires reevaluation of both the diagnosis and treatment. Swelling and tenderness that persist after completion of antimicrobial therapy should be reevaluated to consider other possible diagnoses. These conditions include: testicular tumor, abscess, infarction, tuberculosis, fungal epididymitis, or collagen-vascular disorders (Skoutelas et al, 2000; Kaklamani et al, 2000; Giannopoulos et al, 2001; de Vries et al, 2001). HIV-infected patients with epididymitis should receive the same initial therapy as HIV-negative men. However, fungal infections, atypical mycobacteria, and other opportunistic infections are more likely than in nonimmunosuppressed patients.

**GENITAL ULCER DISEASES**

In the United States, genital herpes (HSV) is the most common cause of genital ulcers. Other important considerations are syphilis and chancroid. In contrast, lymphogranuloma venereum (LGV) and granuloma inguinale (donovanosis) are uncommon causes of genital ulcers in this country. Each of these ulcerative STDs is

**Table 15–3. Epididymitis: Recommended Treatment Regimens.**

<table>
<thead>
<tr>
<th><strong>Gonococcal or chlamydial infection likely</strong></th>
<th><strong>Enteric infection likely</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Ceftriaxone, 250 mg in a single IM dose, plus doxycycline, 100 mg orally twice a day for 10 days</td>
<td>Ofloxacin, 300 mg orally twice a day for 10 days, or levofloxacin, 500 mg orally once a day for 10 days</td>
</tr>
</tbody>
</table>

*According to Centers for Disease Control and Prevention: 2002 Sexually transmitted disease treatment guidelines. MMWR 2002;51:1; and Centers for Disease Control and Prevention: Sexually transmitted diseases treatment guidelines 2006. MMWR51 (No. RR-11).
associated with a two- to fivefold increased risk of HIV transmission.

**Diagnostic Testing**

Diagnosis based solely on the history and physical findings is often inaccurate (Centers for Disease Control and Prevention, 1998, Centers for Disease Control and Prevention, 2006). Patients may be infected with more than one agent simultaneously. Ideally, evaluation of patients with genital ulcers should include testing for the most common causes: HSV, syphilis, and chancroid. These tests include a culture or antigen test for HSV, a darkfield examination or direct immunofluorescence test for *Treponema pallidum* (syphilis), and a culture for *Haemophilus ducreyi* (chancroid). In the future, improved molecular detection tests may become available commercially for these organisms. After thorough diagnostic evaluation, 25% of patients with genital ulcers do not have a laboratory-confirmed diagnosis. HIV testing also should be recommended for patients who have genital ulcers (see below).

Often patients must be treated before test results become available. In this situation, treatment is recommended for both syphilis and chancroid.

**Genital Herpes Virus (HSV) Infection**

Genital HSV is an incurable and recurrent viral infection. Characteristic genital lesions start as painful papules or vesicles. Often the genital lesions have evolved into pustules or ulcers when the patient is seen in the office. With primary genital herpes, the ulcerative lesions persist for 4–15 days until crusting, or reepithelization, or both occur. Pain, itching, vaginal or urethral discharge, and tender inguinal adenopathy are the predominant local symptoms. Primary HSV infection is associated with a high frequency and prolonged duration of systemic and local symptoms. Fever, headache, malaise, and myalgias are common. The clinical symptoms of pain and irritation from genital lesions gradually increase over the first 6–7 days, reach maximum intensity between days 7 and 11 of disease, and then recede gradually during the second or third week.

In contrast to first episodes, recurrent HSV infection is characterized by symptoms, signs, and anatomic sites localized to the genital region. Local symptoms, such as pain and itching, are mild compared with the symptoms of initial infection, and the duration of the usual episode ranges from 8 to 12 days or less.

Two HSV serotypes cause genital ulcers, HSV-1 and HSV-2. Both viruses infect the genital tract. Studies suggest that 5–30% of first-episode cases of genital HSV infection are caused by HSV-1. However, recurrences of HSV-1 infection are substantially less likely than recurrences of HSV-2. Therefore, HSV-2 cases accumulate in the population of patients with recurrent genital lesions. Typing the infecting strain has prognostic importance and is useful for patient counseling. However, most commercially available antibody tests are not accurate enough to distinguish HSV-1 from HSV-2 infection. Better assays should become available in the future.

Serologic studies suggest that 45 million people in the United States are infected with genital HSV-2. Most infections are mild or unrecognized. Therefore, most HIV-infected people do not receive this diagnosis. Such asymptomatic or mildly symptomatic persons shed virus intermittently in their genital tracts and can infect their sex partners. First-episode genital HSV infections are more likely to cause symptoms than recurrent infections. Occasional cases are severe enough to require hospitalization for complications such as disseminated infection, pneumonitis, hepatitis, meningitis, or encephalitis.

**Treatment**

Systemic antiviral therapy results in partial control of symptoms and signs of genital HSV infection. Treatment does not cure the infection or change the frequency or severity of recurrences after discontinuation of treatment. Three antiviral drugs have proved beneficial in randomized clinical trials: acyclovir, valacyclovir, and famciclovir (Table 15–4). Topical treatment with acyclovir has proved substantially less effective than systemic treatment.

Patients with first clinical episodes of genital HSV should receive antiviral treatment to speed healing of genital lesions and shorten the duration of viral shedding. Patients should also be counseled about the natural history of genital herpes, the risks for sexual and perinatal transmission, and methods to reduce transmission. Patients with severe disease should receive intravenous treatment with acyclovir.

Most persons with first clinical episodes of genital HSV-2 experience recurrent episodes. Treatment can shorten the duration of lesions and decrease recurrences. Thus, many patients can benefit from antiviral therapy, and this option should be discussed. There are 2 approaches to antiviral therapy for recurrent HSV: episodic treatment and daily suppressive treatment. Episodic therapy is beneficial for many patients with occasional recurrences. Such therapy is started during the prodrome or first day of lesions. Thus, patients receiving episodic therapy should receive the medication or a prescription so that they may initiate treatment at the first symptom or sign of lesions. Traditionally, 5 days of treatment has been recommended for recurrent HSV (Table 15–4), but recent data suggest that shorter (3-day) courses may work equally well (Leone, Trotter, and Miller, 2002; Wald et al, 2002).

Daily suppressive therapy is useful for patients who experience frequent recurrences (6 or more per year). Therapy reduces the frequency of recurrences by >75%. Such treatment has been shown to be safe and effective for as long as 6 years with acyclovir and for as long as 1 year with both valacyclovir and famciclovir. Daily therapy does
not appear to be associated with clinically significant HSV-drug resistance. After 1 year, discontinuation of treatment should be considered, since the frequency of recurrences often decreases with time.

Syphilis

Syphilis may be the deepest and darkest subject in all of infectious diseases. This complex illness is caused by *T. pallidum*, a spirochete, and holds a special place in the history of medicine as “the great impostor” and “the great imitator.” Sir William Osler in 1897 said, “Know syphilis in all its manifestations and relations, and all other things clinical will be added unto you.”

Syphilis is a systemic disease. Patients may seek treatment for symptoms of signs of primary, secondary, or tertiary infection. Primary infection is characterized by an ulcer, or chancre, at the site of infection. Secondary manifestations include rash, mucocutaneous lesions, and adenopathy. Tertiary infection may present with cardiac, neurologic, ophthalmic, auditory, or gummatous lesions. In addition, syphilis may be diagnosed by serologic testing of asymptomatic patients; this is termed latent syphilis. Latent syphilis acquired within the preceding year is classified as early latent syphilis. All other cases of latent syphilis are classified as late latent or syphilis of unknown duration.

### Table 15–4. Genital Ulcers: Recommended Treatment Regimens.*

<table>
<thead>
<tr>
<th>Condition</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Genital herpes</strong></td>
<td></td>
</tr>
<tr>
<td><strong>First episode</strong></td>
<td>Acyclovir, 400 mg orally 3 times a day for 7–10 days; or acyclovir, 200 mg orally 5 times a day for 7–10 days; or famciclovir, 250 mg orally 3 times a day for 7–10 days; or valacyclovir, 1 g orally twice a day for 7–10 days</td>
</tr>
<tr>
<td><strong>Severe disease</strong></td>
<td>Acyclovir, 5–10 mg/kg body weight IV every 8 hours for 2–7 days or until clinical resolution</td>
</tr>
<tr>
<td><strong>Recurrent episodes</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Episodic recurrences</strong></td>
<td>Acyclovir, 400 mg orally 3 times a day for 5 days; or acyclovir, 200 mg orally 5 times a day for 5 days; or acyclovir, 800 mg orally twice a day for 5 days; or famciclovir, 125 mg orally twice a day for 5 days; or valacyclovir, 500 mg orally twice a day for 3–5 days; or valacyclovir 1 g orally once a day for 5 days</td>
</tr>
<tr>
<td><strong>Daily suppressive therapy</strong></td>
<td>Acyclovir, 400 mg orally twice a day; or famciclovir, 250 mg orally twice a day; or valacyclovir, 250 mg orally twice a day; or valacyclovir, 500 mg orally twice a day; or valacyclovir, 1 g orally once a day</td>
</tr>
<tr>
<td><strong>Syphilis</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Primary and secondary</strong></td>
<td>Benzathine penicillin G, 2.4 million units IM as a single dose</td>
</tr>
<tr>
<td><strong>Tertiary (except neurosyphilis)</strong></td>
<td>Benzathine penicillin G, 2.4 million units IM weekly for 3 weeks</td>
</tr>
<tr>
<td><strong>Neurosyphilis</strong></td>
<td>Aqueous crystalline penicillin G, 3–4 million units IV every 4 hours for 10–14 days; or procaine penicillin, 2.4 million units IM daily for 10–14 days, plus probenecid, 500 mg orally 4 times a day for 10–14 days</td>
</tr>
<tr>
<td><strong>Latent syphilis</strong></td>
<td></td>
</tr>
<tr>
<td>Early</td>
<td>Benzathine penicillin G, 2.4 million units IM as a single dose</td>
</tr>
<tr>
<td>Late or of unknown duration</td>
<td>Benzathine penicillin G, 2.4 million units IM weekly for 3 weeks</td>
</tr>
<tr>
<td><strong>Chancroid</strong></td>
<td>Azithromycin, 1 g as a single oral dose; or ceftriaxone, 250 mg as a single IM dose; or ciprofloxacin, 500 mg orally twice a day for 3 days; or erythromycin base, 500 mg orally 4 times a day for 7 days</td>
</tr>
<tr>
<td><strong>Granuloma inguinale</strong></td>
<td>Trimethoprim-sulfamethoxazole, 1 double-strength tablet orally twice a day for a minimum of 3 weeks; or doxycycline, 100 mg orally twice a day for a minimum of 3 weeks</td>
</tr>
<tr>
<td><strong>Lymphogranuloma venereum</strong></td>
<td>Doxycycline, 100 mg orally twice a day for 21 days</td>
</tr>
</tbody>
</table>

Definitive diagnosis of early syphilis is done by dark-field examination or direct immunofluorescent antibody tests of lesion exudates, because antibodies may not be present. Presumptive diagnosis depends on serologic testing. Serologic tests are either nontreponemal, such as the Venereal Disease Research Laboratory (VDRL) and rapid plasma reagin (RPR) tests, or treponemal, such as the fluorescent treponemal antibody absorption (FTA-ABS) test and microagglutination assay for antibody to T. pallidum (MHA-TP). Use of only one type of serologic test is considered insufficient for diagnosis. False-positive nontreponemal tests occur with a variety of medical conditions. Nontreponemal tests correlate with disease activity, with results reported quantitatively. In general, a fourfold change in titer is considered significant. Most patients with reactive treponemal tests remain reactive for life. Treponemal test titers correlate poorly with disease activity. Thus, combination of both treponemal and nontreponemal tests is necessary for patient management.

Tremendous progress during the last decade has led to sequencing of the entire genome of T. pallidum and to correlation of functional activities with this genetic information (Norris, Cox, and Weinstock, 2001). From an epidemiological perspective, the real news is that eradication of syphilis from the United States (Centers for Disease Control, 2001a,b) and from the world (Rompalo, 2001) has been established as an important public health goal, although this goal may prove difficult to achieve because of high rates of infection in certain at-risk populations (termed “core groups”).

### Treatment

For more than 40 years penicillin has been the treatment of choice for syphilis (Table 15–4). Patients who are allergic to penicillin should receive a 2-week course of doxycycline (100 mg orally twice daily) or tetracycline (500 mg orally 4 times a day). Treatment results in healing of local lesions and prevents sexual transmission and late sequelae. Patients with syphilis should be tested for HIV infection. In areas with a high prevalence of HIV, this test should be repeated after 3 months if the initial HIV test is negative. Patients with syphilis and symptoms or signs of ophthalmic disease should have a slit-lamp examination and those with symptoms or signs of neurologic disease should have cerebrospinal fluid evaluation. Treatment failures occur with any regimen. Thus, serologic testing should be repeated 6 and 12 months after initial treatment.

### Chancroid

Chancroid is an acute ulcerative disease, often associated with inguinal adenopathy ("bubo"). H. ducreyi, a gram-negative facultative bacillus, is the causative agent. The infection is endemic in parts of the United States and the disease also occurs in outbreaks. It is estimated that 10% of patients with chancroid are coinfected with either T. pallidum or HSV. Each of these ulcerative infections is associated with an increased rate of HIV transmission.

Definitive diagnosis of chancroid requires identification of the causative bacterium, H. ducreyi, on specialized culture media that are not widely available. In addition, these media have an estimated sensitivity <80%. In practice, a probable diagnosis of chancroid may be based on the following: the patient has a painful genital ulcer; there is no evidence of T. pallidum by darkfield examination or by a negative syphilis serologic testing at least 7 days before the onset of ulcers; an HSV test is negative; and the clinical appearance is typical. The combination of a painful genital ulcer and tender inguinal adenopathy suggests the diagnosis of chancroid. Unfortunately, this characteristic clinical appearance occurs in only one-third of cases. However, the combination of a painful genital ulcer and suppurative inguinal adenopathy is considered almost pathognomonic. The combination of a human model of infection with new molecular methods has resulted in improved understanding of H. ducreyi genes and virulence factors (Bong et al., 2001; Gelfanova, Humphreys, and Spinola, 2001; Throm and Spinola, 2001; Young et al., 2001).

#### Lymphogranuloma Venereum (LGV)

Lymphogranuloma venereum is caused by the invasive serovars of C. trachomatis (L1, L2, and L3). The disease is a rare cause of genital ulcers in the United States. Tender inguinal or femoral lymphadenopathy or both, often unilateral, is the characteristic clinical presentation in heterosexual men. Women and homosexual men may
present with inflammatory involvement of perirectal and perianal lymphatics, strictures, fistulas, or proctocolitis. The self-limited genital ulcers have usually healed when most patients seek medical care. In most cases, diagnosis is made by serologic testing plus exclusion of other causes of inguinal adenopathy or genital ulcers.

TREATMENT

Therapy causes microbiological cure and prevents ongoing tissue destruction (Table 15–4). Doxycycline is preferred. Erythromycin and azithromycin are alternatives. Prolonged therapy, for a minimum of 3 weeks, is necessary with each of these drugs. However, tissue reaction and scarring can progress after effective treatment. Inguinal adenopathy, known as “bubos,” may require needle aspiration through intact skin or incision and drainage to prevent inguinal or femoral ulcerations. Patients should be followed up until clinical symptoms and signs are resolved.

Granuloma Inguinale (Donovanosis)

Granuloma inguinale is caused by Calymmatobacterium granulomatis, a gram-negative intracellular bacillus that has many similarities to Klebsiella species (Kharsany et al, 1999; O’Farrell, 2001). This infection is rare in the United States. Granuloma inguinale is an important cause of genital ulcers in tropical and developing countries, particularly India, Papua New Guinea, central Australia, and southern Africa.

Clinically, granuloma inguinale presents with painless, progressive genital ulcers. The genital lesions are highly vascular, with a “beefy red” appearance. Patients seldom have inguinal adenopathy. The causative organism cannot be cultured on standard microbiologic media. Diagnosis requires visualization of dark-staining Donovan bodies on tissue crush preparations or biopsy specimens. Molecular diagnostic tests should be available in the near future (O’Farrell, 2001; Behets et al, 1999). Secondary bacterial infections may develop in the lesions. In addition, coinfection with other STD agents may occur.

Treatment

Effective treatment halts progressive tissue destruction (Table 15–4). Trimethoprim-sulfamethoxazole or doxycycline is recommended. Alternative drugs are ciprofloxacin or erythromycin. Azithromycin also appears promising (O’Farrell, 2001; Bowden and Savage, 1998). Prolonged duration of treatment is often necessary to facilitate granulation and reepithelization of the ulcers. Patients should be reevaluated after the first few days of treatment. Addition of an aminoglycoside, such as gentamicin, should be considered if lesions have not responded. Treatment should be continued until all lesions have healed. Relapse can occur 6–18 months after effective initial therapy.

Genital Warts

Genital warts are caused by human papillomavirus (HPV) infection. Of the more than 80 HPV genotypes, more than 20 infect the genital tract. Most of these genital HPV infections are asymptomatic, subclinical, or unrecognized. Depending on their size and anatomic locations, visible external warts can be painful, friable, pruritic, or all three. Most visible genital warts are caused by HPV types 6 or 11. These HPV types can also cause exophytic warts on the cervix and within the vagina, urethra, and anus. HPV types 6 and 11 are only rarely associated with development of invasive squamous cell carcinoma of the external genitalia.

HPV types 16, 18, 31, 33, and 35 are uncommon in visible, external genital warts. These HPV types are associated with cervical dysplasia, as well as vaginal, anal, and cervical squamous cell carcinoma. HPV types 16, 18, 31, 33, and 35 have also been associated with external genital intraepithelial neoplastic lesions, including squamous cell carcinoma, carcinoma in situ, Bowenoid papulosis, erythroplasia of Queyrat, and Bowen’s disease. Patients with external genital warts can be infected simultaneously with multiple HPV types.

Most often, the diagnosis of genital warts can be made by inspection. Diagnosis can be confirmed by biopsy, if necessary, although biopsy is rarely necessary for diagnosis. Biopsy is indicated if the diagnosis is uncertain, if lesions do not respond to standard therapy, if the disease worsens during treatment, if the patient is immunocompromised, or if warts are pigmented, indurated, fixed, or ulcerated. Routine use of type-specific HPV nucleic acid tests is not indicated for diagnosis or management of visible genital warts (Centers for Disease Control and Prevention, 2006).

Treatment

For visible genital warts, the primary goal of treatment is removal of symptomatic lesions. Treatment can induce wart-free periods in most patients. Genital warts are often asymptomatic, and clinical lesions may resolve spontaneously. Currently, there are no data indicating that available therapy can eradicate HPV infection or change the natural history of infection. In theory, removal of exophytic warts may decrease infectivity, but there is no evidence that treatment changes the risk for development of dysplastic or cancerous lesions in the patient or in sexual partners.

Treatment decisions should be guided by the provider’s experience and patient preferences. None of the recommended therapies is superior or ideal for every case. Current treatments can be considered as patient applied or provider administered (Table 15–5). Most patients with visible warts have lesions that respond to most treatment modalities. Many patients require a course of therapy. In general, lesions on moist surfaces or in intertriginous areas respond better to topical treatments, such as trichloroacetic
Table 15–5. External Genital Warts: Recommended Treatment Regimens.*

**Patient-applied**
Podophlox, 0.5% solution of gel to lesions twice a day for 3 days, followed by 4 days off therapy; repeat as needed for up to 4 cycles; or imiquimod, 5% cream to lesions at bedtime 3 times a week for up to 16 weeks; wash off after 6–10 hours

**Provider-administered**
Cryotherapy with liquid nitrogen or cryoprobe; repeat as necessary every 1–2 weeks; or podophyllin resin, 10–25% in tincture of benzoin; repeat weekly as necessary; or trichlor/bichloracetic acid, 80–90%; apply until white “frosting”; repeat weekly as necessary; or surgical removal (laser surgery), or intralesional interferon


Acid, podophyllin, or imiquimod, than do warts on drier surfaces.

Podophlox is an antimitotic drug that results in destruction of warts. Most patients experience pain or local irritation after treatment. Imiquimod is a topically active immune enhancer that stimulates production of cytokines, followed by local inflammation and resolution of warts (Moore et al, 2001; Fife et al, 2001). Effective use of cryotherapy requires training to avoid either overtreatment or undertreatment and poor results. Pain is common after application of the liquid nitrogen, followed by necrosis of the warts. Podophyllin resin contains several antimitotic compounds. Different resin preparations vary in the concentrations of active components and contaminants. Although both trichloroacetic acid and bichloracetic acid are recommended and are used widely, these treatments are associated with several potential problems. The acid can spread rapidly if applied excessively, with damage to normal adjacent tissues. These solutions should be applied sparingly and allowed to dry before the patient stands. If the patient experiences excessive discomfort, the acid can be neutralized by using soap or sodium bicarbonate (baking soda). Recent data suggest that the treatment approach should be changed if a patient has not improved substantially after 3 provider-administered treatments or if warts do not resolve completely after 6 treatments.

Surgical removal offers the advantage of rendering the patient wart free in a single visit. Several approaches are possible, including tangential scissor or shave excision, curettage, electrosurgery, or laser surgery. All of these methods require local anesthesia and are more time consuming and expensive than the methods discussed in the previous paragraph. Surgical approaches are most useful for patients who have a large number or a large volume of genital warts, if the diagnosis is uncertain, or if patients have been unresponsive to other treatments. Patients should be warned that scarring, hypopigmentation, and hyperpigmentation are common after ablative therapies. Occasionally, patients have chronic pain after such treatment.

Recurrence of warts is common after all therapies, with most recurrences occurring within the first 3 months. Women should be counseled about the need for regular cervical cytologic screening. Examination of sex partners is unnecessary for management of external genital warts, because the role of reinfection is probably minimal. However, sex partners of patients with genital warts may benefit from evaluation for genital warts and other STDs. Recent availability of highly effective, multivalent HPV vaccines offer the opportunity to substantially improve the clinical epidemiology of HPV infection by vaccinating adolescents prior to initiation of sexual activity (Koutsky et al, 2002; Garland et al, 2007).

**SUBCLINICAL GENITAL HPV INFECTION**

Subclinical HPV infection (without visible genital warts) is more common than visible genital lesions. Most cases are diagnosed indirectly by cervical cytology, colposcopy, or biopsy of genital skin, or by routine use of acetic acid soaks and examination with magnification for “acetowhite” areas. The consensus of expert opinion is to discourage routine examination for “acetowhiting”, Centers for Disease Control and Prevention, 2006). This test has poor specificity for HPV infection. In addition, the acetowhite test has many false-positive results in low-risk populations. Definitive diagnosis of subclinical HPV infection requires detection of HPV nucleic acid or capsid protein, but these tests are not recommended outside of research settings.

Treatment of subclinical HPV infection is not recommended in the absence of dysplasia. Diagnosis is often questionable because many of the diagnostic tests (i.e., cytology, acetowhiting, colposcopy) correlate poorly with detection of HPV, DNA, or RNA. Furthermore, no therapy has been proved to eradicate infection. HPV has been demonstrated in normal-appearing tissue adjacent to treated areas after aggressive surgical treatment.

**HIV INFECTION: OVERVIEW OF DETECTION, INITIAL EVALUATION, & REFERRAL**

Infection with HIV includes a wide clinical spectrum, ranging from asymptomatic infection to AIDS. The rate of
clinical progression is highly variable. Some persons progress from HIV infection to AIDS within a few months; others remain asymptomatic for decades. Overall, the median time from infection to AIDS is around 10 years. In general, adults with HIV infection remain asymptomatic for prolonged periods. However, HIV viral replication continues during all stages of infection, with substantial increases in the viral burden during later stages of infection, accompanied by marked deterioration in immune functions.

Increasing awareness of risk factors for HIV infection has led to increased testing and earlier diagnosis for many patients. The primary risk factors for HIV infection are sexual contact with an HIV-infected person and sharing injecting-drug equipment.

Early diagnosis is important because treatment can slow the decline in immune function (Centers for Disease Control and Prevention, 2002; Centers for Disease Control and Prevention, 2006). HIV-infected persons with evidence of immune dysfunction are at risk for preventable infections. Prophylactic treatment can substantially reduce the risk for pneumonia (Pneumocystis carinii and bacterial), toxoplasma encephalitis, and mycobacterial disease (tuberculosis and Mycobacterium avium complex). Early diagnosis also facilitates patient counseling, which may reduce transmission. In addition, early diagnosis facilitates planning for referral to a health-care provider/facility experienced in care of HIV-infected persons.

Testing for HIV

Diagnostic testing for HIV should be offered to anyone at risk for infection, especially those seeking evaluation for STDs. Appropriate pre- and posttest counseling and informed consent should be included in the test procedure. Some states required documentation of informed consent.

Usually, HIV infection is documented using HIV-1 antibody tests. HIV antibodies are detected in >95% of infected persons within 6 months of infection. In most laboratories, this is a 2-stage procedure beginning with a sensitive screening test, such as an enzyme immunoassay. Reactive screening test results are then confirmed by a supplemental test, such as the Western blot, or an immunofluorescence assay. Patients with positive results on both the screening and confirmatory tests are infected with HIV. Such infected persons can transmit HIV.

In the United States, almost all HIV infections are caused by HIV-1. Extremely rare cases are caused by a second virus, HIV-2. Thus, routine clinical testing for HIV-2 is not recommended. The only indications are in blood centers or for persons who have specific demographic or behavioral risk factors for HIV-2. These persons include those from countries where HIV-2 is endemic (West Africa, Angola, Mozambique, France, and Portugal) and their sex partners. The possibility of HIV-2 infection should also be considered in situations where there is clinical suspicion of HIV disease in the absence of a positive HIV-1 antibody test.

Acute Retroviral Syndrome

This syndrome occurs in many persons shortly after HIV infection, before antibody tests are positive. The syndrome is characterized by acute symptoms and signs, including fever, malaise, lymphadenopathy, and skin rash. Suspi-

Initial Management of HIV Infection

It is advisable to refer HIV-infected persons to a single clinical resource for comprehensive care (Centers for Disease Control and Prevention, 2006). Because of the limited availability of these facilities, it is often advisable to initiate evaluation and provide access to psychosocial services while planning for referral and continuation of medical care. Thus, brief consideration of initial management is in order.

Recently diagnosed HIV infection may not have been acquired recently. Persons with newly diagnosed HIV infection can be at any of the clinical stages of infection. Thus, it is important to be alert for signs and symptoms that suggest advanced infection, such as fever, weight loss, diarrhea, oral candidiasis, cough, or shortness of breath. These findings suggest the need for urgent referral.

In nonemergent situations, the recommended evaluation of a person with a newly diagnosed HIV infection includes a detailed medical history that emphasizes sexual and substance abuse history, previous STDs, and specific HIV-related symptoms or diagnoses. The physical examination should include a pelvic examination in women, with Pap smear and testing for gonorrhea and chlamydial infection. Recommended blood work includes complete blood count with platelet count; chemistry profile; testing for toxoplasma antibody and hepatitis viral markers; syphilis serologic test; and a CD4+ T-lymphocyte count (Centers for Disease Control and Prevention, 2006). Other evaluations should include a tuberculin skin test and chest x-ray. Finally, provision should be made for evaluation and management of sex and injecting-drug partners.

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Urinary calculi are the third most common affliction of the urinary tract, exceeded only by urinary tract infections and pathologic conditions of the prostate. They are common in both animals and humans. The nomenclature associated with urinary stone disease arises from a variety of disciplines. Struvite stones, for example, composed of magnesium ammonium phosphate hexahydrate, are named in honor of H.C.G. von Struve (1772–1851), a Russian naturalist. Before the time of von Struve, the stones were referred to as guanite, because magnesium ammonium phosphate is prominent in bat droppings. Calcium oxalate dihydrate is frequently referred to as weddellite, because it was commonly found in floor samples collected from the Weddell Sea in Antarctica. The history of the nomenclature associated with urinary stone disease is as intriguing as that of the development of the interventional techniques used in their treatment.

Urinary stones have plagued humans since the earliest records of civilization. The etiology of stones remains speculative. If urinary constituents are similar in each kidney and if there is no evidence of obstruction, why do most stones present in a unilateral fashion? Why don’t small stones pass uneventfully down the ureter early in their development? Why do some people form one large stone and others form multiple small calculi? There is much speculation concerning these and other questions. Advances in the surgical treatment of urinary stones have outpaced our understanding of their etiology. As clinicians we are concerned with an expedient diagnosis and efficient treatment. Equally important is a thorough metabolic evaluation directing appropriate medical therapy and lifestyle changes to help reduce recurrent stone disease. Without such follow-up and medical intervention, stone recurrence rates can be as high as 50% within 5 years. Uric acid calculi can recur even more frequently. Physicians look forward to gaining a better understanding of this multifactorial disease process in hopes of developing more effective prophylaxis.

RENAL & URETERAL STONES

Etiology

Mineralization in all biologic systems has a common theme in that the crystals and matrix are intertwined. Urinary stones are no exception; they are polycrystalline aggregates composed of varying amounts of crystalloid and organic matrix. Theories to explain urinary stone disease are incomplete. Stone formation requires supersaturated urine. Supersaturation depends on urinary pH, ionic strength, solute concentration, and complexation. Urinary constituents may change dramatically during different physiologic states from a relatively acid urine in a first morning void to an alkaline tide noted after meals. Ionic strength is determined primarily by the relative concentration of monovalent ions. As ionic strength increases, the activity coefficient decreases. The activity coefficient reflects the availability of a particular ion.

The role of solute concentrations is clear: The greater the concentration of 2 ions, the more likely they are to precipitate. Low ion concentrations result in undersaturation and increased solubility. As ion concentrations increase, their activity product reaches a specific point termed the solubility product ($K_{sp}$). Concentrations above this point are metastable and are capable of initiating crystal growth and heterogeneous nucleation. As solutions become more concentrated, the activity product eventually reaches the formation product ($K_{fp}$). Supersaturation levels beyond this point are unstable, and spontaneous homogeneous nucleation may occur.

Multiplying 2 ion concentrations reveals the concentration product. The concentration products of most ions are greater than established solubility products. Other factors must play major roles in the development of urinary calculi, including complexation. Complexation influences the availability of specific ions. For instance, sodium complexes with oxalate and decreases its free ionic form, while sulfates can complex with calcium. Crystal formation is modified by a variety of other substances found in the urinary tract, including magnesium, citrate, pyrophosphate, and a variety of trace metals. These inhibitors may act at the active crystal growth sites or as inhibitors in solution (as with citrate).

The nucleation theory suggests that urinary stones originate from crystals or foreign bodies immersed in supersaturated urine. This theory is challenged by the same arguments that support it. Stones do not always form in patients who are hyperexcretors or who are at risk for dehydration. Additionally, many stone formers’ 24-hour urine collections are completely normal with respect to stone-forming ion concentrations.
The crystal inhibitor theory claims that calculi form owing to the absence or low concentration of natural stone inhibitors, including magnesium, citrate, pyrophosphate, and a variety of trace metals. This theory does not have absolute validity since many people lacking such inhibitors may never form stones, and others with an abundance of inhibitors may, paradoxically, form them.

A. CRYSTAL COMPONENT

Stones are composed primarily of a crystalline component. Crystals of adequate size and transparency are easily identified under a polarizing microscope. X-ray diffraction is preferred to assess the geometry and architecture of calculi. A group of stones from the same geographic location or the same historical time period typically have crystalline constituents that are common.

Multiple steps are involved in crystal formation, including nucleation, growth, and aggregation. Nucleation initiates the stone process and may be induced by a variety of substances, including proteinaceous matrix, crystals, foreign bodies, and other particulate tissues. Heterogeneous nucleation (epitaxy), which requires less energy and may occur in less saturated urine, is a common theme in stone formation. It should be suspected whenever an oriented conglomerate is found. A crystal of one type thereby serves as a nidus for the nucleation of another type with a similar crystal lattice. This is frequently seen with uric acid crystals initiating calcium oxalate formation. It takes time for these early nidi to grow or aggregate to form a stone incapable of passing with ease through the urinary tract.

How these early crystalline structures are retained in the upper urinary tract without uneventful passage down the ureter is unknown. The theory of mass precipitation or intranephronic calculosis suggests that the distal tubules or collecting ducts, or both, become plugged with crystals, thereby establishing an environment of stasis, ripe for further stone growth. This explanation is unsatisfactory; tubules are conical in shape and enlarge as they enter the papilla, thereby reducing the possibility of ductal obstruction. Additionally, urine transit time from the glomerulus into the renal pelvis is only a few minutes, making crystal aggregation and growth within the uriniferous tubules unlikely.

The fixed particle theory postulates that formed crystals are somehow retained within cells or beneath tubular epithelium. Randall noted whitish-yellow precipitations of crystalline substances occurring on the tips of renal papillae as submucosal plaques. These can be appreciated during endoscopy of the upper urinary tract. Carr hypothesized that calculi form in obstructed lymphatics and then rupture into adjacent fornices of a calyx. Arguing against Carr’s theory are the grossly visible early stone elements in areas remote from fornices.

B. MATRIX COMPONENT

The amount of the noncrystalline matrix component of urinary stones varies with stone type, commonly ranging from 2% to 10% by weight. It is composed predominantly of protein, with small amounts of hexose and hexosamine. An unusual type of stone called a matrix calculus can be associated with previous kidney surgery or chronic urinary tract infections and has a gelatinous texture (Figure 16–1).

Histologic inspection reveals laminations with scant calcifications. On plain abdominal radiographs, matrix calculi are usually radiolucent and can be confused with other filling defects, including blood clots, upper-tract tumors, and fungal bezoars. Computed tomography (CT) reveals calcifications and can help to confirm the diagnosis. The role of matrix in the initiation of ordinary urinary stones as well as matrix stones is unknown. It may serve as a nidus for crystal aggregation or as a naturally occurring glue to adhere small crystal components and thereby hinder uneventful passage down the urinary tract. Alternatively, the matrix may have an inhibitory role in stone formation or may be an innocent bystander, playing no active role in stone formation.

Urinary Ions

A. CALCIUM

Calcium is a major ion present in urinary crystals. Only 50% of plasma calcium is ionized and available for filtration at the glomerulus. Well over 95% of the calcium filtered at the glomerulus is reabsorbed at both the proximal and distal tubules and limited amounts in the collecting tube. Less than 2% is excreted in the urine. Diuretic medications may exert a hypocalciuric effect by further decreasing calcium excretion. Many factors influence the availability of calcium in solution, including complexation with citrate, phosphate, and sulfate. An increase in monosodium urates and a
decrease in urinary pH further interfere with this complex-
ation and therefore promote crystal aggregation.

B. Oxalate

Oxalate is a normal waste product of metabolism and is relatively insoluble. Normally, approximately 10–15% of oxalate found in the urine originates from the diet; the vast majority is a metabolic by-product. Most of the oxalate that enters the large bowel is consumed by bacterial decomposition. Diet, however, can have an impact on the amount of oxalate found in the urine. Once absorbed from the small bowel, oxalate is not metabolized and is excreted almost exclusively by the proximal tubule. The presence of calcium within the bowel lumen is an important factor influencing the amount of oxalate that is absorbed. The control of oxalate in the urine plays a pivotal role in the formation of calcium oxalate calculi. Normal excretion ranges from 20 to 45 mg/day and does not change significantly with age. Excretion is higher during the day when one eats. Small changes in oxalate levels in the urine can have a dramatic impact on the supersaturation of calcium oxalate. The principal precursors of oxalate are glycine and ascorbic acid; however, the impact of ingested vitamin C (<2 g/day) is negligible.

Hyperoxaluria may develop in patients with bowel disorders, particularly inflammatory bowel disease, small-bowel resection, and bowel bypass. Renal calculi develop in 5–10% of patients with these conditions. Chronic diarrhea with fatty stools results in a saponification process. Intraluminal calcium binds to the fat, thereby becoming unavailable to bind to oxalate. The unbound oxalate is readily absorbed.

Excessive oxalate may occur secondary to the accidental or deliberate ingestion of ethylene glycol (partial oxidation to oxalate). This may result in diffuse and massive deposition of calcium oxalate crystals and may occasionally lead to renal failure.

C. Phosphate

Phosphate is an important buffer and complexes with calcium in urine. It is a key component in calcium phosphate and magnesium ammonium phosphate stones. The excretion of urinary phosphate in normal adults is related to the amount of dietary phosphate (especially in meats, dairy products, and vegetables). The small amount of phosphate filtered by the glomerulus is predominantly reabsorbed in the proximal tubule. Parathyroid hormone inhibits this reabsorption. The predominant crystal found in those with hyperparathyroidism is phosphate, in the form of hydroxyapatite, amorphous calcium phosphate, and carbonate apatite.

D. Uric Acid

Uric acid is the by-product of purine metabolism. The pKa of uric acid is 5.75. Undissociated uric acid predominates with pH values less than this. Elevated pH values increase urate, which is soluble. Approximately 10% of the filtered uric acid finds its way into the urine. Other defects in purine metabolism may result in urinary stone disease. Rarely, a defect in xanthine oxidase results in increased levels of xanthine; the xanthine may precipitate in urine, resulting in stone formation. Unusual alterations in adenine metabolism may result in the production of 2, 8-dihydroxyadeninuria, which is poorly soluble in urine and may develop into a urinary stone. This results from a deficiency of adenine phosphoribosyltransferase (APRT). Pure uric acid crystals and calculi are typically radiolucent in nature and may not be identified on plain abdominal films (Figure 16–2). They are visible on noncontrast CT images. Some uric acid calculi may be partially radiopaque, however, because of associated calcium deposits.

E. Sodium

Although not identified as one of the major constituents of most urinary calculi, sodium plays an important role in regulating the crystallization of calcium salts in urine. Sodium is found in higher than expected concentrations in the core of renal calculi and may play a role in initiating crystal development and aggregation. High dietary sodium intake increases urinary calcium excretion. This reduces the ability of urine to inhibit calcium oxalate crystal agglomeration. These effects are thought to be due to a sodium-induced increase in bicarbonaturia and decrease in serum bicarbonate. Conversely, a reduction in dietary sodium helps to reduce recurrent calcium nephrolithiasis.

F. Citrate

Citrate is a key factor affecting the development of calcium urinary stones. A deficiency commonly is associated
with stone formation in those with chronic diarrhea or renal tubular acidosis type I (distal tubular defect) and in patients undergoing chronic thiazide therapy. Citrate plays a pivotal role in the citric acid cycle in renal cells. Metabolic stimuli that consume this product (as with intracellular metabolic acidosis due to fasting, hypokalemia, or hypomagnesemia) reduce the urinary excretion of citrate. Estrogen increases citrate excretion and may be a factor that decreases the incidence of stones in women, especially during pregnancy. Alkalosis also increases citrate excretion.

G. Magnesium

Dietary magnesium deficiency is associated with an increased incidence of urinary stone disease. Magnesium is a component of struvite calculi. Experimentally, lack of dietary magnesium is associated with increased calcium oxalate stone formation and calcium oxalate crystalluria. The exact mechanism whereby magnesium exerts its effect is undefined. Dietary magnesium supplements do not protect against stone formation in normal people.

H. Sulfate

Urinary sulfates may help prevent urinary calculi. They can complex with calcium. These sulfates occur primarily as components of longer urinary proteins, such as chondroitin sulfate and heparin sulfate.

I. Other Urinary Stone Inhibitors

Inhibitors of urinary stone formation other than citrate, magnesium, and sulfates have been identified. These consist predominantly of urinary proteins and other macromolecules such as glycosaminoglycans, pyrophosphates, and uropontin. Although citrate appears to be the most active inhibitory component in urine, these substances demonstrate a substantial role in preventing urine crystal formation. The N-terminal amino acid sequence and the acidic amino acid content of these protein inhibitors, especially their high aspartic acid content, appear to play pivotal inhibitory roles. Fluoride may be an inhibitor of urinary stone formation.

Stone Varieties

A. Calcium Calculi

Calcifications can occur and accumulate in the collecting system, resulting in nephrolithiasis. Eighty to eighty-five percent of all urinary stones are calcareous. Calcium nephrolithiasis is most commonly due to elevated urinary calcium, elevated urinary uric acid, elevated urinary oxalate, or a decreased level of urinary citrate. Hypercalciiuria is found as a solitary defect in 12% of patients and in combination with other defects in an additional 18%. Hyperuricosuria is identified as a solitary defect in 8% of patients and associated with additional defects in 16%. Elevated urinary oxalate is found as a solitary finding in 5% of patients and as a combined defect in 16%. Finally, decreased urinary citrate is found as an isolated defect in 17% of patients and as a combined defect in an additional 10%. Approximately one-third of patients undergoing a full metabolic evaluation will find no identifiable metabolic defect.

Symptoms are secondary to obstruction, with resultant pain, infection, nausea and vomiting, and rarely culminate in renal failure. Asymptomatic hematuria or repetitive urinary tract infections recalcitrant to apparently appropriate antibiotics should lead one to suspect a urinary stone. Calculifications within the parenchyma of the kidney, known as nephrocalcinosis, rarely cause symptoms, however, and usually are not amenable to traditional therapies appropriate for urinary stone disease (Figure 16–3). Nephrocalcinosis is frequently encountered with renal tubular acidosis and hyperparathyroidism. Nephrolithiasis and nephrocalcinosis frequently coexist. Most patients with nephrolithiasis, however, do not have obvious nephrocalcinosis. Nephrocalcinosis may result from a variety of pathologic states. Ectatic collecting tubules, as seen with medullary sponge kidney, are common. This is frequently a bilateral process. Increased calcium absorption from the small bowel is common with sarcoidosis, milk-alkali syndrome, hyperparathyroidism, and excessive vitamin D intake. Disease processes resulting in bony destruction, including hyperparathyroidism, osteolytic lesions, and multiple myeloma, are a third mechanism. Finally, dystrophic calcifications forming on necrotic tissue may develop after a renal insult.

1. Absorptive hypercalciuric nephrolithiasis—Normal calcium intake averages approximately 900–1000 mg/day. Approximately one-third is absorbed by the small bowel, and of that portion approximately 150–200 mg is
obligatorily excreted in the urine. A large reservoir of calcium remains in the bone. Most dietary calcium is excreted in the stool. Absorptive hypercalcemia is secondary to increased calcium absorption from the small bowel, predominately from the jejunum. This results in an increased load of calcium filtered from the glomerulus. The result is suppression of parathyroid hormone, leading to decreased tubular reabsorption of calcium, culminating in hypercalcemia (>4 mg/kg). This physiologic cascade is in response to the primary defect, an increased absorption of calcium from the small bowel.

Absorptive hypercalcemia can be subdivided into 3 types. Type I absorptive hypercalcemia is independent of diet and represents 15% of all calcareous calculi. There is an elevated urinary calcium level (>150–200 mg/24 h) even during a calcium-restricted diet. Cellulose phosphate is an effective nonabsorbable exchange resin. This effectively binds the calcium in the gut, preventing bowel absorption. Cellulose phosphate has no impact on the calcium transport defect. Urinary calcium excretion returns to normal values with therapy.

Cellulose phosphate must be taken with meals to be available when calcium is ingested. A typical dose is 10–15 g orally in 3 divided doses and is well tolerated. This therapy is relatively contraindicated in postmenopausal women and in children during their active growth cycles. Inappropriate use may lead to a negative calcium balance and a secondary hyperparathyroid state. As with all stone formers, long-term follow-up is required. Cellulose phosphate may bind other cations besides calcium, including magnesium. Secondary hyperoxaluria may develop owing to decreased calcium in the gut. See the section on hyperoxaluria for a more detailed discussion.

Hydrochlorothiazides are an alternative treatment for type I absorptive hypercalcemia. Initially there is a reduction in renal excretion of calcium. The increased absorbed calcium is likely deposited in bone. Eventually the bone reservoir reaches its capacity and the drug becomes ineffective. Hydrochlorothiazides have limited long-term efficacy (approximately 3–5 years). These drugs have no effect on the defective bowel transport system. Hydrochlorothiazides may be alternated with cellulose phosphate as an effective treatment regimen.

Type II absorptive hypercalcemia is dietary dependent and is a common cause of urinary stone disease. There is no specific medical therapy. Calcium excretion returns to normal on a calcium-restricted diet. Patients should limit their calcium intake to 400–600 mg/day. Type II absorptive hypercalcemia is not as severe as type I.

Type III absorptive hypercalcemia is secondary to a phosphate renal leak and accounts for 5% of all urinary calculi. Decreased serum phosphate leads to an increase in 1, 25-dihydroxyvitamin D synthesis. The physiologic cascade culminates in an increased absorption of phosphate and calcium from the small bowel and an increased renal excretion of calcium—hence its classification as absorptive hypercalcemia. Successful treatment replaces bioavailable phosphate. Orthophosphate (Neutra-Phos) inhibits vitamin D synthesis and is administered as 250 mg three to four times daily. It is best taken after meals and before bedtime. Orthophosphates do not alter intestinal calcium absorption.

2. Resorptive hypercalciuric nephrolithiasis—About half the patients with clinically obvious primary hyperparathyroidism present with nephrolithiasis. This group represents less than 5–10% of all patients with urinary stones. Patients with calcium phosphate stones, women with recurrent calcium stones, and those with both nephrocalcinosis and nephrolithiasis should be suspected of having hyperparathyroidism. Hypercalcemia is the most consistent sign of hyperparathyroidism.

Parathyroid hormone results in a cascade of events starting with an increase in urinary phosphorus and a decrease in plasma phosphorus, followed by an increase in plasma calcium and a decrease in urinary calcium. Its action on the kidney and on the bone is independent of each other. Ultimately renal damage is secondary to the hypercalcemia. It limits the concentrating ability of the kidney and impairs the kidney’s ability to acidify urine. Surgical removal of the offending parathyroid adenoma is the only effective way of treating this disease. Attempts at medical management are futile.

3. Renal-induced hypercalciuric nephrolithiasis—Hypercalcemia of renal origin is due to an intrinsic renal tubular defect in calcium excretion. This creates a physiologically vicious cycle. Excessive urinary calcium excretion results in a relative decrease in serum calcium, which leads to a secondarily increased parathyroid hormone level that mobilizes calcium from the bone and increases calcium absorption from the gut. This step completes the pathologic cycle by delivering increased levels of calcium back to the kidney, whereby the renal tubules excrete large amounts of calcium. These patients have an elevated fasting urinary calcium level, normal serum calcium level, and an elevated parathyroid hormone level.

Renal hypercalcemia is effectively treated with hydrochlorothiazides. Unlike their role in type I absorptive hypercalcemia, in this setting hydrochlorothiazides have a durable long-term effect. As a diuretic, they decrease the circulating blood volume and subsequently stimulate proximal tubular absorption of calcium as well as other constituents. They also increase reabsorption at the distal tubule. Both mechanisms correct the secondary hyperparathyroid state.

Hypercalciuric states may result in elevated parathyroid levels. To differentiate primary from secondary hyperparathyroidism in patients with urinary stone disease, one can prescribe a hydrochlorothiazide challenge of 50 mg twice a day for approximately 10 days. Patients with secondary hyperparathyroidism will have normal serum parathyroid
levels, while those with primary hyperparathyroidism will continue to have elevated serum values.

4. Hyperuricosuric calcium nephrolithiasis—Hyperuricosuric calcium nephrolithiasis is due to either an excessive dietary intake of purines or an increase in endogenous uric acid production. In both situations there is an increase in urinary monosodium urates. Monosodium urates absorb and adsorb urinary stone inhibitors and facilitate heterogeneous nucleation.

Patients have elevated urinary uric acid levels (>600 mg/24 h in women and >750 mg/24 h in men) and consistently have a urinary pH >5.5. The urinary pH helps to differentiate hyperuricosuric calcium from hyperuricosuric uric acid stone formation.

Patients with excessive purine intake can be effectively treated by changing their diet to one with low purines. Those with excessive endogenous uric acid production can be successfully treated with allopurinol. Allopurinol is a xanthine oxidase inhibitor. Allopurinol reduces uric acid synthesis and renal excretion of uric acid. It also inhibits uric acid-calcium oxalate crystallization. Allopurinol has many potential side effects, including a variety of skin rashes and liver toxicity, and should be administered with careful monitoring (300 mg daily). Potassium citrate is an alternative treatment, especially when associated with hypocitraturia.

5. Hyperoxaluric calcium nephrolithiasis—Hyperoxaluric calcium nephrolithiasis is secondary to increased urinary oxalate levels (>40 mg/24 h). It is frequently found in patients with inflammatory bowel disease or other chronic diarrheal states that result in severe dehydration. It is rarely associated with excessive oxalate intake, as seen in poisoning with ethylene glycol or endogenous overproduction.

Chronic diarrheal states alter oxalate metabolism. Malabsorption leads to increased luminal fat and bile. Intraluminal calcium readily binds to fat, resulting in a saponification process. Urinary calcium levels are usually low (<100 mg/24 h). The intraluminal gut calcium that normally would have bound to oxalate is decreased. The unbound oxalate is readily absorbed and is unaffected by the usual metabolic inhibitors of energy-dependent pumps. Bile salts may increase the passive bowel absorption of oxalate. A small increase in oxalate absorption and subsequent urinary excretion dramatically increases the formation product of calcium oxalate. This increases the potential for heterogeneous nucleation and crystal growth in this metastable environment. All patients with increased urinary excretion of oxalate do not form calcium oxalate calculi. Other factors must be contributory, therefore, including dehydration, hypocitraturia (associated with an acidosis), decreased excretion of urinary inhibitors including magnesium, and protein malabsorption.

Enteric hyperoxaluric calcium nephrolithiasis is successfully treated with oral calcium supplementation. The calcium binds to the intraluminal oxalate and limits its absorption. It must be given with meals when the oxalate is present. Other oral cations are effective, including magnesium supplements. An alternative therapy includes a diet limited to medium-chain fatty acids and triglycerides; however, it is poorly tolerated by patients. Equally difficult is an attempt to alter oxalate intake. Unless large amounts of specific oxalate-rich foods can be excluded, an alternative diet may result in increased oxalate levels.

Primary hyperoxaluria is a rare hereditary disease. It is associated with calcium oxalate renal calculi, nephrocalcinosis, and other distant deposits of oxalate, culminating in progressive renal failure and eventual death. Type I is associated with an enzyme deficiency of 2-oxoglutarate: glyoxylate carboligase, resulting in elevated urinary levels of glycocolic acid and oxalic acid. Type II has increased excretory levels of D-glyceric acid rather than elevated levels of glycolic acid. It is associated with a D-glycerate dehydrogenase enzyme deficiency. This ultimately results in the accumulation of hydroxypropruvate, which is eventually converted to oxalate. Oxalate crystal deposits develop rapidly in transplanted kidneys. Combined liver and renal transplantation has cured this previously fatal rare disease.

6. Hypocitraturic calcium nephrolithiasis—Citrate is an important inhibitor of urinary stone disease. Increased metabolic demands on the mitochondria of renal cells decrease the excretion of citrate. Such conditions include intracellular metabolic acidosis, hypokalemia (as with thiazide therapy), fasting, hypomagnesemia, androgens, gluconeogenesis, and an acid-ash diet. Citrate may be consumed in the urine by bacteria during a urinary tract infection. The cause of hypocitraturia may be unknown in some cases. In contrast, alkalosis, alkaline-ash diet, estrogen, and vitamin D increase urinary citrate levels.

Citrate has its action in solution. It complexes with calcium, thereby decreasing the ionic calcium concentration and thus the activity product and thereby decreasing the energy for crystallization. Citrate decreases agglomeration, spontaneous nucleation, and crystal growth of calcium oxalate. Citrate also decreases calcium oxalate calculi by decreasing monosodium urates that can absorb inhibitors and facilitate heterogeneous nucleation.

Hypocitraturic (<320 mg/24 h) calcium nephrolithiasis is associated commonly with renal tubular acidosis type I (distal tubule) (Figure 16–4), thiazide therapy (accompanied by potassium wastage), and chronic diarrhea. Treatment is successful with potassium citrate supplementation. Routine dosage is 20–30 mEq two to three times daily and is usually well tolerated. Six to eight glasses of lemonade can increase urinary citrate excretion by approximately 150 mg/24 h and thus either limit or eliminate the need for pharmacologic citrate supplementation.

B. NONCALCIUM CALCULI

1. Struvite—Struvite stones are composed of magnesium, ammonium, and phosphate (MAP). They are found most commonly in women and may recur rapidly. They
frequently present as renal staghorn calculi and rarely present as ureteral stones except after surgical intervention (Figure 16–5). Struvite stones are infection stones associated with urea-splitting organisms, including Proteus, Pseudomonas, Providencia, Klebsiella, Staphylococci, and Mycoplasma. The high ammonium concentration derived from the urea-splitting organisms results in an alkaline urinary pH. The urinary pH of a patient with a MAP stone rarely is <7.2 (normal urinary pH is 5.85). It is only at this elevated urinary pH (>7.19) that MAP crystals precipitate. MAP crystals are soluble in the normal urinary pH range of 5–7. Preoperative bladder cultures do not necessarily reflect the bacteriologic composition found in calculi. Foreign bodies and neurogenic bladders may predispose patients to urinary infections and subsequent struvite stone formation. Massive diuresis does not prevent struvite calculi. Women with recurrent infections despite apparently appropriate antibiotic therapy should be evaluated for struvite calculi with a conventional kidney-ureter-bladder (KUB) film or renal ultrasound, or both. It is impossible to sterilize such calculi with antibiotics. Culture-specific antibiotics can reduce urease levels by 99% and help reduce stone recurrence. Stone removal is therapeutic.

Long-term management is optimized with the removal of all foreign bodies, including catheters of all varieties. A short ileal loop helps decrease the risk of stones in those with supravesical urinary diversion. All stone fragments should be removed with or without the aid of follow-up irrigations. Hemiacidrin (Renacidin) irrigations should be used with caution if at all. Rapid magnesium toxicity can result in death even with a low-pressure irrigation setup (<20 cm water pressure), negative daily urine cultures, and no evidence of upper-tract extravasation. Acetohydroxamic acid inhibits the action of bacterial urease, thereby reducing the urinary pH and decreasing the likelihood of precipitation. Most patients have a difficult time tolerating this medication.

2. Uric acid—Uric acid stones compose <5% of all urinary calculi and are usually found in men. Patients with gout, myeloproliferative diseases, or rapid weight loss, and those treated for malignant conditions with cytotoxic drugs have a high incidence of uric acid lithiasis. Most patients with uric acid calculi, however, do not have hyperuricemia. Elevated uric acid levels are frequently due to dehydration and excessive purine intake. Patients present with a urinary pH consistently <5.5, in contrast to patients with hyperuricosuric calcium nephrolithiasis, who have a urinary pH >5.5. As the urinary pH increases above the dissociation constant pKa of 5.75, it dissociates into a relatively soluble urate ion. Treatment is centered on maintaining a urine volume >2 L/day and a urinary pH >6. Reducing dietary purines or the administration of allopurinol also helps reduce uric acid excretion. Alkalization (with oral sodium bicarbonate, potassium bicarbonate, potassium citrate, or intravenous one-sixth normal sodium lactate) may dissolve calculi and is dependent on the stone surface area. Stone fragments after lithotripsy have a dramatically increased surface area and will dissolve more rapidly. Dissolution proceeds at approximately 1 cm of stone (as seen on KUB) per month, with compliant alkalization.

3. Cystine—Cystine lithiasis is secondary to an inborn error of metabolism resulting in abnormal intestinal (small bowel) mucosal absorption and renal tubular absorption of dibasic amino acids, including cystine, ornithine, lysine, and arginine. The genetic defects associated with cystinuria
have now been mapped to chromosome 2p.16 and more recently to 19q13.1. Cystine lithiasis is the only clinical manifestation of this defect. Classic cystinuria is inherited in an autosomal recessive fashion. Homozygous expression has a prevalence of 1:20,000, while the heterozygous expression is 1:2000. It represents 1–2% of all urinary stones, with a peak incidence in the second or third decade. Homozygous cystinurics excrete more than 250 mg/day, resulting in constant supersaturation. Heterozygous patients usually excrete 100–150 mg/day. Unaffected patients typically excrete <40 mg/day. Approximately 400 mg/L of cystine can remain in solution at a urinary pH of 7. As the urinary pH increases >7, the amount of soluble cystine increases exponentially.

The solubility of cystine is pH-dependent, with a pK of approximately 8.1. There is no difference in the solubility curves in normal versus cystinuric patients. There is no known inhibitor for cystine calculi, and cystine stone formation is completely dependent on excessive cystine excretion. Cystine stones are frequently associated with calcium calculi and their related metabolic abnormalities. They may present as single, multiple, or staghorn stones. The diagnosis is suspected in patients with a family history of urinary stones and the radiographic appearance of a faintly opaque, ground-glass, smooth-edged stone (Figure 16–6). Urinalysis frequently reveals hexagonal crystals. Stone analysis confirms the diagnosis. Quantitative urinary cystine evaluation helps confirm the diagnosis and differentiate heterozygous from homozygous states. It is also important to titrate medical therapy.

Medical therapy includes high fluid intake (>3 L/day) and urinary alkalinization. Patients should monitor their pH with nitrazine indicator paper and keep their pH values >7.5. It is difficult or impossible to maintain levels >8. A low-methionine (precursor to cystine) diet has limited impact, as most of the cystine is endogenous and most of the ingested methionine is incorporated into protein. Glutamine, ascorbic acid, and captopril are effective in some patients. Penicillamine can reduce urinary cystine levels. It complexes with the amino acid, and this complex is dramatically more soluble. Treatment should be titrated with quantitative urinary cystine values. Many patients poorly tolerate penicillamine, reporting skin rashes (discrete or confluent macules with occasional itching), loss of taste, nausea, vomiting, and anorexia. It may inhibit pyridoxine, which should be supplemented during treatment (50 mg/day). Mercaptopropionylglycine (Thiola), 300–1200 mg in divided doses, forms a soluble complex with cystine and can reduce stone formation. Side effects and frequent dosing decrease patient compliance rates. It is better tolerated than penicillamine and is now the first drug of choice in these difficult cases.

Surgical treatment is similar to that for other stones except that most stones are recalcitrant to extracorporeal (outside the body) shock wave lithotripsy (ESWL). One should have a low threshold to proceed with percutaneous stone extraction in symptomatic patients. Two populations of cystine stones have been described, including the rough and smooth varieties, and may reflect subpopulations: those that are effectively treated with ESWL and those that require more invasive therapy. Despite optimum medical therapy, a high stone recurrence rate frequently frustrates both patient and physician. Minimally invasive techniques and optimum medical therapy are paramount.

4. Xanthine—Xanthine stones are secondary to a congenital deficiency of xanthine oxidase. This enzyme normally catalyzes the oxidation of hypoxanthine to xanthine and of xanthine to uric acid. It is of interest that allopurinol, used to treat hyperuricosuric calcium nephrolithiasis and uric acid lithiasis, produces iatrogenic xanthinuria. Blood and urine levels of uric acid are lowered, and hypoxanthine and xanthine levels are increased; however, there are no case reports of xanthine stone formation resulting from allopurinol treatment. It is unlikely that allopurinol completely inhibits xanthine oxidase. Urinary stones develop in approximately 25% of patients with a xanthine
oxidase deficiency. The stones are radiolucent and are tannish yellow in color. Treatment should be directed by symptoms and evidence of renal obstruction. High fluid intake and urinary alkalinization are required for prophylaxis. If stones recur, a trial of allopurinol and a purine-restricted diet is appropriate.

5. Indinavir—Protease inhibitors are a popular and effective treatment in patients with acquired immunodeficiency syndrome. Indinavir is the most common protease inhibitor that results in radiolucent stones in up to 6% of patients who are prescribed this medication. Indinavir calculi are the only urinary stones to be radiolucent on non-contrast CT scans. They may be associated with calcium components and in these situations will be visible on non-contrast CT images. Temporary cessation of the medication with intravenous hydration frequently allows these stones to pass. The stones are tannish red and usually fall apart during basket extraction.

6. Rare—Silicate stones are very rare and are usually associated with long-term use of antacids containing silica. Surgical treatment is similar to that of other calculi.

Triamterene stones are radiolucent and have been identified with an increased frequency. They are associated with antihypertensive medications containing triamterene, such as Dyazide. Discontinuing the medication eliminates stone recurrences. Other medications that may become stone constituents include gafencin and antrafenine.

Rarely, patients arrive at an emergency room at an odd hour feigning signs and symptoms of passing a urinary stone in hopes of obtaining pain medications. They may add blood to their urine and give a believable story of a stone in hopes of obtaining pain medications. The character of the pain depends on the location. Calculi may present with a dull ache or flank discomfort. Frequently present with severe pain, while large staghorn calculi may present with a dull ache or flank discomfort.

The symptoms of acute renal colic depend on the location of the calculus; several regions may be involved: renal calyx, renal pelvis, upper and midureter, and distal ureter. An orderly progression of symptoms as a stone moves down the urinary tract is the exception.

1. Renal calyx—Stones or other objects in calyces or caliceal diverticula may cause obstruction and renal colic. In general, nonobstructing stones cause pain only periodically, owing to intermittent obstruction. The pain is a deep, dull ache in the flank or back that can vary in intensity from severe to mild. The pain may be exacerbated after consumption of large amounts of fluid. Radiographic imaging may not reveal evidence of obstruction despite the patient’s complaints of intermittent symptoms. It remains unclear how much of this pain is related to local mucosal irritation with activation of chemoreceptors. The presence of infection or inflammation in the calyx or diverticulum (eg, milk of calcium) in addition to obstruction may contribute to pain perception. Caliceal calculi occasionally result in spontaneous perforation with urinoma, fistula, or abscess formation.
Figure 16–7. Radiation of pain with various types of ureteral stone. **Upper left:** Ureteropelvic stone. Severe costovertebral angle pain from capsular and pelvic distention; acute renal and urethral pain from hyperperistalsis of smooth muscle of calyces, pelvis, and ureter, with pain radiating along the course of the ureter (and into the testicle, since the nerve supply to the kidney and testis is the same.) The testis is hypersensitive. **Upper right:** Midureteral stone. Same as above but with more pain in the lower abdominal quadrant. **Left:** Low ureteral stone. Same as above, with pain radiating into bladder, vulva, or scrotum. The scrotal wall is hyperesthetic. Testicular sensitivity is absent. When the stone approaches the bladder, urgency and frequency with burning on urination develop as a result of inflammation of the bladder wall around the ureteral orifice.
Caliceal calculi are frequently small and numerous and appear to be able to pass spontaneously. Long-term retention against the flow of urine and against the forces of gravity and antegrade peristalsis suggests a significant element of obstruction. Effective long-term treatment requires stone extraction and elimination of the obstructive component.

Pain relief has been reported in many patients following ESWL for small symptomatic caliceal calculi. Thus, if a patient continues to complain of pain in the face of a small caliceal calculus, ESWL treatment may be justified for both diagnosis and treatment. Percutaneous, retrograde, and laparoscopic techniques have been successful in the management of calculi in calyces or caliceal diverticula.

2. Renal pelvis—Stones in the renal pelvis >1 cm in diameter commonly obstruct the ureteropelvic junction, generally causing severe pain in the costovertebral angle, just lateral to the sacrospinalis muscle and just below the 12th rib. This pain may vary from being dull to excruciatingly sharp and is usually constant, boring, and difficult to ignore. It often radiates to the flank and also anteriorly to the upper ipsilateral abdominal quadrant. It may be confused with biliary colic or cholecystitis if on the right side and with gastritis, acute pancreatitis, or peptic ulcer disease if on the left, especially if the patient has associated anorexia, nausea, or emesis. Acquired or congenital ureteropelvic junction obstruction may cause a similar constellation of symptoms. Symptoms frequently occur on an intermittent basis following a drinking binge or consumption of large quantities of fluid.

Partial or complete staghorn calculi that are present in the renal pelvis are not necessarily obstructive. In the absence of obstruction, these patients often have surprisingly few symptoms such as flank or back pain. Recurrent urinary tract infections frequently culminate in radiographic evaluation with the discovery of a staghorn calculus. If untreated, these “silent” staghorn calculi can often lead to significant morbidity, including renal deterioration, infectious complications, or both.

3. Upper and midureter—Stones or other objects in the upper or midureter often cause severe, sharp back (costovertebral angle) or flank pain. The pain may be more severe and intermittent if the stone is progressing down the ureter and causing intermittent obstruction. A stone that becomes lodged at a particular site may cause less pain, especially if it is only partially obstructive. Stationary calculi that result in high-grade but constant obstruction may allow autoregulatory reflexes and pyelovenous and pyelolymphatic backflow to decompress the upper tract, with diminution in intraluminal pressure gradually easing the pain. Pain associated with ureteral calculi often projects to corresponding dermatomal and spinal nerve root innervation regions. The pain of upper ureteral stones thus radiates to the lumbar region and flank. Midureteral calculi tend to cause pain that radiates caudally and anteriorly toward the mid and lower abdomen in a curved, band-like fashion. This band initially parallels the lower costal margin but deviates caudal toward the bony pelvis and inguinal ligament. The pain may mimic acute appendicitis if on the right or acute diverticulitis if on the left side, especially if concurrent gastrointestinal symptoms are present.

4. Distal ureter—Calculi in the lower ureter often cause pain that radiates to the groin or testicle in males and the labia majora in females. This referred pain is often generated from the ilioinguinal or genital branch of the genitofemoral nerves. Diagnosis may be confused with testicular torsion or epididymitis. Stones in the intramural ureter may mimic cystitis, urethritis, or prostatitis by causing suprapubic pain, urinary frequency and urgency, dysuria, stranguria, or gross hematuria. Bowel symptoms are not uncommon. In women the diagnosis may be confused with menstrual pain, pelvic inflammatory disease, and ruptured or twisted ovarian cysts. Strictures of the distal ureter from radiation, operative injury, or previous endoscopic procedures can present with similar symptoms. This pain pattern is likely due to the similar innervation of the intramural ureter and bladder.

B. Hematuria

A complete urinalysis helps to confirm the diagnosis of a urinary stone by assessing for hematuria and crystalluria and documenting urinary pH. Patients frequently admit to intermittent gross hematuria or occasional tea-colored urine (old blood). Most patients will have at least microhematuria. Rarely (in 10–15% of cases), complete ureteral obstruction presents without microhematuria.

C. Infection

Magnesium ammonium phosphate (struvite) stones are synonymous with infection stones. They are commonly associated with Proteus, Pseudomonas, Providencia, Klebsiella, and Staphylococcus infections. They are rarely if ever associated with Escherichia coli infections. Calcium phosphate stones are the second variety of stones associated with infections. Calcium phosphate stones with a urine pH <6.6 are frequently referred to as brushite stones, whereas infectious apatite stones have a urine pH >6.6. Rarely, matrix stones with minimal crystalline components are associated with urinary tract infections. All stones, however, may be associated with infections secondary to obstruction and stasis proximal to the offending calculus. Culture-directed antibiotics should be administered before elective intervention.

Infection may be a contributing factor to pain perception. Uropathogenic bacteria may alter ureteral peristalsis by the production of exotoxins and endotoxins. Local inflammation from infection can lead to chemoreceptor activation and perception of local pain with its corresponding referral pattern.
1. Pyonephrosis—Obstructive calculi may culminate in the development of pyonephrosis. Unlike pyelonephritis, pyonephrosis implies gross pus in an obstructed collecting system. It is an extreme form of infected hydronephrosis. Presentation is variable and may range from asymptomatic bacteriuria to florid urosepsis. Bladder urine cultures may be negative. Radiographic investigations are frequently nondiagnostic. Renal ultrasonography may be misleading because of the nonspecific and variable appearance of pyonephrosis. Renal urine aspiration is the only way to make the definitive diagnosis. If the condition is noted at the time of a percutaneous nephrolithotomy, the procedure should be postponed to allow for adequate percutaneous drainage and treatment with appropriate intravenous antibiotics (Figure 16–8). If unrecognized and untreated, pyonephrosis may develop into a renocutaneous fistula.

2. Xanthogranulomatous pyelonephritis—Xanthogranulomatous pyelonephritis is associated with upper-tract obstruction and infection. One-third of patients present with calculi; two-thirds present with flank pain, fever, and chills. Fifty percent present with persistent bacteriuria. Urinalysis usually shows numerous red and white cells. This condition is a common imitator of other pathologic states of the kidney. It usually presents in a unilateral fashion. Open surgical procedures, such as a simple nephrectomy for minimal or nonrenal function, can be challenging owing to marked and extensive reactive tissues.

D. ASSOCIATED FEVER

The association of urinary stones with fever is a relative medical emergency. Signs of clinical sepsis are variable and include fever, tachycardia, hypotension, and cutaneous vasodilation. Costovertebral angle tenderness may be marked with acute upper-tract obstruction; however, it cannot be relied on to be present in instances of long-term obstruction. In such instances a mass may be palpable resulting from a grossly hydronephrotic kidney. Fever associated with urinary tract obstruction requires prompt decompression. This may be accomplished with a retrograde catheter (double-J, or an externalized variety to serve as a port for selective urine collections, injection of contrast material, or both). If retrograde manipulations are unsuccessful, insertion of a percutaneous nephrostomy tube is required.

E. NAUSEA AND VOMITING

Upper-tract obstruction is frequently associated with nausea and vomiting. Intravenous fluids are required to restore a euvolemic state. Intravenous fluids should not be used to force a diuresis in an attempt to push a ureteral stone down the ureter. Effective ureteral peristalsis requires coaptation of the ureteral walls and is most effective in a euvolemic state.

Special Situations

A. RENAL TRANSPLANTATION

Urinary stones associated with renal transplantation are rare. Perirenal nerves are severed at the time of renal harvesting. Classic renal colic is not found in these patients. The patients usually are admitted with the presumptive diagnosis of graft rejection. Only after appropriate radiographic and ultrasonic evaluation is the correct diagnosis made (Figure 16–9).

B. PREGNANCY

Renal colic is the most common nonobstetric cause of acute abdominal pain during pregnancy (Figure 16–10). Despite marked hypercalciuria associated with pregnancy, calculi are relatively rare, with an incidence approximating 1:1500 pregnancies. Women with known urinary stone disease do not have an increased risk of stones during pregnancy. The increased filtered load of calcium, uric acid, and sodium from the 25–50% increase in glomerular filtration rate associated with pregnancy has been thought to be a responsible factor in stone development.

The fetus demands special considerations regarding the potential dangers of radiation exposure (especially during the 1st trimester), medications, anesthesia, and surgical intervention. About 90% of symptomatic calculi present during the 2nd and 3rd trimesters. Initial investigations can be undertaken with renal ultrasonography and limited abdominal x-rays with appropriate shielding. Treatment requires balancing the safety of the fetus with the health of the mother. Temporizing measures to relieve upper-tract obstruction with a double-J ureteral stent or a percutaneous nephrostomy tube can be performed under local anesthesia. Treatment usually can be delayed until after delivery.

C. DYSMORPHIA

Patients with severe skeletal dysmorphia that is either congenital (spina bifida, myelomeningocele, cerebral palsy) or
acquired (arthritis, traumatic spinal cord injuries) and concurrent urinary calculi represent a unique clinical situation requiring special considerations (Figure 16–11). These skeletal abnormalities may preclude appropriate positioning for ESWL or percutaneous approaches. Calculi on the concave side in a patient with severe scoliosis may eliminate percutaneous puncture access between the rib and the posterosuperior iliac spine. Retrograde manipulations may need to be performed with flexible endoscopes due to marked contractures, making conventional dorsal lithotomy positioning impossible. Many such patients have undergone supravesical urinary diversion, so that retrograde access may be limited. Risks that need to be addressed include hypercalciuria associated with immobilization, relative dehydration due to patients’ or attendants’ attempts to reduce urinary output into external collecting devices, and the potential inability to drink without assistance. A full metabolic evaluation is even more important because these social and physical restrictions may be difficult or impossible to remedy.

D. OBESITY

Obesity is a risk factor for the development of urinary calculi. Surgical bypass procedures can cause hyperoxaluria. Massive weight gain or loss also may precipitate stone development. Obesity limits diagnostic and treatment
URINARY STONE DISEASE

E. MEDULLARY SPONGE KIDNEY

Medullary sponge kidney is a common condition characterized by tubular ectasia associated with parenchymal cysts and clefts that predispose to nephrolithiasis in 50% of affected patients. It is most often an asymptomatic condition; however it may present with renal colic, hematuria, or urinary tract infection. It is a radiographic diagnosis. The condition can involve select papillae or, more frequently, can be global. A full metabolic evaluation helps direct appropriate medical therapy.

F. RENAL TUBULAR ACIDOSIS

There are 3 main types of renal tubular acidosis: types I, II, and IV. Type I is associated with renal calculi.

Patients with type I renal tubular acidosis present with persistent acidemia with a low serum bicarbonate value unexplained by hyperventilation or known renal failure. The diagnosis should be suspected in those with a known family history, severe hypocitraturia, nephrocalcinosis, medullary sponge kidney, or a fasting urine pH >6 in the absence of infection. Patients usually present with nephrolithiasis (calcium phosphate), nephrocalcinosis, or osteomalacia (or a combination). This disease can be acquired as an adult or inherited with an autosomal dominant pattern. The diagnosis is confirmed by assessing the patient’s response to an acid load. This is frequently produced by a rapid oral ammonium chloride load (0.1 g/kg over 1 hour). The dose can be given before bedtime in the evening; the patient is instructed to fast until a second morning voided urine sample and a serum bicarbonate level are obtained. A normal person responds by eliminating the acid load in the urine, resulting in a urinary pH <5.3. Those who do not respond in this fashion can be said to have type I renal tubular acidosis. Additionally, the diagnosis should be challenged in those with normal citrate values. Treatment is centered on base replacement with potassium citrate or potassium bicarbonate solutions. Urinary citrate levels can be used to monitor effective treatment.

G. ASSOCIATED TUMORS

Squamous cell carcinoma of the upper urinary tract is uncommon but has been associated with calculi in more than 50% of cases. Chronic irritation from calculi or infection may be contributory factors. Upper-tract calculi may predispose patients to transitional cell carcinoma.

H. PEDIATRIC PATIENTS

Urinary calculi are unusual in children. A full and thorough metabolic evaluation should be undertaken. Stone analysis is particularly helpful in directing these investigations. Children born prematurely and given furosemide while in the neonatal intensive care unit are at increased risk of developing urinary stone disease. Treatment may be limited by endoscope size. Preliminary data show no change in renal growth after ESWL.

I. CALICEAL DIVERTICULA

Pyelocaliceal diverticula are cystic urine-containing diverticula of the upper tract lying within the renal parenchyma; they communicate through a narrow channel into the main collecting system (Figure 16–12). These diverticula occur in approximately 0.2–0.5% of the population and are congenital in origin; up to 40% are associated with calculi. Type I diverticula are the most common and are closely related to minor calyces. Type II have a direct communication with the renal pelvis and tend to be larger and symptomatic. Caliceal diverticula are usually asymptom-
atic, but patients may complain of flank pain or recurrent urinary tract infections. Frequently many small calculi, rather than a solitary stone, are found in these obstructed cavities. When intervention was required in the past, treatment was with nephrectomy, heminephrectomy, or open surgical unroofing. Less invasive means are used today. Communications with the collecting system are commonly pinpoint and may be difficult to locate through a retrograde approach. Retrograde access into superior pole diverticula has been successful. Surprisingly, treatment may be successful with ESWL if stone fragments are small enough to pass uneventfully. More commonly, percutaneous access and, more recently, laparoscopic means are used with success. Dilation of the caliceal neck, direct cautery, or sclerosis of the caliceal epithelium, or direct cautery and sclerosis of the caliceal epithelium can help reduce stone recurrence rates.

J. RENAL MALFORMATIONS

Anatomic renal variants such as ectopic kidneys, including the horseshoe kidney, predispose to renal calculi due to impaired urinary drainage. Pain symptoms appear to be no different from those reported in patients with normally positioned kidneys. Radiographic diagnosis may be difficult due to the unexpected location of the ureters and kidneys (Figure 16–13). If calculi can be targeted with ESWL, most stone fragments pass surprisingly uneventfully. Large stone burdens should be approached percutaneously as in normally positioned kidneys. Severe outlet obstruction should be corrected with open surgery, and concurrent calculi can be removed at the same setting. Aberrant vasculature should be appreciated before percutaneous and open procedures are undertaken.

Evaluation

A. DIFFERENTIAL DIAGNOSIS

Urinary stones can mimic other retroperitoneal and peritoneal pathologic states. A full differential diagnosis of the acute abdomen should be made, including acute appendicitis, ectopic and unrecognized pregnancies, ovarian pathologic conditions including twisted ovarian cysts, diverticular disease, bowel obstruction, biliary stones with and without obstruction, peptic ulcer disease, acute renal artery embolism, and abdominal aortic aneurysm, to mention a few. Peritoneal signs should be sought during physical examination.

B. HISTORY

A proper evaluation requires a thorough medical history. The nature of the pain should be evaluated, including its onset, character, potential radiation, activities that exacerbate or ease the pain, associated nausea and vomiting or gross hematuria, and a history of similar pain. Patients with previous stones frequently have had similar types of pain in the past, but not always.

C. RISK FACTORS

1. Crystalluria—Crystalluria is a risk factor for stones. Stone formers, especially those with calcium oxalate stones, frequently excrete more calcium oxalate crystals, and those crystals are larger than normal (>12 mm). The rate of stone formation is proportional to the percentage of large crystals and crystal aggregates. Crystal production is determined by the saturation of each salt and the urinary concentration of inhibitors and promoters. Urine samples should be fresh; they should be centrifuged and examined immediately for optimum results. Cystine crystals are hexagonal; struvite stones appear as coffin lids; brushite (CaHPO₄) stones are splinter-like and may aggregate with a spoke-like center; calcium apatite—(Ca₅(PO₄)₃(OH))—and uric acid crystals appear as amorphous powder because the crystals are so small; calcium oxalate dihydrate stones are bipyramids; and calcium oxalate monohydrate stones are small biconcave ovals that may appear as a dumbbell. Cystine and struvite crystals are always abnormal and require further investigations. Other crystals are frequently found in normal urinalyses.

2. Socioeconomic factors—Renal stones are more common in affluent, industrialized countries. Immigrants from

![Figure 16–13. Scout abdominal radiograph demonstrating horseshoe kidney with lateral ureteral deviation and double-J ureteral stent. Extraosseous calcifications are left lower calyceal stones.](image-url)
less industrialized nations gradually increase their stone incidence and eventually match that of the indigenous population. Use of soft water does not decrease the incidence of urinary stones.

3. Diet—Diet may have a significant impact on the incidence of urinary stones. As per capita income increases, the average diet changes, with an increase in saturated and unsaturated fatty acids, an increase in animal protein and sugar, and a decrease in dietary fiber, vegetable protein, and unrefined carbohydrates. A less energy-dense diet may decrease the incidence of stones. This fact has been documented during war years when diets containing minimal fat and protein resulted in a decreased incidence of stones. Vegetarians may have a decreased incidence of urinary stones. High sodium intake is associated with increased urinary sodium, calcium, and pH, and a decreased excretion of citrate; this increases the likelihood of calcium salt crystallization because the urinary saturation of monosodium urate and calcium phosphate (brushite) is increased. Fluid intake and urine output may have an effect on urinary stone disease. The average daily urinary output in stone formers is 1.6 L/day.

4. Occupation—Occupation can have an impact on the incidence of urinary stones. Physicians and other white-collar workers have an increased incidence of stones compared with manual laborers. This finding may be related to differences in diet but also may be related to physical activity; physical activity may agitate urine and dislodge crystal aggregates. Individuals exposed to high temperatures may develop higher concentrations of solutes owing to dehydration, which may have an impact on the incidence of stones.

5. Climate—Individuals living in hot climates are prone to dehydration, which results in an increased incidence of urinary stones, especially uric acid calculi. Although heat may cause a higher fluid intake, sweat loss results in lowered voided volumes. Hot climates usually expose people to more ultraviolet light, increasing vitamin D3 production. Increased calcium and oxalate excretion has been correlated with increased exposure time to sunlight. This factor has more impact on light-skinned people and may help explain why African Americans in the United States have a decreased stone incidence.

6. Family history—A family history of urinary stones is associated with an increased incidence of renal calculi. A patient with stones is twice as likely as a stone-free cohort to have at least one first-degree relative with renal stones (30% versus 15%). Those with a family history of stones have an increased incidence of multiple and early recurrences. Spouses of patients with calcium oxalate stones have an increased incidence of stones; this may be related to environmental or dietary factors.

7. Medications—A thorough history of medications taken may provide valuable insight into the cause of urinary calculi. The antihypertensive medication triamterene is found as a component of several medications, including Dyazide, and has been associated with urinary calculi with increasing frequency. Long-term use of antacids containing silica has been associated with the development of silicate stones. Carbonic anhydrase inhibitors may be associated with urinary stone disease (10–20% incidence). The long-term effect of sodium- and calcium-containing medications on the development of renal calculi is not known. Protease inhibitors in immunocompromised patients are associated with radiolucent calculi.

D. PHYSICAL EXAMINATION

A detailed physical examination is an essential component of the evaluation of any patient suspected of having a urinary calculus. The patient presenting with acute renal colic typically is in severe pain, often attempting to find relief in multiple, frequently bizarre positions. This fact helps differentiate patients with this condition from those with peritonitis, who are afraid to move. Systemic components of renal colic may be obvious, with tachycardia, sweating, and nausea often prominent. Costovertebral angle tenderness may be apparent. An abdominal mass may be palpable in patients with long-standing obstructive urinary calculi and severe hydronephrosis.

Fever, hypotension, and cutaneous vasodilation may be apparent in patients with urosepsis. In such instances there is an urgent need for decompression of the obstructed urinary tract, massive intravenous fluid resuscitation, and intravenous antibiotics. Occasionally, intensive-care support is needed.

A thorough abdominal examination should exclude other causes of abdominal pain. Abdominal tumors, abdominal aortic aneurysms, herniated lumbar disks, and pregnancy may mimic renal colic. Referral pain may be similar owing to common afferent neural pathways. Intestinal ileus may be associated with renal colic or other intra-peritoneal or retroperitoneal processes. Bladder palpation should be performed because urinary retention may present with pain similar to renal colic. Incarcerated inguinal hernias, epididymitis, orchitis, and female pelvic pathologic states may mimic urinary stone disease. A rectal examination helps exclude other pathologic conditions.

E. RADIOLOGIC INVESTIGATIONS

1. Computed tomography—Noncontrast spiral CT scans are now the imaging modality of choice in patients presenting with acute renal colic. It is rapid and is now less expensive than an intravenous pyelogram (IVP). It images other peritoneal and retroperitoneal structures and helps when the diagnosis is uncertain. It does not depend on an experienced radiologic technician to obtain appropriate oblique views when there is confusion with overlying bowel gas in a nonprepped abdomen. There is no need for intravenous contrast. Distal ureteral calculi can be con-
fused with phleboliths. These images do not give anatomic details as seen on an IVP (for example, a bifid collecting system) that may be important in planning intervention. If intravenous contrast material is used during the study, a KUB film can give additional helpful information. Uric acid stones are visualized no differently from calcium oxalate stones. Matrix calculi have adequate amounts of calcium to be visualized easily by CT.

2. **Intravenous pyelography**—An IVP can document simultaneously nephrolithiasis and upper-tract anatomy. Extraosseous calcifications on radiographs may be erroneously assumed to be urinary tract calculi (Figure 16–14). Oblique views easily differentiate gallstones from right renal calculi. Static hard-copy films can be interpreted by most clinicians. Anecdotally, small ureteral stones have passed spontaneously during such studies. An inadequate bowel preparation, associated ileus and swallowed air, and lack of available technicians may result in a less than ideal study when obtained during acute renal colic. A delayed, planned IVP may result in a superior study.

Acute fornical rupture is not uncommonly associated with a highly obstructive ureteral calculus. It may result in dramatic radiographs but is of no clinical significance, and no intervention is required. The rupture may be precipitated by the osmotic diuresis of the intravenous contrast agent based solely on radiographs.

3. **Tomography**—Renal tomography is useful to identify calculi in the kidney when oblique views are not helpful. It visualizes the kidney in a coronal plane at a set distance from the top of the x-ray table. This study may help identify poorly opacified calculi, especially when interfering abdominal gas or morbid obesity make KUB films suboptimal.

4. **KUB films and directed ultrasonography**—A KUB film and renal ultrasound may be as effective as an IVP in establishing a diagnosis. The ultrasound examination should be directed by notation of suspicious areas seen on a KUB film; it is, however, operator-dependent. The distal ureter is easily visualized through the acoustic window of a full bladder. Edema and small calculi missed on an IVP can be appreciated with such studies.

5. **Retrograde pyelography**—Retrograde pyelography occasionally is required to delineate upper-tract anatomy and localize small or radiolucent offending calculi. Bulb ureterograms frequently leak contrast material back into the bladder, resulting in a suboptimal study. Advancing an angiographic exchange catheter with or without the aid of a guidewire 3–4 cm into the ureter is an alternative technique. Intermittent fluoroscopic images direct appropriate injection volumes and help reduce the likelihood of pyelolymphatic, pyelosinus, and pyelovenous reflux.

6. **Magnetic resonance imaging**—MRI is a poor study to document urinary stone disease.

7. **Nuclear scintigraphy**—Nuclear scintigraphic imaging of stones has recently been appreciated. Bisphosphonate markers can identify even small calculi that are difficult to appreciate on a conventional KUB film (Figure 16–15). Differential radioactive uptake dependent on stone composition appreciated during in vitro studies cannot be appreciated on in vivo studies. Nuclear scintigraphy cannot delineate upper-tract anatomy in sufficient detail to help direct a therapeutic plan.

**Intervention**

**A. Conservative Observation**

Most ureteral calculi pass and do not require intervention. Spontaneous passage depends on stone size, shape, location, and associated ureteral edema (which is likely to depend on the length of time that a stone has not progressed). Ureteral calculus 4–5 mm in size have a 40–50% chance of spontaneous passage. In contrast, calculi >6 mm have a <5% chance of spontaneous passage. This does not mean that a 1-cm stone will not pass or that a 1- to 2-mm stone will always pass uneventfully.
The vast majority of stones that pass do so within a 6-week period after the onset of symptoms. Ureteral calculi discovered in the distal ureter at the time of presentation have a 50% chance of spontaneous passage, in contrast to a 25% and 10% chance in the mid- and proximal ureter, respectively.

B. DISSOLUTION AGENTS

The effectiveness of dissolution agents depends on stone surface area, stone type, volume of irrigant, and mode of delivery. Oral alkalinizing agents include sodium or potassium bicarbonate and potassium citrate. Extra care should be employed in patients susceptible to congestive heart failure or renal failure. Citrate is metabolized to bicarbonate and comes in a variety of preparations. Polycitra contains potassium and sodium citrate and citric acid. Bicitra contains only sodium citrate and citric acid. Food does not alter the effectiveness of these agents. Alternatively, orange juice alkalinizes urine. Intravenous alkalinization is effective with one-sixth molar sodium lactate.

Figure 16–15. A: Scout abdominal radiograph demonstrating large left staghorn renal calculus. B: Nuclear scintigraphic evaluation of renal calculi. Posterior view demonstrating uptake on large left staghorn calculus after furosemide (Lasix) diuresis. Note right kidney with uptake in lower pole. C: Follow-up tomogram confirms calculus (arrow) in right lower pole missed on initial radiograph.
Intrarenal alkalinization may be performed successfully under a low-pressure system (<25 cm water pressure). This may be achieved through a percutaneous nephrostomy tube or an externalized retrograde catheter. A manometer, similar to those used for central venous pressure monitoring, is cheap, available, and practical. Agents include sodium bicarbonate, 2–4 ampules in 1 L of normal saline, producing a urinary pH between 7.5 and 9. Tromethamine-E and tromethamine can produce urinary pHs of 8–10.5 and are especially effective with pH-sensitive calculi as in uric acid and cystine lithiasis.

Cystine calculi can be dissolved with a variety of thiols, including D-penicillamine (0.5% solution), N-acetylcycteine (2–5% solution), and alpha-mercaptopyropionylglycine (Thiola) (5% solution).

Struvite stone dissolution requires acidification and may be achieved successfully with Suby’s G solution and hemiacidrin (Renacidin). Urinary pH may get down to 4. Hemiacidrin must be used with sterile urine and careful monitoring of serum magnesium levels is required. The Food and Drug Administration has not approved hemiacidrin for upper-tract irrigations, and thus appropriate informed consent is required.

C. RELIEF OF OBSTRUCTION

Urinary stone disease may result in significant morbidity and possible mortality in the presence of obstruction, especially with concurrent infection. A patient with obstructive urinary calculi with fever and infected urine requires emergent drainage. Retrograde pyelography to define upper-tract anatomy is logically followed by retrograde placement of a double-J ureteral stent. On occasion such catheters are unable to bypass the offending calculus or may perforate the ureter. In such situations one must be prepared to place a percutaneous nephrostomy tube.

D. EXTRACORPOREAL SHOCK WAVE LITHOTRIPSY

Extracorporeal shock wave lithotripsy has revolutionized the treatment of urinary stones. The concept of using shock waves to fragment stones was noted in the 1950s in Russia. However, it was during the investigation of pitting on supersonic aircraft that Dornier, a German aircraft corporation, rediscovered that shock waves originating from passing debris in the atmosphere can crack something that is hard. It was the ingenious application of a model developed in hopes of understanding such shock waves that ESWL emerged. The first clinical application with successful fragmentation of renal calculi was in 1980. The HM–1 (Human Model–1) lithotriptor underwent modifications in 1982 leading to the HM–2 and, finally, to the widespread application of the HM–3 in 1983 (Figure 16–16). Since then, thousands of lithotriptors have been put into use around the world, with millions of patients successfully treated.

All require an energy source to create the shock wave, a coupling mechanism to transfer the energy from outside to inside the body, and either fluoroscopic or ultrasonic modes, or both, to identify and position the calculi at a focus of converging shock waves. They differ in generated pain and anesthetic or anesthesiologist requirements, consumable components, size, mobility, cost, and durability. Focal peak pressures (400–1500 bar), focal dimensions (6 × 28 mm to 50 × 15 mm), modular design, utilization to help increase mobility of frozen joints, varied distances (12–17 cm) between focus 1 (the shock wave source) and focus 2 (the target), and purchase price differentiate the various machines available today.

1. Shock wave physics—In contrast to the familiar ultrasonic wave with sinusoidal characteristics and longitudinal mechanical properties, acoustic shock waves are unharmonic and have nonlinear pressure characteristics. There is a steep rise in pressure amplitude that results in compressive forces (Figure 16–17). There are 2 basic types of shock wave sources: supersonic and finite amplitude emitters.

Supersonic emitters release energy in a confined space, thereby producing an expanding plasma and an acoustic shock wave. Such shock waves occur in nature—the familiar thunderstorm with lightning (an electrical discharge) followed by thunder (an acoustic sonic boom) is an analogous situation. Under controlled conditions, such an acoustic shock wave can successfully fragment calculi. The initial compression wave travels faster than the speed of sound in water and rapidly slows down to that speed. The traveling pressure wave is reduced in a nonlinear fashion. Medical applications have focused such waves to concentrate energy on a calculus (Figure 16–18).

Finite amplitude emitters, in contrast to point source energy systems, create pulsed acoustic shock waves by displacing a surface activated by electrical discharge. There are 2 major types of finite amplitude emitters: piezoceramic and electromagnetic. The piezoceramic variety results in a shock wave after an electrical discharge causes the ceramic component to elongate in such a manner that the surface is displaced and an acoustic pulse is generated. Thousands of such components placed on the concave side of a spheric surface directed toward a focus result in high stress, strain, and cavitation pressures (Figure 16–19). Electromagnetic systems are similar in concept to a stereo speaker system. An electrical discharge to a slab, adjacent to an insulating foil, creates an electric current that repulses a metal membrane, displacing it and generating an acoustic pulse into an adjacent medium. These waves need to be focused toward the offending stone.

All shock waves, despite their source, are capable of fragmenting stones when focused. Fragmentation is achieved by erosion and shattering (Figure 16–20). Cavitational forces result in erosion at the entry and exit sites of the shock wave. Shattering results from energy absorption with
stress, strain, and shear forces. Surrounding biologic tissues are resilient because they are not brittle nor are the shock waves focused on them.

2. Preoperative evaluation—Physical examination should be as thorough as in preparation for any other surgical procedure. Vital signs including blood pressure should be noted. Body habitus including any gross skeletal abnormalities, contractures, or excessive weight (>300 lb) may severely limit or preclude ESWL. Borderline individuals require simulation before treatment. Pregnant women and patients with large abdominal aortic aneurysms or uncorrectable bleeding disorders should not be treated with ESWL. Individuals with cardiac pacemakers should be thoroughly evaluated by a cardiologist. If ESWL is contemplated, a cardiologist with thorough knowledge and with the ability to override the pacemaker should be present in the lithotripsy suite.

Figure 16–16. Diagrammatic representation of a Dornier HM–3 lithotriptor.

Figure 16–17. Shock wave. Vertical axis represents pressure and horizontal axis represents time.
Figure 16–18. A: Supersonic shock wave emission from a spark gap electrode. B: Reflecting the shock wave from focus 1 to focus 2 allows for stone fragmentation.

Figure 16–19. Piezoceramic finite amplitude emitter. Ceramic components are placed on the concave surface of a sphere and each component is directed to an identified focus.
3. Intraoperative considerations—

a. Stone localization—Proper patient positioning is a prerequisite for successful lithotripsy. Palpating the patient’s ribs and pelvic bony girdle can approximate appropriate positioning. Anterior located kidneys, medial oriented portions of a horseshoe kidney, or transplant kidneys are best treated in the prone position. Understanding positioning options with the various lithotriptors available today is required to optimize therapy.

Small or poorly calcified calculi can be difficult to image with fluoroscopy, irrespective of their location. Placing a ureteral catheter identifies known anatomy and supplies an injection port for radiocontrast agents. A poorly calcified caliceal calculus can be identified by injecting dilute contrast agents into the collecting system and then focusing on the appropriate calyx or filling defect. In patients who cannot have retrograde stents placed, intravenous contrast agents may be used to help localize and thus focus on such stones.

b. Fluoroscopic imaging—The conditions for fluoroscopic imaging include appropriate collimation, dimmed room lighting, and adequate bowel preparation to decrease bothersome bowel gas and thus decrease radiation exposure and improve the quality of the fluoroscopic image. Intermittent fluoroscopy reveals movement of calculi with respiration and is helpful in locating and focusing on offending calculi.

c. Ultrasonic imaging—Ultrasound localization has the advantage of eliminating radiation exposure to the patient or the lithotripsy team. There are 2 basic types: the coaxial unit, aligned with the shock wave generator, and the articulating arm unit with a mobile transducer. Ultrasound can easily identify radiolucent or small calculi that are difficult to visualize with fluoroscopy. However, ureteral or other medially located calculi may be difficult or impossible to identify, especially in nonobstructive collecting systems. Visualization may be difficult or impossible with obese patients. Proficiency in ultrasound localization and assessment of fragmentation has a longer learning curve than that of fluoroscopy. Ultrasound images may be confusing when multiple stones or stone fragments are present.

d. Coupling—Successful fragmentation requires effective coupling. Coupling devices should have properties similar to those of human skin. Optimal systems should prevent pain, ecchymoses, hematomas, or skin breakdown. Interfaces between gas and tissue can result in tissue breakdown. Air bubbles entrapped by hair, by bandages from prior percutaneous procedures, or by inadequately degassed fluid, or air in coupling cushions can significantly impede directed shock waves and result in skin ecchymoses or breakdown. Despite adequate coupling, fragmentation may be inadequate owing to refraction and reflection of shock waves at tissue interfaces, especially with obese patients.

Water bath provides good coupling. Submersion of patients can result in profound hemodynamic changes including peripheral venous compression resulting in increased right atrial pressure, increased pulmonary capillary wedge pressure, and increased cardiac index. These hemodynamic changes should be appreciated and appropriate monitoring should be used in individuals with marginal cardiovascular reserve.

In contrast, water cushion coupling systems have decreased water demands. A coupling gel like those used with ultrasonography provides an excellent interface with skin. The volume of such water cushions can frequently be adjusted to help focus calculi when patients are either extremely thin (eg, children) or obese. Very small patients may require an interposed bag (1–3 L) of normal saline to help with coupling. Both water coupling systems require degassed water to decrease water bubbles.
e. Shock wave triggering—Triggering shock waves with the electrocardiogram was originally performed to decrease cardiac dysrhythmias. The lithotripter would sense the large swing of the QRS complex and initiate the shock wave 20 ms later; this would decrease shock waves during the repolarization phase of the cardiac cycle (myocardium is most sensitive during this time). If cardiac dysrhythmias occur, interruption of the procedure frequently stops them. However, if they continue, standard medical therapy is effective. Conceptually, it makes more sense to trigger the shock waves in response to the respiratory cycle to optimize accurate focusing on the offending calculi that move with respiratory motion. Such systems are available. Many lithotriptors are now triggered without electrocardiogram gating and with unusual associated cardiac dysrhythmias. This can speed up therapy, especially in those with slow heart rates that are not amenable to pharmacologic manipulation.

f. Fragmentation—Safe shock wave dosage is unknown. Shock waves induce trauma, including intrarenal and perirenal hemorrhage and edema, and thus the minimal shocks needed to achieve fragmentation should be given. Casual overtreatment should be avoided since long-term complications are not yet known.

Determination of adequate fragmentation during treatment may be difficult. Initial sharp edges become fuzzy or blurred and have a shotgun-blast-like appearance. Stones that were initially visualized may disappear after successful fragmentation. Intermittent visualization ensures accurate focusing and assessment of progress and eventual termination of the procedure.

Bilateral nephrolithiasis may be treated in the same setting. One must first approach the side that is symptomatic or potentially more troublesome. If there is uncertainty concerning a large stone burden, one or more double-J catheters should be placed to decrease the likelihood of bilateral obstruction.

4. Postoperative care—Patients should be encouraged to maintain an active ambulatory status to facilitate stone passage. Gross hematuria should resolve during the first postoperative week. Fluid intake should be encouraged. Follow-up in approximately 2 weeks for discussion and evaluation of a KUB and renal ultrasonography will help assess success of fragmentation and passage of gravel. Patients may return to work as soon as they feel comfortable in doing so.

Abdominal pain may be related to the shock waves. Severe pain unresponsive to routine intravenous or oral medications should alert the physician for possible rare (0.66%) perirenal hematomas. In such a situation, CT should then be undertaken to stage the injury. The potential association of ESWL with the development of hypertension has not been substantiated. Long-term data are still being collected.

Stone burden correlates with postoperative complications. Steinstrasse (stone street) or columnation of stone gravel in a ureter can be frustrating. It should be specifically ruled out when postoperative radiographs are evaluated. Asymptomatic individuals can be followed up with serial KUBs and ultrasonography. Severe pain or fever requires intervention. Percutaneous nephrostomy drainage is usually uncomplicated owing to the associated hydronephrosis. Decompressing the collecting system allows for effective coaptation of the ureteral walls and encourages resolution of the problem. It is only in the rare patient that steinstrasse does not resolve with the procedures outlined; such cases require retrograde endoscopic manipulations to relieve the obstructed stone fragments. Usually one finds 1 or 2 relatively large fragments that are obstructing. With their removal the columnation of fragments resolves.

Patients with large renal pelvic calculi (>1.5 cm) have a stone-free rate at 3 months approximating 75%, in comparison with those with a similar stone in a lower calyx, which approximates only 50%. Patients with small renal pelvic stones (<1.5 cm) have approximately a 90% stone-free rate in comparison to those with similar stones in a middle calyx (approximately 75%) or lower calyx (approximately 70%). Lower calyceal stone-free rates are increased with a small stone burden, a short and wide infundibulum, and a nonacute infundibulo-pelvic angle. Overall, approximately 75% of patients with renal calculi treated with ESWL become stone-free in 3 months. As stones increase in size, stone-free rates decrease, more so in the lower and middle calyces than in superior calyceal and renal pelvic locations.

E. URETEROSCOPIC STONE EXTRACTION

Ureteroscopic stone extraction is highly efficacious for lower ureteral calculi. The use of small-caliber ureteroscopes and the advent of balloon dilation or ureteral access sheaths have increased stone-free rates dramatically. Even relatively large-caliber endoscopes without balloon dilation are effective in lower ureteral stone retrieval. Stone-free rates range from 66% to 100% and are dependent on stone burden and location, length of time the stone has been impacted, history of retroperitoneal surgery, and the experience of the operator. Complication rates range from 5% to 30%; the rates increase when manipulations venture into the proximal ureter. Ureteral stricture rates are <5%. Postoperative vesicoureteral reflux is extremely rare. Calculi that measure <8 mm are frequently removed intact. Round wire stone baskets can be torqued to help entrap stone or stone fragments. Flat wire baskets should be used with caution; if twisted, they can develop sharp, knifelike edges resulting in ureteral injury. Excessive force with any instrument in the ureter may result in ureteral injury.

A variety of lithotrites can be placed through an ureteroscope, including electrohydraulic, solid and hollow-core
ultrasonic probes, a variety of laser systems, and pneumatic systems such as the Swiss lithoclast. Electrohydraulic lithotrites have power settings as high as 120 V that result in a cavitation bubble, followed by collapse of this bubble causing subsequent shock waves. Care should be taken to keep the tip of the electrode away from surrounding tissue and the tip of the endoscope. Ultrasonic lithotrites have a piezoceramic energy source that converts electrical energy into ultrasonic waves in the range of 25,000 Hz. This vibratory action is effective in fragmenting calculi. Hollow probes can suction stone fragments and debris simultaneously. Laser systems are discussed elsewhere in this book.

**F. PERCUTANEOUS NEPHROLITHOTOMY**

Percutaneous removal of renal and proximal ureteral calculi is the treatment of choice for large (>2.5 cm) calculi; those resistant to ESWL; select lower pole calyceal stones with a narrow, long infundibulum and an acute infundibulo-pelvic angle; and instances with evidence of obstruction; the method can rapidly establish a stone-free status. Needle puncture is directed by fluoroscopy, ultrasound, or both, and is routinely placed from the posterior axillary line into a posterior inferior calyx. Superior caliceal puncture may be required, and in such situations care should be taken to avoid injury to the pleura, lungs, spleen, and liver. Tract dilation is performed by sequential plastic dilators (Amplatz system), telescoping metal dilators (Alken), or balloon dilation with a backloaded Amplatz sheath. Tracts placed during open renal procedures are frequently tortuous and suboptimal for subsequent endourologic procedures.

Percutaneous extraction of calculi requires patience and perseverance. Hardcopy radiographs help to confirm a stone-free status. Remaining calculi can be retrieved with the aid of flexible endoscopes, additional percutaneous puncture access, follow-up irrigations, ESWL, or additional percutaneous sessions. Realistic goals should be established. Patients should be informed that complex calculi frequently require numerous procedures.

Maintenance of body temperature with appropriate blankets during preoperative patient positioning and with warmed irrigation fluids helps to prevent bleeding diatheses associated with hypothermia. The average blood loss during a percutaneous nephrolithotomy is 2–2.8 g/dL of hemoglobin. Multiple percutaneous punctures and renal pelvic perforations are associated with a greater blood loss. Overall, such procedures are safe and effective and have a transfusion rate well <10%.

**G. OPEN STONE SURGERY**

Open stone surgery is the classic way to remove calculi. The morbidity of the incision, the possibility of retained stone fragments, and the ease and success of less invasive techniques have made these procedures relatively uncommon when instruments and surgical experience are available. It is mandatory to obtain a radiograph before the incision is made; calculi frequently move. A variety of incisions to access the kidney are available.

**H. PYELOLITHOTOMY**

Pyelolithotomy is effective, especially with an extrarenal pelvis. A transverse pyelotomy is effective and does not require interruption of the renal arterial blood supply. Inspection with flexible endoscopes helps ensure a stone-free status. Multiple, small renal pelvic calculi and difficult-to-access calceal calculi can be retrieved with the aid of a coagulum. Coagulum was initially produced from pooled human fibrinogen. The risks of hepatitis and other viral infections have made this method unacceptable. Cryoprecipitate can be obtained from rapid freezing of plasma. Autologous plasma may be used to decrease the incidence of bloodborne infections. The tensile strength of cryoprecipitate is approximately 10 times that of a blood clot. Injected into the renal pelvis, endogenous clotting factors result in a Jelly-like coagulum of the collecting system. Small stones are entrapped and removed with the coagulum. A variety of Randall stone forceps help gain access into most of the collecting system.

**I. ANATROPHIC NEPHROLITHOTOMY**

Anatrophic nephrolithotomy is used with complex staghorn calculi. A complete staghorn calculus is a cast of the renal pelvis and calyces (Figure 16–21). A partial staghorn
calculus involves the renal pelvis and extends into at least 2 infundibula. To gain access to the entire collecting system, a longitudinal incision is made on the convex surface of the kidney just posterior to the line of Brödel, taking advantage of the converging anterior and posterior renal blood supplies. Occlusion of the renal artery followed by renal cooling with slushed ice gives a relatively bloodless surgical field. A nerve hook is helpful to tease out calculi. Careful inspection of the entire collecting system helps remove all stones. Repair of narrowed infundibula helps reduce stone recurrence rates. The collecting system is closed followed by the renal capsule. Intraoperative placement of a nephrostomy tube for possible follow-up irrigations or endoscopic inspection or stone retrieval makes hemostasis difficult. Open stone surgery becomes progressively more difficult after the first procedure owing to reactive scar tissue.

J. RADIAL NEPHROTOMY

Radial nephrotomy gives access to limited calyces of the collecting system. An appropriate approach to localized calculi, it is frequently used in blown-out calyces with thin overlying parenchyma. Intraoperative ultrasound helps to localize the calyx and the calculi. Once the kidney has been opened, the introduction of air can make interpretation of subsequent ultrasound scans confusing. A shallow incision of the renal capsule can be followed by puncture into the collecting system. Brain retractors provide excellent exposure. Care should be taken not to force stones through narrow infundibula. Stones may be cut with heavy Mayo scissors, and remaining fragments can be retrieved. Inspection with flexible endoscopes is helpful. Intraoperative radiographs help document a stone-free status.

K. OTHER RENAL PROCEDURES

Partial nephrectomy is appropriate with a large stone burden in a renal pole with marked parenchymal thinning. Caution should be taken with a simple nephrectomy even with a normal contralateral kidney, as stones are frequently associated with a systemic metabolic defect that may recur in the contralateral kidney. What may seem prudent and simple today may be regretted tomorrow.

Other unusual procedures include ileal ureter substitution performed with the hope of decreasing pain with frequent stone passage. Autotransplantation with pyelocystostomy is another option for patients with rare malignant stone disease.

L. URETEROLITHOTOMY

Long-standing ureteral calculi—those inaccessible with endoscopy and those resistant to ESWL—can be extracted with ureterolithotomy. Again, a preoperative radiograph documents stone location and directs an appropriate incision. The proximal ureter may be approached with a dorsal lumbotomy. An incision lateral to the sacrospinalis muscles allows medial retraction of the quadratus lumborum. The anterior fascicle of the dorsal lumbar fascia must be incised to gain proper exposure despite the appearance of potentially opening the peritoneum. Once the ureter is identified, a vessel loop or a Babcock clamp should be placed proximal to the stone to prevent frustrating stone migration. Extension of this incision is limited superiorly by the 12th rib and inferiorly by the iliac crest. A longitudinal incision over the stone with a hooked blade exposes the calculus. The nerve hook is excellent to help tease out the stone. A flank or anterior abdominal muscle splitting incision gives excellent exposure to mid- and distal ureteral stones.

Prevention

In general, 50% of patients experience recurrent urinary stones within 5 years without prophylactic intervention. Appropriate education and preventive measures are best instituted with a motivated patient after spontaneous stone passage or surgical stone removal. Risk factors as described previously should be identified and modified, if possible. Irrespective of the final metabolic evaluation and stone analysis, the patient’s fluid intake should be about 1.6 L/24 h. Fluids should be encouraged during mealtime. Additionally, liquids should be increased approximately 2 hours after meals. Water produced as a metabolic by-product reaches its nadir at this time, and thus the body is relatively dehydrated. Fluid ingestion also should be encouraged to force a nighttime diuresis adequate to awaken the patient to void. Awakening and ambulating to void limit urinary stasis and offer an opportunity to ingest additional fluids. These lifestyle changes are difficult to maintain and should be encouraged during subsequent office visits. Motivated patients who regularly return to a urinary stone clinic have a reduced stone recurrence rate that is probably due to increased compliance.

A. METABOLIC EVALUATION

A systematic metabolic evaluation should be instituted after a patient has recovered from urinary stone intervention or spontaneous stone passage. Stone analysis should be obtained to help direct the workup. An outpatient urine collection during typical activities and fluid intake helps unmask significant abnormalities. An initial 24-hour urine collection for calcium stone formers should include tests for calcium, uric acid, oxalate, citrate, sodium, volume, and pH. An open dialog with local laboratories helps to standardize collection routines and determine whether an outside laboratory is preferred. Baseline serum levels for blood urea nitrogen, creatinine, calcium, phosphorous, and uric acid are appropriate.

Hypercalciuria is the most common abnormality. To differentiate among hypercalciuria types I, II, and III, a patient should be placed on a sodium- and calcium-restricted diet for a few days to a week. This is easily
achieved (100 mEq/day) by eliminating table salt and reducing obviously salty foods. Calcium is restricted (400–500 mg) by excluding dairy products. A repeat 24-hour urine collection is evaluated for calcium. A urinary calcium level <250 mg/day confirms a diagnosis of dietary-dependent hypercalciuria type II. Type I and type III hypercalciuria must be differentiated in patients with urinary calcium levels >250 mg/day. A calcium binder such as cellulose phosphate is prescribed (5 g three times daily with meals) for a few days. This is followed by a repeat 24-hour urine calcium level and parathyroid hormone blood value. Patients who have type I absorptive hypercalciuria have at least a 50% drop in urinary calcium levels and normal parathyroid hormone levels.

Hyperuricosuria, hyperoxaluria, and hypocitraturia calcium stone formers can be treated appropriately and followed with repeat 24-hour urine collections. Many calcium stone formers have multiple defects; although one treatment may reverse one defect, it may exacerbate others. Subsequent 24-hour urine collections are critical for effective long-term follow-up and stone prevention. Treatment of cystinuria should be titrated with repeat 24-hour cystine levels. Repeat urine cultures should be obtained in patients with infectious calculi.

B. ORAL MEDICATIONS

1. Alkalinizing pH agents—Potassium citrate is an oral agent that elevates urinary pH effectively by 0.7–0.8 pH units. Typical dosing is 60 mEq in 3 or 4 divided doses daily. It is available in wax-matrix 10-mEq tablets, liquid preparations, and crystals that must be mixed with fluids. The effect is maintained over many years. Care should be taken in patients susceptible to hyperkalemia, those with renal failure, and those taking potassium-sparing diuretics. Although the medication is usually well tolerated, some patients may complain of abdominal discomfort, especially with tablet preparations. It is indicated in those with calcium oxalate calculi secondary to hypocitruria (<320 mg/day), including those with renal tubular acidosis. Potassium citrate also may be used effectively to treat uric acid lithiasis and nonsevere forms of hyperuricosuric calcium nephrolithiasis.

2. Gastrointestinal absorption inhibitor—Cellulose phosphate binds calcium in the gut and thereby inhibits calcium absorption and urinary excretion. It is a popular drug in the treatment of absorptive hypercalciuria type I with recurrent calcium nephrolithiasis, although it only prevents new stone formation. Patients should have normal parathyroid hormone values, normal serum calcium and phosphate values, no evidence of bone disease, and evidence of increased intestinal calcium absorption. The drug decreases the urinary saturation of calcium phosphate and calcium oxide. It may increase urinary oxalate and urinary phosphate levels. A typical starting dosage is 5 g three times daily with meals; the dosage may be titrated by following 24-hour urinary calcium levels. Urinary magnesium, calcium, oxalate, and sodium levels and serum parathyroid hormone should be monitored one to two times yearly. Magnesium supplements are frequently required and should be taken at least 1 hour before or after cellulose phosphate is taken. Cellulose phosphate is associated with a sodium load and should be used with caution in those with congestive heart failure. Gastrointestinal side effects are infrequent; they include dyspepsia and loose bowel movements.

Cellulose phosphate may be suboptimal treatment for postmenopausal women who are at risk for bone disease. An alternative treatment for such patients would be hydrochlorothiazides supplemented with potassium citrate to offset the potential hypokalemia and hypocitraturia.

3. Phosphate supplementation—Renal phosphate leak is best treated by replacing phosphate. Phosphate absorption may be inhibited in the presence of aluminum-, magnesium-, or calcium-containing antacids. This treatment should be used with caution in digitalized patients and in those with severe renal failure, Addison disease, or severe hepatic dysfunction. It is generally well tolerated. Dosing can begin with 250 mg three to four times daily and may be doubled depending on follow-up serum electrolyte, calcium, and phosphorus levels.

4. Diuretics—Thiazides can correct the renal calcium leak associated with renal hypercalciuria. This prevents a secondary hyperparathyroid state and its associated elevated vitamin D synthesis and intestinal calcium absorption. A rapid decrease in urinary calcium excretion is appreciated and is sustained long-term (>10 years). A starting dose of 25 mg may be titrated based on urinary calcium levels. Side effects are usually well tolerated. Potassium levels should be monitored. Hypokalemia induces a hypocitraturic state; potassium replacement corrects the hypokalemia and its associated hypocitraturia.

Thiazides result in a transient decrease in urinary calcium excretion in absorptive hypercalciurics. Urinary calcium excretion rebounds to pretreatment values in 50% of such patients after 4–5 years of therapy. Dietary changes are not believed to be responsible for this phenomenon. Thiazides do not restore normal intestinal absorption of calcium.

5. Calcium supplementation—Enteric hyperoxaluric calcium nephrolithiasis is effectively treated with calcium supplements. Calcium gluconate and calcium citrate are better absorbed and are more effective in increasing serum calcium availability than are other forms of calcium. Calcium carbonate, calcium phosphate, and oyster shell are forms of calcium that are less efficiently absorbed; they remain in the intestinal lumen, available to bind oxalate,
thus reducing its absorption. These less efficiently absorbed forms of calcium are optimal to treat enteric hyperoxaluric calcium nephrolithiasis and must be given with meals to be effective.

6. Uric acid-lowering medications—Allopurinol is used to treat hyperuricosuric calcium nephrolithiasis with or without hyperuricemia. Unlike uricosuric agents that reduce serum uric acid levels by increasing urinary uric acid excretion, allopurinol is a xanthine-oxidase inhibitor and reduces both serum and urinary levels of uric acid. It has no impact on the biosynthesis of purines; rather, it acts exclusively on purine catabolism. Elevated levels of xanthine and hypoxanthine in the urine secondary to allopurinol have not been associated with nephrolithiasis. Allopurinol is a potentially dangerous drug and should be discontinued at the first appearance of a skin rash, which infrequently may be fatal. Therapy can start at 300 mg/day. It is tolerated best when taken after meals.

7. Urease inhibitor—Acetohydroxamic acid is an effective adjunctive treatment in those with chronic urea-splitting urinary tract infections associated with struvite stones. Acetohydroxamic acid reversibly inhibits bacterial urease, decreasing urinary ammonia levels, and will subsequently acidify urine. It is best used as prophylaxis after removal of struvite stones. It also may be used after unsuccessful attempts at curative surgical removal of calculi or culture-specific antibiotic therapy. Patients with serum creatinine >2.5 mg/dL are unable to achieve therapeutic urinary levels. Acetohydroxamic acid is not effective with non-urease-producing bacteria. Long-term data (>7 years) are unavailable. A significant number of patients complain of side effects, including headaches that are usually short-lived and responsive to aspirin compounds. Other frequent complaints include nausea, vomiting, anorexia, nervousness, and depression. A typical dosing regimen is 1 tablet three or four times daily (total dosage: 10–15 mg/kg/day).

8. Prevention of cystine calculi—Conservative measures, including massive fluid intake and urinary alkalization, are frequently inadequate to control cystine stone formation. Penicillamine, the same drug that is used to chelate excess copper in the treatment of Wilson disease, undergoes a thiol-disulfide exchange with cystine. This reduces the amount of urinary cystine that is relatively insoluble. Cystine solubility is pH-dependent (pH 5: 150–300 mg/L; pH 7: 200–400 mg/L; pH 7.5: 220–500 mg/L). D-Penicillamine is associated with numerous and frequent side effects, including rashes and hematologic, renal, and hepatic abnormalities. An initial dosage of 250 mg daily in 3–4 divided doses may help reduce severe side effects. It may be increased gradually to 2 g/day. Dosage should be titrated with quantitative urinary cystine values. Penicillamine increases the requirement of pyridoxine (vitamin B₆), which should be supplemented with 25–50 mg/day.

Mercaptopropionylglycine (Thiola) is better tolerated by patients than is penicillamine. Mercaptopropionylglycine, a reducing agent, binds to the sulfide portion of cystine, forming a mixed disulfide (Thiola-cysteine) water-soluble compound. It may retard the rate of new stone formation. The dosage should be titrated with repeat 24-hour urinary cystine values. An initial dosage may be 200–300 mg three times daily, either 1 hour before or 2 hours after each meal. Side effects are not infrequent and may include drug fever; nausea, vomiting, and gastrointestinal upset; rash, wrinkling, or friable skin; lupuslike symptoms, decreased taste perception; and a variety of hematologic disorders.

BLADDER STONES

Bladder calculi usually are a manifestation of an underlying pathologic condition, including voiding dysfunction or a foreign body. Voiding dysfunction may be due to a urethral stricture, benign prostatic hyperplasia, bladder neck contracture, or flaccid or spastic neurogenic bladder, all of which result in static urine. Foreign bodies such as Foley catheters and forgotten double-J ureteral catheters can serve as nidi for stones (Figure 16–22). Most bladder calculi are seen in men. In developing countries, they are frequently found in prepubescent boys. Stone analysis frequently reveals ammonium urate, uric acid, or calcium oxalate stones. A solitary bladder stone is the rule, but there are numerous stones in 25% of patients (Figure 16–23). Patients present with irritative voiding symptoms, intermittent urinary stream, urinary tract infections, hematuria, or pelvic pain. Physical examination is unrevealing. A large percentage of bladder stones are radiolucent (uric acid). Ultrasound of the bladder identifies the stone with its characteristic shadowing. The stone moves with changing body position. Stones within a ureterocele do not move with body position (Figure 16–24) as seen on ultrasound examination. They frequently are nonobstructive. Endoscopic incision and stone removal rarely result in vesicoureteral reflux. The mode of stone removal for other bladder stones should be directed by the underlying cause.

Early instruments used to remove bladder calculi were both clever and bizarre. Simple mechanical crushing devices are still used today. Mechanical lithotrites should be used with caution to prevent bladder injury when the jaws are closed. Ensuring partially full bladder and endoscopic visualization of unrestricted lateral movement before forceful crushing of the stones helps reduce this troublesome complication. Cystolitholapaxy allows most stones to be broken and subsequently removed through a cystoscope. Electrohydraulic, ultrasonic, laser, and pneumatic lithotrites similar to those used through a nephroscope are effective. Cystolithotomy can be performed through a small abdominal incision.
PROSTATIC & SEMINAL VESICLE STONES

Prostatic calculi are found within the prostate gland per se and are found uncommonly within the prostatic urethra. They are thought to represent calcified corpora amylacea and are rarely found in boys. Usually small and numerous, they are noted to be tannish gray in color during transurethral resection of the prostate. They are commonly located at the margin of the surgically resected adenoma and are composed of calcium phosphate. Although usually of no clinical significance, rarely they are associated with chronic prostatitis. Large prostatic calculi may be misinterpreted as a carcinoma. The prostate is usually mobile,
however, and a radiograph or transrectal ultrasound helps to confirm the diagnosis.

Seminal vesicle stones are smooth and hard and are extremely rare. They may be associated with hematospermia. Physical examination reveals a stony hard gland, and when multiple stones are present, a crunching sensation may be noted. These stones occasionally are confused with tuberculosis of the seminal vesicle.

URETHRAL & PREPUCIAL STONES

Urethral calculi usually originate from the bladder and rarely from the upper tracts. Most ureteral stones that pass spontaneously into the bladder can pass through the urethra unimpeded. Urethral stones may develop secondary to urinary stasis, secondary to a urethral diverticulum, near urethral strictures, or at sites of previous surgery. Most urethral stones in men present in the prostatic or bulbar regions and are solitary. Patients with recurrent pendulous urethral calculi without evidence of other pathologic conditions should be suspected of self-introduction of such stones in an attempt to obtain pain medications or for attention, as seen in Munchausen syndrome.

Females rarely develop urethral calculi owing to their short urethra and a lower incidence of bladder calculi. Most urethral stones found in women are associated with urethral diverticula.

Symptoms are similar to bladder calculi: intermittent urinary stream, terminal hematuria, and infection. The stones may present with dribbling or during acute urinary retention. Pain may be severe and, in men, may radiate to the tip of the penis. The diagnosis may be confirmed by palpation, endoscopic visualization, or radiographic study.

Treatment should be directed by the underlying cause. Stones associated with a dense urethral stricture or complex diverticula can be removed during definitive open surgical repair. Small stones may be grasped successfully and removed intact. More frequently they need to be fragmented and removed. Long-standing, large impacted stones are best removed through a urethrotomy.

Preputial calculi are rare and usually occur in adults. They develop secondary to a severe obstructive phimosis.

Figure 16–24. A: Scout abdominal radiograph demonstrating extraosseous calcification in the region of the bladder. B: Intravenous pyelogram demonstrates stone to be within a ureterocele.
They may occur secondary to poor hygiene with inspissated smegma. Diagnosis is confirmed by palpation. Treating the underlying cause with a dorsal preputial slit or a formal circumcision prevents recurrent calculi.

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**Phosphate**


**Uric Acid**


**Cystine**


**Xanthine**


**Triamterene**

CHAPTER 16

Silicate


Matrix Urinary Calculi


Urinary Stone Inhibitors

Citrate


Orthophosphates & Pyrophosphates


Urinary Proteins


Trace Elements


2, 8-Dihydroxyadenine Urolithiasis


Renal Tubular Acidoses


Water


Urinary Stone Disease in Uncommon Situations

Spinal Cord Dysfunction


Pregnancy


Renal Transplantation


Obesity


Anatomic Renal Anomalies


**Pediatrics**


**Caliceal Diverticulae**


**Medical Therapy**


**Surgical Therapy**


**Extracorporeal Shock Wave Lithotripsy**


**Percutaneous Nephrolithotomy**


**Ureteroscopy**


Injuries to the Genitourinary Tract

Jack W. McAninch, MD, FACS

EMERGENCY DIAGNOSIS & MANAGEMENT

About 10% of all injuries seen in the emergency room involve the genitourinary system to some extent. Many of them are subtle and difficult to define and require great diagnostic expertise. Early diagnosis is essential to prevent serious complications.

Initial assessment should include control of hemorrhage and shock along with resuscitation as required. Resuscitation may require intravenous lines and a urethral catheter in seriously injured patients. In men, before the catheter is inserted, the urethral meatus should be examined carefully for the presence of blood.

The history should include a detailed description of the accident. In cases involving gunshot wounds, the type and caliber of the weapon should be determined, since high-velocity projectiles cause much more extensive damage.

The abdomen and genitalia should be examined for evidence of contusions or subcutaneous hematomas, which might indicate deeper injuries to the retroperitoneum and pelvic structures. Fractures of the lower ribs are often associated with renal injuries, and pelvic fractures often accompany bladder and urethral injuries. Diffuse abdominal tenderness is consistent with perforated bowel, free intraperitoneal blood or urine, or retroperitoneal hematoma. Patients who do not have life-threatening injuries and whose blood pressure is stable can undergo more deliberate radiographic studies. This provides more definitive staging of the injury.

Special Examinations (Figures 17–1 through 17–3)

When genitourinary tract injury is suspected on the basis of the history and physical examination, additional studies are required to establish its extent.

A. CATHETERIZATION AND ASSESSMENT OF INJURY

Assessment of the injury should be done in an orderly fashion so that accurate and complete information is obtained. This process of defining the extent of injury is termed staging. The algorithms (Figures 17–1 through 17–3) outline the staging process for urogenital trauma.

1. Catheterization—Blood at the urethral meatus in men indicates urethral injury; catheterization should not be attempted if blood is present, but retrograde urethrography should be done immediately. If no blood is present at the meatus, a urethral catheter can be carefully passed to the bladder to recover urine; microscopic or gross hematuria indicates urinary system injury. If catheterization is traumatic despite the greatest care, the significance of hematuria cannot be determined, and other studies must be done to investigate the possibility of urinary system injury.

2. Computed tomography—Abdominal computed tomography (CT) with contrast media is the best imaging study to detect and stage renal and retroperitoneal injuries. It can define the size and extent of the retroperitoneal hematoma, renal lacerations, urinary extravasation, and renal arterial and venous injuries; additionally, it can detect intra-abdominal injuries (liver, spleen, pancreas, bowel). Spiral CT scanning, now common, is very rapid, but it may not detect urinary extravasation or ureteral and renal pelvic injuries. We recommend repeat scanning 10 minutes after the initial study to aid the diagnosis of these conditions.

3. Retrograde cystography—Filling of the bladder with contrast material is essential to establish whether bladder perforations exist. At least 300 mL of contrast medium should be instilled for full vesical distention. A film should be obtained with the bladder filled and a second one after the bladder has emptied itself by gravity drainage. These 2 films establish the degree of bladder injury as well as the size of the surrounding pelvic hematomas.

Cystography with CT is excellent for establishing bladder injury. At the time of scanning, this likewise must be done with retrograde filling of the bladder with 300 mL of contrast media to ensure adequate distention to detect injury.

4. Urethrography—A small (12F) catheter can be inserted into the urethral meatus and 3 mL of water placed in the balloon to hold the catheter in position. After retrograde injection of 20 mL of water-soluble contrast material, the urethra will be clearly outlined on film, and extravasation in the deep bulb area in case of straddle injury—or free extravasation into the retropubic space in case of prostatomembranous disruption—will be visualized.
Figure 17–1. Algorithm for staging blunt trauma in adults.
Figure 17–2. Algorithm for staging penetrating trauma in adults.

Figure 17–3. Algorithm for staging blunt trauma in children.
INJURIES TO THE GENITOURINARY TRACT

5. Arteriography—Arteriography may help define renal parenchymal and renal vascular injuries. It is also useful in the detection of persistent bleeding from pelvic fractures for purposes of embolization with Gelfoam or autologous clot.

6. Intravenous urography—Intravenous urography can be used to detect renal and ureteral injury. This is best done with high-dose bolus injection of contrast media (2.0 mL/kg) followed by appropriate films.

B. CYSTOSCOPY AND RETROGRADE UROGRAPHY

Cystoscopy and retrograde urography may be useful to detect ureteral injury, but are seldom necessary, since information can be obtained by less invasive techniques.

C. ABDOMINAL SONOGRAPHY

Abdominal sonography has not been shown to add substantial information during initial evaluation of severe abdominal trauma.

INJURIES TO THE KIDNEY

Renal injuries are the most common injuries of the urinary system. The kidney is well protected by heavy lumbar muscles, vertebral bodies, ribs, and the viscera anteriorly. Fractured ribs and transverse vertebral processes may penetrate the renal parenchyma or vasculature. Most injuries occur from automobile accidents or sporting mishaps, chiefly in men and boys. Kidneys with existing pathologic conditions such as hydronephrosis or malignant tumors are more readily ruptured from mild trauma.

Etiology (Figure 17–4)

Blunt trauma directly to the abdomen, flank, or back is the most common mechanism, accounting for 80–85% of all renal injuries. Trauma may result from motor vehicle accidents, fights, falls, and contact sports. Vehicle collisions at high speed may result in major renal trauma from rapid deceleration and cause major vascular injury. Gunshot and knife wounds cause most penetrating injuries to the kidney; any such wound in the flank area should be regarded as a cause of renal injury until proved otherwise. Associated abdominal visceral injuries are present in 80% of renal penetrating wounds.

Pathology & Classification (Figure 17–5)

A. EARLY PATHOLOGIC FINDINGS

Lacerations from blunt trauma usually occur in the transverse plane of the kidney. The mechanism of injury is thought to be force transmitted from the center of the

Figure 17–5. Classification of renal injuries. Grades I and II are minor. Grades III, IV, and V are major. A: Grade I—microscopic or gross hematuria; normal findings on radiographic studies; contusion or contained subcapsular hematoma without parenchymal laceration. B: Grade II—nonexpanding, confined perirenal hematoma or cortical laceration less than 1 cm deep without urinary extravasation. C: Grade III—parenchymal laceration extending less than 1 cm into the cortex without urinary extravasation. D: Grade IV—parenchymal laceration extending through the corticomedullary junction and into the collecting system. A laceration at a segmental vessel may also be present. (continued)
Figure 17–5. continued  

E: Grade IV—thrombosis of a segmental renal artery without a parenchymal laceration. Note the corresponding parenchymal ischemia.  
F: Grade V—thrombosis of the main renal artery. The inset shows the intimal tear and distal thrombosis.  
G: Grade V—multiple major lacerations, resulting in a “shattered” kidney.  
H: Grade V—avulsion of the main renal artery or vein or both.
impact to the renal parenchyma. In injuries from rapid deceleration, the kidney moves upward or downward, causing sudden stretch on the renal pedicle and sometimes complete or partial avulsion. Acute thrombosis of the renal artery may be caused by an intimal tear from rapid deceleration injuries owing to the sudden stretch.

Pathologic classification of renal injuries is as follows:

**Grade 1 (the most common)**—Renal contusion or bruising of the renal parenchyma. Microscopic hematuria is common, but gross hematuria can occur rarely.

**Grade 2**—Renal parenchymal laceration into the renal cortex. Perirenal hematoma is usually small.

**Grade 3**—Renal parenchymal laceration extending through the cortex and into the renal medulla. Bleeding can be significant in the presence of large retroperitoneal hematoma.

**Grade 4**—Renal parenchymal laceration extending into the renal collecting system; also, main renal artery thrombosis from blunt trauma, segmental renal vein, or both; or artery injury with contained bleeding.

**Grade 5**—Multiple Grade 4 parenchymal lacerations, renal pedicle avulsion, or both; main renal vein or artery injury from penetrating trauma.

### B. LATE PATHOLOGIC FINDINGS (FIGURE 17–6)

1. **Urinoma**—Deep lacerations that are not repaired may result in persistent urinary extravasation and late complications of a large perinephric renal mass and, eventually, hydronephrosis and abscess formation.

2. **Hydronephrosis**—Large hematomas in the retroperitoneum and associated urinary extravasation may result in perinephric fibrosis engulfing the ureteropelvic junction, causing hydronephrosis. Follow-up excretory urography is indicated in all cases of major renal trauma.

3. **Arteriovenous fistula**—Arteriovenous fistulas may occur after penetrating injuries but are not common.

4. **Renal vascular hypertension**—The blood flow in tissue rendered nonviable by injury is compromised; this results in renal vascular hypertension in less than 1% of cases. Fibrosis from surrounding trauma has also been reported to constrict the renal artery and cause renal hypertension.

### Clinical Findings & Indications for Studies

Microscopic or gross hematuria following trauma to the abdomen indicates injury to the urinary tract. It bears repeating that stab or gunshot wounds to the flank area

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**Figure 17–6.** Late pathologic findings in renal trauma. **Left:** Ureteropelvic stenosis with hydronephrosis secondary to fibrosis from extravasation of blood and urine. **Right:** Atrophy of kidney caused by injury (stenosis) of arterial blood supply.
should alert the physician to possible renal injury whether or not hematuria is present. Some cases of renal vascular injury are not associated with hematuria. These cases are almost always due to rapid deceleration accidents and are an indication for imaging studies.

The degree of renal injury does not correspond to the degree of hematuria, since gross hematuria may occur in minor renal trauma and only mild hematuria in major trauma. However, not all adult patients sustaining blunt trauma require full imaging evaluation of the kidney (Figure 17–1). Miller and McAninch (1995) made the following recommendations based on findings in over 1800 blunt renal trauma injuries: Patients with gross hematuria or microscopic hematuria with shock (systolic blood pressure <90 mm Hg) should undergo radiographic assessment; patients with microscopic hematuria without shock need not. However, should physical examination or associated injuries prompt reasonable suspicion of a renal injury, renal imaging should be undertaken. This is especially true of patients with rapid deceleration trauma, who may have renal injury without the presence of hematuria.

A. SYMPTOMS

There is usually visible evidence of abdominal trauma. Pain may be localized to one flank area or over the abdomen. Associated injuries such as ruptured abdominal viscera or multiple pelvic fractures also cause acute abdominal pain and may obscure the presence of renal injury. Catheterization usually reveals hematuria. Retroperitoneal bleeding may cause abdominal distention, ileus, and nausea and vomiting.

B. SIGNS

Initially, shock or signs of a large loss of blood from heavy retroperitoneal bleeding may be noted. Ecchymosis in the flank or upper quadrants of the abdomen is often noted. Lower rib fractures are frequently found. Diffuse abdominal tenderness may be found on palpation; an "acute abdomen" usually indicates free blood in the peritoneal cavity. A palpable mass may represent a large retroperitoneal hematoma or perhaps urinary extravasation. If the retroperitoneum has been torn, free blood may be noted in the peritoneal cavity but no palpable mass will be evident. The abdomen may be distended and bowel sounds absent.

C. LABORATORY FINDINGS

Microscopic or gross hematuria is usually present. The hematocrit may be normal initially, but a drop may be found when serial studies are done. This finding represents persistent retroperitoneal bleeding and development of a large retroperitoneal hematoma. Persistent bleeding may necessitate operation.

D. STAGING AND X-RAY FINDINGS

Staging of renal injuries allows a systematic approach to these problems (Figures 17–1 through 17–3). Adequate studies help define the extent of injury and dictate appropriate management. For example, blunt trauma to the abdomen associated with gross hematuria and a normal urogram requires no additional renal studies; however, nonvisualization of the kidney requires immediate arteriography or CT scan to determine whether renal vascular injury exists. Ultrasonography and retrograde urography are of little use initially in the evaluation of renal injuries.

Staging begins with an abdominal CT scan, the most direct and effective means of staging renal injuries. This noninvasive technique clearly defines parenchymal lacerations and urinary extravasation, shows the extent of the retroperitoneal hematoma, identifies nonviable tissue, and outlines injuries to surrounding organs such as the pancreas, spleen, liver, and bowel (Figure 17–7). (If CT is not available, an intravenous pyelogram can be obtained [Figure 17–8].)

Arteriography defines major arterial and parenchymal injuries when previous studies have not fully done so. Arterial thrombosis and avulsion of the renal pedicle are best diagnosed by arteriography and are likely when the kidney is not visualized on imaging studies (Figure 17–9). The major causes of nonvisualization on an excretory urogram are total pedicle avulsion, arterial thrombosis, severe contusion causing vascular spasm, and absence of the kidney (either congenital or from operation).

Radionuclide renal scans have been used in staging renal trauma. However, in emergency management, this technique is less sensitive than arteriography or CT.

Differential Diagnosis

Trauma to the abdomen and flank areas is not always associated with renal injury. In such cases, there is no hematuria, and the results of imaging studies are normal.

Figure 17–7. Computed tomography scan of right kidney following knife stab wound. Laceration with urine extravasation is seen. Large right retroperitoneal hematoma is present.
Complications

A. EARLY COMPLICATIONS

Hemorrhage is perhaps the most important immediate complication of renal injury. Heavy retroperitoneal bleeding may result in rapid exsanguination. Patients must be observed closely, with careful monitoring of blood pressure and hematocrit. Complete staging must be done early (Figures 17–1 through 17–3). The size and expansion of palpable masses must be carefully monitored. Bleeding ceases spontaneously in 80–85% of cases. Persistent retroperitoneal bleeding or heavy gross hematuria may require early operation.

Urinary extravasation from renal fracture may show as an expanding mass (urinoma) in the retroperitoneum. These collections are prone to abscess formation and sepsis. A resolving retroperitoneal hematoma may cause slight fever (38.3°C [101°F]), but higher temperatures suggest infection. A perinephric abscess may form, resulting in abdominal tenderness and flank pain.

B. LATE COMPLICATIONS

Hypertension, hydronephrosis, arteriovenous fistula, calculus formation, and pyelonephritis are important late complications. Careful monitoring of blood pressure for several months is necessary to watch for hypertension. At 3–6 months, a follow-up excretory urogram or CT scan should be obtained to be certain that perinephric scarring has not caused hydronephrosis or vascular compromise; renal atrophy may occur from vascular compromise and is detected by follow-up urography. Heavy late bleeding may occur 1–4 weeks after injury.

Treatment

A. EMERGENCY MEASURES

The objectives of early management are prompt treatment of shock and hemorrhage, complete resuscitation, and evaluation of associated injuries.

B. SURGICAL MEASURES

1. Blunt injuries—Minor renal injuries from blunt trauma account for 85% of cases and do not usually require operation. Bleeding stops spontaneously with bed rest and hydration. Cases in which operation is indicated include those associated with persistent retroperitoneal bleeding, urinary extravasation, evidence of nonviable renal parenchyma, and renal pedicle injuries (less than 5% of all renal injuries). Aggressive preoperative staging allows complete definition of injury before operation.

2. Penetrating injuries—Penetrating injuries should be surgically explored. A rare exception to this rule is when staging has been complete and only minor parenchymal injury, with no urinary extravasation, is noted. In 80% of cases of penetrating injury, associated organ injury requires operation; thus, renal exploration is only an extension of this procedure.

C. TREATMENT OF COMPLICATIONS

Retroperitoneal urinoma or perinephric abscess demands prompt surgical drainage. Malignant hypertension requires vascular repair or nephrectomy. Hydronephrosis may require surgical correction or nephrectomy.
INJURIES TO THE GENITOURINARY TRACT

Prognosis
With careful follow-up, most renal injuries have an excellent prognosis, with spontaneous healing and return of renal function. Follow-up excretory urography and monitoring of blood pressure ensure detection and appropriate management of late hydronephrosis and hypertension.

INJURIES TO THE URETER

Ureteral injury is rare but may occur, usually during the course of a difficult pelvic surgical procedure or as a result of stab or gunshot wounds. Rapid deceleration accidents may avulse the ureter from the renal pelvis. Endoscopic basket manipulation of ureteral calculi may result in injury.

Etiology
Large pelvic masses (benign or malignant) may displace the ureter laterally and engulf it in reactive fibrosis. This may lead to ureteral injury during dissection, since the organ is anatomically malpositioned. Inflammatory pelvic disorders may involve the ureter in a similar way. Extensive carcinoma of the colon may involve areas outside the colon wall and directly involve the ureter; thus, resection of the ureter may be required along with resection of the tumor mass. Devascularization may occur with extensive pelvic lymph node dissections or after radiation therapy to the pelvis for pelvic cancer. In these situations, ureteral fibrosis and subsequent stricture formation may develop along with ureteral fistulas. Endoscopic manipulation of a ureteral calculus with a stone basket or ureteroscope may result in ureteral perforation or avulsion.

Pathogenesis & Pathology
The ureter may be inadvertently ligated and cut during difficult pelvic surgery. In such cases, sepsis and severe renal damage usually occur postoperatively. If a partially divided ureter is unrecognized at operation, urinary extravasation and subsequent buildup of a large urinoma will ensue, which usually leads to ureterovaginal or ureterocutaneous fistula formation. Intraperitoneal extravasation of urine can also occur, causing ileus and peritonitis. After partial transection of the ureter, some degree of stenosis and reactive fibrosis develops, with concomitant mild to moderate hydronephrosis.

Clinical Findings

A. SYMPTOMS
If the ureter has been completely or partially ligated during operation, the postoperative course is usually marked by fever of 38.3°C–38.8°C (101°F–102°F) as well as flank and lower quadrant pain. Such patients often experience paralytic ileus with nausea and vomiting. If ureterovaginal or cutaneous fistula develops, it usually does so within the first 10 postoperative days.

Ureteral injuries from external violence should be suspected in patients who have sustained stab or gunshot wounds to the retroperitoneum. The mid portion of the ureter seems to be the most common site of penetrating injury. There are usually associated vascular and other abdominal visceral injuries.

B. SIGNS
The acute hydronephrosis of a totally ligated ureter results in severe flank pain and abdominal pain with nausea and vomiting early in the postoperative course and with associated ileus. Signs and symptoms of acute peritonitis may be present if there is urinary extravasation into the peritoneal cavity. Watery discharge from the wound or vagina may be identified as urine by determining the creatinine concentration of a small sample—urine has many times the creatinine concentration found in serum—and by intravenous injection of 10 mL of indigo carmine, which will appear in the urine as dark blue.

C. LABORATORY FINDINGS
Ureteral injury from external violence is manifested by microscopic hematuria in 90% of cases. Urinalysis and other laboratory studies are of little use in diagnosis when injury has occurred from other causes.

D. IMAGING FINDINGS
Diagnosis is by excretory urography or delayed abdominal spiral CT scan. A plain film of the abdomen may demonstrate a large area of increased density in the pelvis or in an area of retroperitoneum where injury is suspected. After injection of contrast material, delayed excretion is noted with hydronephrosis. Partial transection of the ureter results in more rapid excretion, but persistent hydronephrosis is usually present, and contrast extravasation at the site of injury is noted on delayed films (Figure 17–10).

In acute injury from external violence, the excretory urogram usually appears normal, with very mild fullness down to the point of extravasation at the ureteral transection.

Retrograde ureterography demonstrates the exact site of obstruction or extravasation.

E. ULTRASONOGRAPHY
Ultrasoundography outlines hydroureter or urinary extravasation as it develops into a urinoma and is perhaps the best means of ruling out ureteral injury in the early postoperative period.

F. RADIONUCLIDE SCANNING
Radionuclide scanning demonstrates delayed excretion on the injured side, with evidence of increasing counts owing to accumulation of urine in the renal pelvis. Its great bene-
fit, however, is in the assessment of renal function after surgical correction.

**Differential Diagnosis**

Postoperative bowel obstruction and peritonitis may cause symptoms similar to those of acute ureteral obstruction from injury. Fever, “acute abdomen,” and associated nausea and vomiting following difficult pelvic surgery are definite indications for screening sonography or excretory urography to establish whether ureteral injury has occurred.

Deep wound infection must be considered postoperatively in patients with fever, ileus, and localized tenderness. The same findings are consistent with urinary extravasation and urinoma formation.

Acute pyelonephritis in the early postoperative period may also result in findings similar to those of ureteral injury. Sonography shows normal results, and urography shows no evidence of obstruction.

**Complications**

Ureteral injury may be complicated by stricture formation with resulting hydroureter or hydronephrosis in the area of injury. Chronic urinary extravasation from unrecognized injury may lead to formation of a large retroperitoneal urinoma. Pyelonephritis from hydroureter or urinary infection may require prompt proximal drainage.

**Treatment**

Prompt treatment of ureteral injuries is required. The best opportunity for successful repair is in the operating room when the injury occurs. If the injury is not recognized until 7–10 days after the event and no infection, abscess, or other complications exist, immediate reexploration and repair are indicated. Proximal urinary drainage by percutaneous nephrostomy or formal nephrostomy should be considered if the injury is recognized late or if the patient has significant complications that make immediate reconstruction unsatisfactory. The goals of ureteral repair are to achieve complete debridement, a tension-free spatulated anastomosis, watertight closure, ureteral stenting (in selected cases), and retroperitoneal drainage.

**A. LOWER URETERAL INJURIES**

Injuries to the lower third of the ureter allow several options in management. The procedure of choice is reimplantation into the bladder combined with a psoas-hitch procedure to minimize tension on the ureteral anastomosis. An antireflux procedure should be done when possible. Primary ureteroureterostomy can be used in lower-third injuries when the ureter has been ligated without transection. The ureter is usually long enough for this type of anastomosis. A bladder tube flap can be used when the ureter is shorter.

Transureteroureterostomy may be used in lower-third injuries if extensive urinoma and pelvic infection have developed. This procedure allows anastomosis and reconstruction in an area away from the pathologic processes.

**B. MIDURETERAL INJURIES**

Midureteral injuries usually result from external violence and are best repaired by primary ureteroureterostomy or transureteroureterostomy.

**C. UPPER URETERAL INJURIES**

Injuries to the upper third of the ureter are best managed by primary ureteroureterostomy. If there is extensive loss of the ureter, autotransplantation of the kidney can be done as well as bowel replacement of the ureter.

**D. STENTING**

Most anastomoses after repair of ureteral injury should be stented. The preferred technique is to insert a silicone internal stent through the anastomosis before closure. These stents have a J memory curve on each end to prevent their migration in the postoperative period. After 3–4 weeks of healing, stents can be endoscopically removed from the bladder. The advantages of internal stenting are maintenance of a straight ureter with a constant caliber during early healing, the presence of a conduit for urine during healing, prevention of urinary extravasation, maintenance of urinary diversion, and easy removal.

**Prognosis**

The prognosis for ureteral injury is excellent if the diagnosis is made early and prompt corrective surgery is done.
Delay in diagnosis worsens the prognosis because of infection, hydronephrosis, abscess, and fistula formation.

**INJURIES TO THE BLADDER**

Bladder injuries occur most often from external force and are often associated with pelvic fractures. (About 15% of all pelvic fractures are associated with concomitant bladder or urethral injuries.) Iatrogenic injury may result from gynecologic and other extensive pelvic procedures as well as from hernia repairs and transurethral operations.

**Pathogenesis & Pathology (Figure 17–11)**

The bony pelvis protects the urinary bladder very well. When the pelvis is fractured by blunt trauma, fragments from the fracture site may perforate the bladder. These perforations usually result in extraperitoneal rupture. If the urine is infected, extraperitoneal bladder perforations may result in deep pelvic abscess and severe pelvic inflammation.

When the bladder is filled to near capacity, a direct blow to the lower abdomen may result in bladder disruption. This type of disruption ordinarily is intraperitoneal. Since the reflection of the pelvic peritoneum covers the dome of the bladder, a linear laceration will allow urine to flow into the abdominal cavity. If the diagnosis is not established immediately and if the urine is sterile, no symptoms may be noted for several days. If the urine is infected, immediate peritonitis and acute abdomen will develop.

**Clinical Findings**

Pelvic fracture accompanies bladder rupture in 90% of cases. The diagnosis of pelvic fracture can be made initially in the emergency room by lateral compression on the bony pelvis, since the fracture site will show crepitus and be painful to the touch.

**A. SYMPTOMS**

There is usually a history of lower abdominal trauma. Blunt injury is the usual cause. Patients ordinarily are unable to urinate, but when spontaneous voiding occurs, gross hematuria is usually present. Most patients complain of pelvic or lower abdominal pain.

**B. SIGNS**

Heavy bleeding associated with pelvic fracture may result in hemorrhagic shock, usually from venous disruption of pelvic vessels. Evidence of external injury from a gunshot or stab wound in the lower abdomen should make one suspect bladder injury, manifested by marked tenderness of the suprapubic area and lower abdomen. An acute abdomen may occur with intraperitoneal bladder rupture. On rectal examination, landmarks may be indistinct because of a large pelvic hematoma.

**C. LABORATORY FINDINGS**

Catheterization usually is required in patients with pelvic trauma but not if bloody urethral discharge is noted. Bloody urethral discharge indicates urethral injury, and a urethrogram is necessary before catheterization (Figures 17–1 through 17–3). When catheterization is done, gross or, less commonly, microscopic hematuria is usually present. Urine taken from the bladder at the initial catheterization should be cultured to determine whether infection is present.

**D. X-RAY FINDINGS**

A plain abdominal film generally demonstrates pelvic fractures. There may be haziness over the lower abdomen from blood and urine extravasation. A CT scan should be obtained to establish whether kidney and ureteral injuries are present. Bladder disruption is shown on cystography. The bladder should be filled with 300 mL of contrast material and a plain film of the lower abdomen obtained. Contrast medium should be allowed to drain out completely, and a second film of the abdomen should be obtained. The drainage film is extremely important, because it demonstrates areas of extraperitoneal extravasation of blood and

![Figure 17–11. Mechanism of vesical injury. A direct blow over the full bladder causes increased intravesical pressure. If the bladder ruptures, it will usually rupture into the peritoneal cavity.](image)
urine that may not appear on the filling film (Figure 17–12). With intraperitoneal extravasation, free contrast medium is visualized in the abdomen, highlighting bowel loops (Figure 17–13).

CT cystography is an excellent method for detecting bladder rupture; however, retrograde filling of the bladder with 300 mL of contrast medium is necessary to distend the bladder completely. Incomplete distention with consequent missed diagnosis of bladder rupture often occurs when the urethral catheter is clamped during standard abdominal CT scan with intravenous contrast injection.

Complications
A pelvic abscess may develop from extraperitoneal bladder rupture; if the urine becomes infected, the pelvic hematoma becomes infected too. Intraperitoneal bladder rupture with extravasation of urine into the abdominal cavity causes delayed peritonitis. Partial incontinence may result from bladder injury when the laceration extends into the bladder neck. Meticulous repair may ensure normal urinary control.

Treatment
A. EMERGENCY MEASURES
Shock and hemorrhage should be treated.

B. SURGICAL MEASURES
A lower midline abdominal incision should be made. As the bladder is approached in the midline, a pelvic hematoma, which is usually lateral, should be avoided. Entering the pelvic hematoma can result in increased bleeding from release of tamponade and in infection of the hematoma, with subsequent pelvic abscess. The bladder should be opened in the midline and carefully inspected. After repair, a suprapubic cystostomy tube is usually left in place to ensure complete urinary drainage and control of bleeding.

1. Extraperitoneal bladder rupture—Extraperitoneal bladder rupture can be successfully managed with urethral catheter drainage only. (Typically 10 days will provide adequate healing time.) Large blood clots in the bladder or injuries involving the bladder neck should be managed surgically.

As the bladder is opened in the midline, it should be carefully inspected and lacerations closed from within. Polyglycolic acid or chromic absorbable sutures should be used.

Extraperitoneal bladder lacerations occasionally extend into the bladder neck and should be repaired meticulously. Fine absorbable sutures should be used to ensure complete reconstruction so that the patient will have urinary control after injury. Such injuries are best managed with indwelling urethral catheterization and suprapubic diversion.

2. Intraperitoneal rupture—Intraperitoneal bladder ruptures should be repaired via a transperitoneal approach after careful transvesical inspection and closure of any other perforations. The peritoneum must be closed carefully over the area of injury. The bladder is then closed in separate layers by absorbable suture. All extravasated fluid from the peritoneal cavity should be removed before closure. At the time of closure, care should be taken that the suprapubic cystostomy is in the extraperitoneal position.
INJURIES TO THE GENITOURINARY TRACT

3. Pelvic fracture—Stable fracture of the pubic rami is usually present. In such cases, the patient can be ambulatory within 4–5 days without damage or difficulty. Unstable pelvic fractures requiring external fixation have a more protracted course.

4. Pelvic hematoma—There may be heavy uncontrolled bleeding from rupture of pelvic vessels even if the hematoma has not been entered at operation. At exploration and bladder repair, packing the pelvis with laparotomy tapes often controls the problem. If bleeding persists, it may be necessary to leave the tapes in place for 24 hours and operate again to remove them. Embolization of pelvic vessels with Gelfoam or skeletal muscle under angiographic control is useful in controlling persistent pelvic bleeding.

Prognosis
With appropriate treatment, the prognosis is excellent. The suprapubic cystostomy tube can be removed within 10 days, and the patient can usually void normally. Patients with lacerations extending into the bladder neck area may be temporarily incontinent, but full control is usually regained. At the time of discharge, urine culture should be performed to determine whether catheter-associated infection requires further treatment.

INJURIES TO THE URETHRA
Urethral injuries are uncommon and occur most often in men, usually associated with pelvic fractures or straddle-type falls. They are rare in women. Various parts of the urethra may be lacerated, transected, or contused. Management varies according to the level of injury. The urethra can be separated into 2 broad anatomic divisions: the posterior urethra, consisting of the prostatic and membranous portions, and the anterior urethra, consisting of the bulbous and pendulous portions.

INJURIES TO THE POSTERIOR URETHRA
Etiology (Figure 17–14)
The membranous urethra passes through the pelvic floor and voluntary urinary sphincter and is the portion of the posterior urethra most likely to be injured. When pelvic fractures occur from blunt trauma, the membranous urethra is sheared from the prostatic apex at the prostatomembranous junction. The urethra can be transected by the same mechanism at the interior surface of the membranous urethra.

Clinical Findings
A. Symptoms
Patients usually complain of lower abdominal pain and inability to urinate. A history of crushing injury to the pelvis is usually obtained.

B. Signs
Blood at the urethral meatus is the single most important sign of urethral injury. The importance of this finding cannot be overemphasized, because an attempt to pass a urethral catheter may result in infection of the periprostatic and perivesical hematoma and conversion of an incomplete laceration to a complete one. The presence of blood at the external urethral meatus indicates that immediate urethrography is necessary to establish the diagnosis.

Suprapubic tenderness and the presence of pelvic fracture are noted on physical examination. A large developing pelvic hematoma may be palpated. Perineal or suprapubic contusions are often noted. Rectal examination may reveal a large pelvic hematoma with the prostate displaced superiority. Rectal examination can be misleading, however, because a tense pelvic hematoma may resemble the prostate on palpation. Superior displacement of the prostate does not occur if the puboprostatic ligaments remain intact. Partial disruption of the membranous urethra (currently 10% of cases) is not accompanied by prostatic displacement.

C. X-Ray Findings
Fractures of the bony pelvis are usually present. A urethrogram (using 20–30 mL of water-soluble contrast material) shows the site of extravasation at the prostatomembranous
junction. Ordinarily, there is free extravasation of contrast material into the perivesical space (Figure 17–15). Incomplete prostatomembranous disruption is seen as minor extravasation, with a portion of contrast material passing into the prostatic urethra and bladder.

**D. INSTRUMENTAL EXAMINATION**

The only instrumentation involved should be for urethrography. Catheterization or urethroscopy should not be done, because these procedures pose an increased risk of hematoma, infection, and further damage to partial urethral disruptions.

**Differential Diagnosis**

Bladder rupture may be associated with posterior urethral injuries in approximately 20% of cases. Cystography cannot be done preoperatively, since a urethral catheter should not be passed. Careful evaluation of the bladder at operation is necessary.

**Complications**

Stricture, impotence, and incontinence as complications of prostatomembranous disruption are among the most severe and debilitating mishaps that result from trauma to the urinary system. Stricture following primary repair and anastomosis occurs in about 50% of cases. If the preferred suprapubic cystostomy approach with delayed repair is used, the incidence of stricture can be reduced to about 5%.

The incidence of impotence after primary repair is 30–80% (mean, about 50%). This figure can be reduced to 30–35% by suprapubic drainage with delayed urethral reconstruction.

**Treatment**

**A. EMERGENCY MEASURES**

Shock and hemorrhage should be treated.

**B. SURGICAL MEASURES**

Urethral catheterization should be avoided.

1. **Immediate management**—Initial management should consist of suprapubic cystostomy to provide urinary drainage. A midline lower abdominal incision should be made, with care being taken to avoid the large pelvic hematoma. The bladder and prostate are usually elevated superiorly by large periprostatic and perivesical hematomas. The bladder often is distended by a large volume of urine accumulated during the period of resuscitation and operative preparation. The urine is often clear and free of blood, but gross hematuria may be present. The bladder should be opened in the midline and carefully inspected for lacerations. If a laceration is present, the bladder should be closed with absorbable suture material and a cystostomy tube inserted for urinary drainage. This approach involves no urethral instrumentation or manipulation. The suprapubic cystostomy is maintained in place for about 3 months. This allows resolution of the pelvic hematoma, and the prostate and bladder will slowly return to their anatomic positions.

Incomplete laceration of the posterior urethra heals spontaneously, and the suprapubic cystostomy can be removed within 2–3 weeks. The cystostomy tube should not be removed before voiding cystourethrography shows that no extravasation persists.

2. **Delayed urethral reconstruction**—Reconstruction of the urethra after prostatic disruption can be undertaken within 3 months, assuming there is no pelvic abscess or other evidence of persistent pelvic infection. Before reconstruction, a combined cystogram and urethrogram should be done to determine the exact length of the resulting urethral stricture. This stricture usually is 1–2 cm long and situated immediately posterior to the pubic bone. The preferred approach is a single-stage reconstruction of the urethral rupture defect with direct excision of the strictured area and anastomosis of the bulbous urethra directly to the apex of the prostate. A 16F silicone urethral catheter should be left in place along with a suprapubic cystostomy. Catheters are removed within a month, and the patient is then able to void (Figure 17–16).

3. **Immediate urethral realignment**—Some surgeons prefer to realign the urethra immediately. Persistent bleeding and surrounding hematoma create technical problems. The incidence of stricture, impotence, and incontinence appears to be higher than with immediate cystostomy and

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**Figure 17–15.** Ruptured prostatomembranous urethra shows free extravasation on urethrogram. No contrast medium is seen entering the prostatic urethra.

Total urinary incontinence occurs in <2% of patients and typically is associated with severe sacral fracture and S2-4 nerve injury.
INJURIES TO THE GENITOURINARY TRACT

delayed reconstruction. However, several authors have reported success with immediate urethral realignment.

C. GENERAL MEASURES

After delayed reconstruction by a perineal approach, patients are allowed ambulation on the first postoperative day and usually can be discharged within 3 days.

D. TREATMENT OF COMPLICATIONS

Approximately 1 month after the delayed reconstruction, the urethral catheter can be removed and a voiding cystogram obtained through the suprapubic cystostomy tube. If the cystogram shows a patent area of reconstruction free of extravasation, the suprapubic catheter can be removed; if there is extravasation or stricture, suprapubic cystostomy should be maintained. A follow-up urethrogram should be obtained within 2 months to watch for stricture development.

Stricture, if present (<5%), is usually very short, and urethrotomy under direct vision offers easy and rapid cure. The patient may be impotent for several months after delayed repair. Impotence is permanent in about 10% of patients. Implantation of a penile prosthesis is indicated if impotence is still present 2 years after reconstruction (see Chapter 38). Incontinence after posterior urethral rupture and delayed repair is rare (<2%) and is usually related to the extent of injury rather than to the repair.

Prognosis

If complications can be avoided, the prognosis is excellent. Urinary infections ultimately resolve with appropriate management.

INJURIES TO THE ANTERIOR URETHRA

Etiology (Figure 17–17)

The anterior urethra is the portion distal to the urogenital diaphragm. Straddle injury may cause laceration or contusion of the urethra. Self-instrumentation or iatrogenic instrumentation may cause partial disruption.

Prognosis

If complications can be avoided, the prognosis is excellent. Urinary infections ultimately resolve with appropriate management.
Pathogenesis & Pathology

A. CONTUSION

Contusion of the urethra is a sign of crush injury without urethral disruption. Perineal hematoma usually resolves without complications.

B. LACERATION

A severe straddle injury may result in laceration of part of the urethral wall, allowing extravasation of urine. If the extravasation is unrecognized, it may extend into the scrotum, along the penile shaft, and up to the abdominal wall. It is limited only by Colles’ fascia and often results in sepsis, infection, and serious morbidity.

Clinical Findings

A. SYMPTOMS

There is usually a history of a fall, and in some cases a history of instrumentation. Bleeding from the urethra is usually present. There is local pain into the perineum and sometimes massive perineal hematoma. If voiding has occurred and extravasation is noted, sudden swelling in the area will be present. If diagnosis has been delayed, sepsis and severe infection may be present.

B. SIGNS

The perineum is very tender; a mass may be found, as may blood at the urethral meatus. Rectal examination reveals a normal prostate. The patient usually has a desire to void, but voiding should not be allowed until assessment of the urethra is complete. No attempt should be made to pass a urethral catheter, but if the patient’s bladder is overdistended, percutaneous suprapubic cystostomy can be done as a temporary procedure.

When presentation of such injuries is delayed, there is massive urinary extravasation and infection in the perineum and the scrotum. The lower abdominal wall may also be involved. The skin is usually swollen and discolored.

C. LABORATORY FINDINGS

Blood loss is not usually excessive, particularly if secondary injury has occurred. The white count may be elevated with infection.

D. X-RAY FINDINGS

A urethrogram, with instillation of 15–20 mL of water-soluble contrast material, demonstrates extravasation and the location of injury (Figure 17–18). A contused urethra shows no evidence of extravasation.

Complications

Heavy bleeding from the corpus spongiosum injury may occur in the perineum as well as through the urethral meatus. Pressure applied to the perineum over the site of the injury usually controls bleeding. If hemorrhage cannot be controlled, immediate operation is required.

The complications of urinary extravasation are chiefly sepsis and infection. Aggressive debridement and drainage are required if there is infection. Stricture at the site of injury is a common complication, but surgical reconstruction may not be required unless the stricture significantly reduces urinary flow rates.

Treatment

A. GENERAL MEASURES

Major blood loss usually does not occur from straddle injury. If heavy bleeding does occur, local pressure for control, followed by resuscitation, is required.

B. SPECIFIC MEASURES

1. Urethral contusion—The patient with urethral contusion shows no evidence of extravasation, and the urethra remains intact. After urethrography, the patient is allowed to void; and if the voiding occurs normally, without pain or bleeding, no additional treatment is necessary. If bleeding persists, urethral catheter drainage can be done.

2. Urethral lacerations—Instrumentation of the urethra following urethrography should be avoided. A small midline incision in the suprapubic area readily exposes the dome of the bladder so that a suprapubic cystostomy tube can be inserted, allowing complete urinary diversion while the urethral laceration heals. Percutaneous cystostomy may also be used in such injuries. If only minor extravasation is noted on the urethrogram, a voiding study can be per-
formed within 7 days after suprapubic catheter drainage to search for extravasation. In more extensive injuries, one should wait 2–3 weeks before doing a voiding study through the suprapubic catheter. Healing at the site of injury may result in stricture formation. Most of these strictures are not severe and do not require surgical reconstruction. The suprapubic cystostomy catheter may be removed if no extravasation is documented. Follow-up with documentation of urinary flow rates will show whether there is urethral obstruction from stricture.

3. Urethral laceration with extensive urinary extravasation—After major laceration, urinary extravasation may involve the perineum, scrotum, and lower abdomen. Drainage of these areas is indicated. Suprapubic cystostomy for urinary diversion is required. Infection and abscess formation are common and require antibiotic therapy.

4. Immediate repair—Immediate repair of urethral lacerations can be performed, but the procedure is difficult and the incidence of associated stricture is high.

C. TREATMENT OF COMPLICATIONS

Strictures at the site of injury may be extensive and require delayed reconstruction.

Prognosis

Urethral stricture is a major complication but in most cases does not require surgical reconstruction. If, when stricture resolves, urinary flow rates are poor and urinary infection and urethral fistula are present, reconstruction is required.

INJURIES TO THE PENIS

Disruption of the tunica albuginea of the penis (penile fracture) can occur during sexual intercourse. At presentation, the patient has penile pain and hematoma. This injury should be surgically corrected.

Gangrene and urethral injury may be caused by obstructing rings placed around the base of the penis. These objects must be removed without causing further damage. Penile amputation is seen occasionally, and in a few patients, the penis can be surgically replaced successfully by microsurgical techniques.

Total avulsion of the penile skin may occur from machinery injuries. Immediate debridement and skin grafting are usually successful in salvage. Injuries to the penis should suggest possible urethral damage, which should be investigated by urethrography.

INJURIES TO THE SCROTUM

Superficial lacerations of the scrotum may be debrided and closed primarily. Blunt trauma may cause local hematoma and ecchymosis, but these injuries resolve without difficulty. One must be certain that testicular rupture has not occurred.

INJURIES TO THE TESTIS

Blunt trauma to the testis causes severe pain and, often, nausea and vomiting. Lower abdominal tenderness may be present. A hematoma may surround the testis and make delineation of its margin difficult. Ultrasonography can be used as an aid to better define the organ. If rupture has occurred, the sonogram will delineate the injury, which should be surgically repaired.

REFERENCES

Emergency Diagnosis & Management


Injuries to the Kidney

Injuries to the Bladder


Injuries to the Urethra


Injuries to the Penis


Injuries to the Scrotum


Both experimental and naturally occurring tumors are capable of stimulating a specific antitumor immune response. This observation suggests that there are foreign proteins (antigens) on tumor cells that classically have been described as resulting in humoral and cellular immune responses. However, experimental models suggest that a T-cell (cell-mediated) response may be more important in the killing of tumor cells than a B-cell (humoral) response.

A detailed description of the components of the immune system is beyond the scope of this chapter, but certain features of the immune system as they pertain to diagnostic and therapeutic issues will be reviewed.

**Tumor Antigens**

Tumor antigens can be divided into tumor-specific antigens and tumor-associated antigens. Tumor-specific antigens are not found on normal tissue, and they permit the host to recognize a tumor as foreign. Tumor-specific antigens have been shown to exist in oncogenesis models utilizing chemical, physical, and viral carcinogens but appear to be less common in models of spontaneous tumor development.

The identification of tumor-specific antigens led to the theory of immune surveillance, which suggests that the immune system is continuously trolling for foreign (tumor-specific) antigens. This theory is supported by the observation that at least some cancers are more common in immune-suppressed patients such as transplant patients or human immunodeficiency virus-infected individuals. However, many cancers are not overrepresented in these patient populations. Furthermore, spontaneous tumor models, which more closely resemble human carcinogenesis, appear to have a less extensive repertoire of tumor-specific antigens but instead have been found to express many tumor-associated antigens.

Tumor-associated antigens are found on normal cells but either become less prevalent in normal tissue after embryogenesis (eg, alpha-fetoprotein [AFP]) or remain present on normal tissue but are overexpressed on cancer cells (eg, prostate-specific antigen [PSA]). In either case, the more ubiquitous nature of these antigens appears to cause reduced immune reactivity (also known as tolerance) to the specific antigen. The mechanisms of tolerance are complex and may be due in part to the absence of other required costimulatory molecules (such as B7, a molecule required for T-cell stimulation).

The development of monoclonal (hybridoma) technology has allowed the development of many antibodies against many tumor-associated antigens and has provided insight into the regulation and expression of these antigens. The reexpression or upregulation of these tumor-associated antigens during carcinogenesis may lead to immune response (or loss of tolerance). Many novel therapeutic approaches have sought to break this tolerance, and approaches to enhance a patient’s immune response will be discussed.

**Humoral Immunity**

A large number of monoclonal antibodies have been developed against a variety of tumor-associated antigens. Oncofetal antigens such as AFP and beta-human chorionic gonadotropin (β-hCG) are important markers in germ cell tumors. β-hCG is also expressed in a small percentage of patients with bladder carcinoma. Antibodies directed against specific targets such as vascular endothelial growth factor (vegF) have been correctly developed and are being tested for the treatment of both advanced prostate cancer of RCC.

**Antibodies in Cancer Diagnosis & Detection**

**A. PROSTATE CANCER**

Immunostaining are used to test both body fluids and tissues for the presence of tumor-associated antigens. In the urologic cancers, the most obvious example is the development of monoclonal antibodies against PSA. The utility and limitations of PSA are described elsewhere in this volume. Other antigens that have been tested in prostate cancer include prostatic acid phosphatase, which has largely been replaced by PSA in screening programs and in patients with low tumor burden. Prostatic acid phosphatase may be of some use in detecting or following up bone
metastases and as a predictive marker of response to therapy for metastatic disease, both hormone-sensitive and insensitive. More recently, antibodies to prostate-specific membrane antigen (PSMA) have been used, primarily for immunohistochemistry.

B. RENAL CELL CARCINOMA

Unfortunately, there are as yet no well-established antigens (or antibodies) that can be used to reliably evaluate and monitor renal cell carcinoma, although a variety of target antigens are being evaluated.

C. BLADDER CANCER

Two oncofetal antigens, β-hCG and carcinoembryonic antigen, are expressed by a minority (20% or less) of transitional cell carcinomas. These markers are not routinely used, but in diagnostic dilemmas, measurement of serum levels of β-hCG or staining of tissue for this antigen may be useful.

D. GERM CELL TUMORS

As described in Chapter 23, antibodies to hCG and AFP are routinely used to detect shed antigen from germ cell tumors in the bloodstream. These antigens can also be detected on tissue samples in the setting of some diagnostic dilemmas. While the use of serum markers in germ cell tumors is reviewed elsewhere, it is worth noting that the presence of the oncofetal protein AFP, either in serum or on tissue specimens, is pathognomonic for a nonseminomatous germ cell tumor, regardless of results of routine pathologic evaluation. In addition to their diagnostic utility, AFP and hCG can be used as markers of response to therapy and as predictive factors of outcome. For example, the international germ cell tumor risk classification scheme for patients with metastatic disease relies heavily on AFP and hCG levels as well as levels of a nonspecific marker, lactate dehydrogenase, to assign patients with nonseminomatous germ cell tumors to 1 of 3 risk levels (see Chapter 23).

E. RADIOIMMUNODETECTION

Monoclonal antibodies to a specific antigen can be radiolabeled, and the preferential binding of the monoclonal antibody to tumor cells can be exploited. Theoretically, such an approach could be used for the presurgical evaluation of disease, postsurgical evaluation for minimal residual disease, confirmation of cancer identified by other imaging modalities, and detection of recurrent disease. There are several potential impediments to successful tumor radioimmunodetection. These include dilution of antibody in the bloodstream; metabolism of the antibody; nonspecific binding in liver, reticuloendothelial system, bone marrow, and elsewhere; binding of antibody by circulating or shed antigen; and the development of neutralizing human antihuman antibodies.

The only radioimmunodetection system for urologic cancers at this time is 111In-capromab pendetide (Prostascint), a murine monoclonal antibody to PSMA. Its use has been hampered by a fairly laborious administration process, operator dependence in interpretation of scans, and a less than satisfactory positive predictive value. The use of 111In-capromab pendetide is described in Chapter 10.

Immunotherapy with Monoclonal Antibodies

Immunotherapy with monoclonal antibodies alone (“naked antibodies”) has been fairly extensively evaluated. The use of monoclonal antibodies against tumor-associated antigens has met with only limited success in patients with solid tumors. In lymphoproliferative disorders such as leukemia and lymphoma, some antibodies to tumor-associated surface antigens appear to result in tumor cell death. The mechanism for these effects is certainly multifactorial but may in part be mediated by resultant complement fixation.

Direct antiproliferative effects of antibodies on cancer cells can be achieved by antibodies against functionally important antigens. Thus, the inhibition of growth factors and growth factor receptors and the activation or inhibition of signal transducing molecules are attractive therapeutic targets. In the urologic cancers, while no approved monoclonal antibody therapy exists, trials of antibodies against growth factors, vascular endothelial growth factor (VEGF, an angiogenic molecule), and signal transduction molecules are being undertaken. Kidney cancer is highly dependent on angiogenesis, and bevacizumab (an antibody agent against VEGF) has been shown to prolong time to progression in metastatic disease. Results from a trial of interferon-alpha with and without bevacizumab are awaited. There is, as well, an ongoing trial of chemotherapy with and without bevacizumab in patients with metastatic hormone refractory prostate cancer.

An alternative approach to naked antibodies is to conjugate any of a variety of cytotoxic agents to an antibody. The advantage of this approach is a “bystander effect,” making it unnecessary to use an antibody that binds each and every cell. This can be achieved in a variety of ways. The most straightforward is to use the monoclonal antibody as a means of providing some targeting specificity for the cytotoxic agent used. Cytotoxic agents used include radioisotopes, chemotherapy, and toxins such as ricin. Other means of providing some specificity is to bind a prodrug (with an antibody) to the tumor site and then to activate the bound prodrug. Finally, targeting with bispecific antibodies (eg, to antigen and to an effector T cell, or to antigen and toxin) has been undertaken. These approaches have all been tested in prostate cancer, but all remain investigational at this point.
**Cell-Mediated Immunity**

There is considerable evidence, both clinical and preclinical, that tumor-associated antigens can elicit a cell-mediated immune response. In some models, when carcinogen-induced tumors in mice are resected and then reimplanted, the tumor fails to regrow, suggesting the development of immunity to specific antigens. Specific antigens that are rejected in immunized hosts are termed transplantation antigens. The specificity of tumor rejection has since been demonstrated to reside in T lymphocytes (at a minimum). Lymphocytes of cancer patients can sometimes be stimulated in vitro to recognize specific tumor-associated antigens and consequently demonstrate properties of cytolytic T lymphocytes. Unfortunately, the phenomenon of tumor rejection is by no means universal, either in the laboratory or clinically, and it is unusual to detect cytolytic-T-lymphocyte activity against many tumor-associated antigens.

Nevertheless, there are several clinical scenarios that suggest that cell-mediated antitumor responses exist. These observations have promoted a broad search for the means of enhancing patients’ immune responses to tumor-associated antigens. Renal cell carcinoma (RCC) is in many ways the prototypical immune-mediated tumor and, along with melanoma, has until recently been the primary target of immune manipulation. A dramatic example of such an immune-mediated response is the phenomenon of spontaneous regression of metastatic RCC deposits after nephrectomy. Classically this has been described in less than 1% of patients. The impact of tumor debulking may also explain why a subset of RCC patients with lymph node or renal vein involvement that undergo resection are seemingly cured. The exact mechanism of this phenomenon is not well understood but may involve elimination of inhibitors of cell-mediated immunity. Indeed, tumor-infiltrating lymphocytes in RCC have been shown to exhibit mutant or faulty T-cell-receptor components, and it is unusual to detect cytolytic-T-lymphocyte activity against many tumor-associated antigens.

Nevertheless, there are several clinical scenarios that suggest that cell-mediated antitumor responses exist. These observations have promoted a broad search for the means of enhancing patients’ immune responses to tumor-associated antigens. Renal cell carcinoma (RCC) is in many ways the prototypical immune-mediated tumor and, along with melanoma, has until recently been the primary target of immune manipulation. A dramatic example of such an immune-mediated response is the phenomenon of spontaneous regression of metastatic RCC deposits after nephrectomy. Classically this has been described in less than 1% of patients. The impact of tumor debulking may also explain why a subset of RCC patients with lymph node or renal vein involvement that undergo resection are seemingly cured. The exact mechanism of this phenomenon is not well understood but may involve elimination of inhibitors of cell-mediated immunity. Indeed, tumor-infiltrating lymphocytes in RCC have been shown to exhibit mutant or faulty T-cell-receptor components, and it is unusual to detect cytolytic-T-lymphocyte activity against many tumor-associated antigens.

**Immunotherapy Involving Cell-Mediated Immunity**

Additional evidence of cell-mediated immunity playing a role in tumor rejection lies in the results of a variety of immunotherapeutic interventions. Immunotherapy can be broadly classified as active or passive. This classification refers to the role the host’s immune system plays. Thus, the passive transfer of preformed antibodies is contrasted to a vaccination program in which the host’s immune system must be capable of mounting an immune response. Adoptive therapy refers to a middle ground in which efforts are made to reconstitute, modify, or bolster one of the effector cells involved ex vivo, followed by reinfusion into the patient, where the rest of the immune cascade must then be recruited.

**Active Immunotherapy: Vaccination**

Autologous vaccination programs (the vaccination of patients with their own tumor cells) have been extensively explored. The advantage of autologous vaccination is that the vaccine bears the antigens of the patient’s tumor, although the distinct disadvantage is that not every patient has tumor available for vaccine preparation, and the preparation of each vaccine is tremendously labor intensive. By contrast, allogeneic vaccines (the use of a generic vaccine or “off-the-shelf” antigen) have the benefit of mass production and ease of use, and the identification of specific tumor rejection antigens allows specific antigenic targeting. However, this approach runs the risk of a more narrow shared antigenic spectrum with the patient’s tumor. Both autologous and allogeneic vaccination strategies are under evaluation, both in RCC and prostate cancer.

Several means exist to undertake vaccination. The simplest is to use intact but inactivated tumor cells. Inactivation can be achieved with UV radiation, external beam (photon) radiation, or freeze-thawing. Crude extracts of cells can also be used. The advantage of using cell extracts is that inactivation is not necessary and small particles and proteins that might be more easily phagocytosed are available. One can also enhance the immunogenicity of inactivated cells by growing the cells in cytokines, coinjecting with cytokines (nonspecific active immunotherapy, described below), or transfecting these cells with the genes for immune stimulatory cytokines or the costimulatory molecule B7. Current clinical trials are underway that use prostate cancer cell lines transfected with the GM-CSF gene (GVAX, Cell Genesys, South San Francisco, CA) for vaccination in patients with metastatic hormone refractory prostate cancer. Purified protein or peptides represent a second potential vaccination schema. In prostate cancer, trials of vaccination with PSMA and PSA are under way. Trials of PSA in a vaccine and fowlpox (ProstaVax) are also underway. A third way of undertaking specific vaccination is to attempt to bypass the antigen-presenting function of the immune system and to directly stimulate professional antigen-presenting cells, such as dendritic cells, ex vivo. These cells can be stimulated by pulsing them with protein or peptides of interest or by transfecting them with a gene encoding the antigenic peptide of interest before re-infusion. Initial trials of PAP-pulsed dendritic cells (Provenge, Dendreon Corporation, Seattle, WA) have demonstrated preliminary activity. Confirmatory trials are ongoing.

**Nonspecific Active Immunotherapy: Cytokines & Biologic Response Modifiers**

BCG (Bacillus Calmette-Guérin) is a live attenuated form of tubercle bacillus that appears to have local activity against
Interleukin-2 (IL-2) is a naturally occurring cytokine that has multiple immunoregulatory properties. The observation that exogenously administered IL-2 could result in tumor regression in patients with RCC and melanoma was the first unequivocal indication that cancer regression could be mediated by immune manipulations. IL-2 stimulates lymphocyte proliferation, enhances cytolytic-T-cell activity, induces natural killer cell activity, and induces gamma-interferon and tumor necrosis factor production. IL-2 has no direct cytotoxicity, but when administered endogenously will activate effector cells of the host immune system, including lymphocytes, natural killer cells, lymphokine-activated killer cells, and tumor-infiltrating lymphocytes. The details of immunotherapy for RCC are beyond the scope of this chapter. Nevertheless, in brief, IL-2 has been administered in RCC in several different schemas, including high-dose intravenous bolus (IL-2 is U.S. Food and Drug Administration [FDA] approved with this schedule), continuous intravenous infusion, and at lower doses subcutaneously. The high-dose regimens must be administered on an inpatient basis and are characterized by significant, albeit manageable, toxicities, including fever, malaise; vascular leak syndrome; hypotension; and cardiac, renal, and hepatic dysfunction. Subcutaneous IL-2 is self-administered by patients in the outpatient setting, and while clearly less toxic, still has associated malaise and constitutional symptoms. The optimal dosing regimen is not well established, and overall response proportions rarely exceed 20%. Durable complete responses of 5–8% have been reported with some of the high-dose regimens. IL-2 has also been combined with other active agents such as alpha-interferon and chemotherapy, although it is not clear if these combinations provide additional benefit.

Alpha-interferon is a naturally occurring cytokine that has direct cytotoxic and possibly antiproliferative properties, but also has immunoregulatory properties. It enhances major histocompatibility complex expression, thereby potentially increasing the efficiency of antigen processing and recognition. Alpha-interferon has anticancer activity in both RCC and superficial bladder cancer. Its primary toxicity is fever, malaise, and constitutional symptoms, although at higher doses it can result in bone marrow toxicity, central nervous system toxicity, and hepatic toxicity. In RCC, as a single agent, alpha-interferon can result in clinical responses in up to 20% of patients. In contradistinction to IL-2 as a single agent, durable complete responses are quite rare. Nevertheless, in randomized trials, alpha-interferon appears to confer a modest survival advantage over other agents now known to be largely inactive. Alpha-interferon is also used as an intravesicle treatment in superficial bladder cancer, where it has established activity, and is not infrequently used as second-line therapy after BCG.

Granulocyte macrophage-colony stimulating factor (GM-CSF) is perhaps the most important cytokine in eliciting cellular immune responses. Administered systemically as a subcutaneous injection, GM-CSF has been shown to reduce PSA in patients with both hormone-sensitive and hormone-resistant prostate cancer. However, the use of GM-CSF is neither proven to be of clinical benefit, nor approved for this indication, and must be considered investigational.

**Immunomodulation**

A myriad of immunosuppressive factors exist within cancer patients that may serve to dampen anti-tumor immune responses. Some of these molecules represent natural pathways to inhibit autoimmunity, while some molecules may have been usurped by the tumor to evade immune recognition. Novel approaches are now being developed to target these pathways. For example, CTLA-4 is an inhibitory molecule that blocks binding of B7 to CD28, thereby preventing costimulation and downmodulating T-cell activation. By preventing the action of CTLA-4, an anti-CTLA-4 antibody (ipilimumab) can augment and prolong T-cell immune responses. In animal models, ipilimumab 4 antibody can induce tumor rejection in immunogenic tumors, and in combination with antitumor vaccination, can induce rejection of minimally immunogenic tumors, including in the transgenic adeno carcinoma of mouse/prostate (TRAMP) prostate cancer model. In a phase I study, 14 patients with androgen insensitive prostate cancer were treated with a humanized anti-CTLA-4 antibody (MDX-010, Medarex, Inc., Bloomsbury, NJ). There was no evidence of polyclonal T-cell activation, therapy was well tolerated, and 2 patients had ≥50% decline in their PSA. The combination of CTLA-4 blockade with vaccination is of interest and is under investigation.

**Adoptive Immunotherapy**

Adoptive immunotherapy is the transfer of cellular products (effector cells) to the host or patient in an effort to develop an immune response. The use of adoptive immunotherapy was prompted by the observation that T cells derived from patients with melanoma or RCC had the ability to recognize antigens on the primary tumor. Thus, it was hoped that these cells could be harvested, activated...
ex vivo, and then reinfused into patients. Lymphokine-activated killer cells and tumor-infiltrating lymphocytes have been used to treat patients with metastatic RCC in the investigational setting, frequently along with IL-2. However, randomized trials comparing IL-2 alone with IL-2 plus cellular products have failed to demonstrate an improvement in response proportions or survival. Chapter 22 gives specific details of immunotherapy in RCC.

REFERENCES

Chemotherapy of Urologic Tumors

Eric J. Small, MD

The use of chemotherapy in the treatment of malignant tumors of the genitourinary system serves as a paradigm for a multidisciplinary approach to cancer. The careful integration of surgical and chemotherapeutic treatments has resulted in impressive advances in the management of urologic cancer. By definition, surgical interventions are directed at local management of urologic tumors, whereas chemotherapy and biologic therapy are systemic in nature. While there is no question that there are times in the natural history of genitourinary tumor when only one therapeutic method is required, a multidisciplinary approach is always called for. This chapter details the importance of a joint surgical-medical approach to patients with urologic cancer. A practicing urologist should collaborate closely with a medical oncologist and should feel comfortable speaking with patients about the uses, risks, and benefits of chemotherapy.

PRINCIPLES OF SYSTEMIC THERAPY

A. CLINICAL USES OF CHEMOTHERAPY

Systemic therapy is indicated in the treatment of disseminated cancer when either cure or palliation is the goal. Additionally, chemotherapy may be used as part of a multimodality treatment plan in an effort to improve both local and distant control of the tumor. An understanding of the goals and limitations of systemic therapy in each of these settings is essential to its effective use.

1. Curative intent of metastatic disease—In considering the role of potentially curative chemotherapy in patients with metastatic disease, several factors must be taken into account. The first is the responsiveness of the tumor. Responsiveness is generally defined by the observed partial, complete, and overall responses. It is important to note that a complete response implies complete resolution of abnormal serum tumor markers, if any, and complete radiographic resolution of any abnormalities. This makes the assessment of neoplasms with frequent bony metastases such as prostate cancer, renal cell carcinoma, and transitional cell carcinoma difficult, as a persistently abnormal bone scan does not necessarily imply residual cancer. Patients in whom the only site of disease is bone generally must be considered non-assessable by conventional measures, and if available, intermediate markers of response (such as prostate-specific antigen [PSA]) are required.

If cure is the intent with systemic therapy, the relevant response criterion to consider is the percentage of patients achieving a complete response. This number is less than 10% in patients with metastatic renal cell carcinoma and hormone-refractory prostate cancer, 25% or less in patients with metastatic transitional cell carcinoma, and up to 80% in patients with metastatic germ cell malignancies. Under some circumstances, however (for example, in postchemotherapy residual masses in patients with germ cell carcinoma), an apparent partial response can be converted into a complete response with judicious resection (see Section A. 3.)

The second feature to consider in treating patients with potentially curative systemic therapy is the anticipated toxicity of such therapy. In general, higher levels of toxicity are acceptable if a cure can be achieved, although care must be exercised to avoid a “cure worse than the disease.” This is particularly true in the case of fairly toxic therapies such as interleukin-2 or bone marrow transplantation. These treatments can result in apparent cures of approximately 10% and 30%, respectively, of patients with metastatic renal cell carcinoma or refractory germ cell tumors (GCT). Patients undergoing these rigorous therapies must be carefully selected and must be as fully informed as possible about potential toxicities.

2. Treatment of patients with incurable metastatic cancer—When the goal of systemic therapy is palliation of symptoms rather than cure, the toxicity of the treatment to be offered must be balanced against the cancer-related symptoms the patient is experiencing, and in general, more toxic therapies are not indicated. Nonetheless, an understanding of the potential capabilities of systemic therapy must be understood because even in otherwise incurable disease there may be a role for systemic therapy if there is a likelihood that the patient’s life can be prolonged with its use. In addition, systemic chemotherapy can be associated with a control of pain, and an improvement in quality of life. This appears to be the case for both mitoxantrone and docetaxel in patients with metastatic hormone refractory prostate cancer.

3. Systemic therapy used in conjunction with surgery: adjuvant and neoadjuvant therapy—Systemic therapy administered after a patient has been rendered free of disease surgically is termed adjuvant therapy. Several
important criteria must be met if adjuvant therapy is to be used outside of a research setting. First, an assessment must be undertaken of known risk factors predictive of relapse or development of distant metastases. Patients at low risk of relapse generally should not receive adjuvant therapy because they are unlikely to derive a benefit and will be unnecessarily exposed to the toxicity of therapy. Second, the proposed therapy must have been shown to decrease the rate of relapse and increase the disease-free interval (and, it is hoped, survival) in a randomized, phase III trial. Finally, because patients who are being treated with adjuvant therapy are free of disease and presumably asymptomatic, toxicity must be kept at a minimum. This opens the way to a tailored approach in which patients with high-risk disease, as determined by pathologic review of the surgical specimen, are treated in order to decrease the risk of micrometastatic disease.

By contrast, neoadjuvant therapy is administered before definitive surgical resection. Here, the potential advantages include early therapy of micrometastatic disease and tumor debulking to allow a more complete resection. Patients with known metastatic disease generally do not exhibit high enough response rates to systemic therapy to warrant local surgery following chemotherapy, with the clear exception of patients with GCT. Whether or not patients with metastatic renal cell carcinoma who exhibit a partial response to systemic therapy may benefit from resection of residual masses is not known. As with adjuvant therapy, the proposed therapy must have been demonstrated to impact favorably on rate of relapse, disease-free interval, and survival in a randomized phase III trial.

**B. CHEMOTHERAPEUTIC AGENTS AND THEIR TOXICITY**

The usefulness of antineoplastic agents lies in their therapeutic index or preferential toxicity to malignant cells over normal, nonmalignant cells. The mechanism of action of most chemotherapeutic drugs is based on their toxicity to rapidly dividing cells. Thus, in general, malignancies that have relatively rapid growth, such as GCT, are relatively chemosensitive, whereas slower growing neoplasms such as renal cell carcinoma are less sensitive. Toxicity from chemotherapeutic agents is seen primarily in normal, nonmalignant cells that are also rapidly dividing, such as hematopoietic cells in the bone marrow, gastrointestinal mucosa, and hair follicles, and is manifested in cytopenias, mucositis, and alopecia. Other common toxicities observed with agents frequently used in the treatment of genitourinary malignancies include nephrotoxicity, neurotoxicity, hemorrhagic cystitis, pulmonary fibrosis, and cardiotoxicity. Table 19–1 summarizes the spectrum of activity and primary toxicities of commonly used chemotherapeutic agents.

The development of chemotherapy drug resistance remains an important clinical problem in the field of oncology. Malignant cells develop resistance in a variety of ways, including the induction of transport pumps, which

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**Table 19–1. Commonly Used Chemotherapeutic Agents in Urologic Oncology, and Their Toxicity.**

<table>
<thead>
<tr>
<th>Agent</th>
<th>Activity</th>
<th>Common Toxicities</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cisplatin</td>
<td>Bladder cancer, germ cell tumors, prostate cancer</td>
<td>Renal insufficiency, peripheral neuropathy, auditory toxicity, myelosuppression*</td>
</tr>
<tr>
<td>Carboplatin</td>
<td>Bladder cancer, germ cell tumors</td>
<td>Myelosuppression</td>
</tr>
<tr>
<td>Bleomycin</td>
<td>Germ cell tumors</td>
<td>Fever, chills, pulmonary fibrosis</td>
</tr>
<tr>
<td>Doxorubicin</td>
<td>Bladder cancer, prostate cancer</td>
<td>Myelosuppression, mucositis, cardiomyopathy</td>
</tr>
<tr>
<td>Etoposide (VP-16)</td>
<td>Germ cell tumors, prostate cancer</td>
<td>Myelosuppression</td>
</tr>
<tr>
<td>5-Fluorouracil</td>
<td>Renal cell carcinoma, bladder cancer, prostate cancer</td>
<td>Mucositis, diarrhea, myelosuppression</td>
</tr>
<tr>
<td>Floxuridine (FUdR)</td>
<td>Renal cell carcinoma</td>
<td>Mucositis, diarrhea</td>
</tr>
<tr>
<td>Methotrexate</td>
<td>Germ cell tumors, bladder cancer</td>
<td>Mucositis, myelosuppression, renal toxicity</td>
</tr>
<tr>
<td>Ifosfamide</td>
<td>Germ cell tumors</td>
<td>Myelosuppression, neurologic (CNS) toxicity, cystitis</td>
</tr>
<tr>
<td>Vinblastine</td>
<td>Renal cell carcinoma, bladder cancer, germ cell tumors, prostate cancer</td>
<td>Peripheral, autonomic neuropathy; myelosuppression</td>
</tr>
<tr>
<td>Estramustine</td>
<td>Prostate cancer</td>
<td>Nausea, thromboembolic events</td>
</tr>
<tr>
<td>Paclitaxel (Taxol)</td>
<td>Bladder cancer, germ cell tumors, prostate cancer</td>
<td>Myelosuppression, neuropathy</td>
</tr>
<tr>
<td>Docetaxel (Taxotere)</td>
<td>Bladder cancer, germ cell tumors, prostate cancer</td>
<td>Myelosuppression, neuropathy</td>
</tr>
<tr>
<td>Gemcitabine (Gemzar)</td>
<td>Bladder cancer</td>
<td>Myelosuppression</td>
</tr>
</tbody>
</table>

*Because of recent advances in the treatment of chemotherapy-induced nausea and vomiting, even the most emetogenic agents, such as cisplatin, have virtually no associated nausea and vomiting.

†In combination with estramustine.
actively pump the drug out of the cell and through increased activity of enzymes necessary to inactivate the particular chemotherapeutic agent. While there are several experimental methods of circumventing these mechanisms of drug resistance, one practical approach to this problem is the use of multiagent chemotherapy. Increased tumor cell killing is achieved by exposing neoplastic cells to multiple agents with different mechanisms of action. Furthermore, this approach allows the selection of agents with nonoverlapping toxicity profiles.

The use of increased dose intensity (higher doses of a drug administered over the same time period) as a means of overcoming drug resistance remains experimental in urologic malignancies with one clear exception. A subset of patients with otherwise incurable GCT appear to be curable with high-dose chemotherapy and autologous bone marrow transplant support (see the section Germ Cell Malignancies, following).

C. UNIQUE FEATURES OF GENITOURINARY MALIGNANCIES

The systemic therapy of urologic malignancies offers unique challenges to the practitioner. Renal insufficiency due to obstructive uropathy from local extension of the tumor or postsurgical or postradiotherapy changes is not infrequent and may alter antineoplastic drug clearance. In patients with renal cell carcinoma, previous nephrectomy also may impact on drug clearance. Furthermore, the common use of the nephrotoxic chemotherapeutic agent cisplatin in the treatment of urologic malignancies (most prominently, in bladder and testicular neoplasms) may further diminish renal function. Careful attention must be paid, therefore, to renal function throughout the course of systemic therapy, with appropriate dose adjustments made. Dosing adjustments also must be considered in patients who have undergone cystectomy because ileal conduits or neobladders have the capacity to resorb chemotherapeutic agents that are excreted in the urine in active form (most notably, methotrexate).

Frequent local extension in the pelvis presents additional unique problems. Patients with previous pelvic radiotherapy have markedly diminished bone marrow reserves, which may limit the use of myelosuppressive drugs. Furthermore, local pelvic relapses have the potential to be symptomatic and painful. Particularly in patients who have already received radiotherapy, systemic therapy may be important for palliation.

GERM CELL MALIGNANCIES

A. OVERVIEW

The evolution of therapy for GCT has been deliberate and thoughtful, and has resulted in cures of 80–85% of men with GCT, serving as a model for the treatment of curable cancers. Nonetheless, challenges in the management of GCT remain. Because of their young age, patients who have been cured are at risk of delayed, treatment-induced toxicity. Furthermore, an 80–85% cure rate also implies that 15–20% of patients with GCT will not be cured and ultimately will succumb to their disease. An understanding of staging and risk assessment is crucial if (1) patients with good risk features are not to be overtreated and exposed to undue toxic risks, and (2) patients with poor risk features are to receive adequate (curative) therapy.

The most common multiagent chemotherapy regimen for the treatment of GCT is a 3-drug combination consisting of bleomycin, etoposide, and cisplatin (BEP). The treatment is repeated every 21 days. One cycle consists of cisplatin 20 mg/m² IV day 1–5, etoposide 100 mg/m² IV day 1–5, and bleomycin, 30 units IV, day 2, 9, and 16. Frequently the first 5 days of treatment require hospitalization. The deletion of bleomycin from this regimen results in the PE regimen. The substitution of ifosfamide for bleomycin yield the VIP regime (UP-16, ifosfamide, platinum).

B. USE OF CHEMOTHERAPY FOR PATIENTS WITH STAGE I AND II DISEASE

The standard of care for patients with stage I GCT remains orchiectomy followed by retroperitoneal lymphadenectomy (nonseminoma), radiation therapy (seminoma), or in selected patients, careful surveillance. The use of chemotherapy in stage I GCT in lieu of lymphadenectomy or irradiation remains investigational despite encouraging early results.

Patients with stage II nonseminomatous microscopic disease identified at lymphadenectomy (stage IIA) or patients with low-volume clinical stage II disease (stage IIB) who have undergone retroperitoneal lymphadenectomy may benefit from 2 cycles of adjuvant PE or PEB chemotherapy. The use of adjuvant therapy results in a 96% long-term disease-free survival. While the relapse rate for patients who do not receive adjuvant therapy approaches 40%, the vast majority of relapsing patients can also be cured with either 3 or 4 cycles of chemotherapy, yielding an identical long-term survival rate. The decision about adjuvant chemotherapy after lymphadenectomy must be individualized. Patients at high risk for relapse may choose to undergo 2 cycles of chemotherapy at that point in order to avoid the possibility of 3–4 cycles in the future.

C. USE OF CHEMOTHERAPY IN PATIENTS WITH ADVANCED DISEASE

Patients with advanced GCT should be treated with systemic therapy after completion of their orchiectomy. This group includes some stage IIB nonseminomatous tumors and all stage IIC or higher tumors, both seminomas and nonseminomas. A variety of chemotherapy regimens will result in approximately 80% of patients with advanced GCT achieving a complete response and 70% achieving...
long-term apparent cures (good prognosis). By the same token, however, 20–30% of patients have a poor prognosis and will still ultimately die from their disease. Studies of pretreatment clinical characteristics have sought to identify prognostic features that can be prospectively used to segregate this diverse group of advanced GCT patients into poor- and good-prognostic subsets.

A common classification system has been developed by the International Germ Cell Cancer Collaborative Group (IGCCC). In this system, good-prognosis patients with nonseminomatous GCT have a testis or retroperitoneal primary tumor, no nonpulmonary visceral metastases, and low-serum tumor markers. Intermediate-prognosis patients are the same as good-prognosis patients but have intermediate serum tumor markers. Poor-prognosis patients have a mediastinal primary tumor or nonpulmonary visceral metastases (liver, bone, brain) or high levels of serum tumor markers. Five-year overall survival for the good-, intermediate-, and poor-prognosis categories with current regimens is 92%, 80%, and 48%, respectively. By definition, seminomas are never in the poor-prognosis category. Seminomas are segregated into good-prognosis cases (any primary site, but no nonpulmonary visceral metastases), with an 86% 5-year survival, and intermediate-prognosis cases (any primary site but with the presence of nonpulmonary visceral metastases), with a 72% 5-year survival.

Because it is not likely that the extraordinarily high cure rate for good-prognosis patients can be improved upon, most efforts in the treatment of these patients have been aimed at optimizing treatment with less toxic regimens that will have equal efficacy. Trials evaluating (1) the elimination of bleomycin, (2) a reduction in the number of chemotherapy cycles administered, or (3) the substitution of carboplatin for cisplatin have been undertaken.

The outlook for poor-prognosis patients is grim, with only 38–62% of patients achieving a complete response. Thus, whereas the major concern in good-prognosis patients has been the reduction of toxicity, the major objective of clinical investigation in poor-prognosis patients has been to improve efficacy, with less concern for reducing toxicity. Clinical trials in poor-prognosis patients have by and large relied on one or both of two approaches. The first has been to exploit agents that have been demonstrated to be efficacious in the salvage setting, and the second has been to evaluate the role of dose escalation.

Currently acceptable regimens for good-prognosis patients are fairly well defined and include 3 cycles of PEB or 4 cycles of PE. By contrast, optimal therapy for poor-prognosis patients continues to be investigational. Four cycles of PEB or 4 cycles of VIP (are appropriate options.

D. Adjunctive Surgery and “Salvage” Therapy

Postchemotherapy adjunctive surgery must be integrated into the treatment plan of patients with advanced GCT. Between 10% and 20% of patients with nonseminomato-
B. METASTATIC DISEASE

The development of successful therapy of metastatic bladder TCC has been based on the use of cisplatin. Until recently, two common cisplatin-based regimens are in wide use: (1) cisplatin, methotrexate, and vinblastine (CMV) and (2) the same drugs in a slightly different schedule and dose along with doxorubicin (Adriamycin), in a regimen known as MVAC. These regimens result in overall response rates of approximately 50–60% and complete remission rates in the 20–35% range. Median overall survival for patients with metastatic disease treated with these regimens is in the 8- to 14-month range. Despite early promise, however, long-term survival after MVAC or CMV remains in the single digits. Both CMV and MVAC are intensive regimens, with myelosuppression occurring commonly. The use of hematopoietic growth factors has made it easier to administer full doses on schedule, although this improvement in dose intensity does not appear to translate into a clinical benefit.

More recently, the combination of gemcitabine and cisplatin has been compared to MVAC. This new regimen is far less toxic, is better tolerated, and appears to be equivalent in efficacy to MVAC. As a consequence, gemcitabine/cisplatin can be considered the new standard of care for the treatment of advanced TCC. However, it should be noted that the gemcitabine/cisplatin regimen has been tested in a randomized study only in patients with metastatic disease, and its utility as an adjuvant or neoadjuvant has not been tested. For patients with impaired renal function, agents such as carboplatin and paclitaxel have been utilized.

RENAL CELL CARCINOMA

The treatment of metastatic renal cell carcinoma with chemotherapy remains largely unsatisfactory. The general lack of active agents and the excessive toxicity of many of the agents that exhibit some activity have contributed to the absence of adjuvant or neoadjuvant trials. The only such trials used adjuvant interferon-alpha for patients considered at high risk for relapse after nephrectomy and failed to show an advantage of the adjuvant therapy. Nephrectomy prior to systemic therapy is recommended, particularly in patients in whom the bulk of disease is in the renal mass, and who have a good performance status.

Metastatic renal cell carcinoma is relatively resistant to chemotherapy. The fluoropyrimidines 5-fluorouracil, 5-fluorouracil, and capecitabine have modest activity, as does gemcitabine, with response proportions of 10–15% reported. Renal cell carcinoma is one of very few neoplasms that clearly are responsive to biologic response modifiers. The utility of biologic response modifiers and anti-angiogenic agents in renal cell carcinoma is discussed elsewhere in Chapter 20. In general, these agents are used prior to using chemotherapy.

HORMONE-REFRACTORY PROSTATE CANCER

The systemic therapy of patients with metastatic prostate cancer in whom hormonal therapy has failed generally consists of secondary hormonal manipulations followed by chemotherapy. Approximately 15% of patients who have had progressive disease despite therapy with combined androgen blockade will have a fall in PSA when their antiandrogen is discontinued. This maneuver is mandated, therefore, before initiating other systemic therapy. Furthermore, second-line hormonal maneuvers such as adrenal androgen deprivation with ketoconazole, estrogens, or secondary antiandrogens such as nilutamide clearly have activity and, particularly in asymptomatic patients, should be considered. As noted previously, the evaluation of responses in patients with bone disease only is difficult at best. The use of the PSA in this setting has been fairly extensively evaluated, and it appears to be a reasonable intermediate endpoint. Thus, a decline in PSA of 35–50% appears to be predictive of longer survival for these patients.

Several agents or combinations of agents show promise in the therapy of HRPC. Not only can a significant
decline in PSA be demonstrated in some patients, but also objective responses in patients with soft-tissue disease have been observed. Furthermore, considerable palliation of pain is often possible with chemotherapy in patients in whom narcotics or corticosteroids have failed and for whom palliative irradiation is not an option.

Mitoxantrone is approved in combination with prednisone for the treatment of progressive, symptomatic HRPC. Twenty-nine percent of those treated with the combination experienced decreased pain, compared with 12% receiving prednisone alone. In addition, there were greater improvements in quality-of-life measures. The toxicity of the treatment was mild in both groups; fewer than 2% of patients had infectious episodes. Median survival for both groups was approximately 1 year. Mitoxantrone has modest albeit definable activity in HRPC, although it probably does not significantly prolong survival.

Until recently, chemotherapy for prostate cancer was considered ineffective in prolonging survival. However, the results of 2 phase III trials have established docetaxel-based chemotherapy as the standard-of-care for first-line treatment of metastatic HRPC. SWOG 9916 compared the combination of docetaxel/estramustine with mitoxantrone/prednisone, while Tax 327, a trial conducted by Aventis, tested 2 schedules (weekly and q 3 week) of the combination of docetaxel/prednisone versus mitoxantrone/prednisone. The q 3 week docetaxel regimens in each one of these trials demonstrated a modest but statistically significant (2 month) survival benefit over mitoxantrone/prednisone. The median survival with docetaxel was 18–19 months. In Tax 327, the difference in survival between weekly docetaxel and mitoxantrone did not reach statistical significance. While docetaxel/prednisone was not directly compared with docetaxel/estramustine, the overall survival of the 2 q 3 week docetaxel-based regimens was similar, whether prednisone or estramustine was added, and the use of estramustine was associated with greater toxicity. Thus, every 3-week docetaxel/prednisone has emerged as the FDA-approved, first-line regimen for HRPC. Future directions and the subject of ongoing trials include (1) exploring the addition of novel agents to the docetaxel/prednisone backbone, and (2) using docetaxel in earlier stages of prostate cancer, such as neoadjuvantly prior to prostatectomy, or together with radiation therapy, or for patients with a climbing PSA after definitive local therapy.

REFERENCES


BLADDER CARCINOMAS

Incidence
Bladder cancer is the second most common cancer of the genitourinary tract. It accounts for 7% of new cancer cases in men and 2% of new cancer cases in women. The incidence is higher in whites than in African Americans, and there is a positive social class gradient for bladder cancer in both sexes. The average age at diagnosis is 65 years. At that time, approximately 75% of bladder cancers are localized to the bladder; 25% have spread to regional lymph nodes or distant sites.

Risk Factors & Pathogenesis
Cigarette smoking accounts for 50% of cases in men and 31% in women (Wynder and Goldsmith, 1977). In general, smokers have approximately a twofold increased risk of bladder cancer than nonsmokers, and the association appears to be dose related (Thompson and Fair, 1990). The causative agents are thought to be alpha- and beta-naphthylamine, which are secreted into the urine of smokers.

Occupational exposure accounts for 15–35% of cases in men and 1–6% in women (Matanoski and Elliott, 1981). Workers in the chemical, dye, rubber, petroleum, leather, and printing industries are at increased risk. Specific occupational carcinogens include benzidine, beta-naphthylamine, and 4-aminobiphenyl, and the latency period between exposure and tumor development may be prolonged. Patients who have received cyclophosphamide (Cytoxan) for the management of various malignant diseases are also at increased risk (Fairchild et al, 1979). Ingestion of artificial sweeteners has been proposed to be a risk factor, but several studies have failed to confirm any association (Elcock and Morgan, 1993). Physical trauma to the urothelium induced by infection, instrumentation, and calculi increases the risk of malignancy (Hicks, 1982).

The exact genetic events leading to the development of bladder cancer are unknown, but they are likely to be multiple and may involve the activation of oncogenes and inactivation or loss of tumor suppressor genes (Olumi et al, 1990). Loss of genetic material on chromosome 9 appears to be a consistent finding in patients with both low-grade, low-stage and high-grade, high-stage disease (Tsai et al, 1990; Miyao et al, 1993), which suggests that this may be an early event in bladder cancer development. Loss of chromosome 9 in multiple tumors from an individual patient supports the concept that genetic changes in bladder cancer represent a “field defect” that may occur throughout the urothelium. More recent studies examining p53 tumor suppressor gene mutations in primary, recurrent, and upper tract tumors suggest that these tumors can have a single clonal origin (Dalbagni et al, 2001; Sidransky et al, 1991). Additional genetic changes have been described that are specific for invasive bladder tumors. Chromosome 11p, which contains the c-Ha-ras proto-oncogene, is deleted in approximately 40% of bladder cancers (Olumi et al, 1990). Increased expression of the c-Ha-ras protein product, p21, has been detected in dysplastic and high-grade tumors but not in low-grade bladder cancers. Deletions of chromosome 17p have also been detected in over 60% of all invasive bladder cancers, but 17p deletions have not been described in superficial tumors. This finding is noteworthy because the p53 tumor suppressor gene maps to chromosome 17p. p53 alterations represent the most commonly identified genetic abnormality in human cancers, making deletion of this chromosome an important finding in muscle invasive bladder cancer.

Staging
Currently, the most commonly used staging system allows for a precise and simultaneous description of the primary tumor stage (T stage), the status of lymph nodes (N stage), and metastatic sites (M stage) (American Joint Committee on Cancer, 1997). The T staging system is depicted in Figure 20–1. Nodal (N) stage is defined as Nx – cannot be assessed, N0 – no nodal metastases, N1 – single node <2 cm involved, N2 – single node involved 2–5 cm in size or multiple nodes none >5 cm, N3 – one or more nodes >5 cm in size involved. Metastases (M) stage is defined as Mx – cannot be defined, M0 – no distant metastases, M1 –
distant metastases present. Staging errors exist when one compares the clinical stage (that based on physical examination and imaging) with the pathologic stage (that based on removal of the bladder and regional lymph nodes). Overstaging is relatively uncommon, but clinical understaging may occur in up to 53% of patients (Skinner, 1982; Dutta et al, 2001).

Histopathology

Ninety-eight percent of all bladder cancers are epithelial malignancies, with most being transitional cell carcinomas (TCCs).

**Figure 20–1.** Staging of bladder cancer.

A. NORMAL UROTHELIELUM

The normal urothelium is composed of 3–7 layers of transitional cell epithelium resting on a basement membrane composed of extracellular matrix (collagen, adhesive glycoproteins, glycosaminoglycans) (Figure 20–2A). The epithelial cells vary in appearance: The basal cells are actively proliferating cells resting on the basement membrane; the luminal cells, perhaps the most important feature of normal bladder epithelium, are larger umbrella-like cells that are bound together by tight junctions. Beyond the basement membrane is loose connective tissue, the lamina propria, in which occasionally smooth-muscle fibers can be
identified. These fibers should be distinguished from deeper, more extensive muscle elements defining the true muscularis propria. The muscle wall of the bladder is composed of muscle bundles coursing in multiple directions. As these converge near the bladder neck, 3 layers can be recognized: inner and outer longitudinally oriented layers and a middle circularly oriented layer.

B. PAPILLOMA

The World Health Organization recognizes a papilloma as a papillary tumor with a fine fibrovascular stalk supporting an epithelial layer of transitional cells with normal thickness and cytology (Epstein et al, 1998). Papillomas are a rare benign condition usually occurring in younger patients.

C. TRANSITIONAL CELL CARCINOMA

Approximately 90% of all bladder cancers are TCCs. These tumors most commonly appear as papillary, exophytic lesions (Figure 20–2B); less commonly, they may be sessile or ulcerated. Whereas the former group is usually superficial in nature, sessile growths are often invasive. Carcinoma in situ (CIS) is recognizable as flat, anaplastic epithelium. The urothelium lacks the normal cellular polarity, and cells contain large, irregular hyperchromatic nuclei with prominent nucleoli (Figure 20–2C).

D. NONTRANSITIONAL CELL CARCINOMAS

1. Adenocarcinoma—Adenocarcinomas account for <2% of all bladder cancers. Primary adenocarcinomas of the bladder may be preceded by cystitis and metaplasia. Histologically, adenocarcinomas are mucus-secreting and may have glandular, colloid, or signet-ring patterns. Whereas primary adenocarcinomas often arise along the floor of the bladder, adenocarcinomas arising from the urachus occur at the dome. Both tumor types are often localized at the time of diagnosis, but muscle invasion is usually present. Five-year survival is usually <40%, despite aggressive surgical management (Kramer et al, 1979; Abenoza, Manivel, and Fraley, 1987; Bernstein et al, 1988).

2. Squamous cell carcinoma—Squamous cell carcinoma accounts for between 5% and 10% of all bladder cancers in the United States and is often associated here with a history of chronic infection, vesical calculi, or chronic catheter use. It may also be associated with bilharzial infection owing to *Schistosoma haematobium*, because squamous cell carcinoma accounts for approximately 60% of all bladder cancers in Egypt, parts of Africa, and the Middle East, where this infection is prevalent (El-Bolkainy et al, 1981). These tumors are often nodular and invasive at the time of diagnosis. Histologically they appear as poorly differentiated neoplasms composed of polygonal cells with characteristic intercellular bridges. Keratinizing epithelium is present, although often in small amounts.
3. **Undifferentiated carcinomas**—Undifferentiated bladder carcinomas, which are rare (accounting for <2%), have no mature epithelial elements. Very undifferentiated tumors with neuroendocrine features and small cell carcinomas tend to be aggressive and present with metastases (Quek et al, 2005; Choong et al, 2005).

4. **Mixed carcinoma**—Mixed carcinomas constitute 4–6% of all bladder cancers and are composed of a combination of transitional, glandular, squamous, or undifferentiated patterns. The most common type comprises transitional and squamous cell elements (Murphy, 1989). Most mixed carcinomas are large and infiltrating at the time of diagnosis.

**E. RARE EPITHELIAL & NONEPITHELIAL CANCERS**

Rare epithelial carcinomas identified in the bladder include villous adenomas, carcinoid tumors, carcinosarcomas, and melanomas. Rare nonepithelial cancers of the urinary bladder include pheochromocytomas, lymphomas, choriocarcinomas, and various mesenchymal tumors (hemangioma, osteogenic sarcoma, and myosarcoma) (Murphy, 1989). Cancers of the prostate, cervix, and rectum may involve the bladder by direct extension. The most common tumors metastatic to the bladder include (in order of incidence) melanoma, lymphoma, stomach, breast, kidney, lung and liver (Murphy, 1989; Goldstein, 1967, Franks, 1999).

**Clinical Findings**

**A. SYMPTOMS**

Hematuria is the presenting symptom in 85–90% of patients with bladder cancer. It may be gross or microscopic, intermittent rather than constant. In a smaller percentage of patients, it is accompanied by symptoms of vesical irritability: frequency, urgency, and dysuria. Irritative voiding symptoms seem to be more common in patients with diffuse CIS. Symptoms of advanced disease include bone pain from bone metastases or flank pain from retroperitoneal metastases or ureteral obstruction.

**B. SIGNS**

Patients with large-volume or invasive tumors may be found to have bladder wall thickening or a palpable mass—findings that may be detected on a careful bimanual examination under anesthesia. If the bladder is not mobile, that suggests fixation of tumor to adjacent structures by direct invasion.

Hepatomegaly and supraclavicular lymphadenopathy are signs of metastatic disease. Lymphedema from occlusive pelvic lymphadenopathy may be seen occasionally. On rare occasions, metastases can occur in unusual sites such as the skin presenting as painful nodules with ulceration (Block et al, 2006).

**C. LABORATORY FINDINGS**

1. **Routine testing**—The most common laboratory abnormality is hematuria. It may be accompanied by pyuria, which on occasion may result from concomitant urinary tract infection. Azotemia may be noted in patients with ureteral occlusion owing to the primary bladder tumor or lymphadenopathy. Anemia may be a presenting symptom owing to chronic blood loss, or replacement of the bone marrow with metastatic disease.

2. **Urinary cytology**—Exfoliated cells from both normal and neoplastic urothelium can be readily identified in voided urine. Larger quantities of cells can be obtained by gently irrigating the bladder with isotonic saline solution through a catheter or cystoscope (barbotage). Cytologic examination of exfoliated cells may be especially useful in detecting cancer in symptomatic patients and assessing response to treatment. Detection rates are high for tumors of high grade and stage as well as CIS but not as impressive for low grade superficial tumors.

3. **Other markers**—Several new tests have been developed in order to overcome the shortcomings of urinary cytology such as the low sensitivity for low-grade superficial tumors and inter-observer variability. Commercially available tests include, the BTA test (Bard Urological, Covington, GA), the BTA stat test (Bard Diagnostic Sciences, Inc, Redmond, WA), the BTA TRAK assay (Bard Diagnostic Sciences, Inc), determination of urinary nuclear matrix protein (NMP22; Matriitech Inc, Newton, MA), Immunocyt (Diagnocure, Montreal, Canada) and UroVysion (Abbott Labs, Chicago, IL). These tests can detect cancer specific proteins in urine (BTA/NMP22) or augment cytology by identifying cell surface or cytogenetic markers in the nucleus. Other tests under investigation include identification of the Lewis X antigen on exfoliated urothelial cells, and the determination of telomerase activity in exfoliated cells. Several studies have examined the performance of these voided urinary markers for the detection and follow-up of patients with bladder cancer (summarized in Grossfeld and Carroll, 1998; Grossfeld et al, 2001; Konety and Getzenberg, 2001) (Table 20–1).

These tests have been demonstrated to enhance detection of bladder cancer when used either individually or in combination with cytology. They have been used to detect both new index tumors as well as recurrent tumors. Some of the protein markers lack the specificity of cytology thereby hampering their widespread use. Such exfoliated markers can be expected to play an important and increasing role in the initial evaluation and follow-up of patients with bladder cancer in the future.

**D. IMAGING**

Although bladder cancers may be detected by various imaging techniques, their presence is confirmed by cystoscopy and biopsy. Imaging is therefore used to evaluate the
upper urinary tract and, when infiltrating bladder tumors are detected, to assess the depth of muscle wall infiltration and the presence of regional or distant metastases. Intravenous urography remains one of the most common imaging tests for the evaluation of hematuria. However, intravenous pyelography is increasingly being replaced by computed tomography (CT) urography, which is more accurate, for evaluation of the entire abdominal cavity, renal parenchyma, and ureters in patients with hematuria (Gray Sears et al, 2002). Bladder tumors may be recognized as pedunculated, radiolucent filling defects projecting into the lumen (Figure 20–3); nonpapillary, infiltrating tumors may result in fixation or flattening of the bladder wall. Hydronephrosis from ureteral obstruction is usually associated with deeply infiltrating lesions and poor outcome after treatment (Haleblian et al, 1998).

Superficial (Ta, Tis) bladder cancers staged with a properly performed TUR and examination under anesthesia do not require additional imaging of the bladder or pelvic organs. However, higher stage lesions are often understaged, and the addition of imaging may be useful. Both CT and magnetic resonance imaging (MRI) (Figure 20–4) have been used to characterize the extent of bladder wall invasion and detect enlarged pelvic lymph nodes, with overall staging accuracy ranging from 40% to 85% for CT and from 50% to 90% for MRI (Fisher, Hricak, and Tanngho, 1985; Wood et al, 1988). Both techniques rely on size criteria for the detection of lymphadenopathy: Lymph nodes >1 cm are thought to be suggestive of metastases; unfortunately, small-volume pelvic lymph node metastases are often missed. Because invasive bladder cancers may metastasize to the lung or bones, staging of advanced lesions is completed with chest x-ray and radionuclide bone scan. Bone scans can be avoided if the serum alkaline phosphatase is normal (Berger, 1981).

### E. CYSTOUREROSCOPY & TUMOR RESECTION

The diagnosis and initial staging of bladder cancer is made by cystoscopy and transurethral resection (TUR). Cystoscopy can be done with either flexible or rigid instruments, although the former is associated with less discomfort and only requires local anesthesia. Superficial, low-grade tumors usually appear as single or multiple papillary lesions. Higher grade lesions are larger and sessile. CIS may appear as flat areas of erythema and mucosal irregularity. Use of fluorescent cystoscopy with blue light can enhance the ability to detect lesions by as much as 20% (Jocham, 2005). In this procedure, hematoporphyrin derivatives that accumulate preferentially in cancer cells are instilled into the bladder and fluorescence incited using a blue light. Cancer cells with accumulated porphyrin such as 5-aminolevulenic acid or hexaminolevulinate (HAL) are detected as glowing red under the fluorescent light (Loidl, 2005).

Once a tumor is visualized or suspected, the patient is scheduled for examination under anesthesia and TUR or biopsy of the suspicious lesion. The objectives are tumor diagnosis, assessment of the degree of bladder wall invasion (staging), and complete excision of the low-stage lesions amenable to such treatment.

Patients are placed in the lithotomy position. A careful bimanual examination is performed. The presence of any palpable mass and mobility of the bladder are noted, along with any degree of fixation to contiguous structures. Cystoscopy is repeated with one or more lenses (30° and 70°) that permit complete visualization of the entire bladder surface. A resectoscope is then placed into the bladder, and

### Table 20–1. Exfoliated Markers for the Detection of Bladder Cancer.

<table>
<thead>
<tr>
<th>Marker</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>PPV (%)</th>
<th>NPV (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cytology</td>
<td>35–61</td>
<td>93–100</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>BTA</td>
<td>28–100</td>
<td>40–96</td>
<td>33–80</td>
<td>52–94</td>
</tr>
<tr>
<td>NMP22</td>
<td>47–100</td>
<td>61–99</td>
<td>29–65</td>
<td>60–100</td>
</tr>
<tr>
<td>BTA stat</td>
<td>57–83</td>
<td>33–95</td>
<td>20–56</td>
<td>70–95</td>
</tr>
<tr>
<td>BTA TRAK</td>
<td>62–78</td>
<td>51–98</td>
<td>62</td>
<td>73</td>
</tr>
<tr>
<td>Lewis X antigen</td>
<td>80–97</td>
<td>73–86</td>
<td>72–81</td>
<td>83–98</td>
</tr>
<tr>
<td>Telomerase</td>
<td>62–80</td>
<td>60–99</td>
<td>84</td>
<td>89</td>
</tr>
<tr>
<td>FDP</td>
<td>33–83</td>
<td>66–91</td>
<td>79</td>
<td>78</td>
</tr>
<tr>
<td>Cytokeratin 20</td>
<td>91</td>
<td>85</td>
<td>95</td>
<td>76</td>
</tr>
<tr>
<td>Quantiant</td>
<td>45–59</td>
<td>71–93</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Hyaluronic acid</td>
<td>92</td>
<td>93</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Hyaluronidase</td>
<td>100</td>
<td>89</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>BLCA-4</td>
<td>96</td>
<td>100</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Flow cytometry</td>
<td>45–72</td>
<td>80–87</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

FDP, fibrinogen/fibrin degradation products; PPV, positive predictive value; NPV, negative predictive value.
visible tumors are removed by electrocautery. Suspicious areas may be biopsied with cup biopsy forceps and the areas may be cauterized with an electrode. Some clinicians routinely perform random bladder biopsies of normal-appearing urothelium both close to and remote from the tumor. The value of random bladder biopsies is controversial. Detection of CIS on these biopsies can alter treatment though more recent studies suggest that only 1.5% of low-risk and 3.5% of high-risk patients may have tumor detected on such biopsies. (van der Meijden, 1999; May et al, 2003). Findings of the random biopsy can alter treatment in up to 7% of patients (May et al, 2003).

**Natural History & Selection of Treatment**

**A. STANDARD HISTOPATHOLOGICAL ASSESSMENT**

The natural history of bladder cancers is defined by 2 separate but related processes: tumor recurrence and progression. Progression, including metastasis, represents the greater biologic risk. However, recurrence, even without progression, represents substantial patient morbidity in that it requires periodic reevaluation (cytology, cystoscopy, etc), repeat endoscopic ablation, and often intravesical chemotherapy (which may be costly, uncomfortable, and associated with complications). Treatment decisions are based on tumor stage and grade. Staging is performed using the tumor, node, metastasis (TNM) staging system (Figure 20–1; Table 20-2) while grading has changed from the Ash-Broder system (I–III or I–IV). The new WHO-ISUP system segregates tumors into papillary urothelial neoplasm of low malignant potential (PUNLMP), low grade or high grade.

At initial presentation, approximately 50–70% of bladder tumors are superficial—stage Tis or Ta. Invasion into the lamina propria or muscle wall is identified in a smaller number of patients, approximately 28% and 24%, respectively; regional or distant metastases are found in approxi-
approximately 25%. Unfortunately, 80% of patients with invasive or metastatic disease have no previous history of bladder cancer (Kaye and Lange, 1982). Approximately 43% of tumors are classified as grade I, 25% as grade II, and 32% as grade III (Gilbert et al, 1978). There are strong correlations between tumor grade and stage and tumor recurrence, progression, and survival (Frazier et al, 1993). Patients with low-stage, low-grade disease have a low risk (<5%) of progression to invasive disease, while as many as 40% of patients with low-stage but high-grade disease will progress with extended follow-up (Herr, 2000). Disease-free survival is excellent for patients with pathologically confirmed superficial disease (pT0, pT1, pTIS, 80–88%). However, it falls for patients with pT2 (53–80%), pT3 (39–68%), and pT4 (25–40%) tumors (Stein et al, 2001; Frazier et al, 1993; Thrasher et al, 1994)—owing to the greater likelihood of metastasis in tumors of higher stage. Whereas lymph node metastases are uncommon (5%) in tumors of low stage, they are increasingly more common in higher stage tumors: 10–30% for pT3A, 31–46% for pT3B, and 35–64% for pT4 (Stein et al, 2001; Frazier et al, 1993). In patients with organ-confined disease, the presence of pelvic lymph node metastases appears to be the most important prognostic factor (Vieweg et al, 1999). The presence of lymphovascular invasion even in those with node negative disease may portend a worse prognosis (Lotan et al, 2005).

Although metastasis is less common with superficial bladder cancers, such tumors may progress; most recur and Table 20–2. Initial Treatment Options for Bladder Cancers.

<table>
<thead>
<tr>
<th>Cancer Stage</th>
<th>Initial Treatment Options</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tis</td>
<td>Complete TUR followed by intravesical BCG</td>
</tr>
<tr>
<td>Ta (single, low-to-moderate grade, not recurrent)</td>
<td>Complete TUR</td>
</tr>
<tr>
<td>Ta (large, multiple, high-grade, or recurrent)</td>
<td>Complete TUR followed by intravesical chemo- or immunotherapy</td>
</tr>
<tr>
<td>T1</td>
<td>Complete TUR followed by intravesical chemo- or immunotherapy</td>
</tr>
<tr>
<td>T2–T4</td>
<td>Radical cystectomy</td>
</tr>
<tr>
<td>Any T, N+, M+</td>
<td>Neoadjuvant chemotherapy followed by radical cystectomy</td>
</tr>
<tr>
<td></td>
<td>Radical cystectomy followed by adjuvant chemotherapy</td>
</tr>
<tr>
<td></td>
<td>Neoadjuvant chemotherapy followed by concomitant chemotherapy and irradiation</td>
</tr>
<tr>
<td></td>
<td>Systemic chemotherapy followed by selective surgery or irradiation</td>
</tr>
</tbody>
</table>

TUR, transurethral resection.
require additional treatment. Tumor progression occurs in <6% of patients with Ta disease, but in up to 53% of those with T1 disease, with or without concomitant CIS (Heney et al, 1983; Cookson et al, 1997). Tumor progression occurs in 10–20% of patients with grade I tumors, 19–37% with grade II tumors, and 33–64% with grade III tumors (Torti et al, 1987; Lutzeyer, Rubben, and Dahm, 1982). Using the more recent grading system, progression is observed in 5% of those with low grade tumors, 15–40% with high grade tumors while PUNLMPs almost never demonstrate any risk of progression (Epstein et al, 1998).

Tumor recurrence is related to history of disease and grade, number, and size of the tumor. It is more common in the first 12–24 months after diagnosis (but can become manifest many years later), and patients with one recurrence are more likely to have another. Patients with T1, multiple (>4), large (>3), or high-grade tumors are at greater risk, as are those with either CIS or severe dysplasia in normal-appearing urothelium remote from the tumor site (Heney et al, 1983; Wolf, Olsen, and Hojgaard, 1985). Tumors can be stratified into low- and high-risk categories based on these criteria and this can be used to guide management decisions.

B. MOLECULAR MARKERS

Conventional histopathologic analysis of bladder tumors, including determination of tumor grade and stage, may not reliably predict the behavior of many bladder cancers. Assessment of molecular markers of disease, with immunohistochemical methods, in biopsy specimens, or in cystectomy specimens can yield useful prognostic information.

Tumor growth and metastasis require the growth of new blood vessels, through angiogenesis. Angiogenic stimulators, such as the fibroblastic growth factors and vascular endothelial growth factor, and angiogenic inhibitors, such as thrombospondin-1 and angiostatin regulate angiogenesis. Immunohistochemical quantification of angiogenesis in a given tumor by measuring microvessel density is a useful prognostic indicator for a variety of human malignancies, including bladder cancer. In bladder cancer, microvessel density has been associated with lymph node metastases, disease progression, and overall survival in patients with invasive bladder cancer treated with radical cystectomy (Dickinson et al, 1994; Jaeger et al, 1995; Bochner et al, 1997). The p53 gene is a tumor suppressor gene that plays a key role in the regulation of the cell cycle. When DNA damage occurs, the level of p53 protein increases, causing cell cycle arrest and repair of DNA. Mutations in the p53 gene result in the production of an abnormal protein product, allowing cells with damaged DNA to continue through the cell cycle. The altered p53 protein has a prolonged half-life compared with the wild-type protein, allowing for its detection by immunohistochemical techniques. Patients with altered p53 expression (indicating possible mutation of the p53 gene) appear to have an increased risk for disease recurrence and a decreased overall survival when compared with patients with normal p53 expression (Esrig et al, 1995). Cancers that are p53 positive are associated with recurrence rates of 62% for pT1, 56% for pT2, and 80% for pT3, compared with 7%, 12%, and 11%, respectively, for cancers without p53 reactivity.

Alteration of the retinoblastoma (Rb) gene, a tumor suppressor gene, is associated with high-grade, high-stage bladder cancers. In addition, Rb alteration appears to be significantly associated with decreased overall survival in such patients (Cordon-Cardo et al, 1992; Logothetis et al, 1992). Studies in which both p53 and Rb have been examined in patients with invasive bladder cancer suggest that bladder tumors with alterations in both genes have a poorer prognosis and decreased overall survival when compared with tumors with wild-type p53 and Rb.

Assessment of other markers that may correlate with outcome in patients with bladder cancer includes that of tumor growth fraction (proliferative index) and cellular adhesion molecule expression (E-cadherin) (Okamura et al, 1990; Lipponen and Eskelinen, 1995).

C. TREATMENT SELECTION

Patients with superficial bladder cancers can be treated with TUR followed by selective intravesical chemotherapy or immunotherapy. Patients with initial low-grade small tumors are at low risk of progression and may be treated by TUR alone followed by surveillance or intravesical chemotherapy. Patients with T1, high-grade, multiple, large, recurrent tumors or those associated with CIS on bladder biopsies are at a higher risk of progression and recurrence and should be considered candidates for intravesical chemotherapy or immunotherapy after complete and careful TUR. A second resection of the same area may be required to accurately stage disease and determine treatment (Herr et al, 1999; Grimm, 2003). Repeat resections may also enhance response to intravesical therapy (Herr, 2005). Management of T1 tumors is somewhat controversial; some clinicians advise radical cystectomy, especially for grade III or high grade lesions, which are associated with a high rate of progression. However, progression rates can be reduced by intravesical immunotherapy (Herr et al, 1989; Cookson and Sarosdy, 1992). Recurrence of T1 disease after a trial of intravesical therapy warrants more aggressive therapy (Herr, 1991; Herr and Sogani, 2001).

Patients with more invasive, but still localized, tumors (T2, T3) are candidates for more aggressive local treatment, including partial or radical cystectomy, or a combination of radiation and systemic chemotherapy. Radical TUR alone may be a viable option in select patients with T2 disease particularly if no tumor is found on repeat resection since 10-year survival rates as high as 83% can be achieved (Herr, 2001). However, this approach must be used with caution since there is a substantial risk of leaving
residual disease behind (Solsona et al, 1998). Superficial ductal or acinar in situ carcinoma of the prostatic urethra, not invading the basement membrane or prostatic stroma, may be treated with TUR and intravesical chemotherapy or immunotherapy rather than cystectomy. However, patients with more extensive involvement of the prostatic urethra by TCC, or recurrence after conservative therapy, require more aggressive therapy. Patients with unresectable local tumors (T4B) are candidates for systemic chemotherapy, followed by surgery (or possibly irradiation). Patients with either local or distant metastases should receive systemic chemotherapy followed by the selective use of either irradiation or surgery, depending on the response.

Treatment

A. INTRAVESICAL CHEMOTHERAPY

Immunotherapeutic or chemotherapeutic agents can be instilled into the bladder directly via catheter, thereby avoiding the morbidity of systemic administration in most cases. Intravesical therapy can have a prophylactic or therapeutic objective, either to reduce recurrence in patients whose tumors have been completely resected. Intravesical chemotherapy is used in 2 settings. When instilled immediately following TUR, it acts prophylactically to reduce tumor cell implantation (Solsona et al, 1999). It can also be used therapeutically to reduce risk of recurrence and progression particularly for low-risk superficial tumors. Therefore, intravesical chemotherapy or immunotherapy may be delivered in 3 different fashions to achieve individual goals (Table 20–3). Considerable experience has been gained, but comparison of different agents is difficult owing to the paucity of randomized trials and variations in dose, contact time, patient population, and intervals between treatments. Most agents are administered weekly for 6 weeks except when being used prophylactically where a single dose is administered immediately following TUR. Maintenance therapy (ie, monthly or bimonthly intravesical therapy) may decrease recurrence rates further. Although local toxicity is relatively common—primarily irritative voiding symptoms—systemic toxicity is rare because of the limited absorption of drugs across the lumen of the bladder. Severe systemic complications can be avoided by not administering intravesical chemotherapy in patients with gross hematuria. Efficacy may be improved by increasing contact time and drug concentration (ie, by restricting fluid intake before administration, asking the patient to lie in different positions during treatment, avoiding instillation of air during drug administration, and requiring the patient to avoid urinating for 1–2 hours thereafter). The most common agents in the United States are mitomycin C, thiotepa, and Bacillus Calmette-Guérin (BCG). Patients in whom treatment with one agent fails may respond to another.

1. Mitomycin C—Mitomycin C is an antitumor, antibiotic, alkylating agent that inhibits DNA synthesis. With a molecular weight of 329, systemic absorption is minimal. The usual dose is 40 mg in 40 cc of sterile water or saline given once a week for 6 weeks. The same dose is utilized for a single prophylactic instillation. Between 39% and 78% of patients with residual tumor experience, a complete response to intravesical mitomycin C (Kowalkowski and Lamm, 1988), and recurrence is reduced in 2–33% after complete TUR (Herr, Laudone, and Whitmore, 1987). Side effects are noted in 10–43% of patients and consist largely of irritative voiding symptoms including urinary frequency, urgency, and dysuria. Unique to this drug is the appearance of a rash on the palms and genitalia in approximately 6% of patients, but this effect can be alleviated if patients wash their hands and genitalia at the time of voiding after intravesical administration.

2. Thiotepa—Thiotepa is an alkylating agent with a molecular weight of 189. Although various doses have been used, 30 mg weekly seems to be sufficient. Up to 55% of patients respond completely. Most series show significantly lower recurrence rates in patients taking thiotepa than in those taking a placebo (Herr, Laudone, and Whitmore, 1987; Kowalkowski and Lamm, 1988). Cystitis is not uncommon after instillation but is usually mild and self-limited. Myelosuppression manifested as leukopenia and thrombocytopenia occurs in up to 9% of patients owing to systemic absorption. A complete blood count should be obtained in all patients before successive instillations.

3. BCG—BCG is an attenuated strain of Mycobacterium bovis. Many different strains of BCG exist, and the marketed preparations vary in the number, pathogenicity, viability, and immunogenicity of organisms (Catalona and

<table>
<thead>
<tr>
<th>Use</th>
<th>Timing</th>
<th>Goal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adjunctive</td>
<td>At TUR</td>
<td>Prevent implantation</td>
</tr>
<tr>
<td>Prophylactic</td>
<td>After complete TUR</td>
<td>Prevent or delay recurrence or progression</td>
</tr>
<tr>
<td>Therapeutic</td>
<td>After incomplete TUR</td>
<td>Cure residual disease</td>
</tr>
</tbody>
</table>

TUR, transurethral resection.
The exact mechanism by which BCG exerts its antitumor effect is unknown, but it seems to be immunologically mediated. Mucosal ulceration and granuloma formation are commonly seen after intravesical instillation. Activated helper T lymphocytes can be identified in the granulomas, and interleukin-2 reportedly can be detected in the urine of treated patients (Haaf, Catalona, and Ratliff, 1986). BCG has been shown to be very effective both therapeutically and prophylactically. It appears to be the most efficacious intravesical agent for the management of CIS. Complete responses are recorded in 36–71% of patients with residual carcinoma (Herr, Laudone, and Whitmore, 1987; Catalona and Ratliff, 1990). Recurrence rates are reduced substantially in patients treated after endoscopic resection (11–27% versus a 70% recurrence after endoscopic resection alone) (Catalona and Ratliff, 1990; Herr, Laudone, and Whitmore, 1987; Herr et al, 1985; Lamm, 1985). BCG has been shown to be superior to intravesical chemotherapy in preventing recurrence in patients with high-risk superficial bladder cancer (Lamm et al, 1991). Although BCG appears to be effective in delaying progression of high-risk superficial bladder cancer, 40–50% of these patients will experience disease progression with extended follow-up and many patients will ultimately require cystectomy (Cookson et al, 1997; Herr et al, 1995; Davis et al, 2002). The most commonly recommended induction regimen for BCG is weekly for 6 weeks followed by a period of 6 weeks where no BCG is given. Maintenance therapy should be considered in high-risk patients (Lamm et al, 2000). The utility of maintenance BCG is still under some debate as some randomized studies have not demonstrated a benefit (Badalamenti et al, 1987). The optimal regimen for maintenance therapy is also unclear. Published regimens involve 3 instillations once a week at 3- to 6-month intervals for 3 years following TUR. Only a small proportion (16–32%) of patients received all the treatments in prior studies, which highlights the difficulty of administering maintenance therapy and its side effects (van der Meijden, 2003; Lamm et al, 2000). Maintenance BCG appears to be more effective than intravesical chemotherapy with mitomycin C for intermediate- and high-risk superficial bladder cancer (Bohle, 2003). BCG may be more effective than chemotherapy in preventing progression of superficial cancers (Sylvester et al, 2005). Side effects of intravesical BCG administration are relatively common, although severe complications are uncommon. Most patients experience some degree of urinary frequency and urgency. Hemorrhagic cystitis occurs in approximately 7% of patients, and evidence of distant infection is found in <2%. Patients with mild systemic or moderate local symptoms should be treated with isoniazid (300 mg daily) and pyridoxine (vitamin B6 50 mg/day), and the dosage of BCG should be reduced. Isoniazid is continued while symptoms persist and restarted 1 day before the next instillation.

Patients with severe systemic symptoms should have instillations stopped. Patients with prolonged high fever (>103°F), symptomatic granulomatous prostatitis, or evidence of systemic infection require treatment with isoniazid and rifampin (600 mg daily). Patients with signs and symptoms of BCG sepsis (eg, high fever, chills, confusion, hypotension, respiratory failure, jaundice) should be treated with isoniazid, rifampin, and ethambutol (1200 mg). The addition of cycloserine (500 mg twice daily) or prednisolone (40 mg daily) increases survival rates (Lamm, 1992).

4. New intravesical agents and approaches—The rate of metachronous tumor recurrence is high compared with that of low-grade cancers occurring in other organs (eg, nasopharynx, colon). Recurrence of superficial bladder cancer is related to cancer stage, grade and number of tumors, associated dysplasia, and deoxyribonucleic acid (DNA) content. Recurrent tumors may be due to regrowth of previously resected cancers, growth of new cancers at remote sites, or implantation and subsequent proliferation of cells released into the bladder at the time of endoscopic treatment of the original tumor. Several investigators have studied the efficacy of single-dose therapy delivered at the time of complete TUR (Tolley et al, 1988; Oosterlinck et al, 1993). Such therapy has been shown to decrease recurrence rates, probably by decreasing the risk of tumor cell implantation at the time of initial cancer resection. Studies of interferon-alpha and valrubicin (an anthracycline derivative) suggest that these agents, either alone or perhaps in combination with other agents, may be effective in either high-risk patients or those who fail to respond to first-line therapy (Beldegrun et al, 1998; Sarosdy et al, 1998; Steinberg et al, 2000). Preliminary studies suggest that low-dose BCG, in combination with interferon, may be successful in preventing recurrences up to 24 months in 57% of patients who are BCG naïve and in 42% of those who have failed prior BCG therapy (O’Donnell et al, 2004).

B. SURGERY

1. TUR—TUR is the initial form of treatment for all bladder cancers. It allows a reasonably accurate estimate of tumor stage and grade and the need for additional treatment. Patients with single, low-grade, noninvasive tumors may be treated with TUR alone; those with superficial disease but high-risk features should be treated with TUR followed by selective use of intravesical therapy, as described above. TUR alone has rarely been used in the management of patients with invasive bladder cancer because of a high likelihood of recurrence and progression. Such an approach has been used infrequently for carefully selected patients with comorbid medical conditions and either no residual disease or minimal disease only at restaging TUR of bladder tumor (Herr, 1987; Solsona et al, 1998). Careful follow-up of patients with superficial bladder cancers is mandatory because disease will recur in 30–80% of patients, depend-
ing on cancer grade, tumor stage, and number of tumors. Disease status at 3 months after initial resection is an important predictor of the risk of subsequent recurrence and progression (Holmang and Johansson, 2002; Solsona et al, 2000). For patients who presented initially with solitary, low-grade lesions and who are free of recurrence at 3 months, repeat cystoscopy at 1 year is suggested. Patients who presented initially with multiple or higher grade lesions (or both) and those who have recurrences at 3 months require more careful surveillance. In such patients, cystoscopy at 3-month intervals is necessary. Although periodic cystoscopy is suggested for all patients with a history of bladder cancer, the risk of recurrence decreases as the tumor-free interval increases. After 5 years without recurrence, the risk of recurrence has been estimated to be 22%; the rate is 2% for 10 years (Morris et al, 1995).

2. Partial cystectomy—Patients with solitary, infiltrating tumors (T1–T3) localized along the posterior lateral wall or dome of the bladder are candidates for partial cystectomy, as are patients with cancers in a diverticulum. Disease remote from the primary tumor must be excluded by random bladder biopsies preoperatively. To minimize tumor implantation resulting from contamination of the wound with cancer cells at the time of surgery, short-course, limited-dose (1000–1600 cGy) irradiation can be used, and an intravesical chemotherapeutic agent can be instilled preoperatively (Ojeda and Johnson, 1983). Although survival rates of well-selected patients may approach those for patients with similar stage tumors treated by radical cystectomy, local recurrences are common (Whitmore, 1983; Sweeney et al, 1992). Patients with concomitant CIS and those with lymph node metastases do not respond well to partial cystectomy (Holbeierlein et al, 2004). Given current techniques of bladder replacement surgery, partial cystectomy is rarely indicated in the management of patients with invasive bladder cancer.

3. Radical cystectomy—Radical cystectomy implies removal of the anterior pelvic organs: in men, the bladder with its surrounding fat and peritoneal attachments, the prostate, and the seminal vesicles; in women, the bladder and surrounding fat and peritoneal attachments, cervix, uterus, anterior vaginal vault, urethra, and ovaries. This remains the “gold standard” of treatment for patients with muscle invasive bladder cancer. However, in select female patients, the vaginal vault and urethra can be spared along with the uterus, fallopian tubes, and ovaries, particularly in those who are premenopausal. Sparing of the urethra allows for construction of a neobladder that can be anastomosed to the urethral remnant. Disease-free survival 5 years after surgery is based on tumor stage: 88% for patients with P0, Pa, or PIS disease; 80% for patients with P1 disease; 81% for patients with P2 disease; 68% for patients with P3a and 47% for those with P3b disease; and 44% for patients with P4a disease (Stein et al, 2001). Recurrences after surgery usually occur within the first 3 years. Local pelvic recurrence rates are low (7–10%); most patients who fail therapy have distant disease recurrence.

The risk of urethral tumor occurrence or recurrence in men who undergo radical cystectomy is 6.1–10.6%. Risk factors for urethral tumor involvement in men include infiltration of the prostatic stroma or prostatic urethra with cancer or CIS. Patients with these risk factors are candidates for urethrectomy either at the time of radical cystectomy or as a separate procedure (Zabbo and Montie, 1984). Although prostatic urethral disease is a risk factor for urethral recurrence, recent evidence suggests that urethrectomy may be omitted and orthotopic urinary diversion performed safely in men with only proximal prostatic urethral involvement and a negative urethral margin at radical cystectomy (Iselin et al, 1997).

Urethrectomy was once routinely performed in all women undergoing radical cystectomy. However, recent clinical experience suggests that bladder replacement may be an acceptable procedure in women as well as men. Women with bladder cancer who have an involved urethral margin at the time of cystectomy and whose tumor was not located at the bladder neck are candidates for this procedure. Approximately 66% of women undergoing radical cystectomy for the management of bladder cancer fall into this group (Stein et al, 1995; Stenzl et al, 1995; Stein et al, 1998).

In such women, even the uterus, substantial portion of the vaginal vault, fallopian tubes, and ovaries can be spared. A bilateral pelvic lymph node dissection is usually performed simultaneously with radical cystectomy. Lymph node metastases are identified in approximately 20–35% of patients (Stein et al, 2001)—an incidence that reflects the inability of any imaging mode to identify consistently small-volume lymph node metastases preoperatively. Patients with lymph node metastases have a poorer prognosis. However, some patients (10–33%) with limited disease in regional lymph nodes may be cured by radical cystectomy and lymphadenectomy (Lerner et al, 1993; Vieweg et al, 1999; Stein et al, 2001). Even patients with pathologically negative nodes appear to benefit from an extensive lymphadenectomy (Konety, 2003). Patients with fewer than 5 positive lymph nodes and organ-confined disease in the primary tumor tend to have a better prognosis than patients with more extensive disease. These patients may also benefit from adjuvant chemotherapy (see section Chemotherapy).

Urinary diversion may be accomplished using a variety of techniques. Methods have been developed that allow construction of reservoirs that are continent and do not require the patient to wear an external appliance for collection of urine (see Chapter 24).

C. Radiotherapy

External beam irradiation (5000–7000 cGy), delivered in fractions over a 5- to 8-week period, is an alternative to
radical cystectomy in well-selected patients with deeply infiltrating bladder cancers. Treatment is generally well tolerated, but approximately 15% of patients may have significant bowel, bladder, or rectal complications. Five-year survival rates for stages T2 and T3 disease range from 18% to 41% (Goffinet et al, 1975; Woon et al, 1985; Quilty and Duncan, 1986). Unfortunately, local recurrence is common, occurring in approximately 33–68% of patients. Consequently, radiation as monotherapy is usually offered only to those patients who are poor surgical candidates due to advanced age or significant comorbid medical problems.

D. CHEMOTHERAPY

Approximately 15% of patients who present with bladder cancer are found to have regional or distant metastases; approximately 30–40% of patients with invasive disease develop distant metastases despite radical cystectomy or definitive radiotherapy. Without treatment, survival is limited. Early results with single chemotherapeutic agents and, more recently, combinations of drugs have shown that a significant number of patients with metastatic bladder cancer respond partially or completely (Scher and Sternberg, 1985). The single most active agent is cisplatin, which, when used alone, produces responses in approximately 30% of patients (Yagoda, 1983). Other effective agents include methotrexate, doxorubicin, vinblastine, cyclophosphamide, gemcitabine, and 5-fluorouracil. Response rates improve when active agents are combined. The regimen of methotrexate, vinblastine, doxorubicin (Adriamycin), and cisplatin (MVAC) has been the most commonly used for patients with advanced bladder cancer (Sternberg et al, 1988; Tannock et al, 1989). Approximately 13–35% of patients receiving such regimens attain a complete response. However, the median survival time is approximately 1 year, and the sustained survival rate is 20–25%. Treatment with MVAC is associated with substantial toxicity, including a toxic death rate of 3–4%.

Other newer agents demonstrating activity in this disease include ifosfamide, gemcitabine, paclitaxel, and gallium nitrate (Fagbemi and Stadler, 1998). A recent study demonstrated similar overall survival, time to treatment failure, and response rate for patients treated with MVAC and those treated with the newer combination of gemcitabine and cisplatin (von der Maase et al, 2000). The advantage of gemcitabine and cisplatin over MVAC is significantly lower toxicity and improved tolerability.

E. COMBINATION THERAPY

Once it became apparent that patients with metastatic bladder cancer could benefit from combination chemotherapy, investigators began treating patients with locally invasive (T2–T4), but not metastatic, cancer similarly. Chemotherapy can be given before planned radical cystectomy (neoadjuvant) in an attempt to decrease recurrence rates and, in selected cases, allow for bladder preservation. Approximately 22–43% of patients achieve a complete response to chemotherapy alone (Scher, 1990; Scher et al, 1988). However, additional treatment is still indicated because a substantial number of patients believed to be free of tumors after chemotherapy alone are found to have infiltrating disease at the time of surgery (Scher et al, 1989). Results from a recent randomized trial suggest that neoadjuvant chemotherapy followed by surgery may improve duration of survival when compared with surgery alone for patients with invasive disease. Patients who undergo neoadjuvant chemotherapy are more likely to have no residual tumor in the bladder at cystectomy and this portends a better long-term survival (Grossman, 2003). Alternatively, adjuvant chemotherapy may be offered to selected patients after radical cystectomy because of an increased risk of recurrence due to the presence of locally advanced disease (ie, P3, P4, or N+) (Skinner et al, 1991; Logothetis et al, 1988; Scher, 1990; Stockle et al, 1992; Stockle et al, 1995; Freiha et al, 1996). These studies suggest that patients initially managed with radical cystectomy who are found to be at an increased risk of systemic relapse due to the presence of lymph node metastases or regionally advanced disease are candidates for adjuvant chemotherapy.

Owing to high local and systemic failure rates after definitive irradiation, several investigators have explored the possibility of combining irradiation with systemic chemotherapy to decrease recurrence rates, improve patient survival, and allow bladder preservation. Trials of single-agent chemotherapy and irradiation have shown better local response rates than are found in historical series of irradiation alone (Shipley et al, 1984; Jakse, Fritsch, and Frommhold, 1985; Pearson and Raghaven, 1985).

More recently, investigators have treated patients with invasive bladder cancer with complete TUR followed by concomitant chemotherapy and radiation (Given et al, 1995; Chauvet et al, 1996; Shipley et al, 1997; Zietman et al, 1997; Cervek et al, 1998; Kachnic et al, 1997; Tester et al, 1996; Serretta et al, 1998; Zeitman et al, 2001). Early cystectomy is offered to those who do not tolerate chemotherapy, radiation, or both owing to toxicity and those whose cancers fail to respond to such therapy. Complete response rates to chemoradiation may be as high as 50–70% initially, with 5-year overall survival rates approaching 50–60%. However, local recurrence is common, exceeding 50% in many of these studies. Studies with longer median follow-up of almost 7 years suggest that the rate of superficial disease recurrence may be lower at around 26% (Zeitman et al, 2001). However patients who develop superficial disease recurrence (most commonly CIS) are more likely to require salvage cystectomy with only 34% being alive with a preserved bladder at 8 years compared to 61% of those who do not have such disease recurrence. Owing to invasive local recurrences, only 18–44% of patients may be alive with an intact bladder 5 years
after chemoradiation. Local disease stage and completeness of initial TUR are predictive of response and survival while delivery of radiation therapy by itself is not (Rodel, 2002). Predictors of poor outcome after combined chemoradiation for invasive bladder cancer include hydronephrosis at presentation, advanced clinical tumor stage, inability to complete the entire treatment protocol, and poor performance status. A recent study has suggested that chemoradiation may also be inappropriate for patients with bladder tumors that are p53-positive (Herr et al, 1999). Combined chemotherapy and radiation has also been used successfully to treat high-grade superficially invasive tumors (T1) (Akectin, 2005).

Systemic chemotherapy for locally invasive, but not metastatic, bladder cancer should not yet be considered standard therapy. The durability of the response, ultimate survival rates, and optimal candidates for the treatment regimens described will be determined only after completion of randomized studies.

URETERAL & RENAL PELVIC CANCERS

Incidence
Carcinomas of the renal pelvis and ureter are rare, accounting for only 4% of all urothelial cancers. The ratio of bladder–renal pelvic–ureteral carcinomas is approximately 51:3:1 (Williams and Mitchell, 1973). The mean age at diagnosis is 65 years, and the male–female ratio is 2–4:1 (Babaian and Johnson, 1980). Urothelial cancer often presents as a widespread urothelial abnormality; Patients with a single upper-tract carcinoma are at risk of developing bladder carcinomas (30–50%) and contralateral upper-tract carcinoma (2–4%). Conversely, patients with primary bladder cancer are at low risk (<2%) of developing upper urinary tract cancers (Oldbring et al, 1989). However, patients with multiple, recurrent superficial and in situ bladder cancers that are successfully treated by TUR and BCG are at a substantial lifelong risk of development of upper-tract cancers (Herr, 1998). The cumulative risks of such cancers have been estimated to be 10% at 5 years of follow-up, 26% at 5–10 years, and 34% at >10 years.

Etiology
As with bladder carcinoma, smoking and exposure to certain industrial dyes or solvents are associated with an increased risk of upper urinary tract TCCs. However, these tumors also occur with increased frequency in patients with a long history of excessive analgesic intake, those with Balkan nephropathy, and those exposed to Thorotrast, a contrast agent previously used for retrograde pyelography. Patients with carcinomas associated with analgesic abuse are more likely to be women, have higher stage disease, and be younger than others (Mahoney et al, 1977). All the major constituents of the analgesic compounds consumed (acetaminophen, aspirin, caffeine, and phenacetin) may be associated with an increased risk of upper urinary tract cancer (Ross et al, 1989; Jensen et al, 1989). Balkan nephropathy is an interstitial inflammatory disease of the kidneys that affects Yugoslavians, Rumanians, Bulgarians, and Greeks (Markovic, 1972); associated upper-tract carcinomas are generally superficial and more likely to be bilateral. The exact mechanism of tumor induction in these patients remains unknown.

Pathology
The mucosal lining of the renal pelvis and ureter is similar to that of the urinary bladder, being composed of transitional cell epithelium. Thus, most renal pelvic and ureteral cancers (90% and 97%, respectively) are TCCs. Grading is similar to that for bladder carcinomas. Papillomas account for approximately 15–20% of cases (Grabstald, Whitmore, and Melamed, 1971). They are isolated in just over 50% of patients and multiple in the rest, and in approximately 25% of patients with isolated papillomas and 50% of patients with multiple papillomas, carcinomas eventually develop. Among patients with carcinomas of the ureter, multicentricity approaches 50%. There is a relationship between tumor grade and the likelihood of urothelial abnormalities elsewhere: Low-grade cancers are associated with a low incidence of urothelial atypia or CIS in remote sites; however, these abnormalities are common with high-grade neoplasms (McCarron, Chasko, and Bray, 1982). Most upper urinary tract TCCs are localized at the time of diagnosis; the most common metastatic sites include regional lymph nodes, bone, and lung.

Squamous carcinomas account for approximately 10% of renal pelvic cancers and are much rarer in the ureter. Most carcinomas are usually sessile and infiltrating at the time of diagnosis. Such tumors are commonly identified in patients with a history of chronic inflammation from infection or calculus disease. Adenocarcinomas are very rare tumors of the upper urinary tract and, like squamous carcinomas, tend to be far advanced at the time of diagnosis.

Mesodermal tumors of the renal pelvis and ureter are quite rare. Benign tumors include fibroepithelial polyps (the most common), leiomyomas, and angiomas. Fibroepithelial polyps occur most commonly in young adults and are characterized radiographically by a long, slender, and polyploid filling defect within the collecting system. The most common malignant mesodermal tumors are leiomyosarcomas. The ureter and renal pelvis may be invaded by cancers of contiguous structures, such as primary renal, ovarian, or cervical carcinomas. True metastases to the ureter are rare. The most common metastatic tumors include those of stomach, prostate, kidney, and breast as well as lymphomas.
UROTHELIAL CARCINOMA: CANCERS OF THE BLADDER, URETER, & RENAL PELVIS

Staging & Natural History

Staging of both renal pelvic and ureteral carcinomas (Table 20–4) is based on an accurate assessment of the degree of tumor infiltration and parallels the staging system developed for bladder cancer (Grabstald, Whitmore, and Melamed, 1971; American Joint Committee on Cancer, 1997). Tumor stage and grade correlate with survival (Reitelman et al, 1987). Low-grade and low-stage cancers of the renal pelvis and ureter are associated with survival rates between 60% and 90%, compared with 0% and 33% for tumors of higher grade or those that have penetrated deep into or through the renal pelvic or ureteral wall (Hall et al, 1998). The latter figures reflect a high likelihood of regional or distant metastases—40% and 75% in patients with stages B and C (T2–T4) cancers, respectively.

Clinical Findings

A. SYMPTOMS AND SIGNS

Gross hematuria is noted in 70–90% of patients. Flank pain, present in 8–50%, is the result of ureteral obstruction from blood clots or tumor fragments, renal pelvic or ureteral obstruction by the tumor itself, or regional invasion by the tumor. Irritative voiding symptoms are present in approximately 5–10% of patients. Constitutional symptoms of anorexia, weight loss, and lethargy are uncommon and are usually associated with metastatic disease. A flank mass owing to hydronephrosis or a large tumor is detected in approximately 10–20% (Geerdsen, 1979), and flank tenderness may be elicited as well. Supraclavicular or inguinal adenopathy or hepatomegaly may be identified in a small percentage of patients with metastatic disease.

B. LABORATORY FINDINGS

Hematuria is identified in most patients but may be intermittent. Elevated liver function levels due to liver metastases are noted in a few patients. Pyuria and bacteriuria may be identified in patients with concomitant urinary tract infection from obstruction and urinary stasis.

As with bladder cancers, upper urinary tract cancers may be identified by examining exfoliated cells in the urinary sediment. In addition, specimens may be obtained directly with a ureteral catheter or by passing a small brush through the lumen of an open-ended catheter (Gill, Lu, and Thompson, 1973; Dodd et al, 1997). Detection depends on the grade of the tumor and the adequacy of the specimen obtained: 20–30% of low-grade cancers may be detected by cytologic testing compared with more than 60% of higher grade lesions (McCarron, Mullis, and Vaughn, 1983); using barbotage or a ureteral brush increases diagnostic accuracy. The utility of the newer voided markers, such as the BTA stat test (Bard Diagnostic Sciences, Inc, Redmond, WA), in detecting upper-tract urothelial cancers has not yet been determined (Zimmerman et al, 1998).

C. IMAGING

Findings on intravenous urography in patients with upper urinary tract cancers are usually abnormal. The most common abnormalities identified include an intraluminal filling defect, unilateral nonvisualization of the collecting system, and hydronephrosis (Williams and Mitchell, 1973; Almgard, Freedman, and Ljungqvist, 1973). Ureteral and renal pelvic tumors must be differentiated from nonopaque calculi, blood clots, papillary necrosis, and inflammatory lesions such as ureteritis cystica, fungus infections, or tuberculosis. The intravenous urography is often indeterminate, requiring retrograde pyelography for more accurate visualization of collecting-system abnormalities and simultaneous collection of cytologic specimens. CT urography is being increasingly used as the test of choice for evaluating the upper tract. During retrograde pyelography, contrast material is injected into the ureteral orifice with a bulb or acorn-tip catheter. Intraluminal filling defects may then be identified in the ureter or renal pelvis (Figure 20–5). Ureteral tumors are often characterized by dilation of the ureter distal to the lesion, creating the appearance of a “goblet.” Nonopaque ureteral calculi appear as a narrowing of the ureter distal to the calculus. A ureteral catheter passed up the ureter may coil distal to a ureteral tumor (Bergman’s sign) (Bergman, Friedenberg, and Sayegh, 1961). Ultrasonography, CT, and MRI frequently identify soft-tissue abnormalities of the renal pelvis but may fail to identify ureteral filling defects directly, although they may show hydronephrosis (Figure 20–6). All 3 imaging techniques differentiate blood clot and tumor from nonopaque calculi. In addition, CT and MRI allow simultaneous examination of abdominal and retro-

Table 20–4. Staging of Ureteral and Renal Pelvic Carcinoma.

<table>
<thead>
<tr>
<th>System</th>
<th>Batata*</th>
<th>TNM†</th>
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<tbody>
<tr>
<td>Confined to mucosa</td>
<td>O</td>
<td>Ta, Tis</td>
</tr>
<tr>
<td>Invasion of lamina propria</td>
<td>A</td>
<td>T1</td>
</tr>
<tr>
<td>Invasion of muscularis</td>
<td>B</td>
<td>T2</td>
</tr>
<tr>
<td>Extension through muscularis into fat or renal parenchyma</td>
<td>C</td>
<td>T3</td>
</tr>
<tr>
<td>Spread to adjacent organs</td>
<td>D</td>
<td>T4</td>
</tr>
<tr>
<td>Lymph node metastases</td>
<td>D</td>
<td>N+</td>
</tr>
<tr>
<td>Metastases</td>
<td>D</td>
<td>M+</td>
</tr>
</tbody>
</table>

*Drawn from Batata et al, 1975.
†Drawn from American Joint Committee on Cancer, 1997.
peritoneal structures for signs of regional (lymph node) or more distant metastases.

**D. URETEROPYELOSCOPY**

The use of rigid and flexible ureteropyeloscopes has allowed direct visualization of upper urinary tract abnormalities. These instruments are passed transurethrally through the ureteral orifice; in addition, they (and the similarly constructed but larger nephrosopes) can be passed percutaneously into renal calyces and the pelvis directly. The latter instrument carries with it the theoretic possibility of tumor spillage along the percutaneous tract. Indications for ureteroscopy include evaluation of filling defects within the upper urinary tract and after positive results on cytologic study or after noting unilateral gross hematuria in the absence of a filling defect. Ureteroscopy is also performed as a surveillance procedure in patients who have undergone conservative surgery for removal of a ureteral or renal pelvic tumor. Visualization, biopsy, and, on occasion, complete tumor resection, fulguration, or laser vaporization of the tumor are possible endoscopically. Performance of ureteroscopy with biopsy to establish the diagnosis in a patient with positive urine cytology and an upper tract filling defect may not always be necessary as these patients are presumed to have upper-tract TCC for which nephroureterectomy may be considered. However, any delay that may ensue from first performing a ureteroscopy with biopsy does not appear to jeopardize subsequent patient survival (Boorjian et al, 2005). Ureteroscopic visualization with biopsy is accurate and can identify cancer in a majority of patients. A diagnosis of cancer can be obtained >90% of the time with grade determination possible in >80% of cases (Keeley, 1997). It is harder to obtain lamina propria or muscle in ureteroscopic cup biopsy specimens which limits evaluation for stage of disease. Correlation of grade determined by tumor biopsy to that of the nephroureterectomy specimen is observed in 78% of cases. Biopsies tend to underestimate tumor grade in 22% of patients and stage in 45% of Ta tumors (Guarnizo et al, 2000). Multiple biopsies and biopsy of tumors in the proximal ureter tend to be more reliable in accurately determining stage and grade of ureteric tumors (Guarnizo et al, 2000).
**Treatment**

Treatment of renal pelvic and ureteral tumors should be based primarily on grade, stage, position, and multiplicity. Renal function and anatomy should be assessed. The standard therapy for both tumor types has been nephroureterectomy with excision of a bladder cuff owing to the possibility of multifocal disease within the ipsilateral collecting system. This procedure may be performed using either an open or laparoscopic approach (Landman et al., 2002; Jarrett et al., 2001). When the operation is performed for proximal ureteral or renal pelvic cancers, the entire distal ureter with a small cuff of bladder needs to be removed to avoid recurrence within this segment (Strong et al., 1976; Reitelman et al., 1987). Tumors of the distal ureter may be treated with distal ureterectomy and ureteral reimplantation into the bladder if no proximal defects suggestive of cancer have been noted (Babayan and Johnson, 1980).

Indications for more conservative surgery, including open or endoscopic excision, are not well defined. Absolute indications for renal-sparing procedures include tumor within the collecting system of a single kidney and bilateral urothelial tumors of the upper urinary tract or in patients with 2 kidneys but marginal renal function. In patients with 2 functioning kidneys, endoscopic excision alone should be considered only for low-grade and noninvasive tumors. One must realize that endoscopic examination may fail to detect the degree of infiltration adequately and therefore may understage some tumors. Limited experience with endoscopic resection, fulguration, or vaporization suggests that the procedure is safe in properly selected patients (Blute et al., 1989). However, recurrences have been noted in 15–80% of patients treated with open or endoscopic excision (Maier et al., 1990; Blute et al., 1989; Orihuela and Smith, 1988; Keeley et al., 1997; Stoller et al., 1997). Recurrence may be avoided by treating with instillation of immunotherapeutic or chemotherapeutic agents such as BCG or mitomycin C (Orihuela and Smith, 1988; Keeley and Bagley, 1997; Studer et al., 1989). These agents can be delivered to the upper urinary tract through single or double-J ureteral catheters (Patel and Fuchs, 1998). If patients are treated conservatively, it has been suggested that routine follow-up should include routine endoscopic surveillance because imaging alone may be inadequate for detecting recurrence (Chen et al., 2000).

Radiotherapy plays a limited role in upper urinary tract cancers. Although controversial, postoperative irradiation is believed by some investigators to decrease recurrence rates and improve survival in patients with deeply infiltrating cancers. Patients with metastatic, transitional cell cancers of the upper urinary tract should receive cisplatin-based chemotherapeutic regimens as described for patients with metastatic bladder cancers.

**Future Directions**

Urothelial cancers represent a spectrum of disease defined by various recurrence and progression rates. Further development of biologic markers such as tumor proliferation or antigen expression may permit a better estimate of the biologic potential of individual tumors. More refined visualization techniques such as blue light cystoscopy and use of ferromagnetic particles, as contrast to detect lymph node involvement on MRI, would allow clinicians to identify more advanced disease earlier and thereby select treatment strategies better. Newer intravesical therapies with combinations of chemotherapeutic agents and/or agents directed at molecular targets are being developed. New agents for the management of most patients with metastatic disease who do not respond to conventional chemotherapy need to be developed. Mechanisms of drug resistance and the means to circumvent them need to be investigated.

**REFERENCES**

**Bladder Carcinomas**


Ureteral & Renal Pelvic Cancers


BENIGN TUMORS

With the liberal use of computed tomography (CT) scans and magnetic resonance imaging (MRI), benign renal masses are being detected more frequently. Benign renal tumors include adenoma, oncocytoma, angiomylipoma, leiomyoma, lipoma, hemangioma, and juxtaglomerular tumors.

Renal Adenomas

The adenoma is the most common benign renal parenchymal lesion (Williams, 1992). These are small, well-differentiated glandular tumors of the renal cortex. They are typically asymptomatic and usually identified incidentally. At autopsy, 7–22% of patients are found to have a renal adenoma (Bonsib, 1985). Despite the classification of adenoma as a benign tumor, no clinical, histologic, or immunohistochemical criteria differentiate renal adenoma from renal carcinoma. Previously, all renal tumors <3 cm were considered adenomas. However, even such small tumors can be of high grade and advanced stage and metastasize and are now classified as renal cell carcinoma (RCC) (Remzi, 2006).

Renal Oncocytoma

Renal oncocytoma has a spectrum of behavior ranging from benign to malignant. Composed of large epithelial cells with finely granular eosinophilic cytoplasm (oncocyes), oncocytomas occur in various organs and organ systems including adrenal, salivary, thyroid, and parathyroid glands as well as the kidney. An estimated 3–5% of renal tumors are oncocytomas (Romis, 2004). Men are affected more often than women.

Renal oncocytomas generally occur within a well-defined fibrous capsule, with tumor tissue rarely penetrating the renal capsule, pelvis, collecting system, or perinephric fat. Metastasis is extremely rare though invasion of the lymphovascular spaces has been observed. On cut section, the surface of the tumor is usually tan or light brown with a central stellate scar, but necrosis typical of renal adenocarcinoma is absent. The tumors are usually solitary and unilateral, although several bilateral cases and multiple oncocytomas occurring simultaneously (oncocytomatosis) have been reported (Tickoo et al, 1999).

Oncocytomas can also be associated with benign tumors of hair follicles (fibrofolliculomas), colon polyps/tumors, and pulmonary cysts as part of the Birt-Hogg-Dubé syndrome (Toro et al, 1999). The Familial Renal Oncocytoma syndrome has also been described (Philips, 2001). These patients may have a characteristic genetic abnormality involving a gene located on 17p encoding a protein named folliculin (Nickerson, 2002) Histologically, well-differentiated oncocytomas are made up of large, uniform cells containing an intensely eosinophilic cytoplasm, which on ultrastructural studies is found to be packed with mitochondria. Mitotic activity is absent, and nuclear pleomorphism is uncommon (Figure 21–1). Consistent chromosomal alterations such as loss of chromosome 1 or Y and translocations in the short arm of chromosome 11 have been described in oncocytomas (Philips, 2001; Lindgren et al, 2004). The cellular origin of renal oncocytes has not been fully elucidated, although some early evidence suggested that oncocytes resemble proximal convoluted tubular cells (Merino and Libresi, 1982). Other findings suggest their origin may be a precursor stem cell (Cohen, McCue, and Derose, 1988) or the intercalated cells of the collecting ducts (Storkel et al, 1989).

The diagnosis of oncocytoma is predominantly pathologic because there are no reliable distinguishing clinical characteristics. Gross hematuria and flank pain occur in <20% of patients. No characteristic features of the tumors appear on CT, ultrasound (US), intravenous urography (IVU), or MRI. Angiographic features of oncocytomas include the “spoke-wheel” appearance of tumor arterioles, the “lucent rim sign” of the capsule, and a homogeneous capillary nephrogram phase. Unfortunately, these findings are not invariable, and similar findings have been reported in patients with RCC (Maatman et al, 1984).

High-grade oncocytomas may be intermixed with elements of RCC (Davis et al, 1991) and can be found as coexisting lesions within the same or opposite kidney (Licht et al, 1993). The role of fine-needle aspiration in the preoperative diagnosis of oncocytomas remains controversial and limited, due to a lack of characteristic features that distinguish oncocytoma from RCC.

Angiomyolipoma (Renal Hamartoma)

Angiomyolipoma is a rare benign tumor of the kidney seen in 2 distinct clinical populations. Angiomyolipomas are found in approximately 45–80% of patients with tuberous sclerosis and are typically bilateral and asymptomatic.
Tuberous sclerosis is a familial inherited disorder comprising adenoma sebaceum, mental retardation, and epilepsy. In patients without tuberous sclerosis, renal angiomyolipomas can be unilateral and tend to be larger than those associated with tuberous sclerosis (Anderson and Hatcher, 1990). There is no known histologic difference between the lesions seen in these 2 populations. As many as 25% of cases can present with spontaneous rupture and subsequent hemorrhage into the retroperitoneum (Wong, McGeorge, and Clark, 1981).

Angiomyolipomas are unencapsulated yellow-to-gray lesions, typically round to oval, that elevate the renal capsule, producing a bulging smooth or irregular mass. They are characterized by 3 major histologic components: mature fat cells, smooth muscle, and blood vessels. Renal hamartomas may extend to perirenal or renal sinus fat and involve regional lymphatics and other visceral organs (Ditonno et al, 1992). The presence of renal hamartomas in extrarenal sites is a manifestation of multicentricity rather than metastatic potential, because only one well-documented case of malignant transformation of angiomyolipoma has been reported (Lowe et al, 1992).

Patients with a rare condition termed lymphangioleiomyomatosis may have multiple renal and hepatic angiomyolipomata, multiple pulmonary cysts, enlarged abdominal lymph nodes, and lymphangiomomas (Avila et al, 2000; Urban et al, 1999). The diagnosis of renal angiomyolipoma has evolved with the widespread use of US and CT. Arteriography can reveal neovascularity similar to that of renal cancer and therefore is not helpful in differential diagnosis. Ultrasonography and CT are frequently diagnostic in lesions with high fat content. Fat visualized on US appears as very high intensity echoes. Fat imaged by CT has a negative density, –20 to –80 Hounsfield units, which is pathognomonic for angiomyolipomas when observed in the kidney (Figure 21–2) (Pitts et al, 1980).

The role of MRI as a diagnostic tool has been investigated; as in CT, the high fat content makes this lesion suitable for MRI diagnosis (Uhlenbrock, Fischer, and Beyer, 1988); however, because the presence of bleeding in any renal tumor can mimic the typical pattern of angiomyolipoma, MRI should not be considered the diagnostic method of choice. MRI can however be more useful than CT scanning in distinguishing lipid poor angiomyolipoma with low fat content from other solid renal lesions (Kim, 2006).

The management of angiomyolipomas historically has been correlated with symptoms. Steiner and colleagues (1993) reported a long-term follow-up study of 35 patients with angiomyolipomas. They proposed that patients with isolated lesions <4 cm be followed up with yearly CT or US. Patients with asymptomatic or mildly symptomatic lesions >4 cm should be followed up with semiannual US. Patients with lesions >4 cm with moderate or severe symptoms (bleeding or pain) should undergo renal-sparing surgery or renal arterial embolization. Given the difference in the natural history of angiomyolipomas in patients with tuberous sclerosis, Steiner et al advocate prophylactic intervention in patients with lesions >4 cm irrespective of symptoms, with close follow-up of smaller lesions. Pregnancy may also increase the risk of growth and bleeding from larger renal angiomyolipomas which could be preemptively managed by embolization prior to or early in pregnancy.

Other Rare Benign Renal Tumors

Several other benign renal tumors are quite rare, including leiomyomas, hemangiomas, lipomas, and juxtaglomerular cell tumors. With the exception of juxtaglomerular tumors, there are no features that unequivocally establish the diagnosis before surgery; therefore, the pathologist most often provides the diagnosis after nephrectomy.

Figure 21–1. Histologic section of a grade I (benign) renal oncocytoma (original magnification, X100).

Figure 21–2. Computed tomogram of an angiomyolipoma (arrows).
Leiomyomas are rare small tumors typically found in smooth-muscle-containing areas of the kidney including the renal capsule and renal pelvis. Two groups of renal leiomyomas have been described (Steiner et al, 1990). The more common group comprises cortical tumors that are <2 cm and may be multiple. These tumors are typically found at autopsy and are not clinically significant. A larger, commonly solitary leiomyoma has been described, which may cause symptoms and is confirmed pathologically after nephrectomy.

Hemangiomas are small vascular tumors occurring in the kidney with a frequency second only to that in the liver among visceral organs. Multiple lesions in one kidney occur in approximately 12% of cases; however, they are rarely bilateral. They can occasionally be the elusive source of hematuria in an otherwise well-evaluated patient. The diagnosis may be determined by CT angiography, MR angiography, or by direct visualization by endoscopy (Ekelund and Gothlin, 1975).

Renal lipomas are very uncommon deposits of mature adipose cells without evident mitosis that arise from the renal capsule or perirenal tissue. They are seen primarily in middle-aged females and, owing to the characteristic CT differentiation of fat, are best detected radiographically on CT scanning.

The juxtaglomerular cell tumor is the most clinically significant member of this subgroup of rare benign tumors because it causes significant hypertension that can be cured by surgical treatment. It is a very rare lesion, with <100 reported cases and may have characteristic chromosomal alterations (Brandal, 2005). The tumors occur more commonly in women in their 20s and 30s and are rarely malignant. The tumors originate from the pericytes of afferent arterioles in the juxtaglomerular apparatus and can be shown to contain renin secretory granules. They are typically encapsulated and located in the cortical area. The diagnosis is suspected when there is secondary hyperaldosteronism and is confirmed by selected renal vein sampling for renin. Although complete nephrectomy was advocated in the past, several recent reports indicate that partial nephrectomy can be equally effective (Haab et al, 1995).

ADENOCARCINOMA OF THE KIDNEY (RCC)

In the United States in 2007, an estimated 51,190 new cases of adenocarcinoma of the kidney are expected to be diagnosed, and 12,890 deaths will occur from this disease (Jemal et al, 2007). RCC accounts for roughly 2.8% of adult cancers and constitutes approximately 85% of all primary malignant renal tumors. There appears to be an increase in the incidence of all stages of RCC over the past few decades (Hock, Lynch, and Balaji, 2002; Mindrup et al, 2005). RCC occurs most commonly in the fifth to sixth decade and has a male-female ratio of 2:1. The incidence of renal cancer may vary based on race, with black men demonstrating a higher incidence than in men of all other races. Black men may also have a higher likelihood of a subsequent RCC in the contralateral kidney (Rabbani et al, 2002). Asians appear to have the lowest incidence of RCC (Miller, 1996).

Etiology

The cause of renal adenocarcinoma is unknown. Occupational exposures, chromosomal aberrations, and tumor suppressor genes have been implicated. Cigarette smoking is the only risk factor consistently linked to RCC by both epidemiologic case-control and cohort studies (La Vecchia et al, 1990), with most investigations demonstrating at least a 2-fold increase in risk for the development of RCC in smokers (Yu et al, 1986). Exposure to asbestos, solvents, and cadmium has also been associated with an increased incidence of RCC (Mandel et al, 1995).

RCC occurs in two forms, inherited and sporadic. In 1979, Cohen and colleagues described a pedigree with hereditary RCC in which the pattern of inheritance was consistent with an autosomal dominant gene with a balanced reciprocal translocation between the short arm of chromosome 3 and the long arm of chromosome 8. Subsequent work has documented that both the hereditary and sporadic forms of RCC are associated with structural changes in chromosome 3p (Kovacs et al, 1988; Erlandsen, 1998; Noordzij and Mickisch, 2004).

Two other hereditary forms of RCC have been described. Von Hippel-Lindau disease is a familial cancer syndrome in which affected individuals have a predisposition to have tumors develop in multiple organs, including cerebellar hemangioblastoma, retinal angioma, and bilateral clear cell RCC. In 1993, Latif and colleagues identified the von Hippel-Lindau gene, leading to the detection of a germ line mutation in approximately 75% of families affected by von Hippel-Lindau disease (Chen et al, 1995).

Hereditary papillary renal carcinoma was described in 1994 and is characterized by a predisposition to develop multiple bilateral renal tumors with a papillary histologic appearance (Zbar et al, 1994). In contrast to von Hippel-Lindau patients, the major neoplastic manifestations appear to be confined to the kidney.

Acquired cystic disease of the kidneys is a well-recognized entity of multiple bilateral cysts in the native kidneys of uremic patients (Reichard, Roubidoux, and Dunnick, 1998). The risk of developing RCC has been estimated to be >30 times higher in patients receiving dialysis who have cystic changes in their kidney than in the general population (Brennan et al, 1991). Several series reported in the literature suggest that RCC occurs in 3–9% of patients with acquired cystic disease of the kidneys (Gulanikar et al, 1998). Most RCC cases have been described in patients undergoing hemodialysis, but RCC has been reported in association with peritoneal dialysis (Smith et al, 1987) and
successful renal transplants (Vaziri et al, 1984) and in patients with long-term renal insufficiency not requiring dialysis (Bretan et al, 1986; Fallon and Williams, 1989).

Pathology

RCC originates from the proximal renal tubular epithelium, as evidenced by electron microscopy (Makay, Ordonez, and Khoursland, 1987) and immunohistochemical analysis (Holthöfer, 1990). These tumors occur with equal frequency in either kidney and are randomly distributed in the upper and lower poles. RCCs originate in the cortex and tend to grow out into perinephric tissue, causing the characteristic bulge or mass effect that aids in their detection by diagnostic imaging studies. Grossly, the tumor is characteristically yellow to orange because of the abundance of lipids, particularly in the clear cell type. RCCs do not have true capsules but may have a pseudocapsule of compressed renal parenchyma, fibrous tissue, and inflammatory cells.

Histologically, RCC is most often a mixed adenocarcinoma containing clear cells, granular cells, and, occasionally, sarcomatoid-appearing cells. The classifications of the subtypes of RCC are based on morphology and cytogenetic characteristics. Most RCCs are classified into 1 of the following histologic subtypes: conventional clear cell, papillary (chromophilic), chromophobe, collecting duct, neuroendocrine, and unclassified (Mostofi and Davis, 1998). Benign renal tumors are papillary adenoma, renal oncocytoma, and metanephric adenoma. Clear cells are rounded or polygonal with abundant cytoplasm, which contains cholesterol, triglycerides, glycogen, and lipids (Figure 21–3).

The cells present in the papillary (chromophilic) type contain less glycogen and lipids, and electron microscopy reveals that the granular cytoplasm contains many mitochondria and cytosomes. Chromophobe-type carcinomas contain large polygonal cells with distinct cell borders and reticulated cytoplasm, which can stain diffusely with Hale’s colloidal iron (Theones et al, 1988). Oncocytic RCC or oncocytomas tend to have cytoplasm packed with mitochondria, giving it a granular appearance. Collecting duct tumors tend to have irregular borders and a basophilic cytoplasm with extensive anaplasia and are likely to invade blood vessels and cause infarction of tissue. Sarcomatoid cells are spindle-shaped and form sheets or bundles. This later cell type rarely occurs as a pure form and is most commonly a small component of either the clear cell or papillary cell type (or both).

Pathogenesis

RCCs are vascular tumors that tend to spread either by direct invasion through the renal capsule into perinephric fat and adjacent visceral structures or by direct extension into the renal vein. Approximately 25–30% of patients have evidence of metastatic disease at presentation. The most common site of distant metastases is the lung. However, liver, bone (osteolytic), ipsilateral adjacent lymph nodes and adrenal gland, brain, the opposite kidney, and subcutaneous tissue are frequent sites of disease spread.

Tumor Staging & Grading

A. TUMOR STAGING

The ultimate goal of staging is to select appropriate therapy and obtain prognostic information. Appropriate studies for a complete clinical staging evaluation include history and physical examination, complete blood count, serum chemistries (renal and hepatic function), urinalysis, chest x-ray (chest CT scan for an equivocal exam), CT scan of abdomen and pelvis, and a radionuclide bone scan (with x-rays of abnormal areas).

The original staging system described by Robson (1963) is easy to use, but it does not relate directly to prognosis and hence it is no longer commonly used. The Tumor-Node-Metastasis (TNM) system more accurately classifies the extent of tumor involvement and is currently most often used. The TNM classification system for RCC has undergone multiple revisions with the most recent edition being the 2002 version (Table 21–1). In the most recent AJCC TNM staging, stage T1 disease is further divided into T1a (tumor size <4 cm) and T1b (size 4–7 cm) as there is a difference in long-term survival between stage T1a and T1b (Ficarra, 2005).

B. TUMOR GRADING

Fuhrman grading has become commonly used by pathologists in North America (Fuhrman, Lasky, and Limas, 1982; Goldstein, 1997). The system uses 4 grades based on nuclear size and irregularity and nucleolar prominence. The system is most effective in predicting metastasis (50%...
of high-grade tumors within 5 years). When high-grade, predominantly granular tumors are corrected for grade and stage, there is no apparent difference between clear cell and granular cell tumor prognosis (McNichols, Segura, and DeWeerd, 1981). However, patients presenting with advanced disease do poorly irrespective of tumor grade.

Clinical Findings

A. SYMPTOMS AND SIGNS

The classically described triad of gross hematuria, flank pain, and a palpable mass occurs in only 7–10% of patients and is frequently a manifestation of advanced disease. Patients may also present with hematuria, dysnea, cough, and bone pain which are typically symptoms secondary to metastases. With the routine use of CT scanning for evaluation of nonspecific findings, asymptomatic renal tumors are increasingly detected incidentally (>50%).

B. PARANEoplastIC SYNDROMES

RCC is associated with a wide spectrum of paraneoplastic syndromes including erythrocytosis, hypercalcemia, hypertension, and nonmetastatic hepatic dysfunction. Overall, these manifestations can occur in 10–40% of patients with RCC.

RCC is the most common cause of paraneoplastic erythrocytosis, which is reported to occur in 3–10% of patients with this tumor (Sufrin et al, 1989). In patients with RCC, the elevated erythrocyte mass is physiologically inappropriate and may result either from enhanced production of erythropoietin from the tumor or as a consequence of regional renal hypoxia promoting erythropoietin production from nonneoplastic renal tissue (Hocking, 1987).

Hypercalcemia has been reported to occur in up to 20% of patients with RCC (Muggia, 1990). Hypercalcemia may be due to production of a parathyroid hormone–related peptide that mimics the function of parathyroid hormone (Strewler et al, 1987) or other humoral factors such as osteoclast-activating factor, tumor necrosis factor, and transforming growth factor-alpha (Muggia, 1990).

Hypertension associated with RCC has been reported in up to 40% of patients (Sufrin et al, 1989), and renin production by the neoplasm has been documented in 37%. The excess renin and hypertension associated with RCC are typically refractory to antihypertensive therapy but may respond after nephrectomy (Gold et al, 1996).

In 1961, Stauffer described a reversible syndrome of hepatic dysfunction in the absence of hepatic metastases associated with RCC. Hepatic function abnormalities include elevation of alkaline phosphatase and bilirubin, hypoalbuminemia, prolonged prothrombin time, and
hypergammaglobulinemia. Stauffer’s syndrome tends to occur in association with fever, fatigue, and weight loss and typically resolves after nephrectomy. The reported incidence of Stauffer’s syndrome varies from 3% to 20% (Gold et al, 1996). It may be due to overproduction of granulocyte-macrophage colony stimulating factor by the tumor (Chang et al, 1992).

RCC is known to produce a multitude of other biologically active products that result in clinically significant syndromes, including adrenocorticotropic hormone (Cushing’s syndrome), enteroglucagon (protein enteropathy), prolactin (galactorrhea), insulin (hypoglycemia), and gonadotropins (gynecomastia and decreased libido; or hirsutism, amenorrhea, and male pattern balding) (Sufrin, Golio, and Murphy, 1986).

A paraneoplastic syndrome present at the time of disease diagnosis does not, in-and-of-itself, confer a poor prognosis. However, patients whose paraneoplastic metabolic disturbances fail to normalize after nephrectomy (suggesting the presence of clinically undetectable metastatic disease) have very poor prognoses (Hanash, 1982).

C. LABORATORY FINDINGS

In addition to the laboratory abnormalities associated with the various RCC paraneoplastic syndromes, anemia, hematuria, and an elevated sedimentation rate are frequently observed.

Anemia occurs in about 30% of RCC patients. The anemia typically is not secondary to blood loss or hemolysis and is commonly normochromic. The serum iron and total iron-binding capacity are usually low, as in the anemia of chronic disease. Iron therapy is usually ineffective; however, surgical removal of early-stage tumors usually leads to physiologic correction of the anemia. The potential role of recombinant erythropoietin for patients with unresectable disease represents a potential, but untested, option.

Gross or microscopic hematuria can be seen in up to 60% of patients presenting with RCC. An elevated erythrocyte sedimentation rate is also commonly seen, with a reported incidence as high as 75%. These findings are nonspecific, and normal findings do not rule out a diagnosis of RCC.

D. X-RAY FINDINGS

Although many radiologic techniques are available to aid in the detection and diagnosis of renal masses, CT scanning remains the primary technique with which others must be compared. Other radiologic techniques used include US, MRI, and arteriography. Intravenous pyelography is rarely used for the diagnosis or evaluation of RCC. In this era of cost containment, selecting the appropriate studies for an efficient, cost-effective evaluation is mandatory.

E. ULTRASONOGRAPHY

US examination is a noninvasive, relatively inexpensive technique able to further delineate a renal mass seen on IVU. It is approximately 98% accurate in distinguishing simple cysts from solid lesions. Strict ultrasonographic criteria for a simple cyst include through transmission, a well-circumscribed mass without internal echoes, and adequate visualization of a strong posterior wall (Figure 21–4).

**Figure 21–4.** A: Ultrasound image of a simple renal cyst showing renal parenchyma (long arrows), cyst wall (arrowheads), and a strong posterior wall (short arrows). B: Ultrasound image of a solid renal mass (arrows).
can be helpful in distinguishing the presence and extent of a caval thrombus in equivocal CT studies.

**F. CT Scanning**

CT scanning is more sensitive than US or IVU for detection of renal masses. A typical finding of RCC on CT is a mass that becomes enhanced with the use of intravenous contrast media. In general, RCC exhibits an overall decreased density in Hounsfield units compared with normal renal parenchyma but shows a heterogeneous pattern of enhancement or increased attenuation (slightly decreased from the surrounding parenchyma) when contrast is used (Figure 21–5) (Kosko, Lipuma, and Resnick, 1984). In addition to defining the primary lesion, CT scanning is also the method of choice in staging the patient by visualizing the renal hilum, perinephric space, renal vein and vena cava, adrenals, regional lymphatics, and adjacent organs. In patients with equivocal chest x-ray findings, a CT scan of the chest is indicated. Patients who present with symptoms consistent with brain metastases should be evaluated with either head CT or MRI. Spiral CT with 3-dimensional reconstruction has become useful for evaluating tumors before nephron-sparing surgery to delineate the 3-dimensional extent of the tumor and precisely outline the vasculature, which can aid the surgeon in preventing positive surgical margins (Holmes et al, 1997). Intra-operative ultrasonography is also often used to confirm the extent and number of masses in the kidney at the time of performing a partial nephrectomy.

**G. Renal Angiography**

With the widespread availability of CT scanners, the role of renal angiography in the diagnostic evaluation of RCC has markedly diminished and is now very limited. There remain a very few specific clinical situations in which angiography may be useful; for example, guiding the operative approach in a patient with an RCC in a solitary kidney when attempting to perform a partial nephrectomy may be indicated (Figure 21–6). However, CT angiography or MR angiography can give better information with less risk to the patient.

**H. Radionuclide Imaging**

Determination of metastases to bones is most accurate by radionuclide bone scan, although the study is nonspecific and requires confirmation with bone x-rays of identified abnormalities to verify the presence of the typical osteolytic lesions. There is evidence that patients without bone pain and with a normal alkaline phosphatase level have a very low incidence of bone metastases (Henriksson et al, 1992), and thus a routine bone scan is not necessary in most patients.

**I. Magnetic Resonance Imaging**

MRI is equivalent to CT for staging of RCC (Hricak et al, 1988). Its primary advantage is in the evaluation of patients with suspected vascular extension (Figure 21–7).
Prospective trials have demonstrated that MRI is superior to CT in assessing inferior vena caval involvement (Kabala et al, 1991) and is at least as accurate as venacavography (Horan et al, 1989). In contrast to both CT and cavography, MRI evaluation does not require either iodinated contrast material or ionizing radiation. Recent studies using MRI angiography with gadolinium or CT angiography have improved vascular evaluation of renal neoplasms (Bluemke and Chambers, 1995). MR angiography can also be used to delineate the vascular supply before planned nephron-sparing surgery.

J. Positron Emission Tomography (PET)

This technique allows the measurement of systemically administered biochemical agents such as 18-fluoro-2-deoxyglucose (FDG), which can accumulate in the kidney. Although FDG-PET scanning can yield false-positive results in some patients with RCC (Bachor et al, 1996), it may be useful in monitoring response to systemic therapy in those with metastatic disease (Hoh, Seltzer, and Franklin, 1998). FDG-PET may also be more accurate than routine CT scanning in detecting disease recurrence or progression, which may alter treatment decisions in up to 50% of cases (Ramdave et al, 2001). However, further studies are required before it can be widely used for routine staging of RCC.

K. Fine-Needle Aspiration

Fine-needle aspiration cytology has had a limited role in the evaluation of RCC. Fine-needle aspiration of renal lesions is the diagnostic approach of choice in those patients with clinically apparent metastatic disease who may be candidates for nonsurgical therapy. Other settings in which fine-needle aspiration may be appropriate include establishing a diagnosis in patients who are not surgical candidates, differentiating a primary RCC from a renal metastasis in patients with known primary cancers of non-renal origin, and evaluating some radiographically indeterminate lesions (Renshaw, Granter, and Cibas, 1997). Fine needle aspiration is being increasingly used to confirm the diagnosis of a neoplasm particularly in patients who may undergo observation or percutaneous ablative therapy (Shah, 2005). While core needle biopsies may be able to accurately diagnose malignancy in up to 100% of cases >4 cm and 95% of cases <4 cm this may require multiple cores for accuracy (Wunderlich, 2005). Rare reports of seeding of the needle tract have been reported.

L. Instrumental and Cytologic Examination

Patients presenting with hematuria should also be evaluated with cystoscopy. Blood effluxing from the ureteral orifice identifies the origin of bleeding from the upper tract. Most renal pelvis tumors can be distinguished radiographically from RCC; however, endoscopic evaluation of the bladder, ureters, and renal pelvis is occasionally helpful in making a diagnosis. Additionally, although urine cytologic study is rarely helpful in the diagnosis of RCC, cytologic study of urine with renal pelvis washing is frequently diagnostic in renal pelvis tumors.

Differential Diagnosis

When a patient presents with clinical findings consistent with metastatic disease and is found to have a renal mass, a diagnosis of RCC can be straightforward. Most patients present with a renal mass discovered after an evaluation of hematuria or pain or as an incidental finding during an imaging workup of an unrelated problem. The differential diagnosis of RCC includes other solid renal lesions. The great majority of renal masses are simple cysts. Once the diagnosis of a cyst is confirmed by US, no additional evaluation is required if the patient is asymptomatic. Equivocal findings or the presence of calcification within the mass warrant further evaluation by CT. A wide variety of pathologic entities appear as solid masses on CT scans, and differentiation of benign from malignant lesions is frequently difficult. Findings on CT scan that suggest malignancy include amputation of a portion of the collecting system, presence of calcification, a poorly defined interface between the renal parenchyma and the lesion, invasion into perinephric fat or adjacent structures, and the presence of abnormal periaortic adenopathy or distant metastatic disease (Kosko, Lipuma, and Resnick, 1984). The frequency of benign lesions among renal masses <7 cm in size is as high as 16–20% (Duchene, 2003; Snyder, 2006). Masses >7 cm are rarely benign.

Some characteristic lesions can be defined using CT criteria in combination with clinical findings. Angiomyolipomas (with large fat components) can easily be identified...
by the low-attenuation areas classically produced by substantial fat content. A renal abscess may be strongly suspected in a patient presenting with fever, flank pain, pyuria, and an early needle aspiration and culture should be performed. Other benign renal masses (in addition to those previously described) include granulomas and arteriovenous malformations. Renal lymphoma (both Hodgkin’s disease and non-Hodgkin’s disease), transitional cell carcinoma of the renal pelvis, adrenal cancer, and metastatic disease (most commonly from a lung or breast cancer primary) are additional diagnostic possibilities that may be suspected based on CT and clinical findings.

**Treatment**

**A. SPECIFIC MEASURES**

1. **Localized disease**—Surgical removal of the early-stage lesion remains the only potentially curative therapy available for RCC patients. Appropriate therapy depends almost entirely on the stage of tumor at presentation and therefore requires a thorough staging evaluation. The prognoses of patients with stages T1-T3a disease are similar following radical nephrectomy.

Radical nephrectomy is the primary treatment for localized RCC. Its goal is to achieve the removal of tumor and to take a wide margin of normal tissue. Radical nephrectomy entails en bloc removal of the kidney and its enveloping fascia (Gerota’s) including the ipsilateral adrenal, proximal one-half of the ureter, and lymph nodes up to the area of transection of the renal vessels (Figure 21–8).

Various open incisions provide optimal access for the radical nephrectomy, including an anterior subcostal (unilateral chevron) or thoracoabdominal incision, and, occasionally, a midline incision or the classic flank incision (Droller, 1992).

The likelihood of local recurrence after radical nephrectomy is 2–3% (Itano et al, 2000). Repeat resection of isolated local recurrence can be curative and yield a survival benefit (Itano et al, 2000; Tanguay et al, 1996). The role of regional lymphadenectomy in RCC remains controversial. Between 18% and 33% of patients undergoing radical nephrectomy with lymph node dissection for RCC have metastatic disease identified (Skinner, Lieskovsky, and Pritchett, 1988). Although several retrospective studies (Thrasher and Paulson, 1993) and a prospective, nonrandomized study (Herrlinger et al, 1991) suggest that regional lymphadenectomy can improve survival in patients with T1-T2 RCC. More recent studies including a randomized prospective study as well as a population-based study failed to show any survival benefit that could be obtained by routinely performing regional lymphadenectomy especially in patients with organ-confined disease (Blom et al, 1999; Joslyn et al, 2005). Removal of the adrenal is unnecessary if the tumor is not in the upper pole, because adrenal involvement is uncommon in this instance.

Preoperative renal artery embolization (angioinfarction) has been used in the past as a surgical adjunct to facilitate radical nephrectomy, but because there is no conclusive evidence that preoperative embolization actually decreases blood loss or facilitates surgery, its use should be limited to patients with very large tumors in which the renal artery may be difficult to reach early in the procedure. Additionally, this technique may be useful to palliate patients with nonresectable tumors and significant symptoms such as hemorrhage, flank pain, or paraneoplastic syndromes.

Radiation therapy has been advocated as a neoadjuvant (preoperative) or adjuvant method to radical nephrectomy, but there is no evidence that postsurgical radiation therapy to the renal bed, whether or not residual tumor is present, contributes to prolonged survival.

RCC may invade renal vascular spaces and produce tumor thrombi extending into renal veins, inferior vena cava, hepatic veins, and, occasionally, the right atrium. Between 5% and 10% of patients presenting with RCC have some degree of vena caval involvement (Figure 21–9) (Skinner, Lieskovsky, and Pritchett, 1988). Patients presenting with involvement of the renal vein and vena cava below the hepatic veins (T3bN0M0) but without evidence of regional or distant metastases have a prognosis similar to patients with stage T2 disease when treated by radical exci-
renal parenchymal neoplasms

The surgical approach to the removal of caval thrombi depends entirely on the level of cephalad extension. In general, these thrombi do not invade the wall of the cava and therefore can be removed without resection of the caval wall. For tumor thrombi that have reached the level of the right atrium, the use of cardiopulmonary bypass is typically required.

Laparoscopic radical nephrectomy and partial nephrectomy can also be accomplished successfully and safely. Laparoscopic radical nephrectomy is being used increasingly for patients with localized renal tumors. This approach results in quicker recovery with efficacy comparable to that of open radical nephrectomy and is now the approach of choice in appropriate patients with <10 cm tumors and without local extension or a renal vein or caval thrombus (Portis et al, 2002; Gill et al, 2001).

The approach to the patient with either bilateral RCC or disease in a solitary kidney differs from the standard approach of radical nephrectomy. Bilateral RCC occurs with a frequency as high as 3% (Smith, 1986). Radical nephrectomy in these patients or in those with solitary kidneys obviously commits patients to long-term dialysis or renal transplantation and the morbidities of these conditions. Staging these patients is essentially the same as previously outlined, with the notable exception that either MR or CT angiography is often used to assess the extent of tumor within the kidney and the renal artery anatomy. Surgical alternatives to radical nephrectomy include open or laparoscopic partial nephrectomy, ex vivo partial nephrectomy (bench surgery followed by autotransplantation) (Novick, Stewart, and Straffon, 1980), and enucleation of multiple lesions (Marshall et al, 1986). Given the lack of effective adjuvant therapy and the risk of inadequate excision and subsequent recurrence from various renal-sparing approaches, partial nephrectomy with an adequate parenchymal margin remains the preferred treatment.

Partial nephrectomy and wedge resection with an adequate margin of normal parenchyma is increasingly being used as primary surgical therapy for patients with tumors <4 cm in size, even in the presence of a normal contralateral kidney. Local recurrence of tumor in the same kidney ranges from 0% to 10%, and it is between 0% and 3% for tumors <4 cm in size (Uzzo and Novick, 2001; Morgan and Zincke, 1990; Hafez, Novick, and Campbell, 1997). In patients with multiple small tumors, such as those with von Hippel-Lindau disease, enucleation of the tumor(s) is also an acceptable approach. Long-term follow-up demonstrates that partial nephrectomy has a similar outcome as radical nephrectomy (Herr, 1999). It is now considered the approach of choice in patients with small (<4 cm), incidentally discovered renal tumors that are peripherally located. Laparoscopic partial nephrectomy for these small tumors is currently gaining in use in expert hands as well.

Additional therapeutic approaches being explored for the treatment of small, incidentally discovered renal lesions include the use of cryoablation, high-intensity focused US, and radiofrequency ablation (Murphy and Gill, 2001). Cryoablation with liquid nitrogen or argon gas, either percutaneously using MRI guidance or via laparoscopic probes, has proved to be feasible and effective in selected patients (Shingleton and Sewell, 2002; Gill et al, 2000). Radiofrequency ablation has also been accomplished via the percutaneous approach with minimal morbidity in small groups of patients (Pavlovich et al, 2002). These approaches are particularly attractive in patients with multiple small lesions or larger individuals with many comorbidities. The long-term effectiveness of these emerging technologies remains to be determined.

Observation as treatment should also be mentioned for small (<3.0 cm) lesions, particularly in elderly patients. One recent study noted a growth rate of 0–1.3 cm/year in 40 patients followed up for a mean of 3.5 years (Bosniak, 1995), indicating that with careful follow-up, watchful waiting may be appropriate in selected patients. Only one-third of small (<4 cm) renal masses are observed to increase in size over 2 years with none experiencing disease progression (Volpe, 2004). This further suggests that at least initially observation is a reasonable option particularly for older patients with comorbidities who may not be amenable to surgery.

2. Disseminated disease—Approximately 30% of patients with RCC will present with advanced disease. Metastatic RCC has a natural history that is typically aggressive and rapidly progressive, with 5-year survival rates typically <10% (Motzer et al, 1996). Infrequently,
the disease may have a more protracted course. The biological diversity of RCC is illustrated by the 6.6% response rate (including 3% complete responders) in the placebo arm of a phase III trial of interferon-gamma (IFN-γ) in advanced RCC (Gleave et al, 1998).

a. Surgery—The role of radical nephrectomy in the management of patients with advanced disease has recently been reevaluated based on the results of randomized clinical trials. Historically, radical nephrectomy was primarily used as a palliative procedure in the setting of metastastic disease for managing patients with severe hemorrhage or unremitting pain. Over the past 20 years, retrospective observations of the potential for nephrectomy to improve response rates of patients receiving biologic response modifier therapy prompted a prospective evaluation of this effect. The Southwest Oncology Group performed a randomized phase III trial, randomizing patients with advanced RCC to nephrectomy followed by interferon-alpha (IFN-α) 2b versus interferon alone. About 241 evaluable patients were enrolled. The median survival of patients undergoing nephrectomy followed by interferon was 11.1 months, compared to 8.1 months in those receiving only interferon (P = 0.05) (Flanigan et al, 2001). A similar, smaller, randomized trial conducted in Europe demonstrated similar findings (Mickisch et al, 2001). These 2 studies have prompted a shift in the standard of care for patients with metastatic RCC and good performance status who desire systemic biologic therapy to include an upfront nephrectomy. Nephrectomy in the presence of metastatic disease (cytoreductive nephrectomy) can be performed via the open approach or laparoscopically. Patients who undergo the nephrectomy laparoscopically may have shorter hospital stay, less blood loss, and obtain adjuvant therapy earlier (Rabets, 2004).

Patients presenting with a solitary metastatic site particularly in the lung that is amenable to surgical resection may be candidates for combined nephrectomy and removal of the metastatic foci (Hoffman et al, 2005). This approach can result in 5-year survival rates of 30–40% with patients developing metachronous solitary lung metastases having a better prognosis (Hoffman et al, 2005). In patients who are destined to receive adjuvant therapy, even limited resection of metastases can yield improved survival further emphasizing the potential benefit of tumor debulking (Vogel et al, 2006).

The important role of surgical resection of solitary brain metastases has been highlighted by several randomized trials demonstrating an improvement in survival in patients with solitary brain metastases who undergo both surgical resection and whole-brain radiotherapy compared with patients who receive only radiation therapy (Patchell et al, 1990; Vecht et al, 1993).

b. Radiation therapy—Radiation therapy is an important method in the palliation of patients with metastatic RCC. Despite the belief that RCC is a relatively radioresistant tumor, effective palliation of metastatic disease to the brain, bone, and lungs is reported in up to two-thirds of patients (Fossa, Kjolseth, and Lund, 1982; Onufrey and Mohiuddin, 1985).

c. Biologic response modifiers—The use of metastatic RCC as a model for the investigation of various biologic response modifiers is a consequence of both the lack of effective chemotherapy and the long-recognized biologic “eccentricities” of this tumor. Spontaneous regression of metastatic RCC is a well-recognized, albeit rare, event (Kavoussi et al, 1986; Vogelzang et al, 1992). Although no specific evidence exists, many believe this phenomenon to be immunologically mediated.

Studies using partially purified human leukocyte interferon in renal cancer were first reported in 1983, with subsequent studies using human lymphoblastoid interferon and, more recently, recombinant interferon-alpha (r-IFN-α). Various doses and schedules of r-IFN-α have demonstrated reproducible overall response rates of 10–15% in advanced renal cancer (Pastore et al, 2001). A modest impact on survival has been demonstrated in some randomized trials of IFN-α. The Medical Research Council compared IFN-α to medroxyprogesterone acetate and demonstrated a 2.5-month median survival improvement favoring the IFN-α arm (Medical Research Council Renal Cancer Collaborators, 1999). Other large randomized studies have failed to demonstrate a survival advantage of IFN-α compared to other biologic response modifiers (Negrier et al, 1998; Motzer et al, 2000). IFN-α is commonly administered 3–5 days/week as a subcutaneous injection. Patients most likely to have a clinical benefit from interferon therapy are those who have minimal tumor burden (ie, primary kidney tumor removed), lung or nodal metastases only, and an excellent performance status. Given the modest activity of interferon in patients with advanced disease, trials of interferon administered in the adjuvant setting to patients at high risk of recurrence have been conducted but have not demonstrated clinical benefit (Trump et al, 1996; Pizzocaro et al, 2001). The experience with beta and gamma interferons has been less extensive. In 1989, Aulitzky and colleagues reported a 30% response rate in a trial of low-dose IFN-γ (Aulitzky et al, 1989). Unfortunately, subsequent trials, including a phase III trial, have demonstrated response rates of <10% (Gleave et al, 1998).

Interleukin-2 (IL-2), a T-cell growth factor, was first identified in 1976. Recombinant IL-2 is the only agent approved by the US Food and Drug Administration for patients with advanced renal carcinoma. Despite initial reports of response rates (complete and partial) of 30%, subsequent experience suggests that the overall response rates of IL-2 is in the 15% range. Although most patients do not benefit from IL-2 therapy, a subset of responding patients do very well, with 10–20% alive 5–10 years after therapy (Gitlitz et al, 2001). Controversy persists regarding
hypoxia-inducible factor. This causes increased levels of having mutations of the Von Hippel Lindau gene resulting in RCC. Both inherited and sporadic RCCs appear to inhibitors of tyrosine kinase and other cell cycle activators of interest in evaluating various antiangiogenic agents and inhibitors of interferon-α, IL-2, and IL-2 plus INF-α have demonstrated higher objective response rates to the combination therapy, with no difference in survival and significantly higher toxicity associated with the combination (Negrier et al., 1998). Biochemotherapy regimens have been evaluated, with a recent phase III trial demonstrating no advantage to the combination of IL-2, INF-α, and fluorouracil versus IL-2 and IFN (Negrier et al., 2000).

**d. Newer biologic agents**—There is currently a lot of interest in evaluating various antiangiogenic agents and inhibitors of tyrosine kinase and other cell cycle activators in RCC. Both inherited and sporadic RCCs appear to have mutations of the Von Hippel Lindau gene resulting in loss of the gene product. This causes increased levels of hypoxia-inducible factor α which in turn promotes increased expression of vascular endothelial growth factor (VEGF) and platelet-derived growth factor (PDGF). The hypervascular nature of RCC is ascribed to increased expression of VEGF and PDGF which promote angiogenesis within tumors. Oral agents such as Bevacizumab and Sunitinib can specifically inhibit receptors for VEGF and PDGF thereby halting tumor angiogenesis and tumor progression. Bevacizumab is a monoclonal antibody that binds and inactivates VEGF A. It has shown the ability to yield partial responses, delay disease progression, and improve survival in patients with advanced renal cancer (Yang, 2003). Recent trials have demonstrated that in patients who have already failed cytokine therapy, Sunitinib can result in partial response in a third of the patients with responses lasting a median of 8 months (Motzer, 2006). In direct comparison with INF-α, Sunitinib yields higher response rates and longer progression-free survival (Motzer et al., 2007). The main toxicity appears to be diarrhea. Sorefenib is another agent that can inhibit VEGF and PDGF and has halted disease progression and can yield partial responses in 10% of patients (Escudier, 2007). Patients treated with Sorefenib demonstrated progression-free survival for 5 months compared to the 2.8-month progression-free survival in those treated with placebo. Side effects such as diarrhea, rash, and hand-foot skin reactions were higher in those receiving Sorefenib. Temsirolimus is an agent which inhibits mTOR, a kinase involved in the VEGF pathway to promote angiogenesis. Temsirolimus has been demonstrated to prolong survival in patients with advanced renal cancer when used as first-line therapy either alone or in combination with interferon α (Motzer, 2006). Further trials of all of these agents are ongoing in order to precisely define their utility in treating various stages of RCC. Sunitinib and Sorefenib have been FDA-approved for use in patients with advanced renal cell cancer.

**B. FOLLOW-UP CARE**

There is no universal agreement on the frequency or studies required in the follow-up care of patients with RCC. A stage-specific follow-up schedule is recommended for patients who have undergone radical or partial nephrectomy (Levy et al., 1998; Hafez, Novick, and Campbell, 1998). Patients with stage T1 disease need less stringent follow-up, with yearly chest x-rays and liver and renal function tests. Those with stage T2 or T3 disease require more frequent follow-up of at least 3-month or 6-month intervals in the early postoperative period. Repeat CT scans of the abdomen should also be obtained, especially in those who have undergone partial nephrectomy, to rule out local recurrence. Patients with metastatic disease who are not undergoing therapy need continued follow-up to provide appropriate supportive care.

**Prognosis**

The prognosis of patients is most clearly related to the stage of disease at presentation. Recent studies report 5-year survival rates for patients with stage T1-T2 disease in the 80–100% range, with stage T3 in the 50–60% range. Patients presenting with metastatic disease have a poorer prognosis, with only a 16–32% 5-year survival rate.

**NEPHROBLASTOMA (WILMS TUMOR)**

Nephroblastoma, also known as Wilms tumor, is the most common solid renal tumor of childhood, accounting for roughly 5% of childhood cancers. Approximately 650 new cases are reported annually. The peak age for presentation is during the third year of life, and there is no sex predilection. The disease is seen worldwide with a similar age of onset and sex distribution. Tumors are commonly unicentric, but they occur in either kidney with equal frequency. In 5% of cases the tumors are bilateral.

Wilms tumor occurs in familial and nonfamilial forms. The National Wilms’ Tumor Study (NWTS) group documented the occurrence of a familial Wilms tumor in approximately 1% of cases (Breslow and Beckwith, 1982). Although a relatively rare neoplasm, Wilms tumor has become a very important model for the study of tumorigenesis and has become a prototypical neoplasm for collaborative clinical trials, with approximately 85% of all new cases diagnosed in North America.
enrolled in NWTS group protocols (Beckwith, 1997). Approximately 10% of patients with Wilms tumors have recognized congenital malformations. Among the more common disorders associated with Wilms tumor are the WAGR syndrome (Wilms, aniridia, genitourinary malformation, mental retardation), overgrowth syndromes, such as Beckwith-Wiedemann syndrome and isolated hemihypertrophy, and non-overgrowth disorders, such as isolated aniridia and trisomy 18 (Weiner, Coppes, and Ritchey, 1998). Genitourinary abnormalities such as hypospadias, cryptorchidism, and renal fusion are found in 4.5–7.5% of patients with unilateral Wilms tumor and in up to 13.4% of those with bilateral disease (Breslow et al, 1993). Some of these genetic syndromes are associated with alterations in the WT1 gene but other genes such as the IGF1, H19, and p57 genes can also be implicated (Beckwith-Wiedemann).

**Etiology**

In 1972 Knudson and Strong proposed a 2-hit hypothesis to explain the earlier age of onset and bilateral presentation in children with a familial history of Wilms tumor. In this hypothesis, the pathogenesis of the sporadic form of Wilms tumor results from two postzygotic mutations in a single cell. In contrast, the familial form of the disease arises after one prezygotic mutation and a subsequent postzygotic event. Karyotypic analyses of Wilms tumor patients with various congenital malformations and loss of heterozygosity studies helped identify a region on the short arm of chromosome 11 (11p13) (Riccardi et al, 1978; Huff, 1994). This work ultimately led to the identification of a gene associated with Wilms tumor development (WT1), which maps to chromosome 11p13 (Coppes, Haber, and Grundy, 1994). Although alterations in this gene have been associated with Wilms tumor and genitourinary abnormalities, only 5–10% of sporadic Wilms tumors have been demonstrated to have WT1 mutations (Varanasi et al, 1994). Genetic linkage studies of families with an inherited susceptibility to Wilms tumors suggest that other Wilms tumor genes exist (Weiner, Coppes, and Ritchey, 1998).

**Pathogenesis & Pathology**

In 1990 Beckwith and colleagues proposed a simplified nomenclature and classification of Wilms tumor precursor lesions known as nephrogenic rests (NR). Two distinct categories of NR were identified and designated as perilobar NR and intralobar NR. One concept of Wilms tumor development proposed that some NR remain dormant for many years, with some undergoing involution and sclerosis and others giving rise to Wilms tumors (Beckwith, Kiviat, and Bonadio, 1990; Beckwith, 1997). The typical Wilms tumor consists of blastemal, epithelial, and stromal elements in varying proportions (Figure 21–10). Tumors composed of blastema and stroma or blastema alone have been described. Pure tubular and papillary forms that are very similar to papillary RCC have also been described.

The NWTS correlated pathologic specimens with clinical outcome and divided various histologic features into favorable and unfavorable prognostic groups. The unfavorable subgroup includes tumors that contain focal or diffuse elements of anaplastic cells or two other neoplastic entities considered not to be Wilms tumor variants, clear cell sarcoma of the kidney and rhabdoid tumor of the kidney (Beckwith and Palmer, 1978; Beckwith, 1997). Favorable histology tumors comprise all Wilms tumors without anaplasia. Anaplastic tumors are characterized by extreme nuclear atypia, hyperdiploidy, and numerous complex translocations. Anaplasia occurs in 5% of Wilms tumors and incidence increases with age. It is more common in African-American children and is linked to p53 mutations (Bardesey et al, 1994). Presence of diffuse anaplasia indicates a worse prognosis compared to focal anaplasia.

Grossly, Wilms tumors are generally large, multilobulated, and gray or tan in color with focal areas of hemorrhage and necrosis. A fibrous pseudocapsule is occasionally seen. Tumor dissemination can occur by direct extension through the renal capsule, hematogenously via the renal vein and vena cava, or via lymphatic spread. Metastatic disease is present at diagnosis in 10–15% of patients, with the lungs (85–95%) and liver (10–15%) the most common sites of involvement. Regional lymphatics are involved in as many as 25% of patients. Metastases to liver, bone, and brain are uncommon.
Tumor Staging

The NWTS staging system is most widely used and is based on surgical and pathologic findings. The original classification was used in the first and second NWTS trials and was modified for NWTS III (D’Angio et al, 1989). Further modifications have been introduced to the staging in the NWTS V study.

Stage I:—Tumor limited to kidney and completely excised. No penetration of renal capsule or involvement of renal sinus vessels. Tumor was not ruptured before or during removal. There is no residual tumor apparent beyond the margins of resection.

Stage II:—Tumor extends beyond the kidney but is completely removed. There is either penetration through the outer surface of the renal capsule, invasion of renal sinus vessels, biopsy of tumor before removal or spillage of tumor locally during removal. There is no residual tumor apparent at or beyond the margins of excision and no lymph node involvement.

Stage III:—Residual non-hematogenous tumor confined to abdomen. Any one or more of the following occur: (a) regional lymph node involvement; (b) diffuse peritoneal contamination by tumor, such as spillage of tumor beyond the flank before or during surgery, or by tumor growth that has penetrated through the peritoneal surface; (c) implants are found on the peritoneal surfaces; (d) the tumor extends beyond the surgical margins either microscopically or grossly; (e) the tumor is not completely resectable because of local infiltration into vital structures; (f) tumor spill not confined to the flank occurred either before or during surgery; (g) transected tumor thrombus.

Stage IV:—Hematogenous metastases to lung, liver, bone, and brain.

Stage V:—Bilateral renal involvement at diagnosis. An attempt should be made to stage each side according to the previously given criteria on the basis of extent of disease before biopsy.

Clinical Findings

A. Symptoms and Signs

The diagnosis of Wilms tumor is most commonly made after the discovery of an asymptomatic mass by a family member or a physician during a routine physical examination. Common symptoms at presentation include abdominal pain and distention, anorexia, nausea and vomiting, fever, and hematuria. The most common sign is an abdominal mass. Hypertension is seen in 25–60% of cases and is caused by elevated renin levels (D’Angio et al, 1982; Pizzo et al, 1989). Up to 30% of patients demonstrate hematuria and coagulopathy can occur in 10%.

B. Laboratory Analysis

Urinalysis may show evidence of hematuria, and anemia may be present, particularly in patients with evidence of subcapsular hemorrhage. Patients with liver metastases may have abnormal serum chemistries.

C. X-Ray Imaging

Abdominal US and CT scanning are performed initially to evaluate the mass. CT of the abdomen is performed with suspected Wilms tumor and can be useful in providing information regarding tumor extension, the status of the contralateral kidney, and the presence of regional adenopathy. CT scanning remains an imperfect technique with a relatively high false-positive rate for hepatic invasion in right-sided tumors, and 7% of cases of surgically confirmed synchronous bilateral Wilms tumors were missed by preoperative CT imaging in the NWTS IV (Ritchey et al, 1995).

Abdominal MRI can sometimes be useful to distinguish between NR and Wilms tumor but is otherwise not routinely indicated. MRI can also provide important information in defining the extent of tumor into the inferior vena cava, including those with intracardiac extension. MRI is limited in that there is no bowel contrast agent, and its use in children requires sedation (Babyn et al, 1995).

Chest x-ray remains the initial examination of choice to evaluate for the presence of lung metastases. The role of a chest CT scan is controversial, and it is probably not indicated for routine use in low-risk patients; however, when done concomitantly with an abdominal CT, chest CT may provide clinically useful information in high-risk patients. If pulmonary metastases are seen on chest x-ray, CT of the chest will not alter current therapy. However, the need for chest CT imaging in patients with negative results of chest x-ray remains controversial, because it is not clear whether those lesions detected by CT alone require more aggressive treatment (Weiner, Coppes, and Ritchey, 1998).

D. Needle Biopsy

Preoperative biopsy is indicated routinely only in tumors deemed too large for safe primary surgical resection and for which preoperative chemotherapy or radiation therapy is planned.

Differential Diagnosis

The differential diagnosis of a flank mass in a child includes hydronephrosis, cystic kidneys, intrarenal neuroblastoma, mesoblastic nephroma, and various very rare sarcomas.

Ultrasonography can confirm the presence of hydronephrosis and evaluate for the presence of cystic kidneys. Neuroblastoma, while pathologically distinct from Wilms tumor, frequently presents in the abdomen as a mass arising from the adrenal glands or paraspinal ganglion. Neuroblastosomas are radiographically indistinguishable from Wilms
tumors, but there are several features that may aid in the differentiation. In contrast to Wilms tumors, which are typically confined to one side of the abdomen, neuroblastomas usually cross the midline. Wilms tumors are intrarenal masses and rarely cause a change in the axis of the kidney, while neuroblastomas may cause an outward and downward displacement of the kidney (drooping lily). Children with neuroblastomas are more likely to present with metastatic disease, and these tumors have a higher frequency of calcification observed radiographically. In addition, neuroblastomas may produce various tumor markers including vanillylmandelic acid and other catecholamines that are not seen in patients with Wilms tumor (Pizzo et al, 1989).

Mesoblastic nephromas are benign hamartomas and cannot be preoperatively distinguished from Wilms tumors. They are most commonly seen in the neonatal period and are typically identified by surgical pathology after nephrectomy. The tumor can occur in adults (Truong et al, 1998).

**Treatment**

The goal of therapy is to provide the highest possible cure rate with the lowest treatment-related morbidity. Significant improvements in survival rates for children with Wilms tumor have been achieved by an improved understanding of the disease and a multimodality approach to therapy, advocated by the NWTS, that incorporates surgery, radiation therapy, and chemotherapy.

**A. SURGICAL MEASURES**

For patients with unilateral kidney involvement whose tumors are deemed surgically resectable (tumors not crossing the midline or involving adjacent visceral organs), radical nephrectomy via a transabdominal incision is the procedure of choice.

Retroperitoneal lymph node dissection is not of proven value and is not recommended. However, biopsy of regional lymphatics (renal hilum and para-aortic nodes) and careful examination of the opposite kidney and the remainder of the abdomen provide crucial data for staging and prognosis. Tumor extending into the vena cava should be removed unless there is evidence of total obstruction. Excision of tumor extending into adjacent organs can be attempted if feasible. Complete excision of all tumor would allow downstaging and decrease amount of additional chemotherapy. A major point of emphasis during surgical extirpation is to avoid spillage because there is evidence that this increases abdominal recurrence of disease (Shamberger et al, 1999; Ross and Kay, 1999).

A child with bilateral Wilms tumor, like an adult with bilateral RCC, requires an individualized approach. Patients with favorable histology tumors can frequently be managed with preoperative chemotherapy followed by renal-sparing surgery (Kumar, Fitzgerald, and Breatnach, 1998). In patients for whom preoperative chemotherapy is planned, a biopsy for diagnosis and staging is indicated (Blute et al, 1987). In some centers, needle aspiration biopsy has proved to be a reliable diagnostic tool when evaluated by experienced pathologists (Hanash, 1989). In patients with unfavorable histology tumors, the therapeutic approach consists of aggressive surgery followed by chemotherapy and radiation therapy.

**B. CHEMOTHERAPY**

Wilms tumor has been long recognized as a chemosensitive neoplasm. Consecutive multicenter randomized trials conducted by the National Wilms’ Tumor Study Group (NWTSG) starting in the 1960s have carefully explored various treatment strategies to determine the role of various antineoplastics and the integration of surgery and radiotherapy, with the goal of optimizing response rates and cure while minimizing the toxicity of therapy. Current studies are focused on continued efforts to minimize toxicity (primarily by decreasing chemotherapy treatment duration and removing radiotherapy) in those favorable subgroups with impressive cure rates, and modifying efforts in poor-risk subgroups to improve response and survival.

Patients with stage I favorable or anaplastic histology and stage II favorable histology tumors undergo surgical resection and have adjuvant chemotherapy with vincristine and dactinomycin combinations without radiotherapy. Patients with stage III and IV favorable histology tumors undergo surgical resection and have adjuvant therapy with vincristine, dactinomycin, and doxorubicin, with adjuvant radiotherapy. Patients with stage II–IV focally anaplastic histology tumors receive therapy similar to that for advanced-stage favorable histology tumors. In the NWTS V, patients with stage II–IV anaplastic tumors are receiving vincristine, doxorubicin, cyclophosphamide, and etoposide (Kalapurakal, 2004). Salvage chemotherapy regimens include agents such as cyclophosphamide, ifosfamide, carboplatin and etoposide. For stage V or bilateral Wilms tumors, diagnosis is established by bilateral biopsies followed by chemotherapy. Second-look surgery may be necessary to reassess response 6–8 weeks after chemotherapy. Renal sparing procedures can be attempted but the rate of renal failure is high.

**C. RADIATION THERAPY**

Wilms tumor has long been recognized as a radiosensitive tumor. Despite the proven efficacy of radiation therapy in children, its use is complicated by its potential for growth disturbances and recognized cardiac, pulmonary, and hepatic toxicities. The development of effective chemotherapy combinations has practically replaced radiation therapy in the preoperative setting. The first and second NWTS trials demonstrated that postoperative radiotherapy was not required for patients with favorable histology stage I disease. NWTS III failed to show an advantage for postoperative radiotherapy in patients with favorable stage
II disease and demonstrated that the relapse rate of patients with stage III disease was no different for patients receiving 1000 cGy compared with the traditional 2000 cGy (D’Angio et al, 1989). Postoperative radiation is recommended for patients with stage III or IV disease with favorable histology, stages II–IV with focal anaplasia and clear cell sarcoma, and all stages of rhabdoid tumor of the kidney (Weiner, Coppes, and Ritchey, 1998).

Prognosis

The multimodality approach to the treatment of children with Wilms tumors has significantly improved outcomes. The 4-year survival of patients with favorable histology Wilms tumor now approaches 90% (Weiner, Coppes, and Ritchey, 1998). The most important negative prognostic factors remain the unfavorable histologic subtypes (clear cell sarcoma, rhabdoid, and anaplastic tumors). Although the addition of doxorubicin in NWTS III significantly improved the 2-year survival rate for patients with clear cell sarcomas (61.5–90.3%), it did not affect the survival of children with rhabdoid tumors. Analysis of patients with bilateral Wilms tumors registered with NWTS II and III revealed a 3-year survival rate of 82% (Blute et al, 1987).

Future challenges include improvements in therapy for patients with anaplastic tumors (stages II–IV), clear cell sarcoma, and rhabdoid tumors, and efforts to improve outcomes in favorable histology tumors while decreasing both short-term and late toxicities. Long-term toxicity in these patients include renal failure, cardiac toxicity with congestive heart failure due to chemotherapy, and lung radiation as well as a higher risk for secondary malignancies.

SARCOMA OF THE KIDNEY

Primary sarcomas of the kidney are rare, with a reported incidence ranging from 1% to 3% of all malignant renal neoplasms (Vogelzang et al, 1993; Srinivas et al, 1984). Renal sarcomas are most commonly present in patients in the fifth decade of life, and there is a slight male predominance. Flank or abdominal pain and weight loss are the most frequent presenting symptoms. Primary renal sarcomas may be difficult to distinguish histologically from the sarcomatoid variant of renal carcinoma (Bonsib et al, 1987). Leiomyosarcomas compose approximately 50% of all renal sarcomas and occur with a female predominance of 2:1 (Loomis, 1972). The remaining 40–50% of renal sarcomas consist of fibrosarcomas, liposarcomas, hemangio-pericytomas, osteogenic sarcoma, and malignant schwannomas. Renal sarcomas are typically of renal capsular origin. They present with symptoms analogous to those of other renal masses and tend to exhibit aggressive local spread with distant metastases to lung and liver as late findings.

Radical nephrectomy for localized disease is the only effective therapy. Adjuvant radiotherapy has been demonstrated to decrease the incidence of local recurrence in patients with resectable retroperitoneal sarcomas; however, there is no improvement in overall survival (Kinsella et al, 1988). Various chemotherapeutic agents, including doxorubicin, dacarbazine, and ifosfamide, have activity in the treatment of metastatic disease, but responses are typically partial and of brief duration.

SECONDARY RENAL TUMORS

The kidney is a frequent site for metastatic spread of both solid and hematologic tumors. Wagle, Moore, and Murphy (1975) surveyed 4413 autopsies at a major cancer center and found 81 (18%) cases of secondary carcinoma of the kidney (hematologic tumors were excluded). The most frequent primary site of cancer was lung (20%), followed by breast (12%), stomach (11%), and renal (9%). The authors noted that metastases to the renal parenchyma typically demonstrated capsular and stromal invasion with sparing of the renal pelvis and that bilateral secondary renal involvement was found in approximately 50% of cases.

Albuminuria and hematuria are relatively common findings in patients with secondary renal metastases; however, pain and renal insufficiency are rare (Wagle, Moore, and Murphy, 1975; Olsson, Moyer, and Laferte, 1971). Secondary metastatic disease to the kidneys tends to be a late event, frequently in the setting of widely disseminated disease, which typically portends a poor prognosis. Therapy is dictated by the responsiveness of the primary neoplasm; that is, patients with breast and ovarian cancers for which there is effective therapy are more likely to respond than patients with primary lung or gastrointestinal cancers.

Autopsy series have reported clinically evident renal invasion by lymphoma to be 0.5–7%, with the rates of Hodgkin’s and non-Hodgkin’s lymphoma distributed equally (Goffinet et al, 1977; Weimar et al, 1981). Renal involvement is usually in the form of bilateral, multiple, discrete tumor nodules. Renal involvement by non-Hodgkin’s lymphoma is typically characterized by diffuse, aggressive histologic findings (ie, diffuse large cell) in the setting of extensive disease. Therapy typically consists of combination chemotherapy, with the prognosis of patients similar to that of patients without renal involvement but with widely disseminated, aggressive lymphomas (Geffen et al, 1985).

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Neoplasms of the Prostate Gland

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The prostate gland is the male organ most commonly afflicted with either benign or malignant neoplasms. It comprises the most proximal aspect of the urethra. Anatomically it resides in the true pelvis, separated from the pubic symphysis anteriorly by the retropubic space (space of Retzius). The posterior surface of the prostate is separated from the rectal ampulla by Denonvilliers’ fascia. The base of the prostate is continuous with the bladder neck, and the apex of the prostate rests on the upper surface of the urogenital diaphragm. Laterally, the prostate is related to the levator ani musculature. Its arterial blood supply is derived from branches of the internal iliac artery (inferior vesical and middle rectal arteries). Venous drainage is via the dorsal venous complex, which receives the deep dorsal vein of the penis and vesical branches before draining into the internal iliac veins. Innervation is from the pelvic plexus. The normal prostate measures 3–4 cm at the base, 4–6 cm in cephalocaudad, and 2–3 cm in anteroposterior dimensions.

McNeal has popularized the concept of zonal anatomy of the prostate. Three distinct zones have been identified (Figure 22–1). The peripheral zone accounts for 70% of the volume of the young adult prostate, the central zone accounts for 25%, and the transition zone accounts for 5%. These anatomic zones have distinct ductal systems but, more important, are differentially afflicted with neoplastic processes. Sixty to seventy percent of carcinomas of the prostate (CaP) originate in the peripheral zone, 10–20% in the transition zone, and 5–10% in the central zone (McNeal et al, 1988). Benign prostatic hyperplasia (BPH) uniformly originates in the transition zone (Figure 22–2).

**BENIGN PROSTATIC HYPERPLASIA**

**Incidence & Epidemiology**

BPH is the most common benign tumor in men, and its incidence is age related. The prevalence of histologic BPH in autopsy studies rises from approximately 20% in men aged 41–50, to 50% in men aged 51–60, and to >90% in men older than 80. Although clinical evidence of disease occurs less commonly, symptoms of prostatic obstruction are also age related. At age 55, approximately 25% of men report obstructive voiding symptoms. At age 75, 50% of men complain of a decrease in the force and caliber of their urinary stream.

Risk factors for the development of BPH are poorly understood. Some studies have suggested a genetic predisposition, and some have noted racial differences. Approximately 50% of men under the age of 60 who undergo surgery for BPH may have a heritable form of the disease. This form is most likely an autosomal dominant trait, and first-degree male relatives of such patients carry an increased relative risk of approximately fourfold.

**Etiology**

The etiology of BPH is not completely understood, but it seems to be multifactorial and endocrine controlled. The prostate is composed of both stromal and epithelial elements, and each, either alone or in combination, can give rise to hyperplastic nodules and the symptoms associated with BPH. Each element may be targeted in medical management schemes.

Observations and clinical studies in men have clearly demonstrated that BPH is under endocrine control. Castration results in the regression of established BPH and improvement in urinary symptoms. Additional investigations have demonstrated a positive correlation between levels of free testosterone and estrogen and the volume of BPH. The latter may suggest that the association between aging and BPH might result from the increased estrogen levels of aging causing induction of the androgen receptor, which thereby sensitizes the prostate to free testosterone. However, no studies to date have been able to demonstrate elevated estrogen receptor levels in human BPH.

**Pathology**

As discussed earlier, BPH develops in the transition zone. It is truly a hyperplastic process resulting from an increase in cell number. Microscopic evaluation reveals a nodular growth pattern that is composed of varying amounts of stroma and epithelium. Stroma is composed of varying amounts of collagen and smooth muscle. The differential representation of the histologic components of BPH explains, in part, the potential responsiveness to medical therapy. Thus alpha-blocker therapy may result in excellent responses in patients with BPH that has a significant...
component of smooth muscle, while those with BPH predominantly composed of epithelium might respond better to 5-alpha-reductase inhibitors. Patients with significant components of collagen in the stroma may not respond to either form of medical therapy. Unfortunately, one cannot reliably predict responsiveness to a specific therapy (see below).

As BPH nodules in the transition zone enlarge, they compress the outer zones of the prostate, resulting in the formation of a so-called surgical capsule. This boundary separates the transition zone from the peripheral zone and serves as a cleavage plane for open enucleation of the prostate during open simple prostatectomies performed for BPH.

Pathophysiology

One can relate the symptoms of BPH to either the obstructive component of the prostate or the secondary response of the bladder to the outlet resistance. The obstructive component can be subdivided into the mechanical and the dynamic obstruction.

As prostatic enlargement occurs, mechanical obstruction may result from intrusion into the urethral lumen or bladder neck, leading to a higher bladder outlet resistance. Prior to the zonal classification of the prostate, urologists often referred to the “3 lobes” of the prostate, namely, the median and the 2 lateral lobes. Prostatic size on digital rectal examination (DRE) correlates poorly with symptoms, in part, because the median lobe is not readily palpable.

The dynamic component of prostatic obstruction explains the variable nature of the symptoms experienced by patients. The prostatic stroma, composed of smooth muscle and collagen, is rich in adrenergic nerve supply. The level of autonomic stimulation thus sets a tone to the prostatic urethra. Use of alpha-blocker therapy decreases this tone, resulting in a decrease in outlet resistance.
CHAPTER 22

The irritative voiding complaints (see below) of BPH result from the secondary response of the bladder to the increased outlet resistance. Bladder outlet obstruction leads to detrusor muscle hypertrophy and hyperplasia as well as collagen deposition. Although the latter is most likely responsible for a decrease in bladder compliance, detrusor instability is also a factor. On gross inspection, thickened detrusor muscle bundles are seen as trabeculation on cystoscopic examination. If left unchecked, mucosal herniation between detrusor muscle bundles ensues, causing diverticula formation (so-called false diverticula composed of only mucosa and serosa).

Clinical Findings

A. SYMPTOMS

The symptoms of BPH can be divided into obstructive and irritative complaints. Obstructive symptoms include hesitancy, decreased force and caliber of stream, sensation of incomplete bladder emptying, double voiding (urinating a second time within 2 hours of the previous void), straining to urinate, and post-void dribbling. Irritative symptoms include urgency, frequency, and nocturia.

The self-administered questionnaire developed by the American Urological Association (AUA) is both valid and reliable in identifying the need to treat patients and in monitoring their response to therapy. The AUA Symptom Score questionnaire (Table 22–1) is perhaps the single most important tool used in the evaluation of patients with BPH and is recommended for all patients before the initiation of therapy. This assessment focuses on 7 items that ask patients to quantify the severity of their obstructive or irritative complaints on a scale of 0–5. Thus, the score can range from 0 to 35. A symptom score of 0–7 is considered mild, 8–19 is considered moderate, and 20–35 is considered severe. The relative distribution of scores for BPH patients and control subjects is, respectively, 20% and 83% in those with mild scores, 57% and 15% in those with moderate scores, and 23% and 2% in those with severe scores (McConnell et al., 1994).

A detailed history focusing on the urinary tract excludes other possible causes of symptoms that may not result from the prostate, such as urinary tract infection, neurogenic bladder, urethral stricture, or prostate cancer.

B. SIGNS

A physical examination, DRE, and focused neurologic examination are performed on all patients. The size and consistency of the prostate is noted, even though prostate size, as determined by DRE, does not correlate with severity of symptoms or degree of obstruction. BPH usually results in a smooth, firm, elastic enlargement of the prostate. Induration, if detected, must alert the physician to the possibility of cancer and the need for further evaluation (ie, prostate-specific antigen [PSA], transrectal ultrasound [TRUS], and biopsy).

C. LABORATORY FINDINGS

A urinalysis to exclude infection or hematuria and serum creatinine measurement to assess renal function are required. Renal insufficiency may be observed in 10% of patients with prostatism and warrants upper-tract imaging. Patients with renal insufficiency are at an increased risk of developing postoperative complications following surgical intervention for BPH. Serum PSA is considered optional, but most physicians will include it in the initial evaluation. PSA, compared with DRE alone, certainly increases the ability to detect CaP, but because there is much overlap between levels seen in BPH and CaP, its use remains controversial (see Screening for CaP).

D. IMAGING

Upper-tract imaging (intravenous pyelogram or renal ultrasound) is recommended only in the presence of concomitant urinary tract disease or complications from BPH (eg, hematuria, urinary tract infection, renal insufficiency, history of stone disease).

E. CYSTOSCOPY

Cystoscopy is not recommended to determine the need for treatment but may assist in choosing the surgical approach in patients opting for invasive therapy.

F. ADDITIONAL TESTS

Cystometrograms and urodynamic profiles are reserved for patients with suspected neurologic disease or those who
Table 22–1. Questionnaire for American Urological Association Symptom Score.

<table>
<thead>
<tr>
<th>URINARY SYMPTOMS (SYMPTOM SCORE CRITERIA)</th>
<th>AUA Score</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Not at all</td>
</tr>
<tr>
<td>1. Incomplete emptying</td>
<td>0</td>
</tr>
<tr>
<td>Over the past month, how often</td>
<td></td>
</tr>
<tr>
<td>have you had a sensation of</td>
<td></td>
</tr>
<tr>
<td>not emptying your bladder</td>
<td></td>
</tr>
<tr>
<td>completely after you finished urinating?</td>
<td></td>
</tr>
<tr>
<td>2. Frequency</td>
<td>0</td>
</tr>
<tr>
<td>Over the past month, how often</td>
<td></td>
</tr>
<tr>
<td>have you had to urinate again</td>
<td></td>
</tr>
<tr>
<td>less than two hours after you</td>
<td></td>
</tr>
<tr>
<td>finished urinating?</td>
<td></td>
</tr>
<tr>
<td>3. Intermittency</td>
<td>0</td>
</tr>
<tr>
<td>Over the past month, how often</td>
<td></td>
</tr>
<tr>
<td>have you found you stopped</td>
<td></td>
</tr>
<tr>
<td>and started again several times when you</td>
<td></td>
</tr>
<tr>
<td>urinate?</td>
<td></td>
</tr>
<tr>
<td>4. Urgency</td>
<td>0</td>
</tr>
<tr>
<td>Over the past month, how often</td>
<td></td>
</tr>
<tr>
<td>have you found it difficult to</td>
<td></td>
</tr>
<tr>
<td>postpone urination?</td>
<td></td>
</tr>
<tr>
<td>5. Weak stream</td>
<td>0</td>
</tr>
<tr>
<td>Over the past month, how often</td>
<td></td>
</tr>
<tr>
<td>have you had a weak urinary</td>
<td></td>
</tr>
<tr>
<td>stream?</td>
<td></td>
</tr>
<tr>
<td>6. Straining</td>
<td>0</td>
</tr>
<tr>
<td>Over the past month, how often</td>
<td></td>
</tr>
<tr>
<td>have you had to push or strain</td>
<td></td>
</tr>
<tr>
<td>to begin urination?</td>
<td></td>
</tr>
<tr>
<td>7. Nocturia</td>
<td>None</td>
</tr>
<tr>
<td>Over the past month, how many</td>
<td></td>
</tr>
<tr>
<td>times did you most typically get</td>
<td></td>
</tr>
<tr>
<td>up to urinate from the time you</td>
<td></td>
</tr>
<tr>
<td>went to bed at night until the</td>
<td></td>
</tr>
<tr>
<td>time you got up in the morning?</td>
<td></td>
</tr>
</tbody>
</table>

AUA Symptom Score = sum of questions A1 to A7

QUALITY OF LIFE DUE TO URINARY PROBLEMS

<table>
<thead>
<tr>
<th>Delighted</th>
<th>Pleased</th>
<th>Mostly satisfied</th>
<th>Mixed—about equally satisfied and unsatisfied</th>
<th>Mostly dissatisfied</th>
<th>Unhappy</th>
<th>Terrible</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
<td>6</td>
</tr>
</tbody>
</table>

If you were to spend the rest of your life with your urinary condition just the way it is now, how would you feel about that?

have failed prostate surgery. Measurement of flow rate, determination of post-void residual urine, and pressure-flow studies are considered optional.

**Differential Diagnosis**

Other obstructive conditions of the lower urinary tract, such as urethral stricture, bladder neck contracture, bladder stone, or CaP, must be entertained when evaluating men with presumptive BPH. A history of previous urethral instrumentation, urethritis, or trauma should be elucidated to exclude urethral stricture or bladder neck contracture. Hematuria and pain are commonly associated with bladder stones. CaP may be detected by abnormalities on the DRE or an elevated PSA (see below).

A urinary tract infection, which can mimic the irritative symptoms of BPH, can be readily identified by urinalysis and culture; however, a urinary tract infection can also be a complication of BPH. Although irritative voiding complaints are also associated with carcinoma of the bladder, especially carcinoma in situ, the urinalysis usually shows evidence of hematuria. Likewise, patients with neurogenic bladder disorders may have many of the signs and symptoms of BPH, but a history of neurologic disease, stroke, diabetes mellitus, or back injury may be present as well. In addition, examination may show diminished perineal or lower extremity sensation or alterations in rectal sphincter tone or the bulbocavernosus reflex. Simultaneous alterations in bowel function (constipation) might also alert one to the possibility of a neurologic origin.

**Treatment**

After patients have been evaluated, they should be informed of the various therapeutic options for BPH. It is advisable for patients to consult with their physicians to make an educated decision on the basis of the relative efficacy and side effects of the treatment options.

Specific treatment recommendations can be offered for certain groups of patients. For those with mild symptoms (symptom score 0–7), watchful waiting only is advised. On the other end of the therapeutic spectrum, absolute surgical indications include refractory urinary retention (failing at least one attempt at catheter removal), recurrent urinary tract infection from BPH, recurrent gross hematuria from BPH, bladder stones from BPH, renal insufficiency from BPH, or large bladder diverticula (McConnell et al, 1994).

**A. WATCHFUL WAITING**

Very few studies on the natural history of BPH have been reported. The risk of progression or complications is uncertain. However, in men with symptomatic BPH, it is clear that progression is not inevitable and that some men undergo spontaneous improvement or resolution of their symptoms. Retrospective studies on the natural history of BPH are inherently subject to bias, related to patient selection and the type and extent of follow-up. Very few prospective studies addressing the natural history of BPH have been reported. Recently, a large randomized study compared finasteride with placebo in men with moderately to severely symptomatic BPH and enlarged prostates on DRE (McConnell et al, 1998). Patients in the placebo arm of the study had a 7% risk of developing urinary retention over 4 years.

As mentioned earlier, watchful waiting is the appropriate management of men with mild symptom scores (0–7). Men with moderate or severe symptoms can also be managed in this fashion if they so choose. Neither the optimal interval for follow-up nor specific end points for intervention have been defined.

**B. MEDICAL THERAPY**

1. **Alpha-blockers**—The human prostate and bladder base contains alpha-1-adrenoreceptors, and the prostate shows a contractile response to corresponding agonists. The contractile properties of the prostate and bladder neck seem to be mediated primarily by the subtype alpha-1a receptors. Alpha-blockade has been shown to result in both objective and subjective degrees of improvement in the symptoms and signs of BPH in some patients. Alpha-blockers can be classified according to their receptor selectivity as well as their half-life (Table 22–2).

Phenoxybenzamine and prazosin have comparable efficacy with respect to symptomatic relief, but the higher side-effect profile of phenoxybenzamine, associated with its

<table>
<thead>
<tr>
<th>Classification</th>
<th>Oral Dosage</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Alpha-blockers</strong></td>
<td></td>
</tr>
<tr>
<td>Nonselective</td>
<td></td>
</tr>
<tr>
<td>Phenoxybenzamine</td>
<td>10 mg twice a day</td>
</tr>
<tr>
<td>Alpha-1, short-acting</td>
<td></td>
</tr>
<tr>
<td>Prazosin</td>
<td>2 mg twice a day</td>
</tr>
<tr>
<td>Alpha-1, long-acting</td>
<td></td>
</tr>
<tr>
<td>Terazosin</td>
<td>5 or 10 mg daily</td>
</tr>
<tr>
<td>Doxazosin</td>
<td>4 or 8 mg daily</td>
</tr>
<tr>
<td>Alpha-1a selective</td>
<td></td>
</tr>
<tr>
<td>Tamsulosin</td>
<td>0.4 or 0.8 mg daily</td>
</tr>
<tr>
<td>Alfuzosin</td>
<td>10 mg daily</td>
</tr>
</tbody>
</table>

| **5-alpha-reductase inhibitors**      |                   |
| Finasteride                           | 5 mg daily        |
| Dutasteride                           | 0.5 mg daily     |
| Subcutaneous implant                  | Yearly           |
| Triptorelin pamoate                   | 3.75 mg every month|
lack of alpha-receptor specificity, precludes its use in BPH patients. Dose titration is necessary with prazosin, with typical therapy started at 1 mg at bedtime for 3 nights, then increased to 1 mg twice a day, which is titrated up to 2 mg twice a day if necessary. At higher doses, little additional symptomatic improvement is observed and side-effect profiles worsen. Typical side effects include orthostatic hypotension, dizziness, tiredness, retrograde ejaculation, rhinitis, and headache.

Long-acting alpha-blockers make once-a-day dosing possible, but dose titration is still necessary. Terazosin is initiated at 1 mg daily for 3 days and increased to 2 mg daily for 11 days and then to 5 mg/day. Dosage can be escalated to 10 mg daily if necessary. Therapy with doxazosin is started at 1 mg daily for 7 days and increased to 2 mg daily for 7 days, and then to 4 mg daily. Dosage can be escalated to 8 mg daily if necessary. Side effects are similar to those described for prazosin.

The most recent advance in alpha-blocker therapy is related to the identification of subtypes of alpha-1-receptors. Selective blockade of the alpha-1a receptors, which are localized in the prostate and bladder neck, results in fewer systemic side effects (orthostatic hypotension, dizziness, tiredness, rhinitis, and headache), thus obviating the need for dose titration. Tamsulosin is initiated at 0.4 mg daily and can be increased to 0.8 mg daily if necessary. Alfuzosin is a functionally uroselective alpha-1-adrenergic antagonist. Like tamsulosin, no dosage titration is needed for an extended release preparation (10 mg), and this agent has fewer adverse cardiovascular effects compared to nonspecific alpha-blocker therapy.

Several randomized, double-blind, placebo-controlled trials, individually comparing alpha-blockers with placebo, have demonstrated the safety and efficacy of all of these agents.

2. **5-Alpha-reductase inhibitors**—Finasteride is a 5-alpha-reductase inhibitor that blocks the conversion of testosterone to dihydrotestosterone. This drug affects the epithelial component of the prostate, resulting in a reduction in the size of the gland and improvement in symptoms. Six months of therapy are required to see the maximum effects on prostate size (20% reduction) and symptomatic improvement.

Several randomized, double-blind, placebo-controlled trials have compared finasteride with placebo. Efficacy, safety, and durability are well established. However, symptomatic improvement is seen only in men with enlarged prostates (>40 cm³). Side effects include decreased libido, decreased ejaculate volume, and impotence. Serum PSA is reduced by approximately 50% in patients being treated with finasteride, but individual values may vary.

Dutasteride differs from finasteride as it inhibits both isoenzymes of 5-alpha-reductase. Similar to finasteride, it reduces serum prostatic specific antigen and total prostate volume. Randomized, placebo-controlled trials have shown the efficacy of dutasteride in symptomatic relief, symptoms scores, peak urinary flow rate, and reduced risk of acute urinary retention and the need for surgery. The main side effects are erectile dysfunction, decreased libido, gynecomastia, and ejaculation disorders.

3. **Combination therapy**—The first randomized, double-blind, placebo-controlled study investigating combination alpha-blocker and 5-alpha-reductase inhibitor therapy was a four-arm Veterans Administration Cooperative Trial comparing placebo, finasteride alone, terazosin alone, and combination finasteride and terazosin (Lepor et al, 1996). Over 1200 patients participated, and significant decreases in symptom score and increases in urinary flow rates were seen only in the arms containing terazosin. However, one must note that enlarged prostates were not an entry criterion; in fact, prostate size in this study was much smaller than that in previous controlled trials using finasteride (32 versus 52 cm³). McConnell and colleagues conducted a long-term, double-blind trial involving 3047 men to compare the effects of placebo, doxazosin, finasteride, and combination therapy on measures of the clinical progression of BPH (McConnell et al, 2003). The risk of overall clinical progression—defined as an increase above baseline of at least 4 points in the AUA Symptom Score, acute urinary retention, urinary incontinence, renal insufficiency, or recurrent urinary tract infection—was significantly reduced by doxazosin (39% risk reduction) and finasteride (34% risk reduction), as compared with placebo. The reduction in risk associated with combination therapy (66% risk reduction) was significantly greater than that associated with doxazosin or finasteride alone. Patients most likely to benefit from combination therapy are those in whom baseline risk of progression is very high, generally patients with larger glands and higher PSA values.

4. **Phytotherapy**—Phytotherapy refers to the use of plants or plant extracts for medicinal purposes. The use of phytotherapy in BPH has been popular in Europe for years, and its use in the United States is growing as a result of patient-driven enthusiasm. Several plant extracts have been popularized, including the saw palmetto berry, *(Serenoa repens)* the bark of *Pygeum africanum*, the roots of *Echinacea purpurea* and *Hypoxis rooperi*, pollen extract, and the leaves of the trembling poplar. *S. repens* has been the most well-studied agent usually at 320 mg/day. Although a favorable effect on symptom score and flow rate has been noted in some studies, such findings have not been consistently demonstrated (Wilt et al, 2002). A recent published prospective, randomized clinical trial of saw palmetto showed no benefit beyond placebo for symptom improvement or urinary flow rate. The mechanisms of action of these phyotherapies are unknown, and the efficacy and safety of these agents have not been well tested in multicenter, randomized, double-blind, placebo-controlled studies.
C. Conventional Surgical Therapy

1. Transurethral resection of the prostate—Ninety-five percent of simple prostatectomies can be done endoscopically. Most of these procedures involve the use of a spinal anesthetic and require a 1- to 2-day hospital stay. Symptom score and flow rate improvement with transurethral resection of the prostate (TURP) is superior to that of any minimally invasive therapy. The length of hospital stay of patients undergoing TURP, however, is greater. Much controversy revolves around possible higher rates of morbidity and mortality associated with TURP in comparison with those of open surgery, but the higher rates observed in one study were probably related to more significant comorbidities in the TURP patients than in the patients undergoing open surgery. Several other studies could not confirm the difference in mortality when results were controlled for age and comorbidities. Risks of TURP include retrograde ejaculation (75%), impotence (5–10%), and incontinence (<1%). Complications include bleeding, urethral stricture or bladder neck contracture, perforation of the prostate capsule with extravasation, and if severe, TUR syndrome resulting from a hypervolemic, hyponatremic state due to absorption of the hypotonic irrigating solution. Clinical manifestations of the TUR syndrome include nausea, vomiting, confusion, hypertension, bradycardia, and visual disturbances. The risk of the TUR syndrome increases with resection times >90 minutes. Treatment includes diuresis and, in severe cases, hypertonic saline administration.

2. Transurethral incision of the prostate—Men with moderate to severe symptoms and a small prostate often have posterior commissure hyperplasia (elevated bladder neck). These patients will often benefit from an incision of the prostate. This procedure is more rapid and less morbid than TURP. Outcomes in well-selected patients are comparable, although a lower rate of retrograde ejaculation with transurethral incision has been reported (25%). The technique involves two incisions using the Collins knife at the 5- and 7-o’clock positions. The incisions are started just distal to the urethral orifices and are extended outward to the verumontanum.

3. Open simple prostatectomy—When the prostate is too large to be removed endoscopically, an open enucleation is necessary. What constitutes “too large” is subjective and will vary depending upon the surgeon’s experience with TURP. Glands >100 g are usually considered for open enucleation. Open prostatectomy may also be initiated when concomitant bladder diverticulum or a bladder stone is present or if dorsal lithotomy positioning is not possible.

Open prostatectomies can be done with either a suprapubic or retropubic approach. A simple suprapubic prostatectomy is performed transvesically and is the operation of choice in dealing with concomitant bladder pathology. After the bladder is opened, a semicircular incision is made in the bladder mucosa, distal to the trigone. The dissection plane is initiated sharply, and then blunt dissection with the finger is performed to remove the adenoma. The apical dissection should be done sharply to avoid injury to the distal sphincteric mechanism. After the adenoma is removed, hemostasis is attained with suture ligatures, and both a urethral and a suprapubic catheter are inserted before closure.

In a simple retropubic prostatectomy, the bladder is not entered. Rather, a transverse incision is made in the surgical capsule of the prostate, and the adenoma is enucleated as described above. Only a urethral catheter is needed at the end of the procedure.

D. Minimally Invasive Therapy

1. Laser therapy—Many different techniques of laser surgery for the prostate have been described. Two main energy sources of lasers have been utilized—Nd:YAG and holmium:YAG.

Several different coagulation necrosis techniques have been described. Transurethral laser-induced prostatectomy (TULIP) is done with TRUS guidance. The TULIP device is placed in the urethra, and TRUS is used to direct the device as it is slowly pulled from the bladder neck to the apex. The depth of treatment is monitored with ultrasound.

Most urologists prefer to use visually directed laser techniques. Under cystoscopic control, the laser fiber is pulled through the prostate at several designated areas, depending upon the size and configuration of the prostate. Four quadrant and sextant approaches have been described for lateral lobes, with additional treatments directed at enlarged median lobes. Coagulative techniques do not create an immediate visual defect in the prostatic urethra, but rather tissue is sloughed over the course of several weeks and up to 3 months following the procedure.

Visual contact ablative techniques are more time-consuming procedures because the fiber is placed in direct contact with the prostate tissue, which is vaporized. An immediate defect is obtained in the prostatic urethra, similar to that seen during TURP.

Interstitial laser therapy places fibers directly into the prostate, usually under cystoscopic control. At each puncture, the laser is fired, resulting in submucosal coagulative necrosis. This technique may result in fewer irritative voiding symptoms, because the urethral mucosa is spared and prostate tissue is resorbed by the body rather than sloughed.

Advantages of laser surgery include (1) minimal blood loss, (2) rare instances of TUR syndrome, (3) ability to treat patients receiving anticoagulation therapy, and (4) ability to be done as an outpatient procedure. Disadvantages include (1) lack of availability of tissue for pathologic examination, (2) longer postoperative catheterization time, (3) more irritative voiding complaints, and (4) high cost of laser fibers and generators.
Large-scale, multicenter, randomized studies with long-term follow-up are needed to compare laser prostate surgery with TURP and other forms of minimally invasive surgery.

2. Transurethral electrovaporization of the prostate—Transurethral electrovaporization uses the standard resectoscope but replaces a conventional loop with a variation of a grooved rollerball. High current densities cause heat vaporization of tissue, resulting in a cavity in the prostatic urethra. Because the device requires slower sweeping speeds over the prostatic urethra, and the depth of vaporization is approximately one-third of a standard loop, the procedure usually takes longer than a standard TURP. Long-term comparative data are needed.

3. Hyperthermia—Microwave hyperthermia is most commonly delivered with a transurethral catheter. Some devices cool the urethral mucosa to decrease the risk of injury. However, if temperatures are not >45°C, cooling is unnecessary. Improvement in symptom score and flow rate is obtained, but as with laser surgery, large-scale, randomized studies with long-term follow-up are needed to assess durability and cost effectiveness.

4. Transurethral needle ablation of the prostate—Transurethral needle ablation uses a specially designed urethral catheter that is passed into the urethra. Interstitial radio frequency needles are then deployed from the tip of the catheter, piercing the mucosa of the prostatic urethra. The use of radio frequencies to heat the tissue results in a coagulative necrosis. This technique is not adequate treatment for bladder neck and median lobe enlargement. Subjective and objective improvement in voiding occurs, but as mentioned above, comparative long-term randomized studies are lacking.

5. High-intensity focused ultrasound—High-intensity focused ultrasound is another means of performing thermal tissue ablation. A specially designed dual-function ultrasound probe is placed in the rectum. This probe allows transrectal imaging of the prostate and also delivers short bursts of high-intensity focused ultrasound energy, which heats the prostate tissue and results in coagulative necrosis. Bladder neck and median lobe enlargement are not adequately treated with this technique. Although ongoing clinical trials demonstrate some improvement in symptom score and flow rate, the durability of response is unknown.

6. Intraurethral stents—Intraurethral stents are devices that are endoscopically placed in the prostatic fossa and are designed to keep the prostatic urethra patent. They are usually covered by urothelium within 4–6 months after insertion. These devices are typically used for patients with limited life expectancy who are not deemed to be appropriate candidates for surgery or anesthesia. With the advent of other minimally invasive techniques requiring minimal anesthesia (conscious sedation or prostatic blocks), their application has become more limited.

Prostate cancer is the most common cancer detected in American men. Approximately 230,000 American men will be diagnosed with prostate cancer this year (Jemal et al, 2005). Although prostate cancer is the second leading cause of cancer death for men, mortality rates have been declining since the mid-1990s. However, because of reductions in death due to cardiovascular disease and the aging of American men, prostate cancer will continue to be a major health-care concern unless more effective forms of prevention and treatment are identified (Chan, Jou et al, 2004). Of all cancers, the prevalence of CaP increases the most rapidly with age. However, unlike most cancers, which have a peak age of incidence, the incidence of CaP continues to increase with advancing age. The lifetime risk of a 50-year-old man for latent CaP (detected as an incidental finding at autopsy, not related to the cause of death) is 40%; for clinically apparent CaP, 9.5%; and for death from CaP, 2.9%. Thus, many prostate cancers are indolent and inconsequential to the patient while others are virulent, and if detected too late or left untreated, they result in a patient’s death. This broad spectrum of biological activity can make decision making for individual patients difficult.

Several risk factors for prostate cancer have been identified. As discussed above, increasing age heightens the risk for CaP. Which of the factors associated with the aging process are responsible for this observation is unknown. The probability of CaP developing in a man under the age of 40 is 1 in 10,000; for men 40–59 it is 1 in 103, and for men 60–79 it is 1 in 8. African Americans are at a higher risk for CaP than whites. In addition, African American men tend to present at a later stage of disease than whites. Controversial data have been reported suggesting that mortality from this disease may also be higher for African Americans. A positive family history of CaP also increases the relative risk for CaP. The age of disease onset in the family member with the diagnosis of CaP affects a patient’s relative risk. If the age of onset is 70, the relative risk is increased fivefold; and if the age of onset is 50, the relative risk is increased fourfold; if the age of onset is 60, the relative risk is increased fivefold; and if the age of onset is 50, the relative risk is increased sevenfold. Although diagnostic biases exist between countries, differences in the incidence of prostate cancer are real. These differences may be related to differences in diet (Chan et al, 2005). Epidemiologic studies have shown that the incidence of clinically significant prostate cancer is much lower in parts of the world where people eat a predominantly low fat, plant-based diet. In addition, migrant studies demonstrate that when men from a low-risk country move to the United States and begin eating a westernized diet, their rates of prostate cancer increase severalfold and approach that of the host country. Total fat intake, animal fat intake, and red meat intake are associated with an increased risk of prostate can-
cancer, whereas intake of fish is associated with a decreased risk. There is considerable controversy on the impact of obesity on prostate cancer. Some studies suggest that obesity is associated with an increased risk of more advanced disease and a higher recurrence rate after treatment. Additionally, lycopene, selenium, omega-3 fatty acids (fish), and vitamin E intake have been shown to be protective, whereas vitamin D and calcium increase risk. Previous vasectomy has been suggested as a factor that heightens the risk for CaP, but these data are controversial.

MOLECULAR GENETICS & PATHOBIOLOGY

Molecular profiling of human tissues has identified differential expression of specific genes and proteins in the progression from normal precursor tissue to preneoplastic lesions to cancer (both androgen dependent and independent). In doing so diagnostic, prognostic, and therapeutic markers have been discovered. Chromosomal rearrangements or copy number abnormalities at 8p, 10q, 11q, 13q, 16q, 17p, and 18q have been described in prostate cancers. Some of these such as specific loss at 8p23.2 and/or gain at 11q13.1 are predictive of prostate cancer progression.

The entire prostate microenvironment, not just the epithelial compartment, is important for both normal and neoplastic growth as significant epithelial-mesenchymal/stromal interactions occur (Cunha et al, 2004; Chung et al, 2005). Molecular events may not always be spontaneous, but the product of environmental influences. For instance, both epidemiologic and molecular data suggest that inflammation may be related to prostate cancer development (Nelson et al, 2004). RNASEL, encoding an interferon inducible ribonuclease, and MSR1, encoding subunits of the macrophage scavenger receptor, are candidate-inherited susceptibility genes for prostate cancer, including familial cancer. Proliferative inflammatory atrophy lesions containing activated inflammatory cells and proliferating epithelial cells appear likely to be precursors to prostatic intraepithelial neoplasia (PIN) lesions and prostatic carcinomas.

Using a novel bioinformatics approach, Tomlins and colleagues identified 2 transcription factors ERG and EtV1 that were overexpressed in prostate cancer tissue (Tomlins et al, 2005). Furthermore TMPRSS2 was fused to these genes suggesting that fusion accounted for overexpression. This genetic rearrangement appears to be the most common identified in prostate cancer. Some of the overexpressed genes or combinations of genes may be important biomarkers capable of not only identifying cancer in equivocal biopsy samples (alpha-methylacyl coenzyme A racemase or AMACR and EPCA) but in predicting response to treatment and progression (Rubin, 2004).

The number of prostate cancers attributable to heritable factors may be greater than once thought (Lichtenstein et al, 2000). Although the loci 1p, 3q, 5q, and 22q have been identified as harboring potential predisposition genes in those with a family history of prostate cancer, a multigene model may best explain familial clustering of this disease. In addition, men with a family history of breast and/or ovarian cancer may be offered a predictive genetic test to determine whether or not they carry the family specific BRCA1/2 mutations as they are at increased risk of breast and prostate cancers.

Pathology

Over 95% of the cancers of the prostate are adenocarcinomas. Of the other 5%, 90% are transitional cell carcinomas, and the remaining cancers are neuroendocrine (“small cell”) carcinomas or sarcomas. This discussion will address only adenocarcinomas.

The cytologic characteristics of CaP include hyperchromatic, enlarged nuclei with prominent nucleoli (Figure 22–3). Cytoplasm is often abundant; thus, nuclear-to-cytoplasmic ratios are not often helpful in making a diagnosis of CaP, unlike their usefulness in diagnosing many other neoplasms. Cytoplasm is often slightly blue tinged or basophilic, which may assist in the diagnosis. The diagnosis of CaP is truly an architectural one. The basal cell layer is absent in CaP, although it is present in normal glands, BPH glands, and the precursor lesions of CaP. If the diagnosis of CaP is in question, high-molecular-weight keratin immunohistochemical staining is useful, as it preferentially stains basal cells. Absence of staining is thus consistent with CaP. Those biopsies that remain equivocal could be stained with new markers such as AMACR or EPCA, which appear to identify those with the disease, but who have equivocal or negative biopsies based on standard tissue staining.

PIN and atypical small acinar proliferation (ASAP) are thought to be precursor lesions. Men found to have either lesion may be at an increased risk of prostate cancer and warrant repeat biopsy certainly if an extended-core biopsy was not performed initially. High-grade PIN (HGPIN) is characterized by cellular proliferations within preexisting ducts and glands, with nuclear and nucleolar enlargement similar to prostate cancer. However, unlike cancer, HGPIN retains a basal cell layer identifiable by immunohistochemistry.

Approximately, 60–70% of cases of CaP originate in the peripheral zone, while 10–20% originate in the transition zone, and 5–10% in the central zone. Although prostate cancer is most often thought to be multifocal, the use of widespread screening and extended biopsy techniques has resulted in the increasing detection of unifocal and smaller cancers.

The histology of the remaining 5% of prostate cancer is heterogenous, arising from stromal, epithelial, or ectopic cells. Nonadenocarcinoma variants can be cate-
NEOPLASMS OF THE PROSTATE GLAND

Epithelial variants consist of endometrioid, mucinous, signet-ring, adenoid cystic, adenosquamous, squamous cell, transitional cell, neuroendocrine, and comedocarcinoma. Nonepithelial variants include rhabdomyosarcoma, leiomyosarcoma, osteosarcoma, angiosarcoma, carcinosarcoma, malignant lymphoma, and metastatic neoplasms among others. It is becoming increasingly evident that neuroendocrine differentiation may occur in response to prolonged androgen deprivation. This can be recognized by staining such tissue for neuroendocrine markers (chromogranin A, neuron-specific enolase) and/or by measuring such markers in serum.

Grading & Staging

The Gleason grading system is the most commonly employed grading system in the United States. It is truly a system that relies upon the low-power appearance of the glandular architecture under the microscope. In assigning a grade to a given tumor, pathologists assign a primary grade to the pattern of cancer that is most commonly observed and a secondary grade to the second...
most commonly observed pattern in the specimen. Grades range from 1 to 5 (Figure 22–3). If the entire specimen has only one pattern present, then both the primary and secondary grade are reported as the same grade. The **Gleason score** or **Gleason sum** is obtained by adding the primary and secondary grades together. As Gleason grades range from 1 to 5, Gleason scores or sums thus range from 2 to 10. Well-differentiated tumors have a Gleason sum of 2–4, moderately differentiated tumors have a Gleason sum of 5–6, and poorly differentiated tumors have a Gleason sum of 8–10. Historically, tumors having a Gleason sum of 7 have sometimes been grouped with the moderately differentiated tumors and at other times with the poorly differentiated tumors. One point that needs to be clarified is that the primary Gleason grade is perhaps the most important with respect to placing patients in prognostic groups. This is most important in assessing patients with a Gleason sum of 7. Patients with a Gleason sum of 7 who have a primary Gleason grade of 4 (4 + 3) tend to have a worse prognosis than those who have a primary Gleason grade of 3 (3 + 4). Many clinical series have failed to distinguish between these two populations and, therefore, caution must be exercised in reviewing these series.

Gleason grades 1 and 2 are characterized by small, uniformly shaped glands, closely packed with little intervening stroma. Gleason grade 3 is characterized by variable-sized glands that percolate between normal stroma. A variant of Gleason grade 3 is referred to as a cribriform pattern. Here a small mass of cells is perforated by several gland lumens with no intervening stroma. This results in a cookie-cutter-like appearance of cell nests. The border of these cribriform glands is smooth. Gleason grade 4 has several histologic appearances. The characteristic observation common to all Gleason grade 4 patterns is **incomplete gland formation**. Sometimes glands appear fused, sharing a common cell border. At other times sheets of cell nests are seen or long cords of cells are observed. Cribriform glands can also occur in Gleason grade 4, but the cell masses are large and borders tend to appear ragged, with infiltrating finger-like projections. Gleason grade 5 usually has single infiltrating cells, with no gland formation or lumen appearance. Comedocarcinoma is an unusual variant of Gleason grade 5 carcinoma that has the appearance of cribriform glands with central areas of necrosis.

The TNM staging system for CaP is presented in Table 22–3 (American Joint Committee on Cancer, 1997). Note that with respect to the primary tumor categorization (T stage), the clinical staging system uses results of the DRE and TRUS, but not the results of the biopsy. Some examples to illustrate this staging system follow. If a patient has a palpable abnormality on one side of the prostate, even though biopsies demonstrate bilateral disease, his clinical stage remains T2a. If a patient has a normal DRE,

<table>
<thead>
<tr>
<th><strong>Table 22–3.</strong> TNM Staging System for Prostate Cancer.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>T—Primary tumor</strong></td>
</tr>
<tr>
<td>Tx</td>
</tr>
<tr>
<td>T0</td>
</tr>
<tr>
<td>Tis</td>
</tr>
<tr>
<td>T1a</td>
</tr>
<tr>
<td>T1b</td>
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<tr>
<td>T1c</td>
</tr>
<tr>
<td>T2a</td>
</tr>
<tr>
<td>T2b</td>
</tr>
<tr>
<td>T3a</td>
</tr>
<tr>
<td>T3b</td>
</tr>
<tr>
<td>T4</td>
</tr>
<tr>
<td><strong>N—Regional lymph nodes (obturator, internal iliac, external iliac, presacral lymph nodes)</strong></td>
</tr>
<tr>
<td>Nx</td>
</tr>
<tr>
<td>N0</td>
</tr>
<tr>
<td>N1</td>
</tr>
<tr>
<td><strong>M—Distant metastasis</strong></td>
</tr>
<tr>
<td>Mx</td>
</tr>
<tr>
<td>M0</td>
</tr>
<tr>
<td>M1a</td>
</tr>
<tr>
<td>M1b</td>
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<tr>
<td>M1c</td>
</tr>
</tbody>
</table>

DRE, digital rectal examination; PIN, prostatic intraepithelial neoplasia; PSA, prostate-specific antigen; TRUS, transrectal ultrasound.  
with TRUS demonstrating a lesion on one side and a biopsy confirming cancer, his clinical stage is also T2a (using results of DRE and TRUS). A T1c cancer must have both a normal DRE and a normal TRUS.

PROSTATE CANCER CHEMOPREVENTION

There is much interest in prostate cancer prevention due to the disease’s high prevalence, slowly progressive nature, long latency period, and high absolute mortality. The ideal therapeutic intervention would arrest disease progression during the latency period and decrease the incidence of clinical disease. The ideal agent should be nontoxic and of low cost and the ideal patient would be one at high risk of the disease. To date, several promising chemopreventive agents have been identified and are under laboratory and clinical investigation. In one important study, 18,882 men aged 55 or over with normal DRE and PSA <3 ng/mL were randomized to either placebo or the 5-alpha-reductase inhibitor, finasteride (5 mg/day) (Thompson et al, 2003). Prostate cancer was detected in 18.4% of those on finasteride and 24.4% on placebo, for a 24.8% reduction in prevalence. Cancers of Gleason grades 7, 8, 9, or 10 were more common in the finasteride group than in the placebo group (37.0% versus 22.2%). This apparent increase in the risk of high-grade cancers may be artifactual for several reasons. Further studies on this issue are ongoing. Sexual side effects were more common in those receiving finasteride, whereas urinary symptoms were more common in those receiving placebo. Other agents currently being studied in clinical trials include vitamin E, selenium, dutasteride (another 5-alpha-reductase inhibitor), cyclooxygenase-2 inhibitors, dietary supplements, and selective estrogen receptor modulators (toremifene).

Patterns of Progression

The pattern of CaP progression has been well defined. The likelihood of local extension outside the prostate (extracapsular extension) or seminal vesicle invasion and distant metastases increases with increasing tumor volume and more poorly differentiated cancers. Small and well-differentiated cancers (grades 1 and 2) are usually confined to the prostate, whereas large-volume (>4 cm³) or poorly differentiated (grades 4 and 5) cancers are more often locally extensive or metastatic to regional lymph nodes or bone. Penetration of the prostatic capsule by cancer is a common event and often occurs along perineural spaces. Seminal vesicle invasion is associated with a high likelihood of regional or distant disease. Locally advanced CaP may invade the bladder trigone, resulting in ureteral obstruction. Of note, rectal involvement is rare as Denovilliers’ fascia represents a strong barrier.

Lymphatic metastases are most often identified in the obturator lymph node chain. Other sites of nodal involvement include the common iliac, presacral, and periaortic lymph nodes. The axial skeleton is the most usual site of distant metastases, with the lumbar spine being most frequently implicated (Figure 22–4). The next most common sites in decreasing order are proximal femur, pelvis, thoracic spine, ribs, sternum, skull, and humerus. The bone lesions of metastatic CaP are typically osteoblastic. Involvement of long bones can lead to pathologic fractures. Vertebral body involvement with significant tumor masses extending into the epidural space can result in cord compression. Visceral metastases most commonly involve the lung, liver, and adrenal gland. Central nervous system involvement is usually a result of direct extension from skull metastasis.

Clinical Findings

A. SYMPTOMS

Most patients with early-stage CaP are asymptomatic. The presence of symptoms often suggests locally advanced or metastatic disease. Obstructive or irritative voiding complaints can result from local growth of the tumor into the urethra or bladder neck or from its direct extension into
the trigone of the bladder. Metastatic disease to the bones may cause bone pain. Metastatic disease to the vertebral column with impingement on the spinal cord may be associated with symptoms of cord compression, including paresthesias and weakness of the lower extremities and urinary or fecal incontinence.

B. SIGNS

A physical examination, including a DRE, is needed. Induration, if detected, must alert the physician to the possibility of cancer and the need for further evaluation (ie, PSA, TRUS, and biopsy). Locally advanced disease with bulky regional lymphadenopathy may lead to lymphedema of the lower extremities. Specific signs of cord compression relate to the level of the compression and may include weakness or spasticity of the lower extremities and a hyperreflexic bulbocavernous reflex.

C. LABORATORY FINDINGS

Azotemia can result from bilateral ureteral obstruction either from direct extension into the trigone or from retroperitoneal adenopathy. Anemia may be present in metastatic disease. Alkaline phosphatase may be elevated in the presence of bone metastases. Serum acid phosphatase may be elevated with disease outside the confines of the prostate.

D. TUMOR MARKERS—PROSTATE-SPECIFIC ANTIGEN

PSA is a serine protease produced by benign and malignant prostate tissues. It circulates in the serum as uncomplexed (free or unbound) or complexed (bound) forms. Normal PSA values are those \( \leq 4 \) ng/mL.

Current detection strategies include the efficient use of the combination of DRE, serum PSA, and TRUS with systematic biopsy. Unfortunately, PSA is not specific for CaP, as other factors such as BPH, urethral instrumentation, and infection can cause elevations of serum PSA. Although the last two factors can usually be clinically ascertained, distinguishing between elevations of serum PSA resulting from BPH and those related to CaP remains the most problematic. Serum PSA concentrations are decreased by treatment with agents that lower serum testosterone such as LHRH agonists and antagonists used to treat prostate cancer as well as with 5-alpha-reductase inhibitors used to treat BPH. Interestingly, serum PSA levels are decreased in men with high body mass indexes compared to normal weight men.

The positive predictive value of a serum PSA between 4 and 10 ng/mL is approximately 20–30%. For levels in excess of 10 ng/mL, the positive predictive value increases from 42% to 71.4%. Given that most men with elevated serum PSA levels do not have prostate cancer, there is great interest in identifying makers with greater sensitivity and/or specificity. Candidate markers include novel autoantibodies or other prostate antigens (early prostate cancer antigen) (Paul et al, 2005; Wang et al, 2005).

Numerous strategies to refine PSA for cancer detection have been explored. Their common goal is to decrease the number of false-positive test results. This would increase the specificity and positive predictive value of the test and lead to fewer unnecessary biopsies, lower costs, and reduced morbidity of cancer detection. Attempts at refining PSA have included PSA velocity (change of PSA over time), PSA density (standardizing levels in relation to the size of the prostate), age-adjusted PSA reference ranges (accounting for age-dependent prostate growth and occult prostatic disease), and PSA forms (free versus protein-bound molecular forms of PSA).

1. PSA velocity—PSA velocity refers to the rate of change of serum PSA. A retrospective study has shown that men with prostate cancer have a more rapidly rising serum PSA in the years before diagnosis than do men without prostate cancer. Patients whose serum PSA increases by 0.75 ng/mL/y appear to be at an increased risk of harboring cancer. However, PSA velocity must be interpreted with caution. An elevated PSA velocity should be considered significant only when several serum PSA assays are carried out by the same laboratory over a period of at least 18 months. As a screening tool in large population-based studies, PSA velocity appeared to add little to enhance early detection of prostate cancer. However, it is becoming increasingly clear that those with rapid PSA velocity (ie, PSA doubling times \( \leq 6 \) months) both before diagnosis and/or after treatment are at an increased risk of failure of initial treatment, the development of metastases and prostate cancer specific mortality.

2. PSA density—PSA levels are elevated approximately 0.12 ng/mL/g of BPH tissue. Thus, patients with enlarged glands due to BPH may have elevated PSA levels. The ratio of PSA to gland volume is termed the PSA density. Some investigators advocate prostate biopsy only if the PSA density exceeds 0.1 or 0.15, while others have not found PSA density to be useful. Problems with this approach include the facts that (1) epithelial-stromal ratios vary from gland to gland and only the epithelium produces PSA, and (2) errors in calculating prostatic volume may approach 25%. The positive predictive value of PSA density is slightly higher than the use of a PSA level \( >4 \) ng/mL in several series (30–40% versus 20–30%). Instead of adjusting the PSA to total prostate volume, some have advocated adjusting it to transition zone volume (PSA transition zone density). However, like PSA density, such calculations are subject to error, require TRUS, and do not seem to be superior to the use of PSA in most patients.

3. Age-adjusted reference ranges for PSA—Age-adjusted PSA values for normal men are presented in Table 22–4 (Oesterling JE et al, 1993). It is thought that the rise in PSA with increasing age results from prostate gland growth from BPH, the higher incidence of subclinical prostatitis, and the growing prevalence of microscopic,
Prostate biopsy should be considered in men with an elevated serum PSA, a DRE, or a combination of the two. The most recent refinement in PSA has been the recognition of the various molecular forms of PSA—free and protein-bound. Approximately 90% of the serum PSA is bound to alpha-1-antichymotrypsin, and lesser amounts are free or are bound to alpha-2-macroglobulins. In the latter form, no epitopes to the antibodies used in the current assays are available, while PSA bound to alpha-1-antichymotrypsin may have 3 of its 5 epitopes masked. Early studies suggest that prostate cancer patients demonstrate a lower percentage of free PSA than do patients with benign disease. A large multicenter study has reported that in men with a normal DRE and a total PSA level between 4 and 10 ng/mL, a 25% free PSA cutoff would detect 95% of cancers while avoiding 20% of unnecessary biopsies. The cancers associated with >25% free PSA were more prevalent in older patients and generally were less threatening in terms of tumor grade and volume (Catalona et al, 1998).

**E. Prostate Biopsy**

Prostate biopsy is best performed under TRUS guidance using a spring-loaded biopsy device coupled to the imaging probe. Biopsies are taken throughout the peripheral zone of the prostate, rather than just sampling an area abnormal on the basis of DRE or TRUS. Traditionally, 6 (sextant) biopsies were taken along a parasagittal line between the lateral edge and the midline of the prostate at the apex, midgland, and base bilaterally. However, several investigators have shown that increasing the number (≥10) and performing more laterally directed biopsies of the peripheral zone will increase detection rates 14–20% over the more traditional sextant technique. Although a small number of prostate cancers will originate in the transition zone, specific transition zone biopsies add little to overall cancer detection rates when an extended-pattern biopsy is performed. There is ongoing interest in the use of even more extended biopsy schemes (“saturation biopsy”) or use of a transperineal approach to improve cancer detection, usually in those who have had a negative biopsy, but are thought to be at an increased risk of prostate cancer based on a persistently abnormal serum PSA.

Prostate biopsy is usually performed using local anesthesia and preprocedure antibiotic prophylaxis. Although prostate biopsy is usually very well tolerated by patients, approximately 10–24% of those undergoing the procedure will find it very painful. The use of local anesthesia, either applied topically along the anterior rectal wall, injected into or adjacent to the prostate, or a combination of the two, decreases pain associated with the procedure. Hematospermia and hematuria are common occurring in approximately 40–50% of patients. Minor rectal bleeding may occur, as well. High fever is rare occurring in 2.9–4.2% of patients.

**F. Combined Modality Risk Assessment**

Nomograms and probability tables incorporating serum PSA, cancer grade, T stage, and cancer volume as assessed by extent of biopsies involved with cancer and patient age have been published, are widely used, and allow a better assessment of disease natural history (with or without treatment) and more appropriate selection of initial treatment (Table 22–5). It is likely that identification and incorporation of novel genotypic and phenotypic markers will further enhance the predictive value of these nomograms and probability tables.

**G. Imaging**

1. TRUS—TRUS is useful in performing prostatic biopsies and in providing some useful local staging information if cancer is detected. Almost all prostate needle biopsies are performed under TRUS guidance. This allows uniform spatial separation and sampling of the regions of the prostate and also makes lesion-directed biopsies possible. If visible, CaP tends to appear as a hypoechoic lesion in the peripheral zone.
TRUS provides more accurate local staging than does DRE. The sonographic criteria for extracapsular extension are bulging of the prostate contour or angulated appearance of the lateral margin. The criteria for seminal vesicle invasion are a posterior bulge at the base of the seminal vesicle or asymmetry in echogenicity of the seminal vesicle associated with hypoechoic areas at the base of the prostate.

TRUS also enables measurement of the prostate volume, which is needed in the calculation of PSA density. Typically, a prolate ellipsoid formula is used: \( \frac{\pi}{6} \times (\text{anterior-posterior diameter}) \times (\text{transverse diameter}) \times (\text{sagittal diameter}) \). TRUS is also used in the performance of cryosurgery and brachytherapy (see below). Color or power Doppler TRUS assesses blood flow through prostatic vessels. As cancers may have increased vascularity, such technology may improve the sensitivity and specificity of U.S. imaging. 3D color Doppler permits a three-dimensional image to be constructed from a series of 2D images by a computer algorithm. Use of a microbubble, intravenous contrast agent may also improve visualization of cancers.

2. Endorectal magnetic resonance imaging (MRI)—Use of an endorectal coil improves cancer detection and staging compared to the use of a standard body coil. While rendering high image quality, the use of an endorectal coil appears to be operator dependent, requiring education and expertise. Routine use of this technology may not alter treatment decisions compared to the information gained by assessment of more standard clinopathologic information. Use of magnetic resonance spectroscopy (MRS) in conjunction with MRI may improve the accuracy of imaging. Prostate cancer is associated with proportionately lower levels of citrate and higher levels of choline and creatine compared to BPH or normal prostate tissue. The combined metabolic and anatomic information provided by MRI and MRS may allow for a more accurate assessment of cancer location and stage. The reported staging accuracy of endorectal MRI varies from 51% to 92%. It appears to add novel information to the assessment of some patients over the use of nomograms alone, but may be best utilized in high-risk patients where it is most accurate and helpful (Hricak 2005; Wang et al, 2005).

3. Axial imaging (CT, MRI)—Cross-sectional imaging of the pelvis in patients with CaP is selectively performed to exclude lymph node metastases in high-risk patients who are thought to be candidates for definitive local therapy, whether it be surgery or irradiation. Both MRI and computed tomography (CT) are used for this purpose. Patients identified as having lymphadenopathy on imaging may undergo CT-guided fine-needle aspiration. If lymph node metastases are confirmed, such patients may be candidates for alternative treatment regimens. However, the incidence of lymph node metastases in contemporary radical prostatectomy series is low (<10%). In addition, imaging is costly and its sensitivity is limited (30–40%). Various criteria can be used to identify patients for axial imaging, including negative bone scans and either T3 cancers or a PSA >20 ng/mL and primary Gleason grade 4 or 5 cancers. Intravenous administration of superparamagnetic nanoparticles, which gain access to lymph nodes by means of interstitial-lymphatic fluid transport, at the time of high-resolution MRI, appears to improve visualization of small nodal metastases.

Analyses of several contemporary series of patients with clinically localized prostate cancer suggest that the risk of lymph node metastases is low and that its risk can be quantified on the basis of serum PSA, local tumor stage, and tumor grade.

4. Bone scan—When prostate cancer metastasizes, it most commonly does so to the bone (Figure 22–4). Soft tissue metastases (eg, lung and liver) are rare at the time of initial presentation. Although a bone scan has been considered a standard part of the initial evaluation of men with newly

<table>
<thead>
<tr>
<th>Risk Group</th>
<th>Criteria</th>
<th>Recurrence after Local Therapy (Ranges)</th>
<th>Risk Lower</th>
<th>Risk Higher</th>
<th>Need for Routine Radiographic Imaging</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>PSA &lt;10 ng/mL, Gleason ≤6 (No grade 4/5) and T1, T2a</td>
<td>6–20%</td>
<td>Gleason 2–4</td>
<td>&gt;50% positive biopsies</td>
<td>None</td>
</tr>
<tr>
<td>Intermediate</td>
<td>PSA 10–20 ng/mL, Gleason 7, and/or T2b, T3a</td>
<td>26–60%</td>
<td>&lt;50% positive biopsies, PSA &lt;6 ng/mL</td>
<td>&gt;50% positive biopsies, Gleason 4/3</td>
<td>Bone scan for PSA &gt;15 ng/mL</td>
</tr>
<tr>
<td>High</td>
<td>PSA &gt;20 ng/mL, Gleason 8–10, and/or T3b</td>
<td>31–100%</td>
<td>PSA, 10 ng/mL, T1/T2 disease</td>
<td>7113 Variables</td>
<td>Bone scan, CT/MRI of pelvis</td>
</tr>
</tbody>
</table>

Table 22–5. Pretreatment Risk Assessment.
diagnosed prostate cancer, good evidence has been accumulated that it can be excluded in most of these men on the basis of serum PSA. Several investigators have shown that bone scans can be omitted in patients with newly diagnosed, untreated prostate cancer who are asymptomatic, have T1 and T2 disease, and have serum PSA concentrations <15 ng/mL. However, patients with PSA 15 ng/mL or greater, locally advanced disease (T3B, T4) are at higher risk for bone metastases and should be considered for bone scan.

5. Antibody imaging—ProstaScint is a murine monoclonal antibody to an intracellular component of the prostate-specific membrane antigen (PSMA), which is conjugated to 111 indium. After infusion of the antibody, single photon emission computed tomography (SPECT) images are usually obtained at 30 minutes to access vasculature and at 72–120 hours. It has been approved by the U.S. Food and Drug Administration (FDA) for use in the evaluation of patients prior to treatment and for detecting the site of recurrent disease in patients who have biochemical relapse after initial treatment. However, this antibody recognizes the intracellular domain of PSMA; only soft tissues are imaged and the test may suffer from both false-positive and negative results in both clinical situations described above. Use of new antibodies, which recognize the extracellular domain of PSMA, appear to allow for recognition of both bone and soft tissue metastases and could be used as agents for therapy, not only improved imaging (Bander et al, 2005).

Differential Diagnosis

Not all patients with an elevated PSA concentration have CaP. Other factors that elevate serum PSA include BPH, urethral instrumentation, infection, prostatic infarction, or vigorous prostate massage. Induration of the prostate is associated not only with CaP, but also with chronic granulomatous prostatitis, previous TURP or needle biopsy, or prostatic calculi. Sclerotic lesions on plain x-ray films and elevated levels of alkaline phosphatase can be seen in Paget’s disease and can often be difficult to distinguish from metastatic CaP. In Paget’s disease, PSA levels are usually normal and x-ray findings demonstrate subperiosteal cortical thickening.

Screening for CaP

The case for CaP screening is supported by the following: The disease is burdensome in this country; PSA improves detection of clinically important tumors without significantly increasing the detection of unimportant tumors; most PSA-detected tumors are curable; prostate cancer mortality is declining in regions where screening occurs; and curative treatments are available. If screening is undertaken, it appears that the use of both DRE and serum PSA is preferable to either one used alone. Although many recommend that screening be undertaken at age 50, some have advocated for earlier screening starting at age 40. Although annual screening is most often recommended. Some feel that men with very low serum PSA level (≤1 ng/mL) may be able to be screened at less frequent intervals (every 2 or 3 years).

What constitutes a serum PSA at which biopsy is recommended is a matter of debate. Although a normal PSA is considered to be 4 ng/mL or less, this value was set in men of all ages and prostate volumes. As mentioned earlier, younger men, with less BPH, should have lower levels. In addition, recent information suggests that many men with serum PSA concentrations in the normal range, even with a normal DRE, may harbor significant disease. In the PCPT trial of finasteride for chemoprevention of prostate cancer discussed earlier, the prevalence of prostate cancer was 6.6% among men with a PSA level of up to 0.5 ng/mL, 10.1% among those with values of 0.6–1 ng/mL, 17% among those with values of 1.1–2 ng/mL, 23.9% among those with values of 2.1–3 ng/mL, and 26.9% among those with values of 3.1–4 ng/mL (Thompson et al, 2003; Thompson et al, 2004; Thompson et al, 2005). PSA cutoff values of 1.1, 2.1, 3.1, and 4.1 ng/mL yielded sensitivities of 83.4%, 52.6%, 32.2%, and 20.5%, and specificities of 38.9%, 72.5%, 86.7%, and 93.8%, respectively. Even more importantly, the prevalence of high-grade cancers varied from 12.5% to 25% in these low ranges. Therefore, there is no PSA cutpoint where cancer can be excluded. Based on these results, some have suggested lowering the PSA cutpoint for biopsy to 2 or 2.5 ng/mL.

Screening to date has resulted in considerable stage migration (more lower stage cancers being detected) and mortality rates are falling. The reasons for this later finding are a matter of much debate. Although several investigators have shown a reduction in prostate cancer mortality related to early detection efforts, none have done so on the basis of a well-conducted randomized trial. In the Prostate, Lung, Colon and Ovarian Trial (PLCOT), 74,000 men have been randomized to annual DRE and PSA screening versus no screening. Results are anticipated in 2006. A larger trial involving 190,000 men in the European Randomized Study for Prostate Cancer Screening is anticipated to provide results in 2008.

One concern related to screening is the fact that some cancers may be detected which would never result in clinically significant disease in the patient if left untreated, a phenomenon called overdetection. Some have estimated that between 29% and 48% of cancers detected by an aggressive screening program are such cancers (Eziozie et al, 2002; Draisma et al, 2003). This underscores the importance of informed consent before screening is undertaken and the need to discuss all treatment options, including active surveillance, in those found to have the disease. Screening should be undertaken in men who are healthy enough to benefit from it. Screening may be highly encouraged in certain populations with a higher disease
prevalence and/or mortality such as African American men and those with a strong family history of the disease.

**Treatment**

**A. Localized Disease**

1. **General considerations**—The optimal form of therapy for all stages of CaP remains a subject of great debate. Treatment dilemmas persist in the management of localized disease (T1 and T2) because of the uncertainty surrounding the relative efficacy of various modalities, including radical prostatectomy, radiation therapy, and surveillance. Currently, treatment decisions are based on the grade and stage of the tumor, the life expectancy of the patient, the ability of each therapy to ensure disease-free survival, its associated morbidity, and patient and physician preferences. Until recently, there was little information to be sure that treatment of early stage disease had an important impact on overall and cancer-specific survival. A well-conducted randomized trial of radical prostatectomy versus surveillance in men with early stage prostate cancer was conducted in Scandinavia (Bill-Axelson et al, 2005). Men who underwent radical prostatectomy were less likely to die, die of prostate cancer (risk reduction 0.56), develop metastases (risk reduction 0.60), or suffer local cancer progression (risk reduction 0.33) compared to men who underwent initial surveillance. The advantage to surgery was most apparent in younger patients.

2. **Watchful waiting and active surveillance**—Although local cancer progression may occur, with watchful waiting for early stage prostate cancer, disease-specific mortality at 10 years is low varying generally between 4% and 15%. However, in further follow-up from 15 to 20 years, a substantial increase in the risk of local and systemic progression and death from prostate cancer may be seen (Johansson, Andren et al, 2004). The risk of progression is related significantly to cancer grade. The risk of progression is low in those with Gleason grades 2–6 (no pattern 4 or 5), but increases significantly for those with Gleason grades 7 through 10. Most of the men, in these previously reported series of men managed with watchful waiting, had palpable disease and, therefore, larger, more significant cancer than most of those detected currently based on serum PSA. In addition, most men were not followed carefully with periodic clinical, radiographic, and laboratory (PSA) reevaluation. They were treated, usually with androgen deprivation, when symptomatic metastatic disease was detected. A more modern concept of watchful waiting is better termed “active surveillance” where men with very well-characterized, early stage, and low to intermediate grade cancer are followed very carefully and treated at the first sign of subclinical progression based on serial and regular physical examinations, serum PSA measurements, and repeat prostatic biopsy (Klotz 2005). Although between 20% and 41% of men on such regimens may require treatment at 5 years of follow-up, treatment at progression appears to be as effective as it would have been if delivered at the time of diagnosis for most men. However, optimal surveillance strategies, end points for intervention and exact risks of surveillance have not been well defined, as yet.

3. **Radical prostatectomy**—The first radical perineal prostatectomy was performed by Hugh Hampton Young in 1904, and Millin first described the radical retropubic approach in 1945. However, the procedure remained unpopular because of frequent complications of incontinence and impotence. The rebirth of radical prostatectomy has resulted from a better understanding of the surgical anatomy of the pelvis. Description of the anatomy of the dorsal vein complex resulted in modifications in the surgical technique leading to reduced operative blood loss. In addition, improved visualization made possible a more precise apical dissection, allowing better reconstruction of the urinary tract and improved continence. Eversion of the bladder mucosa before anastomosis ensures a mucosa-to-mucosa apposition. Anatomical dissections have led to a better understanding of the prostate apex anatomy and its relationship to the distal urethral sphincteric mechanism. Description of the course of the cavernous nerves enabled modifications of the surgical technique, resulting in preservation of potency. Lymph node dissection, once done routinely, may be performed only in those at significant risk of lymph node metastases. Such men can be identified with use of probability tables and nomograms. Previously, only limited node dissections were performed harvesting lymph nodes from the obturator fossa. However, results from extended dissections showed that more than half of lymph node metastases are found outside this region. Therefore, a more extended and meticulous dissection is advised. Some feel that this may not only have diagnostic value, but also could have a therapeutic impact in those with limited nodal disease (Bader et al, 2003; Allaf et al, 2004).

Considerable experience has been gained recently using a laparoscopic approach to radical prostatectomy. This can be performed through an extra- or transperitoneal approach. Also, a robotic interface is thought, by some, to improve and/or facilitate the procedure. The procedure is associated with reduced blood loss, shorter hospitalization, and return to normal activity. Whether it results in improved long-term outcomes (ie, biochemical relapse-free survival, continence, potency, etc.) compared to open techniques is not known, as yet. The procedure is more costly to perform and does require special expertise to perform well.

The prognosis of patients treated by radical prostatectomy correlates with the pathologic stage of the specimen. Distant metastasis is inevitable in patients with positive lymph nodes. A high percentage of patients with seminal vesical involvement at radical prostatectomy are destined
to distant failure. Fortunately, the number of patients with these adverse prognostic factors undergoing surgery is decreasing because of better candidate selection based on appropriate use of preoperative clinical parameters. Several investigators have established nomograms to predict final pathologic stage at radical prostatectomy based on the serum PSA level, clinical DRE stage, and Gleason sum derived from the biopsy.

Patients with organ-confined cancer have 10-year disease-free survival ranging from 70% to 85% in several series. Those with focal extracapsular extension demonstrate 85% and 75% disease-free survival at 5 and 10 years, respectively. Patients with more extensive extracapsular extension demonstrate 70% and 40% disease-free survival at 5 and 10 years, respectively. High-grade tumors (Gleason sum >7) have a higher risk of progression than do low-grade tumors. Disease-free survival at 10 years for patients with Gleason sum 2–6 tumors is in excess of 70%; for Gleason sum 7, 50%; and for Gleason sum >8, 15%. Positive surgical margins significantly affect only tumors with extensive extracapsular extension. Neoadjuvant androgen deprivation, studied by several investigators, reduces the risk of positive surgical margins, but does not appear to impact long-term biochemical relapse-free survival.

The management of patients with positive surgical margins at radical prostatectomy remains controversial. Not all such patients relapse, but the identification of appropriate candidates for adjuvant radiation therapy remains problematic. A large multicenter randomized trial was completed to determine whether adjuvant radiation therapy in this setting is superior to radiation therapy at the time of relapse. Results of this study will be available shortly. Patients with positive surgical margins are at an increased risk of cancer recurrence. Adjuvant radiation therapy appears to reduce the risk of biochemical relapse, but its impact on overall and disease specific survival is not known, as yet. The risk of extracapsular extension, and therefore, the risk of a positive surgical margin can be predicted preoperatively based on cancer grade (Gleason score >7), cancer volume (extent of core involved and total number of cores involved with cancer), and imaging (endorectal MRI/MRS and/or TRUS). Those at significant risk of extracapsular excision should undergo wide surgical excision with sacrifice of the neurovascular bundle on that side to better ensure complete cancer excision.

Morbidity associated with radical prostatectomy can be significant and is in part related to the experience of the surgeon. Immediate intraoperative complications include blood loss, rectal injury, and ureteral injury. Blood loss is more common with the retropubic approach than with the perineal approach because in the former, the dorsal venous complex must be divided. Rectal injury is rare with the retropubic approach and more common with the perineal approach but usually can be immediately repaired without long-term sequelae. Ureteral injury is exceedingly rare with any technique. Perioperative complications include deep venous thrombosis, pulmonary embolism, lymphocele formation, and wound infection. Late complications include urinary incontinence and impotence. Although total urinary incontinence tends to be rare (<3%), stress urinary incontinence may be seen in up to 20% of patients. The return of continence after surgery is gradual, with 50% of patients continent at 3 months, 75% at 6 months, and the remainder at 9–12 months. Age is the single most important factor in the restoration of continence. Preservation of one or both neurovascular bundles may allow maintenance of erectile function in men who are potent and sexually active before the procedure. However, the nerve-sparing procedure should be used selectively, for extracapsular extension may be a common finding in patients with presumed localized CaP. If extracapsular extension is present, preservation of the neurovascular bundle may increase the likelihood that the tumor will recur. Preservation of potency varies as a function of age, preoperative sexual function, and preservation of one or both neurovascular bundles. Reported rates of potency preservation vary from 40% to 82% in men under the age of 60 when both nerves are preserved and drops to 20–60% when only one nerve is preserved. For men between the ages of 60 and 69, comparable rates are 25–75% with bilateral nerve sparing and 10–50% with unilateral nerve sparing. Recovery of sexual function generally occurs within 6–18 months following surgery. Potency can be improved with early use of PDE-5 inhibitors. (See Chapter 38 for more information on impotence.)

4. Radiation therapy—external beam therapy—
Traditional external beam radiotherapy (XRT) techniques allow the safe delivery of 6500–7000 cGy to the prostate. Standard XRT techniques depend upon bony landmarks to define treatment borders or a single CT slice to define the target volume. These standard XRT techniques generally involve the use of open square or rectangular fields with minimal to no blocking and are characterized by the use of relatively small boost fields. Often, these XRT techniques fail to provide adequate coverage of the target volume in as many as 20–41% of patients with CaP irradiated.

Improved imaging and use of novel treatment planning (three-dimensional, conformal, and intensity-modulated radiation therapy) allow for better targeting, conforming or shaping radiation volume more closely around the prostate, and the use of higher doses without exceeding tolerance of surrounding normal tissues. Such radiotherapy has resulted in dramatic reductions in acute and late toxicity of radiation treatment and improved tumor control compared with conventional dose radiotherapy. Doses ≥72 cGy appear to result in improved biochemical outcomes compared to lower doses. Day to day variations in patient/prostate position can be accounted for by the use of daily online CT scanning, transabdominal ultrasound imaging, and insertion of an endorectal balloon or imaging of radio-
opaque markers placed before treatment. Although some feel that whole pelvic radiation to include regional lymph nodes, especially when combined with androgen deprivation, results in improved outcomes in those with intermediate and high-risk prostate cancer as defined by serum PSA, T stage, and cancer grade, not all agree.

In addition to the use of dose escalation and improved tumor targeting, several investigators have shown that the results of radiation therapy may be improved with the use of neoadjuvant, concurrent, and adjuvant androgen deprivation. On the basis of numerous randomized trials, androgen deprivation improves the outcome of radiation in those with intermediate (PSA 10–20 ng/mL, T2b, or Gleason score 7) or high-risk (PSA >20 ng/mL, T3, or Gleason score 8, 9, or 10) disease. The use of short-term (3–4 months) neoadjuvant and concurrent androgen deprivation is recommended for those with intermediate risk disease, whereas those with high-risk disease should receive neoadjuvant, concurrent, and long-term adjuvant (24 months) androgen deprivation (Bolla et al., 2002; Roach, 2003).

Like radical prostatectomy, men who receive radiation may experience side effects especially those related to urinary, bowel, and sexual function. Most such side effects are limited in extent. Whereas men who undergo surgery are more likely to suffer incontinence, men who undergo radiation are more likely to suffer obstructive or irritative voiding or bowel symptoms (diarrhea, rectal bleeding, tenesmus). Whereas the impact of surgery on sexual function occurs early and may improve with time, the impact of radiation on sexual function may not be seen for 18–24 months. Sexual side effects may be exacerbated with the concurrent use of androgen deprivation, especially if used long term.

Readers are referred to Chapter 25 for a more detailed discussion of XRT in CaP.

5. Radiation therapy—brachytherapy—A resurgence in the interest in brachytherapy has occurred because of the technologic developments making it possible to place radioactive seeds under TRUS guidance. Previously, free-hand seed placement techniques were used; however, very high failure rates were observed and the technique was virtually abandoned. Currently, with the use of computer software, one can preplan a precise dose of radiotherapy to be delivered by TRUS guidance. Implants can be permanent (iodine 125 or palladium 103) in that the seeds are placed in the prostate and the radiation dose is delivered over time or temporary in that the seeds are loaded into hollow-core catheters and both the seeds (iridium 192) and catheters are removed after a short period of hospitalization and radiation exposure. Permanent implants have a lower dose rate, but a higher total dose delivered compared to temporary implants which have a higher dose rate, but deliver a lower total dose. External beam radiation can be given to those with intermediate and high-risk cancers who receive permanent brachytherapy and is routinely given to all those who undergo temporary or high dose rate brachytherapy. Some clinicians are determining whether men with low-risk disease can be treated effectively with high dose rate brachytherapy alone without the use of neoadjuvant external beam radiation.

As opposed to external beam radiation, androgen deprivation does not appear to improve the outcomes of men with intermediate disease who are treated with brachytherapy.

Men with high-risk disease who choose brachytherapy receive external beam radiation and long-term adjuvant androgen deprivation as described for those managed with external beam techniques alone.

Based on contemporary information, radiation therapy delivered well, in appropriate doses by any means, and when combined appropriately with androgen deprivation appears to result in similar long-term, relapse-free outcomes as does surgery. Only well-designed randomized trials will determine whether or not there may be small to moderate differences in outcomes between these two forms of treatment.

Readers are referred to Chapter 25 for a more detailed discussion of brachytherapy in CaP.

6. Cryosurgery and high-intensity focused ultrasound (HIFU)—There has been a resurgence of interest in cryosurgery as a treatment for localized CaP in the past several years. This is due to an increased interest in less invasive forms of therapy for localized CaP as well as several recent technical innovations, including improved percutaneous techniques, expertise in TRUS, improved cryotechnology, and better understanding of cryobiology.

Freezing of the prostate is carried out by using a multi-probe cryosurgical device. Multiple hollow-core probes are placed percutaneously under TRUS guidance. Most surgeons routinely perform 2 freeze-thaw cycles in all patients, and if the iceball does not adequately extend to the apex of the prostate, the cryoprobes are pulled backward into the apex and additional freeze-thaw cycles are undertaken. The temperature at the edge of the iceball is 0°C to −2°C, while actual cell destruction requires −25°C to −50°C. Therefore, actual tissue destruction occurs a few millimeters inside the iceball edge and cannot be monitored precisely by ultrasound imaging. Double freezing creates a larger tissue destruction area and theoretically brings the iceball edge and destruction zone edge closer together.

Studies to date indicate that, in the short term at least, cryosurgery can result in negative posttreatment prostatic biopsies and low or undetectable serum PSA levels. However, the morbidity of cryosurgery is significant and the long-term results are unknown. More recently, further refinement in the cryo technology has resulted in the development of smaller cryoprobes, which may result in a more controlled freezing. Some have advocated more focal cryosurgery where only the side/site of cancer is presumed to be in order to minimize side effects of treatment. Further refinements in imaging and tumor site identification
are required before such treatment can be done with confidence in larger numbers of patients. HIFU can be delivered to the prostate using a rectal probe. This technology induces coagulative necrosis of benign and malignant prostate tissue. Clinical experience, largely from Europe and Japan, suggests that such treatment is associated with clinical outcomes similar to those seen with cryotherapy. As urinary retention is one of the most common complications of this technique, some have performed transurethral resection of the prostate at the time of HIFU to reduce the risk of postoperative urinary retention.

C. Recurrent Disease

1. Overview—A substantial number of men who are treated with either surgery or radiation for presumed clinically localized prostate cancer will relapse based on evidence of a detectable or rising serum PSA after treatment, respectively. Although a persistently detectable serum PSA after surgery is considered a failure, what constitutes biochemical failure after radiation is a matter of some debate. The American Society for Therapeutic Radiology and Oncology (ASTRO) adopted the definition of 3 consecutive rises in serum PSA above nadir. However, this has been modified, by some, to improve its specificity by defining failure as a rise of at least 2 ng/mL greater than the nadir level. It must be remembered that up to one-third of patients will experience a “PSA bounce” following radiation (especially brachytherapy), which is defined by a rise in serum PSA followed by a decline. Such patients are not at an increased risk of cancer recurrence and repeat prostate biopsy should be deferred in such patients. Biochemical failure may have a variable natural history after any kind of initial treatment and may signify localized disease, systemic disease, or a combination of the two. After either form of treatment, an interval to PSA failure <3–6 years or a posttreatment PSA doubling time <3 months place a man at increased risk for metastases and subsequent prostate cancer-specific mortality.

2. Following radical prostatectomy—The likelihood of recurrence following radical prostatectomy is related to cancer grade, pathologic stage, and the extent of extracapsular extension. Cancer recurrence is more common in those with positive surgical margins, established extracapsular extension, seminal vesicle invasion, and high-grade disease. For those patients in whom a detectable PSA level develops after radical prostatectomy, the site of recurrence (local versus distant) can be established with reasonable certainty based on the interval from surgery to the detectable PSA concentration, PSA doubling time, and selective use of imaging studies.

Patients with persistently detectable serum PSA levels immediately after surgery, those with PSA levels that become detectable in the early postoperative period, and those with serum PSA levels that double rapidly are more likely to have systemic relapse.

Those patients thought to have recurrent localized disease based on a long time from surgery to biochemical failure, prolonged PSA doubling times (>10–12 months), and presence of positive surgical margins at the time of surgery are most likely to benefit from salvage radiation (approximately 77% freedom from subsequent relapse). Those with high-grade disease or seminal vesicle involvement at the time of surgery, who fail early or with rapid PSA kinetics after surgery, are less likely to respond and should be considered for systemic therapy.

3. Following radiation therapy—A rising PSA level following definitive radiotherapy is indicative of cancer recurrence. For those who undergo radiation and experience biochemical failure as defined earlier, the site of recurrence may be identified using PSA kinetics, time to failure as noted above, prostate biopsies, and selective use of imaging. Most patients who fail radiation therapy, irrespective of the site of recurrence, currently are managed with androgen deprivation. However, those with local recurrence only may be candidates for brachytherapy, cryosurgery, or salvage prostatectomy. However, morbidity can be high with these forms of treatment as is subsequent relapse.

D. Metastatic Disease

1. Initial endocrine therapy—Since death due to CaP is almost invariably a result of failure to control metastatic disease, a great deal of research has concentrated on efforts to improve control of distant disease. It is well known that most prostatic carcinomas are hormone dependent and that approximately 70–80% of men with metastatic CaP respond to various forms of androgen deprivation. Testosterone, the major circulating androgen, is produced by the Leydig cells in the testes (95%), with a smaller amount being produced by peripheral conversion of other steroids. Although 98% of serum testosterone is protein bound, free testosterone enters prostate cells and is converted to dihydrotestosterone (DHT), the major intracellular androgen. DHT binds to a cytoplasmic receptor protein and the complex moves to the cell nucleus, where it modulates transcription. Androgen deprivation may be induced at several levels along the pituitary-gonadal axis using a variety of methods or agents (Table 22–6). Use of a class of drugs (LHRH agonists) has allowed induction of androgen deprivation without orchiectomy or administration of diethylstilbestrol. There are 4 LHRH agonists currently approved by the FDA for the treatment of prostate cancer: goserelin acetate, triptorelin pamoate, histrelin acetate, and leuprolide acetate. These can be delivered by injection monthly or as depot preparations lasting 3–6 months. A subcutaneous implant that releases leuprolide acetate at a constant rate for 1 year is also available. An LHRH agonist (Abarelix, Praecis Pharmaceuticals) has been developed, as well. However, concerns about allergic reactions associated with administration have limited its development. Agonists would avoid the “flare” phenomenon associated with
antagonists where serum testosterone concentrations increase before falling. Such an increase could cause symptoms in those with advanced cancer. Currently, administration of LHRH agonists is the most common form of primary androgen blockade used. Orchiectomy, once common, is much less commonly performed today. Estrogens achieve castration by feedback inhibition of the hypothalamic-pituitary axis and, perhaps, by a direct cytotoxic effect. Although effective, their use is limited due to an increased risk of negative cardiovascular effects. Transdermal preparations are under investigation. Because of its rapid onset of action, ketoconazole should be considered in patients with advanced prostate cancer who present with spinal cord compression or disseminated intravascular coagulation. Although testosterone is the major circulating androgen, the adrenal gland secretes the androgens dehydroepiandrosterone, dehydroepiandrosterone sulfate, and androstenedione. Some investigators believe that suppressing both testicular and adrenal androgens (complete androgen blockade) allows for a better initial and a longer response compared with those methods that inhibit production of only testicular androgens. Complete androgen blockade can be achieved by combining an antiandrogen with the use of an LHRH agonist or orchiectomy. Antiandrogens appear to act by competitively binding the receptor for DHT, the intracellular androgen responsible for prostatic cell growth and development. When patients with metastatic prostate cancer are stratified with regard to extent of disease and performance status, those patients with limited disease and a good performance status who are treated with combined androgen blockade (an LHRH agonist and antiandrogen agent) seem to survive longer than those treated with an LHRH agonist alone (Crawford et al, 1989). However, another study comparing the use of an antiandrogen with and without an orchiectomy failed to demonstrate a survival difference between the two arms (Eisenberger et al, 1998). A meta-analysis of monotherapy and complete androgen blockade for the treatment of men with advanced prostate carcinoma suggested that there might be a small survival advantage to complete androgen blockade. This advantage must be balanced against an increased risk of side effects in those on combined therapy (Samson 2002). Ongoing trials are studying the use of intermittent androgen deprivation to determine whether this might result in a delay in the appearance of the hormone-refractory state. Intermittent therapy, compared to continuous therapy, may be associated with improved quality of life as serum testosterone levels may normalize during periods off therapy. High-dose antiandrogen monotherapy (bicalutamide 150 mg/day) is an alternative to castration both in patients with locally advanced and metastatic disease who are interested in maintaining libido and erectile function. In those with locally advanced disease, no significant difference in overall survival has been demonstrated between bicalutamide monotherapy and castration. However in those with metastatic disease, castration is associated with better survival.

The timing of initial endocrine therapy in CaP has been an area of great debate for many years. Data from the Veterans Administration Cooperative Studies from the 1960s did not demonstrate a clear survival advantage for early intervention with androgen ablation therapy in patients with advanced CaP. However, a randomized study from the Medical Research Council comparing early with delayed endocrine therapy in patients with advanced CaP demonstrated improved survival as well as lower com-

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plication rates (cord compression, ureteric obstruction, bladder outlet obstruction, and pathologic fractures) in patients treated with early endocrine therapy (Medical Research Council, 1997). In patients who undergo radical prostatectomy and are found to have microscopic lymph node involvement, early endocrine therapy has also resulted in a survival advantage (Messing et al, 1999). Most would agree that androgen deprivation should be instituted in all those with metastatic disease, whether symptomatic or not. In addition, there may be an advantage to early therapy in those without radiographic evidence of cancer, but who relapse after initial therapy and are found to have rapid PSA doubling times as such patients are at great risk of developing metastatic disease early and dying of their disease. Androgen deprivation is not without side effects including hot flashes, anemia, loss of libido and sexual function, loss of bone mineral density, increased weight and body fat, and cognitive changes. In addition, increases in total cholesterol, low- and high-density lipoproteins, and serum triglycerides have been reported. Men on androgen deprivation should be monitored for such side effects as treatment for most is readily available. Many men diagnosed with prostate cancer suffer from low bone mineral density, which can be exacerbated with androgen deprivation therapy. Many agents may prevent generalized and localized bone loss, including calcium and vitamin D supplements and, if significant, bisphosphonates. Anemia is usually mild, but may be managed with recombinant erythropoietin. Although there are a number of treatments for men with hot flashes that are especially troublesome, medroxyprogesterone acetate (300–400 mg IM monthly) is an effective treatment with limited side effects.

2. Manipulations for hormone refractory prostate cancer—Patients receiving complete androgen blockade therapy who demonstrate a rise in serum PSA levels are currently managed by discontinuing the antiandrogen. Approximately 20–30% of such patients have a secondary PSA response. Responses are not just serologic, as regression of soft tissue disease has been reported. The pathophysiologic underlying this secondary response, referred to as the antiandrogen withdrawal syndrome, is not understood. Some investigators have postulated that emergence of the hormone-refractory state results from mutations in the androgen receptor. Typically, antiandrogens competitively inhibit the androgen receptor, but it is possible that these agents actually stimulate a mutant androgen receptor. Removal of this stimulus (stopping the antiandrogen), thus leads to a secondary response.

Patients receiving monotherapy (LHRH agonist or orchiectomy) whose PSA level starts rising may respond to the addition of an antiandrogen. Response rates are approximately 20–30% in this setting. In addition, use of ketoconazole, aminogluthethimide, corticosteroids, and estrogenic compounds should be considered, as a significant number of patients who have failed initial forms of androgen deprivation will respond to these agents. Although chemotherapy was once not thought to be very effective men with hormone refractory prostate cancer, two recent trials demonstrated a survival benefit for docetaxel-based therapy in men with hormone refractory prostate cancer. Earlier use of chemotherapy and novel agents are currently under investigation.

Readers are referred to Chapter 19 for a detailed discussion of the therapy for hormone-refractory disease.

**SUGGESTED READINGS**

**Prostate Gland Anatomy**


**Benign Prostatic Hyperplasia**


Bent S et al. Saw Palmetto for benign prostatic hyperplasia. NEJM 2006;354. (pages released on Feb 9th)


Prostate Cancer

Epidemiology


Genetics, Pathobiology, & Pathology


Chemoprevention


Serum Tumor Markers, Staging, & Nomograms


Parsons JK et al: Complexed prostate specific antigen (PSA) reduces unnecessary prostate biopsies in the 2.6-4.0 ng/mL range of total PSA. BJU Int 2004;94(1):47–50.


Imaging & Staging


Radical Prostatectomy


**Radiation**


**Active Surveillance & Watchful Waiting**


**Cryotherapy & HIFU**


**Quality of Life**


**Relapse & Androgen Deprivation**


Wirth MP et al: Bicalutamide 150 mg in addition to standard care in patients with localized or locally advanced prostate cancer: results from the second analysis of the early prostate cancer program at median followup of 5.4 years. J Urol 2004;172(5 Pt 1):1865–70.

**Secondary Therapy & Chemotherapy**


TUMORS OF THE TESTIS

GERM CELL TUMORS OF THE TESTIS

Epidemiology & Risk Factors

Malignant tumors of the testis are rare, with approximately 9 new cases per 100,000 males reported in the United States each year. Of all primary testicular tumors, 90–95% are germ cell tumors (seminoma and nonseminoma), while the remainder are nongerminal neoplasms (Leydig cell, Sertoli cell, gonadoblastoma). The lifetime probability of developing testicular cancer is 0.2% for a white male in the United States. Survival of patients with testicular cancer has improved dramatically in recent years, reflecting the development and refinement of effective combination chemotherapy. Of the 8000 new cases of testicular cancer in the United States in 2005, less than 400 deaths are expected.

The incidence of testicular cancer shows marked variation among different countries, races, and socioeconomic classes. Scandinavian countries report up to 6.7 new cases per 100,000 males annually; Japan reports 0.8 per 100,000 males. In the United States, the incidence of testicular cancer in blacks is approximately one-fourth that in whites. Within a given race, individuals in the higher socioeconomic classes have approximately twice the incidence of those in the lower classes.

Testicular cancer is slightly more common on the right side than on the left, which parallels the increased incidence of cryptorchidism on the right side. Of primary testicular tumors, 1–2% are bilateral, and about 50% of these tumors occur in men with a history of unilateral or bilateral cryptorchidism. Primary bilateral tumors of the testis may happen synchronously or asynchronously but tend to be of the same histologic type. Seminoma is the most common germ cell tumor in bilateral primary testicular tumors, while malignant lymphoma is the most common bilateral tumor of the testis.

Although the cause of testicular cancer is unknown, both congenital and acquired factors have been associated with tumor development. The strongest association has been with the cryptorchid testis. Approximately 7–10% of testicular tumors develop in patients who have a history of cryptorchidism; seminoma is the most common form of tumor these patients have. However, 5–10% of testicular tumors occur in the contralateral, normally descended testis. The relative risk of malignancy is highest for the intra-abdominal testis (1 in 20) and is significantly lower for the inguinal testis (1 in 80). Placement of the cryptorchid testis into the scrotum (orchiopexy) does not alter the malignant potential of the cryptorchid testis; however, it does facilitate examination and tumor detection.

Exogenous estrogen administration to the mother during pregnancy has been associated with an increased relative risk for testicular tumors in the fetus, ranging from 2.8 to 5.3 over the expected incidence. Other acquired factors such as trauma and infection-related testicular atrophy have been associated with testicular tumors; however, a causal relationship has not been established.

Classification

Numerous classification systems have been proposed for germ cell tumors of the testis. Classification by histologic type proves to be the most useful with respect to treatment. The 2 major divisions are seminoma and nonseminomatous germ cell tumors (NSGCT), which include embryonal, teratoma, choriocarcinoma, and mixed tumors.

Tumorigenic Hypothesis for Germ Cell Tumor Development

During embryonal development, the totipotential germ cells can travel down normal differentiation pathways and become spermatocytes. However, if these totipotential germ cells travel down abnormal developmental pathways, seminoma or embryonal carcinomas (totipotential tumor cells) develop. If the embryonal cells undergo further differentiation along intraembryonic pathways, teratoma will result. If the embryonal cells undergo further differentiation along extraembryonic pathways, either choriocarcinoma or yolk sac tumors are formed (Figure 23–1). This model helps to explain why specific histologic patterns of testicular tumors produce certain tumor markers. Note that yolk sac tumors produce alpha-fetoprotein (AFP) just as the yolk sac produces AFP in normal development. Likewise, choriocarcinoma produces human chorionic
gonadotropin (hCG) just as the normal placenta produces hCG.

**Pathology**

**A. SEMINOMA (35%)**

Three histologic subtypes of pure seminoma have been described. However, stage for stage, there is no prognostic significance to any of these subtypes. Classic seminoma accounts for 85% of all seminomas and is most common in the fourth decade of life. Grossly, coalescing gray nodules are observed. Microscopically, monotonous sheets of large cells with clear cytoplasm and densely staining nuclei are seen. It is noteworthy that syncytiotrophoblastic elements are seen in approximately 10–15% of cases, an incidence that corresponds approximately to the incidence of hCG production in seminomas.

Anaplastic seminoma accounts for 5–10% of all seminomas. Diagnosis requires the presence of 3 or more mitoses per high-power field, and the cells demonstrate a higher degree of nuclear pleomorphism than the classic types. Anaplastic seminoma tends to present at a higher stage than the classic variety. When stage is taken into consideration, however, this subtype does not convey a worse prognosis.

Spermatocytic seminoma accounts for 5–10% of all seminomas. Microscopically, cells vary in size and are characterized by densely staining cytoplasm and round nuclei that contain condensed chromatin. More than half the patients with spermatocytic seminoma are over the age of 50.

**B. EMBRYONAL CELL CARCINOMA (20%)**

Two variants of embryonal cell carcinoma are common: the adult type and the infantile type, or yolk sac tumor (also called endodermal sinus tumor). Histologic structure of the adult variant demonstrates marked pleomorphism and indistinct cellular borders. Mitotic figures and giant cells are common. Cells may be arranged in sheets, cords, glands, or papillary structures. Extensive hemorrhage and necrosis may be observed grossly.

The infantile variant, or yolk sac tumor, is the most common testicular tumor of infants and children. When seen in adults, it usually occurs in mixed histologic types and possibly is responsible for AFP production in these tumors. Microscopically, cells demonstrate vacuolated cytoplasm secondary to fat and glycogen deposition and are arranged in a loose network with large intervening cystic spaces. Embryoid bodies are commonly seen and resemble 1- to 2-week-old embryos consisting of a cavity surrounded by syncytiotrophoblastic elements.

**C. TERATOMA (5%)**

Teratomas may be seen in both children and adults. They contain more than one germ cell layer in various stages of maturation and differentiation. Grossly, the tumor appears lobulated and contains variable-sized cysts filled with gelatinous or mucinous material. Mature teratoma may have elements resembling benign structures derived from ectoderm, mesoderm, and endoderm, while immature teratoma consists of undifferentiated primitive tissue. In contrast to its ovarian counterpart, the mature teratoma of the testis does not attain the same degree of differentiation as teratoma of the ovary. Microscopically, ectoderm may be represented by squamous epithelium or neural tissue; endoderm by intestinal, pancreatic, or respiratory tissue; and mesoderm by smooth or skeletal muscle, cartilage, or bone.
D. CHORIOCARCINOMA (<1%)

Pure choriocarcinoma is rare. Lesions tend to be small within the testis and usually demonstrate central hemorrhage on gross inspection. Microscopically, syncytiotrophoblasts must be visualized. The syncytial elements are typically large, multinucleated cells with vacuolated, eosinophilic cytoplasm; the nuclei are large, hyperchromatic, and irregular. Cytotrophoblasts are uniform cells with distinct cell borders, clear cytoplasm, and a single nucleus.

Clinically, choriocarcinomas behave in an aggressive fashion characterized by early hematogenous spread. Paradoxically, small intratesticular lesions can be associated with widespread metastatic disease.

E. MIXED CELL TYPE (40%)

Within the category of mixed cell types, most (up to 25% of all testicular tumors) are teratocarcinomas, which are a combination of teratoma and embryonal cell carcinoma. Up to 6% of all testicular tumors are of the mixed cell type, with seminoma being one of the components. Treatment for these mixtures of seminoma and NSGCT is similar to that of NSGCT alone.

F. CARCINOMA IN SITU (CIS)

In a series of 250 patients with unilateral testicular cancer, Berthelsen et al (1982) demonstrated the presence of CIS in 13 (5.2%) of the contralateral testes. This is approximately twice the overall incidence of bilateral testicular cancer. The presence of contralateral atrophy or ultrasonographic microlithiasis in patients with testicular tumors warrants contralateral biopsy. If diagnosed, CIS is usually treated by external beam radiation therapy.

Patterns of Metastatic Spread

With the exception of choriocarcinoma, which demonstrates early hematogenous spread, germ cell tumors of the testis typically spread in a stepwise lymphatic fashion. Lymph nodes of the testis extend from T1 to L4 but are concentrated at the level of the renal hilum because of their common embryologic origin with the kidney. The primary landing site for the left testis is the para-aortic area at the level of the right renal hilum. Stepwise spread, in order, is to the precaval, preaortic, paracaval, right common iliac, and right external iliac lymph nodes. The primary landing site for the left testis is the para-aortic area at the level of the left renal hilum. Stepwise spread, in order, is to the preaortic, left common iliac, and left external iliac lymph nodes. In the absence of disease on the left side, no crossover metastases to the right side have ever been identified. However, right-to-left crossover metastases are common. These observations have resulted in modified surgical dissections to preserve ejaculation in selected patients (see section on Treatment, following).
A testicular mass or diffuse enlargement is found in most cases. The mass is typically firm and nontender and the epididymis should be easily separable from it. A hydrocele may accompany the testicular tumor and help to camouflage it. Transillumination of the scrotum can help to distinguish between these entities.

Palpation of the abdomen may reveal bulky retroperitoneal disease; assessment of supraclavicular, scalene, and inguinal nodes should be performed. Gynecomastia is present in 5% of all germ cell tumors but may be present in 30–50% of Sertoli and Leydig cell tumors. Its cause seems to be related to multiple complex hormonal interactions involving testosterone, estrone, estradiol, prolactin, and hCG. Hemoptysis may be seen in advanced pulmonary disease.

### C. Laboratory Findings and Tumor Markers

Anemia may be detected in advanced disease. Liver function tests may be elevated in the presence of hepatic metastases. Renal function may be diminished (elevated serum creatinine) if ureteral obstruction secondary to bulky retroperitoneal disease is present. The assessment of renal function (creatinine clearance) is mandatory in patients with advanced disease who require chemotherapy.

Several biochemical markers are of importance in the diagnosis and management of testicular carcinoma, including AFP, hCG, and LDH. Alpha-fetoprotein is a glycoprotein with a molecular mass of 70,000 daltons and a half-life of 4–6 days. Although present in fetal serum in high levels, beyond the age of 1 year it is present only in trace amounts. While present to varying degrees in many NSGCTs (Table 23–2), it is never found in seminomas.

Human chorionic gonadotropin is a glycoprotein with a molecular mass of 38,000 daltons and a half-life of 24 hours. It is composed of 2 subunits: alpha and beta. The alpha subunit is similar to the alpha subunits of luteinizing hormone (LH), follicle-stimulating hormone (FSH), and thyroid-stimulating hormone (TSH). The beta subunit conveys the activity to each of these hormones and allows for a highly sensitive and specific radioimmunoassay in the determination of hCG levels. A normal man should not have significant levels of beta-hCG. While more commonly elevated in NSGCTs, hCG levels may be elevated in up to 7% of seminomas.

Lactic acid dehydrogenase (LDH) is a cellular enzyme with a molecular mass of 134,000 daltons that has 5 isoenzymes; it is normally found in muscle (smooth, cardiac, skeletal), liver, kidney, and brain. Elevation of total serum LDH and in particular isoenzyme-I was shown to correlate with tumor burden in NSGCTs. LDH may also be elevated in seminoma.

Other markers have been described for testis cancer, including placental alkaline phosphatase (PLAP) and gamma-glutamyl transpeptidase (GGT). These markers, however, have not contributed as much to the management of patients as those mentioned previously.

### D. Imaging

The primary testicular tumor can be rapidly and accurately assessed by scrotal ultrasonography. This technique can determine whether the mass is truly intratesticular, can be used to distinguish the tumor from epididymal pathology, and may also facilitate testicular examination in the presence of a hydrocele.

Once the diagnosis of testicular cancer has been established by inguinal orchectomy, careful clinical staging of disease is mandatory. Chest radiographs (posteroanterior and lateral) and computed tomography (CT scan) of the abdomen and pelvis are used to assess the 2 most common sites of metastatic spread, namely, the lungs and retroperitoneum. The role of CT scanning of the chest remains...
controversial because of its decreased specificity. Of note is the fact that routine chest x-rays detect 85–90% of pulmonary metastases. Pedal lymphangiography (LAG) is rarely used owing to its invasiveness as well as low specificity, although it may be warranted in patients undergoing a surveillance protocol (see section on treatment).

Differential Diagnosis

An incorrect diagnosis is made at the initial examination in up to 25% of patients with testicular tumors and may result in delay in treatment or a suboptimal surgical approach (scrotal incision) for exploration. Epididymitis or epididymoorchitis is the most common misdiagnosis in patients with testis cancer. Early epididymitis should reveal an enlarged, tender epididymis that is clearly separable from the testis. In advanced stages, the inflammation may spread to the testis and result in an enlarged, tender, and indurated testis and epididymis. A history of acute onset of symptoms including fever, urethral discharge, and irritative voiding symptoms may make the diagnosis of epididymitis more likely. Ultrasonography may identify the enlarged epididymis as the cause of the scrotal mass.

Hydrocele is the second most common misdiagnosis. Transillumination of the scrotum may readily distinguish between a translucent, fluid-filled hydrocele and a solid testicular tumor. Since 5–10% of testicular tumors may be associated with hydroceles, if the testis cannot be adequately examined a scrotal ultrasound examination is mandatory. Aspiration of the hydrocele should be avoided because positive cytologic results have been reported in hydroceles associated with testicular tumors.

Other diagnoses to be considered include spermatocele, a cystic mass most commonly found extending from the head of the epididymis; hematocoele associated with trauma; granulomatous orchitis, most commonly resulting from tuberculosis and associated with beading of the vas deferens; and varicocele, which is engorgement of the pampiniform plexus of veins in the spermatic cord and should disappear when the patient is in the supine position.

Although most intratesticular masses are malignant, one benign lesion, an epidermoid cyst, may be seen on rare occasions. Usually these cysts are very small benign nodules located just underneath the tunica albuginea; however, on occasion they can be large. The diagnosis is usually made following inguinal orchietomy; as frozen sections, the larger lesions are often difficult to distinguish from teratoma.

### Treatment

Inguinal exploration with cross-clamping of the spermatic cord vasculature and delivery of the testis into the field is the mainstay of exploration for a possible testis tumor. If cancer cannot be excluded by examination of the testis, radical orchietomy is warranted. Scrotal approaches and open testicular biopsies should be avoided. Further therapy depends on the histologic characteristics of the tumor as well as the clinical stage.

#### A. Low-Stage Seminoma

Seminoma is exquisitely radiosensitive. About 95% of all stage I seminomas are cured with radical orchietomy and retroperitoneal irradiation (usually 2500–3000 cGy). This low dose of radiation is usually well tolerated, with minimal, if any, gastrointestinal side effects.

Low-volume retroperitoneal disease also can be treated effectively with retroperitoneal irradiation with an average 5-year survival rate of 87%. Prophylactic mediastinal radiation is no longer employed because this may cause considerable myelosuppression and thus compromise the patient’s ability to receive chemotherapy if required. Chemotherapy should be used as salvage therapy for patients who relapse following irradiation.

#### B. High-Stage Seminoma

Patients with bulky seminoma and any seminoma associated with an elevated AFP should receive primary chemotherapy. Seminomas are also sensitive to platinum-based regimens, as are their NSGCT counterparts. Some of the successful regimens include cisplatin, etoposide, and bleomycin (PEB); vinblastine, cyclophosphamide, dactinomycin, bleomycin, and cisplatin (VAB-6); and cisplatin and etoposide. All seminomas receive low-risk chemotherapy regimens, which currently consist of cisplatin and etoposide (4 cycles) or 3 cycles of PEB.

Ninety percent of patients with advanced disease achieve a complete response with chemotherapy. Residual retroperitoneal masses following chemotherapy are often fibrosis (90%) unless the mass is well circumscribed and in excess of 3 cm, under which circumstances approximately 40% of patients harbor residual seminoma. In such cases surgical excision is warranted.

#### C. Low-Stage Nonseminomatous Germ Cell Tumors

Standard treatment for stage I disease in the United States has included retroperitoneal lymph node dissection (RPLND). However, because three-fourths of patients

<table>
<thead>
<tr>
<th>Histologic Type</th>
<th>hCG (%)</th>
<th>AFP (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Seminoma</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>Teratoma</td>
<td>25</td>
<td>38</td>
</tr>
<tr>
<td>Teratocarcinoma</td>
<td>57</td>
<td>64</td>
</tr>
<tr>
<td>Embryonal</td>
<td>60</td>
<td>70</td>
</tr>
<tr>
<td>Choriocarcinoma</td>
<td>100</td>
<td>0</td>
</tr>
</tbody>
</table>
with clinical stage I disease are cured by orchiectomy alone and the morbidity of RPLND is not negligible, other alternatives have been explored. These options include surveillance and modified RPLND.

Surveillance in stage I NSGCT was proposed because, as mentioned previously, 75% of patients with clinical stage I disease have, in fact, pathologic stage I disease. In addition, infertility related to disruption of sympathetic nerve fibers is common following RPLND. Clinical staging has been markedly improved in the presence of CT scanning and LAG. And finally, effective chemotherapy regimens have been developed for relapse. Patients are considered candidates for surveillance if the tumor is an NSGCT confined within the tunica albuginea, the tumor does not demonstrate vascular invasion, tumor markers normalize after orchiectomy, radiographic imaging shows no evidence of disease (chest x-ray [CXR], CT), and the patient is considered reliable.

Surveillance should be considered an active process on the part of both the physician and the patient. Patients are followed monthly for the first 2 years and bimonthly in the third year. Tumor markers are obtained at each visit, and CXR and CT scans are obtained every 3–4 months. Follow-up continues beyond the initial 3 years. Most relapses occur, however, within the first 8–10 months. With rare exceptions, patients who relapse can be cured by chemotherapy or surgery, or both.

Retroperitoneal lymph node dissection has been the preferred treatment of low-stage NSGCTs in the United States until recently. A thoracoabdominal or midline transabdominal approach may be used, and all nodal tissue between the ureters from the renal vessels to the bifurcation of the common iliac vessels is removed. Patients with negative nodes or N1 disease do not require adjuvant therapy, whereas the recommendation for those with N2 disease is to receive 2 cycles of chemotherapy because their relapse rate approaches 50%.

While effective in surgically staging and potentially curing a subset of patients, RPLND is associated with significant morbidity, especially with respect to fertility in young men. With a standard RPLND, sympathetic nerve fibers are disrupted, resulting in loss of seminal emission. Currently a modified RPLND has been developed that preserves ejaculation in up to 90% of patients. By modifying the dissection below the level of the inferior mesenteric artery to include only the nodal tissue ipsilateral to the tumor, important sympathetic fibers from the contralateral side are preserved, thus maintaining emission.

An alternative approach to patients with clinical stage I disease and vascular invasion in the primary is 2 cycles of chemotherapy. While obviating the need for surgery, such an approach is associated with neurotoxicity and fertility issues for these young patients.

D. HIGH-STAGE NONSEMINOMATOUS GERM CELL TUMORS

Patients with bulky retroperitoneal disease (>3-cm nodes or 3 or more 1-cm cuts on CT scan) or metastatic NSGCT are treated with primary platinum-based combination chemotherapy following orchiectomy. If tumor markers normalize and a residual mass is apparent on imaging studies, resection of that mass is mandatory, because 20% of the time it will harbor residual cancer, 40% of the time it will be teratoma, and 40% of the time it will be fibrosis (Figure 23–2). In patients with residual cancer in the resected tissue, the histologic picture is usually embryonal cell carcinoma, but malignant teratoma is seen in less than 5% of cases. Malignant teratoma is unresponsive to chemotherapy, and only 15% of patients survive following surgical resection. If tumor markers fail to normalize following primary chemotherapy, salvage chemotherapy is required (cisplatin, etoposide, bleomycin, ifosfamide). Even if patients

Figure 23–2. Upper: Computed tomography scan of patient with bulky retroperitoneal mass after radical orchiectomy for embryonal carcinoma. Lower: Residual cystic mass after chemotherapy; it was resected and found to be teratoma.
attain a complete response after chemotherapy (normal tumor markers, no mass on CT scan or CXR), some investigators advocate an RPLND because viable germ cell tumor may be seen in up to 10% of cases.

Although the treatment plan described cures up to 70% of patients with high-volume disease, there are patients who fail to respond. Also, the potential complications from chemotherapy including sepsis, neuropathy, renal toxicity, and death must be considered. It is thus apparent that it is important to be able to discriminate between patients who are likely to respond to standard chemotherapy (low risk) and those who may require more aggressive regimens (high risk). High-risk patients are only patients with NSGCT that have one of the following features: mediastinal primary tumor, nonpulmonary visceral metastases, or S3 marker levels. The rate of decline of serum tumor markers during chemotherapy has also been used to predict response in patients with advanced disease.

Follow-up Care

All patients with testicular cancer require regular follow-up care. As discussed previously, patients on a surveillance protocol require vigorous follow-up. Those who have undergone surgery (RPLND) or radiotherapy are followed at 3-month intervals for the first 2 years, then every 6 months until 5 years, and then yearly. Follow-up visits should include careful examination of the remaining testis, the abdomen, and the lymph node areas. Laboratory investigation should include AFP, hCG, and LDH levels. A CXR and an abdominal film (if an LAG was performed) should also be included at each visit. Abdominal CT scans are used less frequently as risk of relapse in the retroperitoneum is low following RPLND.

Prognosis

Survival in testicular cancer has improved dramatically over the past several years, reflecting the continuing improvement and refinement in combination chemotherapy. For seminoma treated by orchiectomy and radiotherapy, the 5-year disease-free survival rate is 98% for stage I and 92–94% for stage II-A in several recent series. Higher-stage disease treated by orchiectomy and primary chemotherapy has a 5-year disease-free survival rate of 35–75%, yet the lower value comes from older series in which more crude chemotherapy regimens were employed.

Survival in patients with NSGCTs treated by orchiectomy and RPLND for stage I disease ranges from 96 to 100%. For low-volume stage II disease treated with chemotherapy plus surgery, greater than 90% 5-year disease-free survival rates are attainable. Patients with bulky retroperitoneal or disseminated disease treated with primary chemotherapy followed by surgery have a 5-year disease-free survival rate of 55–80%.

Currently much work is being done to stratify patients into “high-risk” and “low-risk” groups so that treatment regimens may be modified in order to increase survival and decrease morbidity.

NON–GERM CELL TUMORS OF THE TESTIS

Approximately 5–6% of all testis tumors are non–germ cell tumors of the testis. Three types will be considered, namely, Leydig cell tumors, Sertoli cell tumors, and gonadoblastomas.

1. Leydig Cell Tumors

Epidemiology & Pathology

Leydig cell tumors are the most common non–germ cell tumors of the testis and account for 1–3% of all testicular tumors. They follow a bimodal age distribution: the 5- to 9-year-old and the 25- to 35-year-old age groups. Twenty-five percent of these tumors occur in childhood. Bilaterality is seen in 5–10% of cases. The cause of these tumors is unknown; unlike germ cell tumors, there is no association with cryptorchidism.

Pathologic examination reveals a small, yellow, well-circumscribed lesion devoid of hemorrhage or necrosis. Microscopically, hexagonally shaped cells with granular, eosinophilic cytoplasm containing lipid vacuoles are seen. Reinke crystals are fusiform-shaped cytoplasmic inclusions that are pathognomonic for Leydig cells.

Clinical Findings

Prepubertal children usually present with virilization, and tumors are benign. Adults are usually asymptomatic, although gynecomastia may be present in 20–25%. Ten percent of tumors in adults are malignant. Laboratory findings include elevated serum and urinary 17-ketosteroids as well as estrogens.

Treatment & Prognosis

Radical orchiectomy is the initial treatment for Leydig cell tumors. Clinical staging is similar to that for germ cell tumors, and levels of the 17-ketosteroids can be helpful in distinguishing between benign and malignant lesions. Elevations of 10–30 times normal are typical of malignancy. RPLND is recommended for malignant lesions. Because of the rarity of these lesions, the role of chemotherapy remains to be defined. Prognosis is excellent for benign lesions, while it remains poor for patients with disseminated disease.

2. Sertoli Cell Tumors

Epidemiology & Pathology

Sertoli cell tumors are exceedingly rare, composing less than 1% of all testicular tumors. A bimodal age
distribution is seen: 1 year old or younger and the 20- to 45-year-old age group. Approximately 10% of the lesions are malignant. Gross examination reveals a yellow or gray-white lesion with cystic components. Benign lesions are well circumscribed, while malignant lesions show ill-defined borders. Microscopically, tumors appear heterogeneous with mixed amounts of epithelial and stromal components. Sertoli cells are columnar or hexagonal cells with a large nucleus and solitary nucleolus and contain vacuolated cytoplasm.

**Clinical Findings**

A testicular mass is the most common presentation. Virilization is often seen in children, and gynecomastia may be present in 30% of adults. Because of the rarity of these tumors, minimal endocrine data are available on these patients.

**Treatment**

Radical orchiectomy is the initial procedure of choice. In cases of malignancy, RPLND is indicated; however, the roles of chemotherapy and radiotherapy remain unclear.

3. **Gonadoblastomas**

**Epidemiology & Pathology**

Gonadoblastomas comprise 0.5% of all testicular tumors and are almost exclusively seen in patients with some form of gonadal dysgenesis. Most of these tumors occur in patients under 30 years of age, although the age distribution ranges from infancy to beyond 70 years. Gross examination reveals a yellow or gray-white lesion that can vary in size from microscopic to greater than 20 cm and may exhibit calcifications. Microscopically, 3 cell types are seen: Sertoli cells, interstitial cells, and germ cells.

**Clinical Findings**

The clinical manifestations are predominantly related to the underlying gonadal dysgenesis and are discussed elsewhere in this book. It is noteworthy that four-fifths of patients with gonadoblastomas are phenotypic females. Males typically have cryptorchidism or hypospadias.

**Treatment & Prognosis**

Radical orchiectomy is the primary treatment of choice. In the presence of gonadal dysgenesis, a contralateral gonadectomy is recommended because the tumor tends to be bilateral in 50% of cases in this setting. Prognosis is excellent.

**SECONDARY TUMORS OF THE TESTIS**

Secondary tumors of the testis are rare. Three categories are considered: lymphoma, leukemia, and metastatic tumors.

1. **Lymphoma**

**Epidemiology & Pathology**

Lymphoma is the most common testicular tumor in a patient over the age of 50 and is the most common secondary neoplasm of the testis, accounting for 5% of all testicular tumors. It may be seen in 3 clinical settings: (1) late manifestation of widespread lymphoma; (2) initial presentation of clinically occult disease; and (3) primary extranodal disease. Gross examination reveals a bulging, gray or pink lesion with ill-defined margins. Hemorrhage and necrosis are common. Microscopically, diffuse histiocytic lymphoma is the most common type.

**Clinical Findings**

Painless enlargement of the testis is common. Generalized constitutional symptoms occur in one-fourth of patients. Bilateral testis involvement occurs in 50% of patients, usually asynchronously.

**Treatment & Prognosis**

Fine needle aspiration should be considered in patients with a known or suspected diagnosis of lymphoma while radical orchiectomy is reserved for those with suspected primary lymphoma of the testicle. Further staging and treatment should be handled in conjunction with the medical oncologist. Prognosis is related to the stage of disease. Some reports support adjuvant chemotherapy for primary testicular lymphoma, with improved survival rates of up to 93% after 44 months of follow-up.

2. **Leukemic Infiltration of the Testis**

The testis is a common site of relapse for children with acute lymphocytic leukemia. Bilateral involvement may be present in one-half of cases. Testis biopsy rather than orchiectomy is the diagnostic procedure of choice. Bilateral testicular irradiation with 20 Gy and reinstitution of adjuvant chemotherapy constitute the treatment of choice. Prognosis remains guarded.

3. **Metastatic Tumors**

Metastasis to the testis is rare. These lesions are typically incidental findings at autopsy. The most common primary site is the prostate, followed by the lung, gastrointestinal tract, melanoma, and kidney. The typical pathologic finding is neoplastic cells in the interstitium with relative sparing of the seminiferous tubules.
EXTRAGONADAL GERM CELL TUMORS

Epidemiology & Pathology
Extragonadal germ cell tumors are rare, accounting for approximately 3% of all germ cell tumors. Debate continues over whether these lesions originate from “burned-out” testicular primaries or originate de novo. Most retroperitoneal tumors have their origin from a testicular primary, whereas mediastinal germ cell tumors are truly ectopic.

The most common sites of origin in decreasing order are mediastinum, retroperitoneum, sacrococcygeal area, and pineal gland. All germ cell types may be observed. Seminoma composes more than half of the retroperitoneal and mediastinal tumors.

Clinical Findings
Clinical presentation depends on the site and volume of disease. Mediastinal lesions may present with pulmonary complaints. Retroperitoneal lesions may present with abdominal or back pain and a palpable mass. Sacrococcygeal tumors are most commonly seen in neonates and may present with a palpable mass and bowel or urinary obstruction. Pineal tumors may present with headache, visual or auditory complaints, or hypopituitarism.

Metastatic spread is to regional lymph nodes, lung, liver, bone, and brain. Metastatic workup is similar, therefore, to that of testicular germ cell tumors. A careful testicular examination is mandatory along with ultrasonography to exclude an occult testicular primary.

Treatment & Prognosis
Treatment of extragonadal germ cell tumors parallels that of testicular tumors. Low-volume seminoma can be managed with radiotherapy. High-volume seminoma should receive primary chemotherapy. Prognosis parallels that of testicular seminoma. Primary chemotherapy should be employed for nonseminomatous elements with surgical excision of residual masses; however, prognosis remains poor for these patients.

TUMORS OF THE EPIDIDYMIS, PARATESTICULAR TISSUES, & SPERMATIC CORD

Primary tumors of the epididymis are rare and are most commonly benign. Adenomatoid tumors of the epididymis are the most common and typically occur in the third and fourth decade of life. They are typically asymptomatic, solid lesions that arise from any portion of the epididymis. Leiomyomas are the second most common tumor of the epididymis. These lesions tend to be painful and are often associated with a hydrocele. Cystadenomas are benign lesions of the epididymis that are bilateral in 30% of cases and are frequently seen in association with von Hippel-Lindau disease. Histologically these lesions are difficult to distinguish from renal cell carcinoma. Malignant lesions of the epididymis are extremely rare. In general, an inguinal approach should be used, and if frozen section confirms a benign lesion, epididymectomy should be performed. If a malignant tumor is diagnosed, radical orchietomy must be performed.

Tumors of the spermatic cord are typically benign. Lipomas of the cord account for most of these lesions. Of the malignant lesions, rhabdomyosarcoma is the most common, followed by leiomyosarcoma, fibrosarcoma, and liposarcoma.

Clinical diagnosis of tumors of the spermatic cord can be difficult. Differentiating between a hernia and a spermatic cord tumor may be possible only at exploration. In general, these lesions should be approached through an inguinal incision. The cord should be occluded at the internal ring and frozen sections obtained. If malignancy is diagnosed, attention should be directed toward performing wide local excision to avoid local recurrence. Staging of disease is similar to that of testicular tumors. For rhabdomyosarcoma, RPLND should be performed with adjuvant radiotherapy and chemotherapy. The value of RPLND for the other malignant spermatic cord tumors remains to be determined. Prognosis relates to the histologic status, stage, and site of disease.

TUMORS OF THE PENIS

Epidemiology & Risk Factors
Carcinoma of the penis accounts for less than 1% of cancers among males in the United States, with approximately 1–2 new cases being reported per 100,000 men. There is marked variation in incidence with geographic location. In areas such as Africa and regions of South America, penile carcinoma may compose 10–20% of all malignant lesions. Penile carcinoma occurs most commonly in the sixth decade of life, although rare case reports have included children.

The one etiologic factor most commonly associated with penile carcinoma is poor hygiene. The disease is virtually unheard of in males circumcised near birth. One theory postulates that smegma accumulation under the phim-
otic foreskin results in chronic inflammation leading to carcinoma. A viral cause has also been suggested as a result of the association of this tumor with cervical carcinoma.

Pathology

A. Precancerous Dermatologic Lesions

Leukoplakia is a rare condition that most commonly occurs in diabetic patients. A white plaque typically involving the meatus is seen. Histologic examination reveals acanthosis, hyperkeratosis, and parakeratosis. This lesion may precede or occur simultaneously with penile carcinoma.

Balanitis xerotica obliterans is a white patch originating on the prepuce or glans and usually involving the meatus. The condition is most commonly observed in middle-aged diabetic patients. Microscopic examination reveals atrophic epidermis and abnormalities in collagen deposition.

Giant condylomata acuminata are cauliflower-like lesions arising from the prepuce or glans. The cause is believed to be viral (human papillomavirus). These lesions may be difficult to distinguish from well-differentiated squamous cell carcinoma.

B. Carcinoma in Situ (Bowen Disease, Erythroplasia of Queyrat)

Bowen disease is a squamous cell carcinoma in situ typically involving the penile shaft. The lesion appears as a red plaque with encrustations. Erythroplasia of Queyrat is a velvety, red lesion with ulcerations that usually involve the glans. Microscopic examination shows typical, hyperplastic cells in a disordered array with vacuolated cytoplasm and mitotic figures.

C. Invasive Carcinoma of the Penis

Squamous cell carcinoma composes most penile cancers. It most commonly originates on the glans, with the next most common sites, in order, being the prepuce and shaft. The appearance may be papillary or ulcerative.

Verrucous carcinoma is a variant of squamous cell carcinoma composing 5–16% of penile carcinomas. This lesion is papillary in appearance and on histologic examination is noted to have a well-demarcated deep margin unlike the infiltrating margin of the typical squamous cell carcinoma.

Patterns of Spread

Invasive carcinoma of the penis begins as an ulcerative or papillary lesion, which may gradually grow to involve the entire glans or shaft of the penis. Buck’s fascia represents a barrier to corporal invasion and hematogenous spread. Primary dissemination is via lymphatic channels to the femoral and iliac nodes. The prepuce and shaft skin drain into the superficial inguinal nodes (superficial to fascia lata), while the glans and corporal bodies drain to both superficial and deep inguinal nodes (deep to fascia lata). There are many cross-communications, so that penile lymphatic drainage is bilateral to both inguinal areas. Drainage from the inguinal nodes is to the pelvic nodes. Involvement of the femoral nodes may result in skin necrosis and infection or femoral vessel erosion and hemorrhage. Distant metastases are clinically apparent in less than 10% of cases and may involve lung, liver, bone, or brain.

Tumor Staging

The staging system used most commonly in the United States was proposed by Jackson (1966), as follows: In stage I, the tumor is confined to the glans or prepuce. Stage II involves the penile shaft. Stage III has operable inguinal node metastasis. In stage IV, the tumor extends beyond the penile shaft, with inoperable inguinal or distant metastases. The TNM classification of the American Joint Committee (1996) is given in Table 23–3.

Clinical Findings

A. Symptoms

The most common complaint at presentation is the lesion itself. It may appear as an area of induration or erythema, an ulceration, a small nodule, or an exophytic growth. Phimosis may obscure the lesion and result in a delay in seeking medical attention. In fact, 15–50% of patients delay

<table>
<thead>
<tr>
<th>TNM Classification of Tumors of the Penis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>T</strong>—Primary tumor</td>
</tr>
<tr>
<td>TX: Cannot be assessed</td>
</tr>
<tr>
<td><strong>T</strong>0: No evidence of primary tumor</td>
</tr>
<tr>
<td><strong>T</strong>is: Carcinoma in situ</td>
</tr>
<tr>
<td><strong>Ta</strong>: Noninvasive verrucous carcinoma</td>
</tr>
<tr>
<td><strong>T</strong>1: Invades subepithelial connective tissue</td>
</tr>
<tr>
<td><strong>T</strong>2: Invades corpus spongiosum or cavernosum</td>
</tr>
<tr>
<td><strong>T</strong>3: Invades urethra or prostate</td>
</tr>
<tr>
<td><strong>T</strong>4: Invades other adjacent structures</td>
</tr>
<tr>
<td><strong>N</strong>—Regional lymph nodes</td>
</tr>
<tr>
<td><strong>NX</strong>: Cannot be assessed</td>
</tr>
<tr>
<td><strong>N</strong>0: No regional lymph node metastasis</td>
</tr>
<tr>
<td><strong>N</strong>1: Metastasis in single superficial inguinal node</td>
</tr>
<tr>
<td><strong>N</strong>2: Metastasis in multiple or bilateral superficial inguinal nodes</td>
</tr>
<tr>
<td><strong>N</strong>3: Metastasis in deep inguinal or pelvic nodes</td>
</tr>
<tr>
<td><strong>M</strong>—Distant metastasis</td>
</tr>
<tr>
<td><strong>MX</strong>: Cannot be assessed</td>
</tr>
<tr>
<td><strong>M</strong>0: No distant metastasis</td>
</tr>
<tr>
<td><strong>M</strong>1: Distant metastasis present</td>
</tr>
</tbody>
</table>

Source: American Joint Committee on Cancer: *TNM Classification—Genitourinary Sites*, 1996.
for at least 1 year in seeking medical attention. Other symptoms include pain, discharge, irritative voiding symptoms, and bleeding.

B. Signs

Lesions are typically confined to the penis at presentation. The primary lesion should be characterized with respect to size, location, and potential corporal body involvement. Careful palpation of the inguinal area is mandatory because more than 50% of patients present with enlarged inguinal nodes. This enlargement may be secondary to inflammation or metastatic spread.

C. Laboratory Findings

Laboratory evaluation is typically normal. Anemia and leukocytosis may be present in patients with long-standing disease or extensive local infection. Hypercalcemia in the absence of osseous metastases may be seen in 20% of patients and appears to correlate with volume of disease.

D. Imaging

Metastatic workup should include CXR, bone scan, and CT scan of the abdomen and pelvis. Disseminated disease is present in less than 10% of patients at presentation.

Differential Diagnosis

In addition to the dermatologic lesions discussed previously, carcinoma of the penis must be differentiated from several infectious lesions. Syphilitic chancre may present as a painless ulceration. Serologic and darkfield examination should establish the diagnosis. Chancroid typically appears as a painful ulceration of the penis. Selective cultures for *Haemophilus ducreyi* should identify the cause. Condylomata acuminata appear as exophytic, soft, “grape cluster” lesions anywhere on the penile shaft or glans. Biopsy can distinguish this lesion from carcinoma if any doubt exists.

Treatment

A. Primary Lesion

Biopsy of the primary lesion is mandatory to establish the diagnosis of malignancy. Treatment varies depending on the pathology as well as the location of the lesion. Carcinoma in situ may be treated conservatively in reliable patients. Fluorouracil cream application or neodymium:YAG laser treatment is effective for CIS and is preserving of the penis. Patients must come for frequent follow-up examinations to monitor response.

The goal of treatment in invasive penile carcinoma is complete excision with adequate margins. For lesions involving the prepuce, this may be accomplished by simple circumcision. For lesions involving the glans or distal shaft, partial penectomy with a 2-cm margin to decrease local recurrence has traditionally been suggested. Less aggressive surgical resections such as Mohs micrographic surgery and local excisions directed at penile preservation have gained popularity. For lesions involving the proximal shaft or when partial penectomy results in a penile stump of insufficient length for sexual function or directing the urinary stream, total penectomy with perineal urethrostomy has been recommended.

B. Regional Lymph Nodes

As discussed previously, penile carcinoma spreads primarily to the inguinal lymph nodes. However, enlargement of inguinal nodes at presentation does not necessarily imply metastatic disease. In fact, up to 50% of the time this enlargement is caused by inflammation. Thus patients who present with enlarged inguinal nodes should undergo treatment of the primary lesion followed by a 4- to 6-week course of oral broad-spectrum antibiotics. Persistent adenopathy following antibiotic treatment should be considered to be metastatic disease, and sequential bilateral ilioinguinal node dissections should be performed. If lymphadenopathy resolves with antibiotics, observation in low-stage primary tumors (Tis, T1) is warranted. However, if lymphadenopathy resolves in higher-stage tumors, more limited lymph node samplings should be considered, such as Mohs micrographic surgery and local excisions directed at penile preservation.
as the sentinel node biopsy described by Cabanas (1977) or a modified (limited) dissection as suggested by Catalona (1988) (Figure 23–3). If positive nodes are encountered, bilateral ilioinguinal node dissection should be performed. A decision tree for penile carcinoma is presented in Figure 23–4. Patients who initially have clinically negative nodes but in whom clinically palpable nodes later develop should undergo a unilateral ilioinguinal node dissection.

Patients who have inoperable disease and bulky inguinal metastases are treated with chemotherapy (cisplatin and 5-fluorouracil). In some cases regional radiotherapy can provide significant palliation by delaying ulceration and infectious complications and alleviating pain.

**C. SYSTEMIC DISEASE**

Four chemotherapeutic agents demonstrate activity against penile carcinoma: bleomycin, methotrexate, cisplatin, and 5-fluorouracil. However, no long-term responders have been reported. The rarity of the disease in the United States has resulted in limited clinical trials.

**Prognosis**

Survival in penile carcinoma correlates with the presence or absence of nodal disease. Five-year survival rates for patients with node-negative disease range from 65% to 90%. For patients with positive inguinal nodes, this rate decreases to 30–50% and with positive iliac nodes decreases to less than 20%. In the presence of soft-tissue or bony metastases, no 5-year survivors have been reported.

**Other Penile Tumors**

Squamous cell carcinoma accounts for 98% of penile cancers. Sporadic cases of melanoma, basal cell carcinoma, and Paget disease have been reported. The incidence of Kaposi sarcoma of the penis is increasing with the increasing prevalence of the human immunodeficiency virus. It appears as a painful papule on the glans or shaft with bluish-purple discoloration. These lesions tend to be radiosensitive.
TUMORS OF THE SCROTUM

Tumors of the scrotal skin are rare. The most common benign lesion is a sebaceous cyst. Squamous cell carcinoma is the most common malignant tumor of the scrotum, although rare cases of melanoma, basal cell carcinoma, and Kaposi sarcoma have been reported. In the past, squamous cell carcinoma of the scrotum most commonly resulted from exposure to environmental carcinogens including chimney soot, tars, paraffin, and some petroleum products. Today, most cases result from poor hygiene and chronic inflammation.

Biopsy of the scrotal lesion must be performed to establish a histologic diagnosis. Wide excision with a 2-cm margin should be performed for malignant tumors. Surrounding subcutaneous tissue should be excised with the primary tumor; however, resection of the scrotal contents is rarely necessary. Primary closure using the redundant scrotal skin is usually possible. The management of inguinal nodes should be similar to that of penile cancer.

Prognosis correlates with the presence or absence of nodal involvement. In the presence of inguinal node metastasis, the 5-year survival rate is approximately 25%; there are virtually no survivors if iliac nodes are involved.

REFERENCES

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American Joint Committee on Cancer: TNM Classification—Genitourinary Sites. 1996.

Tumors of the Penis
American Joint Committee on Cancer: TNM Classification—Genitourinary Sites. 1996.

Recommended Readings
Selected patients with lower urinary tract cancers or severe functional or anatomic abnormalities of the bladder may require urinary diversion. Although this can be accomplished by establishing direct contact between the urinary tract and the skin surface, it is most often performed by incorporating various intestinal segments into the urinary tract. Virtually every segment of the gastrointestinal tract has been used to create urinary reservoirs or conduits. No single technique is ideal for all patients and clinical situations. A decision is based on a patient’s underlying disease and its method of treatment as well as on renal function, individual anatomy, and personal preference. An ideal method of urinary diversion would most closely approximate the normal bladder: it would be nonrefluxing, low pressure, continent, and nonabsorptive.

Individual methods of urinary diversion can be categorized in various ways, such as (1) by the segment of intestine used, and (2) by whether the method provides complete continence or simply acts as a conduit carrying urine from the renal pelvis or ureter to the skin, where the urine is collected in an appliance attached to the skin surface. Continent forms of urinary diversion can be categorized further according to whether they are attached to the urethra (ie, as a bladder substitute) or are placed in the abdomen and rely on another mechanism for continence (continent urinary reservoir).

Preoperative Counseling & Preparation

All candidates for urinary diversion or bladder substitution should undergo careful preoperative counseling and preparation, including a detailed discussion of the objectives and potential complications of each method. Any potential impact of a procedure on sexual function, body image, and lifestyle should be discussed. Overall satisfaction of most patients undergoing urinary diversion appears to be high (Dutta et al, 2002; Hara et al, 2002; Allareddy et al, 2006; Fujisawa et al, 2000a). However, because they allow freedom from an external appliance, continent forms of urinary diversion, especially bladder substitution, may be of great psychological and functional benefit to well-selected patients (Okada et al, 1997; Bjerre, 1995).

A careful history taken from the patient should note any previous abdominal or pelvic surgery, irradiation, or systemic disease. A history of intestinal resection or irradiation, renal failure, diverticulitis, regional enteritis, or ulcerative colitis would be especially important to consider when selecting a method of urinary diversion or bladder substitution. A complete blood count and measurement of serum electrolytes, urea nitrogen, and creatinine should be performed. The upper urinary tract should be imaged with intravenous urography, ultrasound, or computed tomography to determine whether hydronephrosis, renal parenchymal scarring, or calculous disease exists. Contrast imaging of the small or large bowel or colonoscopy should be considered preoperatively for patients with a history of significant intestinal irradiation, occult bleeding, or other gastrointestinal diseases. Patients with benign bladder diseases—such as a reduced bladder capacity due to neurologic disorders or irradiation, bladder fistulas, or interstitial cystitis—are occasionally considered candidates for urinary diversion or bladder substitution to manage urinary incontinence; however, with such patients, careful evaluation of bladder function and anatomy is required, as adequate urinary function can often be restored by urinary tract reconstruction, pharmacologic manipulation, or intermittent catheterization.

Patients undergo a standard mechanical and oral antibiotic bowel cleansing program beginning 1 or 2 days before surgery. Much of patients’ dissatisfaction with urinary diversion can be attributed to poor stomal construction (Fitzgerald et al, 1997). The patient should be preoperatively evaluated in the lying, sitting, and standing positions. The stoma should preferably be located above or below the belt line. The most common site for stoma placement is along a line between the anterior superior iliac spine and umbilicus and through the rectus abdominis muscle. The surface area should be flat and able to support an appliance. In patients with a short mesentery location of the stoma closer to the level of the umbilicus may be required.
INTESTINAL CONDUIT

URINARY DIVERSION

Ileal Conduit

Ureteroileal urinary diversion is the most common method of urinary diversion in the United States. The conduit is constructed using a segment of ileum 18–20 cm long and located approximately 15–20 cm proximal to the ileocecal valve (Figure 24–1). Longer conduits may be required in obese patients but a short segment minimizes the absorptive surface of the bowel in contact with urine. Once the appropriate length of bowel is selected and isolated, the mesentery is divided proximally and distally and individual mesenteric blood vessels are ligated. The bowel is divided, thus isolating the segment selected for conduit construction. The continuity of the small intestine is reestablished, allowing for normal bowel function. The conduit is usually positioned in the right lower quadrant of the abdomen in an isoperistaltic direction. The posterior closed end of the conduit can be fixed to the posterior peritoneum or the sacral promontory to prevent volvulus of the conduit. The ureters are reimplanted individually in an end to side fashion or joined together (Wallace technique) and anastomosed in an end-to-end fashion, creating a refluxing ureteroileal anastomosis. Ureteral stents (7–8F, single-J, silastic) are usually placed through the ureteral anastomosis and conduit into the renal pelvis to facilitate urinary drainage while the anastomosis is healing. The preselected stoma site is identified and a small circle of skin and underlying fat excised. The fascia is incised in a cruciate fashion. The end of the conduit is brought through the rectus abdominis muscle and anchored to the fascia, and the stoma is then formed. The stoma should protrude, without tension, approximately 1–1.5 in above the skin surface.

Jejunal Conduit

Jejunal conduit urinary diversion is used rarely and mainly in cases where other segments cannot be used due to significant ileal and colonic disease caused by previous irradiation or inflammatory bowel disease. Electrolyte disturbances are more common when the jejunum is used for conduit construction and hence this bowel segment is used only if none of the other intestinal segments can be used to fashion a conduit.

Colon Conduit

There are several advantages to using the large bowel in construction of urinary conduits: nonrefluxing ureterointestinal anastomoses are easily performed, possibly abrogating the deleterious effects of reflux on the upper urinary tracts (Richie and Skinner, 1975); stoma stenosis is uncommon because of the wide diameter of the large bowel; limited absorption of electrolytes occurs; and the blood supply to the transverse and sigmoid colon is abundant. Either the transverse or the sigmoid colon can be used, allowing for placement of the conduit high or low in the abdomen, depending on the integrity and condition of the ureters. Use of the transverse colon for conduit construction is especially well suited for patients who have received extensive pelvic irradiation or when the middle and distal ureters are absent.

The blood supply of the transverse colon is based on the middle colic artery. The greater omentum is separated from the superior surface of the transverse colon, and a segment of bowel, usually 15 cm in length, is selected for the conduit (Figure 24–2). Short mesenteric incisions are made, and the colon is divided proximally and distally. Once the conduit is isolated, bowel continuity is reestablished. The proximal end of the conduit is closed and fixed in the midline posteriorly. The ureters are brought through small incisions in the posterior peritoneum and reimplanted into the base of the conduit. The stoma may be positioned on either the patient’s right or left side.

A sigmoid conduit is constructed in a similar manner. Great care should be taken to preserve the blood supply by carefully selecting a segment with a good blood supply and by making short mesenteric incisions. The conduit is positioned lateral to the reapproximated sigmoid colon. Ureteral reimplantation and stoma construction are completed. The ureters can be reimplanted into the large intestine either in a nonrefluxing submucosal tunnel or in a refluxing end to side manner. A submucosal tunnel can be created by incising the tenia up to the mucosa of the large bowel for a distance of 3–4 cm. A “button” of mucosa is removed, and the ureter is anastomosed to the mucosa of the bowel. The muscularis of the tenia is reapproximated over the ureter to complete the tunnel (Figure 24–3).

CONTINENT URINARY DIVERSION & BLADDER SUBSTITUTION

General Considerations

Various procedures have been developed for construction of bladder substitutes or continent urinary reservoirs that preclude the need for an external urine collection appliance. They offer psychological and functional advantages for selected patients who require urinary diversion. Such reservoirs or bladder substitutes are composed of 3 segments: ureterointestinal (afferent limb) anastomosis, the reservoir itself, and the conduit carrying the urine from the reservoir to the surface (effenter continence mechanism). Bladder substitutes rely on the intact urethra and sphincter to provide outlet resistance and carry urine to the urethral meatus. In men and women whose urethrae are involved by cancer or are not functional owing to benign diseases, an effenter continence mechanism can be constructed with...
the appendix or short segments of tapered, intussuscepted, or reimplanted intestine.

The decision to proceed with bladder replacement is dependent on the risk of urethral recurrence and the continence of the patient. Both men and women who have low risk of urethral recurrence and those who have intact external urinary sphincters should be considered for bladder replacement rather than construction of a continent urinary reservoir. The risk of urethral occurrence or recurrence in men who undergo radical cystectomy is 6.1–
Figure 24–2. Transverse colon conduit.
10.6%. Carcinoma in situ and tumor multifocality appear to be risk factors for prostatic urethral tumor involvement in men with bladder cancer (Nixon et al, 2002). Although prostatic urethral disease is a risk factor for urethral recurrence, recent evidence suggests that orthotopic diversion may be considered in those with only proximal prostatic urethral involvement and negative urethral margins at cystectomy (Iselin et al, 1997). Although orthotopic bladder replacement was once reserved for men, recent clinical and laboratory experience suggests that it may be an acceptable procedure in women as well (Stein et al, 1997). Women with bladder cancer whose tumors are not located at the bladder neck and have a clear urethral margin at the time of cystectomy are candidates for this procedure. Approximately 66% of women undergoing radical cystectomy for the management of bladder cancer fall into this group (Stein et al, 1995, 1998a; Stenzl et al, 1995). Intraoperative inspection and frozen section assessment of the bladder neck limits the risk of urethral recurrence.

Although various segments of the gastrointestinal tract can be used, the principles for success are standard. The bowel segments should be opened and refashioned (detubularized) to interrupt the normal high-pressure contractions of the intact intestine (Hinman, 1988). A large radius is preferred, as results in a reservoir with a larger geometric capacity and lower pressure. Continental reservoirs and bladder substitutes may be made of small intestine, large intestine, or a combination of both. Even though bladder substitution is perceived by some as a more complex procedure, prior studies have demonstrated complication and reoperation rates similar to those of ileal conduit urinary diversion in the hands of experienced surgeons (Gburek, Lieber, and Blute, 1998; Parekh et al, 2000). Long-term results of bladder replacement demonstrate excellent functional outcomes (Elmajian et al, 1996; Hautmann et al, 1999; Abol-Enein and Ghoneim, 2001; Steven and Poulsen, 2000; Stein et al, 1997; Stenzl et al, 2001). Daytime continence can be expected in 87–100% of men and 82–100% of women. Nighttime continence is achieved in 86–94% of men and 72–82% of women. Nearly all men are able to void to completion, while approximately 15–20% of women require intermittent catheterization to completely empty the neobladder. It is beyond the scope of this chapter to review all techniques and minor modifications; instead, the most common techniques and the general principles of continent diversion are reviewed.

**Ureterosigmoidostomy**

The first direct anastomosis of the ureters into the intact colon was performed by Smith in 1878 (Smith, 1879). Peritonitis (from fecal spillage) and pyelonephritis (result-
ing from ascending infection and stricturing of the ureteral anastomosis) led initially to very high surgical mortality rates. Recognizing that ascending infection from the rectum into the kidney was a major problem, surgeons developed techniques to reimplant the ureters into the colon in an antirefluxing fashion. Since patients will retain large amounts of urine and fecal material simultaneously in the rectum, assurance of adequate rectal sphincter function is important preoperatively. Since ammonia may be absorbed across the bowel surface, patients with liver disease who may be at an increased risk of hyperammonemic encephalopathy should not undergo this procedure. Patients who have primary diseases of the colon or have received extensive pelvic irradiation should undergo alternative forms of diversion.

The ureters are identified at or below the common iliac arteries. The overlying peritoneum is incised and the ureters are mobilized carefully to preserve their blood supply. A site low in the sigmoid is selected for ureteral reimplantation. The ureters are reimplanted separately into the respective ipsilateral tenia coli, using the antirefluxing techniques described earlier. The peritoneum is sutured over the completed ureteral anastomosis (Figure 24–4).

One particularly worrisome complication of ureterosigmoidostomy is development of adenocarcinoma at the site where the ureters have been reimplanted into the large intestine. The incidence of anastomosis site adenocarcinoma is uncertain, but there seems to be a several thousand-fold increase in its incidence in patients who have undergone ureterosigmoidostomy over individuals who have not undergone this type of surgery. The induction period varies but may be approximately 20 years; thus, the risk is especially high in young patients. Experimental studies have shown that the development of adenocarcinoma seems dependent on the presence of urine, feces, urothelium, and colonic epithelium in close approximation. All those who undergo ureterosigmoidostomy should be counseled to undergo yearly sigmoidoscopy starting 5 years after the procedure and at any time occult or gross gastrointestinal bleeding or a major change in bowel habits is noted.

Reservoirs Constructed of Small Intestine

Nils Kock was responsible for the early and ongoing development of a continent urinary reservoir fashioned entirely of small intestine (Kock et al, 1982; Nieh, 1997). Sixty to seventy centimeters of small intestine is selected for reservoir construction. The proximal and distal 15-cm segments are preserved for construction of nipple valves, allowing for antireflux ureteroileal anastomoses (inlet) and a continent, catheterizable abdominal stoma (outlet) (Figure 24–5). The middle 40 cm is opened along the antimesenteric border and folded into a U; the back wall is sutured together. The proximal and distal nipple valves are constructed by intussuscepting the bowel, stapling it in place, and further securing it with a mesh anchoring collar. This fixes the intussuscepted bowel segments in place and allows for high-pressure zones, which prevent ureteral reflux proximally and incontinence distally. The reservoir is closed by folding the middle segment and suturing it in place. The ureters are sutured to the proximal nipple valve. The distal nipple valve is brought to the skin as a “flush,” catheterizable stoma. Revision of the nipple valve may be necessary in some patients to correct valve slippage or eversion. In men who have an intact urethra and external urinary sphincter, the distal nipple valve can be omitted and the reservoir attached directly to the urethra (Elmajian et al, 1996). The Koch reservoir is no longer commonly used due to a high incidence of complications.

In an attempt to decrease the complication rate associated with the afferent intussuscepted antireflux nipple of the Kock ileal neobladder, Stein and colleagues (1998b) have described an innovative antireflux technique (Figure 24–6). This new reservoir, the T pouch, has several important advantages over the Kock ileal neobladder: (1) It requires a smaller segment of ileum to create the antireflux technique, (2) the serosal-lined antireflux mechanism eliminates the need for intussusception, (3) blood supply is better preserved, and (4) urine is not in contact with the implanted portion of the ileum.

Camey described a technique of bladder substitution whereby an intact segment of ileum was anastomosed directly to the urethra (Lilien and Camey, 1984). A 40-cm segment of ileum was isolated from the gastrointestinal tract, and its midpoint was anastomosed, without tension, directly to the urethra. The ureters were reimplanted into

![Figure 24–4. Ureterosigmoidostomy.](image-url)
either end of the ileal segment in an antirefluxing fashion. However, failure to detubularize the ileal segment led to a high incidence of urinary incontinence compared with substitutes fashioned from detubularized small-bowel segments (Studer and Zingg, 1997; Hautmann et al, 1999). Forty- to sixty-centimeter segments of ileum can be detubularized and folded into U-, S-, or W-shaped reservoirs, which can be connected directly to the urethra (Figure 24–7). The ureters are reimplanted directly into a short segment of ileum at the proximal end of the bowel segment, isolated to fashion the diversion, which is not detubularized. End to side ureteric anastomosis into this isoperistaltic segment of ileum prevents reflux. With the Hautmann type of ileal neobladder (W configuration), non-detubularized segments of ileum can be left intact at either end of the W and the ureters individually implanted into each of these segments. This would make it easier to dissect the ureter if a nephroureterectomy is required at a later date for upper tract tumor recurrence. The larger diameter and lower pressure of these reservoirs compared with those of non-detubularized bowel have led to better continence rates.

**Reservoirs Constructed of Large Intestine**

Various investigators have described using segments of the large intestine alone or combinations of large- and small-bowel segments to fashion urinary reservoirs (Lampel et al, 1996; Bihrl, 1997). Bladder substitutes have been constructed using detubularized bowel segments from the ileocecal region or sigmoid colon. Use of

**Figure 24–5.** Kock Pouch urinary reservoir. **A:** 60 cm of small intestine selected. **B:** Afferent (nonrefluxing) limb for ureteral reimplantation and efferent limb for stoma fashioned using stapling devices. **C:** Completed reservoir.
the ascending colon and terminal ileum to construct continent urinary reservoirs has gained great popularity. Cecum and ascending colon are detubularized and refashioned or augmented with small intestine to provide for a spherical reservoir. The ureters are reimplanted in a nonrefluxing fashion (Figure 24–8). The reservoir can be anastomosed directly to the urethra. In women or in men who require a urethrectomy, a continent, catheterizable stoma can be fashioned of appendix or tapered terminal ileum (Figure 24–9).

**Postoperative Care**

Postoperative care varies depending on the method of urinary diversion or bladder substitution. As with all patients who have undergone major abdominal surgery, early ambulation, use of intermittent compression stockings, and incentive spirometry may be used to prevent pulmonary emboli or respiratory complications. If a nasogastric tube is used, it is left in place until there are signs of intestinal peristalsis. Serum electrolytes and creatinine should be monitored postoperatively for the development of metabolic abnormalities. If ureteral stents have been placed, they are usually removed sometime after postoperative day 5; they may stay in place for up to 3 weeks in patients undergoing bladder replacement. Continent urinary reservoirs and bladder substitutes produce mucus. They should be irrigated regularly in the early postoperative period to prevent mucus accumulation. Mucus production decreases over time, and irrigation ultimately becomes unnecessary. Upper urinary tract surveillance for hydronephrosis should be performed on a regular basis using ultrasound, intrave-
nous urography, or computed tomography. An initial assessment of the upper urinary tract is made in the early postoperative period; if upper urinary tract dilation is not found, the patient undergoes repeat imaging on a yearly basis.

**COMPLICATIONS**

Complications occurring after urinary diversion, bladder substitution, or continent urinary diversion are generally a product of surgical technique, the underlying disease pro-
cess and its treatment, the age of the patient, and the length of follow-up (Carlin, Rutchik, and Resnick, 1997; Gburek et al, 1998). Early complications, which are uncommon (occurring in approximately 10–20% of patients), include excessive bleeding, intestinal obstruction, urinary extravasation, and infection. Late complications (10–20% of patients) include metabolic disorders, stomal problems (stenosis or hernia), pyelonephritis, and calculi. Rarely, spontaneous neobladder rupture may occur presumably due to overdistention or blunt abdominal trauma (Nippgen et al, 2001). Such patients usually present with peritonitis and acute abdominal pain. Perforation of continent cutaneous diversions can occur and can be related to vigorous catheterization.

**Metabolic & Nutritional Disorders**

Fluid, electrolyte, nutrient, and waste product excretion or absorption normally occurs across the intestinal wall. The extent of absorption or excretion is dependent on the concentration of these substances in the lumen or blood and on which segment of bowel is in contact with them. Metabolic abnormalities may occur when intestinal segments are interposed into the urinary tract. As pointed out previously, use of the jejunum may result in hyponatremic, hypochloremic, hyperkalemic metabolic acidosis in up to 40% of patients. The jejunum is unable to maintain large solute gradients, so large amounts of water and solute pass through the jejunal wall. Sodium and chloride are rapidly excreted into the conduit, and potassium is passively absorbed. Aldosterone is produced, resulting in reabsorption of hydrogen and excretion of potassium into the distal tubule of the kidney and consequent acidosis and movement of potassium from the body's intracellular stores. As water is lost into the conduit, extracellular fluid volume is reduced, as is the glomerular filtration rate. The renin-angiotensin system is activated which further stimulates aldosterone secretion. Urea may be absorbed from the jejunal lumen which (with dehydration) contributes to azotemia. This syndrome is often characterized clinically by nausea, vomiting, anorexia, and muscle weakness.

The pathogenesis and nature of metabolic abnormalities occurring after incorporation of the ileum or colon into the urinary tract differ from those associated with jejunal conduits. When such segments are used, sodium and chloride are absorbed across the bowel surface. Chloride is absorbed in slight excess of sodium, resulting in a net loss of bicarbonate into the bowel lumen. Preexisting renal failure contributes to the development and severity of the disorder, as does a large bowel surface area and long contact time. Hyperchloremic acidosis is more common in patients who undergo ureterosigmoidostomy than in patients who undergo simple conduit construction using either the ileum or the colon, because of the larger surface area and longer contact time with urine associated with ureterosigmoidostomies. Hyperchloremic metabolic acido-
Figure 24–9. Continent urinary reservoir constructed from the ileocecal segment (A). B: Bowel detubularized. Stoma constructed by tapering the terminal ileum with intestinal stapling instrument (shown in C) or sutures. D: Completed stoma.
sore may manifest clinically as weakness, anorexia, vomiting, Kussmaul breathing, and coma. One potential long-term complication of chronic acidosis may be decreased bone calcium content and osteomalacia (McDougal et al, 1988; Kawakita et al, 1996).

Bile salts are important for fat digestion and uptake of vitamins A and D. Bile salt metabolism may be altered after ileal resection (Olofsson et al, 1998). Resection of small segments of the ileum may be associated with mild malabsorption and steatorrhea owing to increased concentrations of bile salts delivered to the colon. The increased concentration of such salts leads to decreased colonic absorption of water and electrolytes. Resection of large segments of ileum may reduce bile salt reabsorption to very low levels, leading to severe fat malabsorption. Bowel transit time may be reduced further by resection of the ileocecal valve. Cholestyramine may be used to treat secretory bile salt diarrhea. If diarrhea persists, addition of agents such as loperamide that decrease bowel motility and increase transit time may be effective. Cholelithiasis may be more common after ileal resections as well.

Vitamin B<sub>12</sub> deficiency may occur as a result of either gastric or ileal resection (Terai et al, 1997; Fujisawa et al, 2000b). Because B<sub>12</sub> stores are likely to last for years, its deficiency may not become apparent for several years after intestinal surgery. B<sub>12</sub> deficiency results in megaloblastic anemia and peripheral nerve paresthesias. The levels of this vitamin can be assessed in patients after ileal or gastric resection. Alternatively, yearly replacement by injection of Vitamin B<sub>12</sub> is adequate in those in whom deficiency is noted.

**Stoma**

A good deal of patients’ dissatisfaction with urinary diversion can be attributed to stomal complications. Stoma construction is especially important in procedures in which a pouch must be worn. Careful stomal construction as outlined earlier is essential (Fitzgerald et al, 1997). Complications related to the stoma can be separated into (1) stomal complications: necrosis, stenosis, hernia, retraction, and prolapse and (2) peristomal complications: dermatitis related to fungal infections, contact allergic reactions, mechanical trauma, and contact dermatitis from urine on the skin over time. They are often related as a stenotic stoma may be difficult to pouch leading to chronic skin irritation. Early stomal complications may occur in as much as 45.6% in conduits or late complications as much as 18.9% (Bass, 1997). The likelihood of stomal stenosis increases with time. Stomal stenosis can lead to conduit elongation and upper-tract obstruction. This condition can be diagnosed relatively easily by catheterizing the stoma and measuring the residual urine volume. It is corrected by revision of the stoma. Parastomal herniation tends to occur as a delayed complication in 2–6.5% of patients (Franks and Hrebinko, 2001). One cause for this is the location of the stoma lateral to the rectus muscle which eliminates the circumferential support provided by the muscle cuff around the stoma. The hernia typically tends to occur around the superolateral aspect of the stoma. Placing the stoma such that the conduit is traversing the rectus muscle may not always decrease the likelihood of parastomal hernia (Gray, 2005). Some degree of stomal bulge without a hernia is commonly noted in most patients with conduits. Repair of a stomal hernia requires reinforcing the rectus fascia with mesh, either intraperitoneally through the previous midline incision or through an incision lateral to the stoma. In some cases it may be necessary to move the stoma to a new site, preferably to the contralateral side of the abdominal wall. Recurrence rates following hernia repair were high previously but have declined considerably to 0–7% in recent series (Franks and Hrebinko, 2001; Ho and Fawcett, 2004).

Skin irritation or infections are most common in procedures in which an appliance is worn and there is prolonged contact of the skin with urine. Some patients’ skin may be sensitive to adhesive agents.

Stomal or skin problems are minimized by selecting an appropriate pouching system. Extended wear pouches provide better skin protection because of better adhesion and delayed moisture absorption (Colwell, 2005). The treatment of parastomal conditions is summarized in Table 24–1.

The most troublesome stomal complication of external urinary appliances is unexpected pouch leakage. This is often a result of improper application of the pouch but may be caused by poor stomal construction. With a properly constructed and located stoma, the choice of the pouch is relatively simple and is often determined by patient preference. Factors that must be taken into account when selecting a pouch include patient dexterity and preference, available resources, and stomal construction. Pouches have an adhesive skin barrier that attaches to the skin and a bag that collects urine. An opening is made in the middle of the skin barrier, this opening should be of a diameter just larger than the stoma to avoid erosion of the stoma or prolonged contact of the skin with urine, typically one-eighth of an inch on each side unless the stoma is retracted or at skin level then the opening may be more secure if there is one-fourth-inch clearance on each side to avoid undermining (Doughty, 2005).

Most patients favor using disposable urinary collection pouches. Disposable, one-piece pouches are single use. Two-piece systems allow the patient to remove the pouch, leaving the skin barrier intact. A larger collection device is often attached to the pouch at night.

**Continence & Urinary function**

One of the common late complications of continent urinary diversion is urinary incontinence that can
occur following both cutaneous diversion and in neo-
bladders. Nighttime incontinence is particularly a
problem with neobladders and can occur in about 20–
50% of all patients (Meyer et al, 2004; Lee et al,
2003). Daytime incontinence is significantly lower
and ranges from <10% to 33%. Reported continence
rates are higher with the T pouch at around 87%
(daytime) and 72% (nighttime) (Stein, 2004). The wide
variations in reported continence rates stem from varying
definitions of continence. Improvement in continence is
observed with time and increasing neobladder capac-
ity reaching 92% for daytime and 80% at nighttime
by 2 years (Studer, 1996). Bladder capacity increases
gradually and stabilizes to reach >450 cc and contin-
ence also improves over time in these patients (Per-
menis, 2004). Incomplete emptying can occur in a
small percentage of patients requiring long-term self-
catheterization. Women who undergo neobladder con-
struction are especially at risk for hypercontinence,
which can occur in 12% of patients and must be
counseled preoperatively in this regard (Stenzl et al,
2001). Day- and nighttime continence rates appear to
be similar between male and female patients and the
most common types of neobladders reconstruction
(Studer versus Hautmann) (Stenzl et al, 2001; Lee et
al, 2003). Sphincteric damage and low urethral resting
pressures are believed to be the reasons for inconti-
ence while urethral denervation, poor pouch contrac-
tility, and/or exaggerated acute angulation of the neo-
bladder urethral junction are believed to be the
reasons for hypercontinence and retention. Frequent
bladder emptying, urethral collagen injection, artifi-
cial urinary sphincter placements, and urethral slings
are all potential treatment options for incontinence
while catheterization is the main therapy for urinary
retention.

**Pyelonephritis & Renal Deterioration**

Pyelonephritis occurs in approximately 10% of patients
who have undergone urinary diversion. Treatment is
based on a properly collected urine sample for culture.
A urine sample should not be collected from the pouch;
rather, the pouch should be removed, the stoma

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<table>
<thead>
<tr>
<th>Problem/Description</th>
<th>Cause</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fungal infections/yeast dermatitis</td>
<td>• Yeast overgrowth on skin exposed to excess moisture (barrier opening too large, pouch leakage)</td>
<td>• Antifungal powder (Nystatin) or systemic therapy with fluconazole (Diflucan).</td>
</tr>
<tr>
<td>Allergic contact dermatitis</td>
<td>• Allergic reaction in sensitive patient.</td>
<td>• Identify irritant (skin patch test) and discontinue product.</td>
</tr>
<tr>
<td>Mechanical trauma</td>
<td>• Frequent or excessive pouch changing leading to skin stripping</td>
<td>• Apply hydrocortisone cream for short period to reduce skin reaction.</td>
</tr>
<tr>
<td>Pseudoverrucous skin lesions</td>
<td>• Urine contact with peristomal skin over extended period</td>
<td>• Topical antihistamines (diphenhydramine gel).</td>
</tr>
<tr>
<td>Peristomal hernia</td>
<td>• Not clear at this time—may be related to abdominal aperture size created at time of surgery</td>
<td>• Ensure correct opening in pouch.</td>
</tr>
</tbody>
</table>

The patient should see a Wound, Ostomy, Continence Nurse (WOCN), if available.
cleansed with an antiseptic, and a catheter advanced gently through the stoma. If infection has occurred in a patient with a simple conduit, the volume of residual urine within the conduit should be recorded. Obstruction and stasis of urine within the reconstructed urinary tract are risk factors for the development of infection.

Although many patients with preexisting dilation of the upper urinary tract show improvement or resolution of the dilation after urinary diversion or bladder substitution, progressive renal deterioration as manifested by hydronephrosis or an increasing serum creatinine level (or both) occurs in some patients who undergo these procedures. The incidence of either complication increases after 10 years. The presence of hydronephrosis, particularly in patients with a conduit diversion, may indicate the presence of ureteric reflux or obstruction at the uretero-vesical junction. A loopgram will help demonstrate if there is retrograde flow of contrast from conduit to the ureter and renal pelvis. In some situations it may be necessary to obtain a radionuclide renal scan (MAG3) to determine if there is antegrade flow of urine from the kidney through the ureters. Pyelographic evidence of upper urinary tract deterioration has been noted in up to 50% of patients who have undergone urinary diversion at an early age. Recurrent upper urinary tract infection and high-pressure ureteral reflux and obstruction, usually in combination, contribute to the likelihood of renal deterioration. Upper-tract deterioration is less likely following ileal continent diversion. While a majority of kidneys will demonstrate mild dilation, 97% of patients show no change in renal parenchymal size and 100% show no change in serum creatinine 10–15 years following the Studer type of ileal neobladder (Perimenis, 2004).

Calculi

Calculi occur in approximately 8% of patients who undergo urinary diversion or bladder substitution (Cohen and Streem, 1994; Terai et al, 1996). Such patients have several risk factors for the development of various calculi. Nonabsorbable staples, mesh, or suture material used to construct conduits or reservoirs may act as a nidus for stone formation. Colonization in either conduits or reservoirs is common, whereas symptomatic infection is much rarer. Certain bacteria can contribute to stone formation; some bacteria commonly found in the urinary tract, including Proteus, Klebsiella, and Pseudomonas species, produce urease, a urea-splitting enzyme that contributes to the formation of ammonia and carbon dioxide. Hydrolysis of these products results in an alkaline urine supersaturated with magnesium ammonium phosphate, calcium phosphate, and carbonate apatite crystals. Management of such infection-related stones requires stone removal, resolution of infection, and, often, use of adjunctive agents to complete stone dissolution.

The likelihood of stone formation is increased by the development of systemic acidosis, as described previously (Terai et al, 1995). Prolonged contact of the urine with the intestinal surface facilitates the exchange of chloride for bicarbonate. Bicarbonate loss results in systemic acidosis and hypercalciuria. The combination of hypercalciuria and alkaline urine predisposes a patient to the development of calcium calculi. In addition, the terminal ileum is responsible for bile salt absorption; if this portion of the intestine is used for conduit or bladder reservoir construction, excess bile salts in the intestine may bind calcium and result in increased absorption of oxalate, which may lead to the development of oxalate-containing calculi. Hypocitraturia may also be a risk factor for stone disease in patients undergoing bladder replacement (Osther, Poulsen, and Steven, 2000). Excess conduit length, urine stasis, and dehydration make the development of calculi more likely.

REFERENCES

General


Colon Conduit

Continent Urinary Diversion & Bladder Substitution


Ureterosigmoidostomy

Smith T: An account of an unsuccessful attempt to treat extroversion of the bladder by a new operation. St Barth Hosp Rep 1879:15:29.

Complications


**Ostomy Care**


The primary management of genitourologic malignant diseases has been tied to the use of radiation for more than 100 years. In 1895 Roentgen described x-rays; by 1899 a patient with skin cancer was cured with radiation; and within 10 years, radiation was used to treat prostate cancer (Pasteau and Degrais, 1914). Radiotherapy became a mainstay for treatment of bladder and testicular cancers and later prostate cancer as supervoltage sources became available. Although chemotherapy and aggressive surgery have supplanted some of the uses of radiotherapy, radiation continues to play a major role in the management of carcinomas of the penis, urethra, prostate, and bladder. In this chapter we review general principles and the indications for using radiation as a component in the primary management of urologic malignant diseases. The role of radiation as an agent of palliation has been well documented elsewhere and is excluded from this chapter.

GENERAL PRINCIPLES OF RADIOTHERAPY

Mechanisms of Cytotoxicity

The effects of radiation on tumor and surrounding normal tissues are thought to be mediated primarily through the induction of unrepaired double-strand breaks in DNA. Excited electron species generated in the presence of oxygen form peroxide radicals, which fix chemical lesions and result in the generation of either repairable or nonrepairable DNA double-strand breaks. High linear energy transfer radiation (including neutrons and heavy-charged particles) is associated with less repairable DNA damage. Classically, the expression of radiation damage is not seen until the target cells enter mitosis. Differentiated normal tissues with low mitotic activity, such as the heart and spinal cord, tend to express the effects of radiation much later than cells from more kinetically active tissues, such as the epithelial cells lining the rectum, bladder, or urethra. However, differentiated normal tissues with low mitotic activity are more sensitive to the use of high dose per fraction or high linear energy transfer radiotherapy. In organs in which the functional stromal cells are postmitotic, such as muscle cells and neurons, the damage is expressed by slowly dividing support cells such as endothelial cells.

In addition to the classic mechanism described above, radiation has been shown to induce programmed cell death (apoptosis). Androgen-independent human prostate cancer cells activate a genetic program of apoptotic cell death in response to exposure to ionizing radiation, in a dose-dependent fashion (Sklar, 1993). Results from animal models suggest that it is better to achieve maximal androgen suppression before starting radiation treatment (Zietman, 2000).

Radiation Sensitivity & Tolerance

Radiation tolerance levels have been determined for nearby normal tissues that are likely to be affected during conventional fractionated radiotherapy treatment of tumors arising from the urinary tract. The term conventional fractionation generally refers to the delivery of a single daily dose of 180 cGy (1.8 Gy) to 200 cGy (2.0 Gy). When used alone, cumulative doses of at least 65 Gy are necessary for local control of gross disease (adenocarcinomas, transitional cell carcinomas [TCCs], and squamous carcinomas) arising from the prostate, bladder, urethra, or ureters. When used prophylactically for presumed microscopic disease (lymph nodes) or postoperatively, doses of 45–50 Gy are generally sufficient. For testicular seminomas, doses of 25 Gy are usually adequate.

Total dose, dose per fraction, and volume of normal tissue irradiated are the major risk factors for radiation-induced complications. The presence of several comorbid conditions such as previous surgery, diabetes, inflammatory bowel diseases, or old age is also associated with an increased risk of radiation-induced complications. Accurate estimates of the “true tolerance” of surrounding normal tissues have been hampered until recently by our inability to reconstruct the actual relationship between normal tissue doses and volumes in 3 dimensions. Early reports assumed that organ movement and day-to-day treatment setup error had an insignificant impact on the doses of radiation delivered to surrounding normal tissues. Recent studies have demonstrated that these assumptions are inaccurate and probably result in an underestimate of the true tolerance of surrounding normal tissues to radiation (Langen and Jones, 2001).
Dose per Fraction Considerations

Dose per fraction, the total dose, volume and overall treatment time are all critical determinants of chronic genitourinary toxicity. The linear quadratic equation (L-Q equation) has been adopted by many clinical investigators as the most useful model for comparing various doses and fractionation schemes (Fowler et al, 2001). This equation can be written as follows:

\[ \text{Effect} = E = n (\alpha d + \beta d), \]
where:
- \(d\) = Dose per fraction
- \(\alpha\) = Nonrepairable effects
- \(\beta\) = Repairable effects
- \(n\) = Number of identical fractions

For comparing two different fractionation schemes, assuming a similar overall treatment time, the L-Q equation also can be written as follows:

\[ \frac{D_2}{D_1} = \frac{(1 + d_1 \beta / \alpha)}{(1 + d_2 \beta / \alpha)} \]
where:
- \(D\) = Total dose and \(D_1 = n_1 d_1\)
- \(d\) = Dose/fraction

For most clinical circumstances, it is assumed that the \(\alpha/\beta\) ratio for late-reacting normal tissues such as the bladder or rectum is 3. For early-responding normal tissues and for tumor, it is assumed that the \(\alpha/\beta\) ratio is 10.

Altered Fractionation Schedules

Radiobiologic modeling using the \(\alpha/\beta\) ratios as described previously has been used to develop “altered fractionation schedules” to improve the therapeutic ratio between efficacy and toxicity. Accelerated hyperfractionation is the most frequently used altered fractionation schedule. With accelerated hyperfractionation, more than one treatment is given per day with a minimum of 6 hours between treatments, using a decreased dose per fraction. Since most radiation-induced damage is repaired within 6 hours, the use of multiple treatments per day in theory should allow greater doses of radiation to be given over a shorter period of time, reducing the opportunity for tumor repopulation. Using \(\alpha/\beta\) modeling, the late effects predicted using 1.2 Gy fractions twice daily (separated by 6 hours) to 69.6 Gy would be expected to be equivalent to those using conventional fractionation to 58 Gy. In contrast, early-responding tissues such as the epithelium of the bladder would be expected to respond as though they had been treated with a dose of 65 Gy using conventional fractionation. This model predicts an increase in acute effects, including tumor response, but a decrease in late effects, such as fibrosis.

Clinical use of hyperfractionation schemes in the management of urologic tumors has been limited. Preliminary results with hyperfractionation for prostate tumors have been mixed. Acceptable toxicity was reported among the first 20 patients treated with 1.3 Gy twice daily to 78 Gy (Forman et al, 1996). Because of practical considerations (such as time and cost), there appears to be more interest in hypofractionated regimens than in hyperfractionated ones.

Hypofractionation involves the use of larger than conventional fraction sizes. Such an approach results in nearly a 10% reduction in the total dose to the tumor without sparing late complications (assuming that the values chosen for the \(\alpha/\beta\) ratio are correct for normal tissues and for tumors) (Fowler et al, 2001). Several recent investigators have argued that the \(\alpha/\beta\) ratio for prostate cancer is much lower than previously thought and that this formed a strong rationale for the use of high dose rate (HDR) to treat prostate cancer (Brenner and Hall, 1999; King and Fowler, 2001; Fowler et al, 2001). For patients being treated with external-beam radiotherapy (EBRT), a growing number of investigators are studying the use of larger doses per fraction in an attempt to shorten overall time and cost (Pickett et al, 1999). Because of day-to-day setup variation, some type of monitoring will probably be required to ensure that this can be carried out safely. A number of recent studies suggest that the use of hypofractionated regimens hold promise but the only Phase III Trial completed to date raises concerns about the selection of the most appropriate dose to yield equal or better outcomes (Kupelian et al, 2007; Lukka, 2003; Pollack et al, 2006; Tang et al, 2006).

Brachytherapy

The term brachytherapy refers to a treatment technique that places radioactive sources in close proximity to or directly into the tumor. Brachytherapy can be classified as either interstitial or intracavity. Interstitial brachytherapy involves the placement of radioactive needles, afterloaded needles or catheters, or radioactive seeds directly into the prostate, bladder, penis, or perirethral soft tissues. Intracavitary brachytherapy includes placement of radioactive catheters into a lumen or orifice, such as in the urethra, to treat urethral and penile tumors. Permanent implants involve the use of radioactive seeds that are left in the patient. Nowadays, temporary implants generally involve the use of needles or catheters that act as conduits for delivering radiation from high-activity seeds attached to a long wire. Most modern temporary implants are delivered using HDR brachytherapy systems to deliver moderately high doses of radiation over a relatively short period of time (minutes). HDR brachytherapy is usually delivered over 2 or more treatment sessions to reduce the risk of late complications. Low-dose rate brachytherapy is usually delivered in a continuous fashion over days to weeks via temporary or permanent implants, respectively. Figure 25–1 depicts an example of a transrectal ultrasound-based iodine-125 permanent interstitial implant of the prostate. Figure 25–2 shows an example of intracavitary brachytherapy for a urethral tumor in a female patient.
CHAPTER 25

SPECIFIC UROLOGIC SITES

Prostate Cancer

Many urologists are aware that the radical perineal prostatectomy (RPP) approach was first used to completely remove the prostate in the 1800s (Bilroth, 1869). Unfortunately, the first 2 patients so treated died in 14 months or less. Few urologists are aware that radiotherapy was first used to successfully treat prostate cancer 100 years ago and that it was not until 1949 that Memmelaar first reported the successful use of the retropubic approach—the radical retropubic approach (RRP)—for treatment of prostate cancer (Memmelaar, 1949; Reiner and Walsh, 1979). The basic RRP is very much unchanged since his description and is probably still the most common approach used today. In the 1950s and 1960s, it was believed that radiotherapy provided comparable results with less morbidity. However, in 1979 Walsh and Donker demonstrated the potential value of the RRP to preserve nerves (Walsh and Donker, 1982). Radical perineal prostatectomies became less fashionable in the 1980s after Walsh and others reported improvements in the RRP that promised to reduce blood loss, minimize the risk of incontinence, and preserve potency (Reiner and Walsh, 1979). With this advancement, the RRP became the surgical approach of choice in the eyes of most urologists.

A. CONVENTIONAL TREATMENT

Conventional EBRT has been used in the United States for treating prostate cancer for more than 35 years. From the late 1960s through the mid-1980s, definitive prostate radiation was delivered to the whole pelvis by using the “4-field box” technique, followed by a “cone down” prostate boost, popularized by Bagshaw and coworkers from Stanford University (Bagshaw et al, 1988). The placement of the treatment fields was based on bony anatomic landmarks. It is now known that this technique resulted in inadequate coverage of the target volume in nearly one-third of patients. Computed tomography has improved...
our ability to localize and reconstruct pelvic anatomy, allowing more accurate design of treatment portals.

Several retrospective and prospective studies support the efficacy of radiation therapy in the management of localized prostate cancer (Roach et al, 1999). Based on these studies, the local control rate following EBRT has been estimated to be between 70% and 90%. The end points of these studies were based primarily on a clinical assessment of “local control.” These end points are now known to underestimate the true incidence of local failures because of the presence of occult cancer in clinically controlled patients. Many patients whose disease was once thought to be controlled locally, according to physical examination, do in fact have persistent disease when assessed by prostate-specific antigen (PSA) or biopsy, or both (Crook et al, 2000). Although the posttreatment PSA is more sensitive, it is a less specific end point for treatment failure than is biopsy, because distant failures can contribute to a rising level. PSA has been long recognized as an acceptable end point for the assessment of disease status, but only recently has it been validated as a predictor of disease-specific survival (DSS) and overall survival (OS) (Roach et al, 2003b; 2006). Based on a Consensus Conference held in January 2005 it was determined that a rise of more than 2 ng/mL was strongly suggestive of recurrences that were subsequently associated with clinical signs and symptoms. The transition from using the so-called American Society of Therapeutic Radiology and Oncology (ASTRO definition) (Roach, 2006) was stimulated by the recognition that the use of androgen derivation therapy complicated a definition based on three rises because PSAs frequently rise after discontinuing ADT due to the recovery of testosterone (ASTRO, 1997).

Table 25–1 summarizes patient characteristics and treatment outcomes for selected surgical and conventional radiotherapy series. It should be kept in mind, however, that in general patients treated with surgery tend to have lower Gleason scores and T-stages, and many surgical series excluded lymph node-positive patients. When matched for pretreatment PSA, freedom from biochemical relapse at 5 years is not significantly different, suggesting similar efficacy for surgery and EBRT in the management of prostate cancer. There is a trend, however, for surgically treated patients to have a slightly higher freedom from biochemical failure. This could be explained in part by the better parameters for surgical patients at the outset.

Table 25–2 summarizes some of the factors contributing to the risk of local failure following external-beam irradiation. Traditional techniques used until recently were associated with inadequate coverage of the target volume in 20–41% of patients treated. Early series also routinely used total doses in the range of 65–70 Gy because it was believed that this dose was sufficient and close to the maximum dose allowable by the surrounding normal tissues. Another factor contributing to the data regarding biochemical relapse is that many of the patients had disease that was underestimated by physical examination, so they were incurable at the time of diagnosis.

An analysis of the long-term results from approximately 1500 men treated with EBRT alone between 1980 and 1992 in one of four Phase III randomized trials conducted by the Radiation Therapy Oncology Group (RTOG) has been completed (Roach, 2000a). In a multivariate analysis, Gleason score, clinical stage, and pathologic node status were correlated with OS and DSS. By combining patient groups with similar risk for DSS, 4 distinct prognostic subgroups have been defined with varying risk of death due to prostate cancer. Table 25–3 summarizes the estimated 5-, 10-, and 15-year DSS for each subgroup. These outcome data are likely to represent a worst-case scenario, since the vast majority of these patients were treated before PSA screening. In addition, the patients for whom PSA levels were available had levels in the range of 20–30 ng/mL. Patients with lower PSAs treated with 3-dimensional technology to higher doses are likely to do substantially better

### Table 25–1. Biochemical Failure from Selected Surgical or Radiotherapy Series.

<table>
<thead>
<tr>
<th>Pretreatment PSA</th>
<th>Surgical Aver. % (range)</th>
<th>XRT Aver. % (range)</th>
<th>Surgical b-NED 5 y</th>
<th>XRT b-NED 5 y</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–4</td>
<td>26 (15–43)</td>
<td>21 (11–33)</td>
<td>85–95</td>
<td>80–86</td>
<td>Patients treated with surgery usually were not node-positive and tended to have lower grade and lower stage tumors than those treated with XRT.</td>
</tr>
<tr>
<td>4.1–10</td>
<td>41 (35–52)</td>
<td>30 (27–32)</td>
<td>55–93</td>
<td>42–67</td>
<td></td>
</tr>
<tr>
<td>10–20</td>
<td>26 (16–32)</td>
<td>21 (18–22)</td>
<td>56</td>
<td>30–75</td>
<td></td>
</tr>
<tr>
<td>&gt;20</td>
<td>13 (6–20)</td>
<td>35 (18–50)</td>
<td>—</td>
<td>45</td>
<td></td>
</tr>
</tbody>
</table>


b-NED, no evidence of disease; PSA, prostate-specific antigen; XRT, external-beam radiotherapy.

* Most series exclude node-positive patients and PSA failures >0.2–0.6.

† Clinical staging includes only node-positive and T3–T4 patients and PSA failures >1.0–4.0 or rising.
Large multi-institutional series are now available for patients treated with EBRT and permanent seed implants as well (Kuban, 2003; Zelefsky, 2007). Early attempts at developing risk stratification systems to predict OS were validated using contemporary patients (Roach et al, 2003a). Since this approach, numerous other risk-stratification schemes have gained increasing popularity among radiation oncologists managing prostate cancer (D’Amico et al, 2000; Kattan et al, 2001; Kattan et al, 2000; Ross and Scardino, 2001). More recent classification schemes have emphasized more clinically meaningful endpoints than PSA recurrences alone (Roach et al, 2006). The application of such approaches may improve our ability to design and implement Phase III trials. Reasonable treatment guidelines using the most common risk-stratification systems and incorporating exceptions and considerations in selecting treatment options have become well established (Speight, 2005). In future it is anticipated that the staging system for prostate cancer will be modified to reflect these risk-stratification systems (Roach et al, 2007).

Regarding hormonal therapy and radiotherapy, there currently exists no uniform consensus as to who should receive hormonal therapy and how long it should last. However, the best evidence suggests that low-risk patients do not benefit, and when treated with EBRT at doses of 65–70 Gy intermediate-risk patients benefit from short-term hormonal therapy, and high-risk patients benefit from long-term hormonal therapy (Speight, 2005). The standard for “short-term” hormonal therapy consists of combined androgen blockade using a luteinizing hormone-releasing hormone agent and an anti-androgen for a duration of as few as 3 months to as long as 8 months, with little or no obvious advantage to OS for intermediate-risk patients confirmed to date (Roach, 2004). Patients with intermediate- to high-risk disease receiving hormonal therapy should probably also receive whole-pelvic radiotherapy to the L5/S1 level (Roach, 2006). Patients with high-risk disease (T3, GS 7–10 or GS = 8–10 or very high PSAs) should receive neoadjuvant and concurrent hormonal therapy, but they should also receive long-term adjuvant hormonal therapy consisting of a luteinizing hormone-releasing hormone for 2–3 years or more (Bolla et al, 2002; Hanks et al, 2003; Roach et al, 2000, 2003).

**Table 25–2.** Issues in Risk of Local Failure following External-Beam Irradiation.

<table>
<thead>
<tr>
<th>Explanations for the high local failure rate:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Inadequate target definitions.</td>
</tr>
<tr>
<td>a. Field sizes too small or inferior border too high.</td>
</tr>
<tr>
<td>b. Seminal vesicles not covered.</td>
</tr>
<tr>
<td>2. Inadequate doses of radiation.</td>
</tr>
<tr>
<td>a. Doses limited by the belief that 65–70 Gy is sufficient for cure.</td>
</tr>
<tr>
<td>b. Dose limited by technical difficulties and normal tissue tolerances.</td>
</tr>
<tr>
<td>3. Understanding of patients: patients with metastatic disease.</td>
</tr>
</tbody>
</table>

**Solutions for the problems discussed:**

1. Adequate target definitions: field designed conformally using 3-dimensional reconstructions of CT-based tumor volumes and urethromgrams.
2. Adequate dose: higher doses of radiation to improve local control rate.
3. Better pre-staging of patients: use of pretreatment PSA and Gleason score.

CT, computed tomography; PSA, prostate-specific antigen.

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<table>
<thead>
<tr>
<th>Group*</th>
<th>Deaths/No.</th>
<th>5-Year (%)†</th>
<th>10-Year (%)†</th>
<th>15-Year (%)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>63/474</td>
<td>97 (95–99)</td>
<td>85 (81–89)</td>
<td>71 (61–81)</td>
</tr>
<tr>
<td>2</td>
<td>69/335</td>
<td>91 (88–94)</td>
<td>75 (69–81)</td>
<td>59 (49–69)</td>
</tr>
<tr>
<td>3</td>
<td>89/336</td>
<td>82 (78–86)</td>
<td>60 (52–68)</td>
<td>38 (21–55)</td>
</tr>
<tr>
<td>4</td>
<td>138/314</td>
<td>66 (60–72)</td>
<td>34 (26–42)</td>
<td>28 (19–37)</td>
</tr>
</tbody>
</table>


*Group 1 had patients with a Gleason score (GS) = 2–5, any T stage, or T1–2Nx and GS = 6; group 2 had stage T3Nx, GS = 6, or any T stage, N+, GS = 6, or T1–2Nx, GS = 7; group 3 had T3Nx, GS = 7, or any T stage, N+, GS = 7, or T1–2Nx, GS = 8–10; group 4 had T3Nx, GS = 8–10, or any T stage, N+, GS = 8–10.

†95% confidence intervals in parentheses.
B. THREE-DIMENSIONAL CONFORMAL RADIOTHERAPY (3DCRT) AND INTENSITY MODULATED RADIOTHERAPY (IMRT)

The technical problems mentioned in Table 25–2 have been addressed in several centers by the use of computed tomography–assisted localization and reconstruction of the pelvic anatomy. Beginning in the early 1990s, 3DCRT began to be established as the new standard of care for treating clinically localized prostate cancer. This technology takes advantage of sophisticated imaging and computerized treatment planning software that allows high-dose radiation to conform to the target volume with greater sparing of the surrounding normal tissues. Figure 25–3 is an example of the dose distribution associated with the use of several different types of IMRT. IMRT is a more sophisticated form of 3DCRT that allows higher doses of radiation to be given with less toxicity (Zelefsky, 2002). Figure 25–4A demonstrates a digitally reconstructed radiograph (DRR) from the treatment planning computed tomography and Figure 25–4B is a lateral port (treatment) film. The advantages provided by using improved imaging modality and treatment planning have resulted in the adoption of 3DCRT and more recently IMRT by major centers as the new standard of care (Table 25–4) (Speight, 2005, 2007; Boyer, 2001).

Although there appears to be no reduction in acute toxicity in patients treated with 3DCRT compared to those receiving conventional radiotherapy, chronic rectal toxicity was reduced in patients treated with conformal radiotherapy and appears to be lower with 3DCRT when the same dose is delivered (Dearnaley et al, 1999). When higher doses are given to the patients receiving 3DCRT, late complications may be increased if the technique used is not sophisticated enough to compensate for the higher doses used (Pollack et al, 2002b). Biochemical failure rates appear to be improved, but no significant difference in local control or OS has been seen so far, although the follow-up is relatively short. To date there have been six Phase III Trials addressing the issue of radiation dose. Despite a variety of approaches, doses, and differences in patient selection, a consistent pattern of outcomes supporting higher doses can be seen (Table 25–5). A much larger Phase III trial, being conducted by the RTOG (RTOG 0126), comparing results using 72.9 or 83.2 Gy for “intermediate”risk patients has recently closed to accrual. The results of this study will be available in a few years.

C. BRACHYTHERAPY

Alternative forms of radiation for the treatment of prostate cancer are growing in popularity. The most common of these alternative forms of radiation is brachytherapy. The major theoretic advantage with this form of radiation is the ability to deliver a very high dose of radiation to a localized area with a decreased number of treatment visits. The use of modern-era imaging techniques for visualizing the placement of radioactive seeds has obviated the need for open surgical procedures. Transrectal ultrasound–guided closed techniques are currently the standard. Permanent implants involve a lower dose rate of delivery but a higher

Figure 25–3. Example of the dose distribution associated with the use of several different types of IMRT.
**Figure 25-4.** Example of a 7-field conformal external-beam irradiation technique for the treatment of prostate cancer. **A:** A digitally reconstructed radiograph (DRR) from the treatment planning CT. **B:** A lateral port (treatment) film.

**Table 25–4.** 3DCRT and IMRT: Selected Major Prospective Trials.

<table>
<thead>
<tr>
<th>First Author (year)</th>
<th>Institution(s)</th>
<th>Type of Study</th>
<th>Conclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dearnaley et al (1999)</td>
<td>United Kingdom, Medical Research Council</td>
<td>Prospective randomized phase III trial (n = 225)</td>
<td>Less late GI and GU toxicity with 3D compared to non-3D, but only 64 Gy delivered and 3D technology used for only a portion of the treatment</td>
</tr>
<tr>
<td>Pollack et al (2002b)</td>
<td>MD Anderson</td>
<td>Prospective randomized phase III trial (n = 305) T1–3 randomized: 70 vs. 78 Gy</td>
<td>Patients with a PSA &gt; 10 ng/mL benefited the most from the 78 Gy arm vs. 70 Gy (62% vs. 43%; P = .01). Rectal side effects were also greater with Grade 2 or higher toxicity rates at 6 years; rates were 12% and 26% for the 70 Gy and 78 Gy arms, respectively (P = .001).</td>
</tr>
<tr>
<td>Zelefsky et al (2002)</td>
<td>Memorial Sloan Kettering</td>
<td>Phase I–II including more (n = 772) patients treated with IMRT</td>
<td>After 81–86.4 Gy with IMRT, the 3-year risk of ≥ late Grade 2 rectal or urinary toxicity was 4% and 15%. The 3-year PSA relapse-free rates for low, intermediate, and high-risk patients were 92%, 86%, and 81%.</td>
</tr>
<tr>
<td>Ryu et al (2002)</td>
<td>RTOG (Radiation Therapy Oncology Group) Multi-institutional</td>
<td>Prospective phase I–II trial (n = 173)</td>
<td>Feasible to treat safely to higher than conventional dose of 79.2 Gy in a multi-institutional setting</td>
</tr>
</tbody>
</table>

3DCRT, three-dimensional conformal radiotherapy; IMRT, Intensity Modulated Radiotherapy.

Temporary implants generally involve a lower total dose but at a higher dose rate; hence the term “high
dose rate” brachytherapy. An example of an ultrasound-
based iodine-125 permanent seed implant of the prostate,
performed at our institution, is shown in Figure 25–1. The
failure rates reported in several older studies (done in the
late 1960s and 1970s) suggested that the permanent
implants are less effective than EBRT. More recent series
suggest that the results of permanent implants may be
improved with the use of transrectal ultrasound, with long-
term results equal to or better than other treatments
(Blasko et al, 2002; Pickett et al, 2006).

At many centers, intermediate- and high-risk patients
(groups 2, 3, and 4, Table 25–3) are treated with a combina-
tion of EBRT and interstitial implant with or without
hormonal therapy. Some clinicians routinely add EBRT
for all patients undergoing permanent seed implants (Critz
et al, 2000). However, most brachytherapists agree that
low-risk patients can be equally well treated without the
additional cost or morbidity of EBRT (Frank et al, 2007;
Potters et al, 2000; Robinson et al, 2002). Although pro-
ponents of prostate brachytherapy commonly believe that
the morbidity associated with interstitial brachytherapy is
less than that associated with 3DCRT, most recent studies
suggest that acute morbidity is greater with permanent
implants, whereas long-term morbidity tends to be similar.

**Table 25–5. Major Phase III Randomized Trials Addressing the Issue of Dose of Radiation.**

<table>
<thead>
<tr>
<th>First Author</th>
<th>Trial Design</th>
<th>Estimated Control Rate at 5 years Low-Dose Arm/High-Dose Arm</th>
<th>T-Stages and PSA (Median)**</th>
<th>Impact of Higher Doses of Radiation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pollack et al (2002)</td>
<td>70 vs 78 Gy</td>
<td>60/90%</td>
<td>T1-3</td>
<td>Improved PSA control rates and trend for reduced DM</td>
</tr>
<tr>
<td>Sathya et al, 2005</td>
<td>66 Gy EBRT versus 40 Gy EBRT + 35 Gy iridium192 boost</td>
<td>~40/70%</td>
<td>T2 and T3</td>
<td>Better outcomes with iridium implant (higher biologic dose)</td>
</tr>
<tr>
<td>Lukka et al, 2005</td>
<td>66 Gy in 33 fractions versus 52.5 Gy in 20 fractions</td>
<td>53/60%</td>
<td>T1c to T2c</td>
<td>Patients receiving lower doses did worse</td>
</tr>
<tr>
<td>Zietman et al, 2005</td>
<td>70 Gy versus 79 GyE (Protons)</td>
<td>60/82%</td>
<td>T1-2, PSA 6.3 ng/mL</td>
<td>Improved PSA control rates, no difference in OS, CSS, DM</td>
</tr>
<tr>
<td>Peeters et al, 2006</td>
<td>68 versus 78 Gy</td>
<td>64/74%</td>
<td>T1-4</td>
<td>Improved PSA control rates (not by Phoenix definition) no difference in OS, CSS, DM</td>
</tr>
<tr>
<td>Dearnaley et al, 2007</td>
<td>64*** versus 74 Gy***</td>
<td>60/71%</td>
<td>T1-3</td>
<td>Improved PSA control rates, no difference in OS, CSS, DM</td>
</tr>
</tbody>
</table>

*OS, overall survival, CSS, cause-specific survival, DM, distant metastasis.
**Estimated from papers.
***Neoadjuvant androgen deprivation therapy used on each arm.

Temporary implants have the advantage of decreasing radiation exposure to hospital personnel and greater flexibility because of the ability to compensate for less than optimal needle placement. Temporary implants tend to be used for patients with more advanced disease in part because they are usually combined with EBRT and because HDR can be used to cover disease that is thought to be outside the gland. Iridium-192 is the only widely used isotope for temporary prostate implants. Based on the available data, in the hands of experts, it appears that HDR represents an excellent treatment option.

**D. NEUTRONS, PROTONS, AND HEAVY-CHARGED PARTICLES**

Eradication of tumor by radiation is believed to be dose dependent. Unfortunately, the dose beyond which no additional benefit is likely is unknown. Conformal treatment approaches have 3 interrelated goals: (1) more accurate tumor targeting, (2) the safe delivery of higher doses of radiation to the tumor, and (3) diminishing of the normal tissue toxicity associated with high-dose radiation. Particle beam radiation is an alternative form of EBRT. This class of radiation involves the use of heavy particles (eg, neutrons), charged particles (eg, protons), or heavy-charged particles (eg, neon). The theoretic advantage of proton-based radiotherapy is the potential for a more con-
formal dose distribution. The largest and most recently completed prospective randomized trial to date, addressing the use of proton beam radiotherapy, showed a significant improvement in biochemical control, but no obvious advantage to the outcomes that would be expected if conventional x-rays had been used (Zietman et al, 2005).

The attractiveness of neutron or carbon ion-based radiotherapy relates to the relative lack of oxygen dependence (Forman et al, 1996; Chuba, 1999). Heavy-charged particles (e.g. carbon ions) are thought to have the advantages of both neutrons and protons. Early studies using this technology have been encouraging, but the series are small, follow-up is relatively short, and this equipment has limited availability (Russell et al, 1994). Longer follow-up studies will be required to assess the impact of these alternative types of radiation on long-term survival.

E. POSTOPERATIVE RADIOTHERAPY

The objective of postoperative radiotherapy is to improve local-regional control by eliminating microscopic residual tumor in the surgical bed, periprostatic tissues, and regional lymph nodes. As such, there are several indications for the use of adjuvant radiation, including (1) positive surgical margins, (2) seminal vesicle involvement, (3) lymph node involvement, (4) extracapsular extension, (5) increasing PSA, and (6) biopsy-proven recurrence. The presence of any of these variables is associated with a higher incidence of local recurrence. Based on the results of two Phase III randomized trials, adjuvant, EBRT appears to reduce the incidence of local recurrence in patients with postsurgical microscopic residual tumor after radical prostatectomy (Bolla et al, 2005; Thompson et al, 2006). Patients who are treated before clinically manifesting a local recurrence appear to have improved disease-free survival, time to distant metastasis, freedom from biochemical relapse, and the need for androgen deprivation therapy (ADT) compared with patients undergoing salvage treatment. With the use of modern equipment and contemporary planning techniques, the incidence of complications is quite low. The addition of ADT with or without pelvic nodal postoperative radiotherapy may be of value in selected patients, but prospective randomized trials are needed to confirm this assertion supported by retrospective studies (Katz et al, 2003; Stephenson et al, 2004; Spiotto, 2007).

F. COMPLICATIONS OF RADIOTHERAPY FOR PROSTATE CANCER

Most patients experience urinary frequency and dysuria during the course of their treatment. In patients receiving whole-pelvic irradiation, mild diarrhea may develop, but moderate-to-severe late complications are similar (Roach, 2003a). Mild, self-limited rectal bleeding occurs in approximately 10% of patients and is dose and volume related. Urinary incontinence is usually associated with a history of a prior transurethral resection of the prostate. Hematuria and ureteral strictures occur in <2–10% of patients and are usually mild and self-limited. Fecal incontinence is uncommon, but rectal urgency due to reduction in rectal distensibility may occur in 10% of patients (Litwin, 1994). Loss of erectile function is the most worrisome, the most common, and the most permanent sequela of radiotherapy. Impotence is reported in 35–40% of patients who were potent before treatment. Sexual dysfunction may be slightly lower after brachytherapy than after EBRT (Robinson et al, 2002). However, most patients experience a decrease in the frequency and quality of intercourse, and most note a decrease in the volume of ejaculate. Potency diminishes further with time owing to aging and late radiation-induced normal tissue injury. Sparing proximal penile structures appears to be beneficial in reducing the risk of radiation-induced impotency (Roach et al, 2004; Merrick et al, 2002).

Acute urinary toxicity associated with brachytherapy is more common and longer lasting than that seen with 3DCRT. The incidence of strictures is also higher. Acute obstruction occurs in 2–20% of patients. Although incontinence is quite uncommon following radiotherapy, it can occur in up to 50% of patients if they have had a previous transurethral resection of the prostate. The frequency of rectal toxicity following brachytherapy is generally believed to be less than that with external beam treatment 3DCRT (Roach and Hsu, 2002).

NON-PROSTATE GENITOURINARY CANCERS

Urinary Tract Tumors

Urothelial cancers (UC) can occur along the entirety of the urinary tract, from the kidney to the urethra. Most occur in the bladder, though up to 5% of UC occur in the upper urinary tract. The majority of these involve the renal pelvis (Munoz, 2000). The role for radiation therapy in the management of UC varies by site, ranging from a palliative role, as in the management of renal cell cancers, to a joint role with surgery and chemotherapy, as in organ-sparing approaches for the management of muscle-invasive bladder cancer, to a primary, role as can be considered for the management of penile cancer. This section will review the common uses of EBRT and brachytherapy in the management of urinary tract malignancies.

Bladder Cancer

In the absence of durable local control, the natural history of bladder cancer is that of progressive growth and invasion with the eventual development of distant metastases. At diagnosis, the majority of patients (85%) with TCC of the bladder have superficial mucosal lesions (Ta, T1), however 70% of patients recur locally following trans-
urethral resection of the bladder tumor (TURBT). About 50–65% of these patients will progress to muscle-invasive disease (Brake, 2000). The presence of TCC in situ is associated with a more aggressive natural history, with higher probability of recurrence and progression to muscle invasive disease (Wolf, 1994). The addition of intravesicle immunotherapy (Bacille Calmette-Guerin; BCG) or chemotherapy decreases the overall recurrence rate by approximately 30% compared with TURBT alone. Nonetheless, within the first 5 years, tumor progression is noted in 20–40% of patients despite this additional treatment (Smith, 1999; Soloway, 2002). The development of muscle invasion (T2–T4) is accompanied by a significant increase in the incidence of metastatic spread and cause-specific death. Unfortunately, more than half of the patients diagnosed with muscle-invasive TCC have disseminated disease, often occult, at diagnosis. Five-year survival rates of up to 60% are reported for early invasive lesions (T1/T2a, N0); however, rates fall to ≤40% for more advanced tumors (T2b/T4, N+). Late systemic disease recurrence, most frequently pulmonary metastases, with or without local recurrence accounts for the decline in survival (Stein, 2001; Dalbagni et al, 2001), emphasizing the importance of adjuvant cytotoxic chemotherapy in the management of TCC. Following decades of unsuccessful single and bimodal therapies, contemporary management plans utilize combinations of cytotoxic chemotherapy, radiotherapy, and/or surgery in an attempt to improve survival and if possible, achieve organ preservation.

A. EBRT MANAGEMENT OF BLADDER CANCER

EBRT has had no role in the management of in situ (Tis) or superficial (T1) bladder cancer. Recently, investigators at the University of Erlangen, Germany proposed a role for post-TURBT EBRT or EBRT with chemotherapy (EBRT/CT; cisplatin or carboplatin with 5-fluorouracil) for high-risk (T1G3, T1G1-2 associated with Tis, multifocality or tumor diameter >5 cm) or multiply recurrent superficial bladder cancers (Weiss, 2006). Eighty-eight percent of (121/137) patients treated with EBRT or EBRT/CT 4–6 weeks after initial TURBT were found to have a complete response (CR) at restaging TURBT. Patients not achieving a CR (16/137; 12%) were managed with immediate cystectomy. Five- and 10-year DSS and OS rates for patients with CR were 89% and 75%, and 79% and 53%, respectively. When the evaluation was limited to patients with T1G3 tumors, 5- and 10-year DSS and OS rates were 80% and 64% and 71% and 47%. These rates are comparable to those seen in primary cystectomy series in T1 bladder cancer (Amling, 1994; Freeman, 1995; Malkowicz, 1990). Of note, patients receiving EBRT/CT had significantly lower 5-year DSS rates than patients treated with EBRT only. These findings are provocative, however a randomized trial comparing EBRT or EBRT/CT with BCG will be necessary to fully investigate the utility of this organ-sparing approach. A Phase III randomized trial by Harland et al reported that adjuvant EBRT provided no benefit over observation alone for time to progression, progression-free survival or OS for T1G3 bladder tumors (Harland et al, 2007).

The primary use of EBRT has been in muscle-invasive TCC, however many oncologists have felt that the role for radiotherapy in the management of TCC has been limited. Surgical and medical oncologists typically recommend EBRT only for those patients who are medically unfit for or refuse cystectomy, or as palliation for locally advanced, unresectable tumors. Radical cystectomy remains the “gold standard” for the management of recurrent superficial and primary muscle-invasive TCC in the United States, despite the absence of robust evidence supporting its superiority. In fact, the “optimum” management approach remains undetermined.

In earlier studies, neither radiotherapy monotherapy or pre-cystectomy radiotherapy have shown DSS or OS benefits versus radical cystectomy (Huncharek, 1998). However, most of these studies had small sample sizes, compared pathologically and clinically staged patients, and used inadequate radiotherapy techniques by current standards. Radiotherapy monotherapy yields poorer local control rates, but comparable 5-year survival rates to radical cystectomy. For muscle-invasive TCC, three of four randomized trials comparing EBRT (≤50 Gy) plus immediate cystectomy versus primary EBRT (60 Gy) and delayed (salvage) cystectomy demonstrated equivalent long-term survival rates with either treatment; only one trial demonstrated a significant benefit associated with immediate cystectomy (Bloom, 1982; Miller, 1977; Sell, 1991). In addition, no significant difference in 5- and 10-year survival rates or rates of development of distant metastases is seen with delayed or salvage cystectomy (Horwich, 1995; Petrovich, 2001). The use of combined modality treatments to achieve organ preservation without compromising treatment outcome has become a management approach of choice for many malignancies, including breast, esophageal, laryngeal, and anorectal cancers. Demonstration of equivalent outcomes with salvage surgery has made organ preservation a reasonable and appropriate treatment choice for some patients with muscle-invasive TCC.

B. COMBINED MODALITY MANAGEMENT OF MUSCLE-INVASIVE BLADDER CANCER (TRANSURETHRAL BLADDER RESSECTION, CHEMOTHERAPY, AND EBRT) AND ORGAN PRESERVATION

Several prospective randomized trials evaluating combined modality therapy for bladder preservation have been completed. In general, each of the trials has followed a common bladder-preservation algorithm including maximal TURBT, followed by induction chemoradiation, and an
assessments of treatment response. 

Individuals with clinically complete response continued with bladder-sparing therapy; all others were recommended for extirpative surgery. Completeness of TURBT (visibly complete versus not visibly complete) is associated with significantly lower salvage cystectomy rates. The key features of the contemporary bladder-sparing trials are summarized in Table 25–6.

As a body of work, nearly 1000 patients have been enrolled on these trials. Various cytotoxic agents have been evaluated for efficacy and safety when administered with EBRT. The timing of chemotherapy delivery has also been addressed. Concurrent chemoradiation schedules offer higher complete response rates compared with sequential administration (Shipley, 1998; 2005). Combinations of cisplatin-based chemotherapy with 5-fluorouracil (5-FU), Paclitaxel, or gemcitabine appear well suited for multimodality treatment for tolerability, radiosensitizing, and complementary cell-killing effects (Kaufman, 2000; von der Maase, 2005). Cisplatinum-based chemotherapy regimens administered concurrently with EBRT were well tolerated and resulted in a significant increase in freedom from distant metastases and OS. However, additional cycles of neoadjuvant chemotherapy were found not beneficial with regard to complete response rates, metastasis-free or overall survival, and were associated with higher morbidity and mortality (Tester, 1993; Shipley, 1998). Gemcitabine and the taxanes have also demonstrated significant single-agent activity against TCC. Assessment of whether or not concurrent administration of these agents with EBRT will yield acceptable toxicity and improved outcomes is ongoing. Preliminary data is promising, with 87% CR rates and acceptable toxicity with cisplatin and paclitaxel (Kaufman, 2000).

For patients achieving a complete response to the induction phase of treatment, this approach, followed by consolidation chemotherapy, yields long-term disease-free, overall, and metastases-free survival rates equivalent to those achieved with radical cystectomy. Five-year survival rates range from 50% to 62%, with nearly two-thirds of the surviving patients maintaining a well-functioning bladder. The incidence of cystectomy performed for palliation of treatment-related morbidity is low (Zeitman, 2001). Overall survival and metastases-free survival rates realized from these organ-preserving strategies approximate those achieved with primary radical cystectomy (Zeitman 2000; Nichols et al, 2000; Stein, 2001), suggesting that overall survival is driven by the presence or absence of occult distant disease at diagnosis. Limited overall survival, driven by high rates of distant metastases highlights the need to optimize systemic therapy and better select for patients who are likely to benefit from local treatment.

Several studies have evaluated epidermal growth factor receptor (EGFR) and Her-2/neu expression in bladder cancer. Immunohistochemical staining has revealed Her-2/neu overexpression in 40–80% of tumors. Data regarding the relationship between expression and treatment response and outcome are conflicting. One report evaluated the use of

Table 25–6. Contemporary Combined Modality Bladder Preservation Trials.

<table>
<thead>
<tr>
<th>Series (yr)</th>
<th>Induction Treatment*</th>
<th>CR Rate**</th>
<th>5-yr OS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shipley: RTOG 85-12 (1987)</td>
<td>CDDP+EBRT</td>
<td>66%</td>
<td>52%</td>
</tr>
<tr>
<td>Tester: RTOG 88-02 (1993) (Tester, 1993)</td>
<td>MCV+ CDDP + EBRT</td>
<td>75%</td>
<td>51%</td>
</tr>
<tr>
<td>Sauer et al (1998)</td>
<td>CDDP/Carbo + EBRT</td>
<td>71%</td>
<td>56%</td>
</tr>
<tr>
<td>Shipley: RTOG 89-03 (1998) (Shipley, 1998)</td>
<td>+/- neoadj MCV then CDDP+EBRT</td>
<td>59%</td>
<td>49%</td>
</tr>
<tr>
<td>Arias (2000)</td>
<td>Neoadj MVAC then CDDP + EBRT</td>
<td>68%</td>
<td>48%</td>
</tr>
<tr>
<td>Hussain: SWOG (2001) (Hussain, 2001)</td>
<td>CDDP + 5FU + EBRT</td>
<td>72%</td>
<td>45%</td>
</tr>
<tr>
<td>Rodel: Erlangen (2002) (Rodel, 2002)</td>
<td>CDDP/Carbo + EBRT</td>
<td>72%</td>
<td>50%</td>
</tr>
<tr>
<td>Kaufman: RTOG 99-06 (2005)</td>
<td>TAX + CDDP + bid EBRT</td>
<td>87%</td>
<td>N/A</td>
</tr>
<tr>
<td>RTOG 0223</td>
<td>TAX +CDDP + bid EBRT vs 5FU + CDDP + bid EBRT</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>RTOG 0524</td>
<td>PAX +TMaB +EBRT vs PAX + EBRT</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

*All patients underwent TURBT before induction treatment.
**Complete response rate at time of post-induction cystoscopy. CR rate, complete response rate; OS, overall survival; RTOG, Radiation Therapy Oncology Group; CDDP, cisplatin; EBRT, external beam radiotherapy; neoadj, neoadjuvant; MCV, methotrexate, cisplatin, vinblastine; 5FU, 5-fluorouracil; TAX, Taxotere; bid EBRT, twice daily EBRT; GEM, gemcitabine; MVAC, methotrexate, vinblastine, Adriamycin, cisplatin; Carbo, carboplatin; Pax, paclitaxel; TMaB, trastuzumab.
EGFR and/or Her-2 with chemotherapy and radiotherapy resistance and treatment outcomes (Chakravarti et al, 2005). EGFR expression appears to be a favorable prognostic factor for muscle-invasive TCC, and correlates with significantly higher absolute and disease-specific survival (p=0.044 and p=0.42, respectively). A trend toward decreased incidence of distant metastases was also associated with EGFR expression. Her-2 expression was significantly correlated with reduced response rates to chemoradiation. Unlike other studies, p53, p16 had no prognostic significance (del Muro, 2004). In vitro, a synergistic effect between EGFR and ionizing radiation has been shown to increase apoptosis when compared to EGFR alone (Maddineni et al, 2005). The potential diagnostic and therapeutic implications of these findings remain to be clarified.

Following chemoradiation, residual tumor will be found in 20–30% of patients at restaging cystoscopy and TURBT. In addition, 20–30% of patients who achieve a complete response develop a new or recurrent TCC. Typically, half of these tumors are superficial and half are muscle invasive. Persistent and superficial recurrences of TCC are successfully managed with TURBT with or without intravesical chemotherapy. Treatment outcomes for patients with superficial recurrences are comparable to those for patients who achieve a complete response. Invasive recurrences are managed with prompt cystectomy. Salvage surgery is not associated with compromised in overall survival (Rodel, 2002; Dunst et al, 2001; Zeitman, 2001). Although no difference in overall survival is seen in patients who subsequently develop a superficial disease relapse, the 5-year survival rate is less for patients with a native bladder than for patients who do not develop a recurrence.

C. IMPROVING TREATMENT OUTCOMES

It appears that an EBRT dose–response relationship exists for TCC. The ability to deliver higher radiotherapy doses requires sophisticated treatment planning and delivery techniques that can spare the small bowel and rectum. Anatomy-based, image-guided EBRT has multiple goals including accurate tumor targeting, the ability to safely deliver higher radiation doses, and minimizing normal tissue toxicity. The importance of image-guided conformal EBRT has been discussed in the section on prostate cancer. Here again, treatment precision including accommodation for organ motion and patient positioning is particularly important. As with prostate and other pelvic malignancies, external pressure from surrounding bowel and rectum as well as changes in the volume of urine within the bladder, lead to considerable variation in the position of the bladder (Pos et al, 2003; Langen, 2001). Fiducial marker placement and real-time imaging has been used and appears promising (Shimizu, 2000). For bladder-sparing treatment of TCC, this also preserves the option for later creation of continent diversions for patients who have an incomplete response to induction chemoradiation.

Other approaches to dose intensification include brachytherapy and altered fractionation regimens. Select European centers have used interstitial brachytherapy, usually in addition to EBRT, to treat TCC with reported local control rates of 70–90%, excellent preservation of bladder function, and low treatment-related toxicity. However in the absence of randomized prospective trials comparing treatment outcome and toxicity, interstitial brachytherapy cannot be considered a standard of care for TCC. As discussed in the section on general principles of radiotherapy, dose escalation via altered fractionation schedules (Housset et al, 1993; Hagan, 2003; Sangar et al, 2005; Kaufman, 2000) enhances the therapeutic ratio by delivering a higher effective radiotherapy dose. Aggressive dose-intensified regimens yield higher complete response rates though at the cost of moderate severe toxicity. Longer follow-up data is needed to comprehensively assess efficacy and safety.

Bladder preservation therapy provides an alternative treatment option for select patients with invasive TCC, without compromised survival. Patient selection for bladder-sparing approaches is of prime importance. Only those patients that would be suitable primary surgical candidates should be considered for this approach, since salvage cystectomy may be indicated. The optimal regimen of combined radiotherapy and chemotherapy remains to be determined.

D. TOXICITY OF RADIOTHERAPY TREATMENT FOR BLADDER CANCER

Treatment-related toxicities during and after chemoradiation primarily effect the bladder, rectum, and small bowel. Acute enteritis and cystitis are frequent complaints occurring in the majority of patients. There are usually mild and are managed symptomatically. Severe marrow-related toxicity is reported in ≤10% of organ-preservation patients. Rates of chronic bladder dysfunction of up to 10% have been reported, but symptomatic reduction of the bladder capacity is rare. Chronic, moderately severe rectal and small-bowel injuries are reported in 3–4% and 1–2%, respectively. Mortality rates are ≤1% (Chao, 1995). With wider use of conformal radiotherapy techniques, continued declines in toxicity may be anticipated.

CANCERS OF THE KIDNEY, RENAL PELVIS, & URETER

EBRT has had limited use in the management of primary renal cell adenocarcinoma. In vivo and in vitro experiments have demonstrated variable though low radiosensitivity to
conventionally fractionated EBRT (Ning et al, 1997). Randomized trials have failed to show a survival or relapse-free survival benefit from preoperative or postoperative radiotherapy (van der werf Messing, 1981; van der werf Messing, 1973; Juusela et al, 1977; Finney, 1973; Kjaer, 1987). More contemporary retrospective studies with better patient selection and using contemporary EBRT techniques have suggested a benefit of postoperative EBRT in select patients with a high-risk of local-regional failure (T3a and T3c) (Makarewicz, 1998; Kao et al, 1994; Stein, 1992). Similarly, scant data exists supporting a benefit from EBRT for renal pelvis or ureteral carcinoma. Some studies have shown a local control benefit from post-nephroureterectomy irradiation in T3-T4, N0 or node-positive patients (Maulard-Durdux et al, 1996; Cozad, 1992, 1995).

A role exists for palliative EBRT for metastatic renal cell renal pelvis and ureteral carcinomas. Palliative radiotherapy is effective at relieving pain from bone metastases, palliation of neurologic sequelae from brain metastases, spinal cord and nerve root compression, or invasion (Sheehan et al, 2003; Huguenin et al, 1998; Onufrey and Mohiuddin, 1985; Wronski et al, 1997).

**URETHRAL CANCERS**

Primary urethral cancers are very rare in both men and women. The National Cancer Institute Surveillance, Epidemiology and End Results (SEER) database in the United States identified only 1615 cases in the time period between 1973 and 2002 (www.seer.cancer.gov). As a result, knowledge regarding risk factors is extremely limited and no consensus exists regarding optimal management. Squamous cell histologies appear to be the most common followed by adenocarcinoma and TCC. Distal or anterior lesions appear to have a more favorable prognosis than proximal or posterior lesions.

**Cancer of the Female Urethra**

Urethral adenocarcinoma comprises <0.003% of urogenital tract cancers in women (Meis, 1987). The treatment strategy for each patient is largely based upon tumor size and location; however, prognosis remains relatively poor, regardless of treatment approach. Surgical excision has been used for small lesions of the distal urethra with limited success (Bracken, 1976). Both EBRT and brachytherapy are alternatives to surgical resection of early-stage urethral cancers <1 cm in size. For larger lesions, or lesions that extend into surrounding structures, preoperative EBRT radiation to the inguinal, external iliac, and hypogastric nodes is recommended (Grigsby, 1998b). Doses of 45–50 Gy are delivered to clinically uninvolved nodes, with an additional 10–15 Gy boost to any involved nodes. A total dose of 60–70 Gy is delivered to the tumor through reduced fields.

Tumors involving the posterior urethra often involve the bladder and have a high incidence of nodal involvement. Locally advanced tumors may be managed with preoperative radiotherapy and exenteration. If feasible, early lesions may be treated with surgical resection and postoperative EBRT or EBRT alone. Local control rates of 20–30% and correspondingly low 5-year survival rates are noted (Grigsby, 1998b). Preoperative EBRT doses are typically in the range of 45–50 Gy. Definitive and postoperative doses deliver 45–50 Gy to the pelvis and clinically uninvolved nodes, with an additional 10–15 Gy boost to involved nodes. A total dose of 60 Gy is delivered to the entire surface of the vaginal mucosa and brachytherapy is employed to deliver a final dose of 70–80 Gy (Grigsby, 1998a) to the primary tumor.

Urethral strictures are the most frequently reported complication of radiotherapy treatment. Urinary incontinence, cystitis, and vaginal atrophy and stenosis may also occur. Fistulas and small-bowel obstruction due to radiation or tumor necrosis are uncommon.

**Cancer of the Penis & Male Urethra**

Surgery has been the primary management choice for penile cancer. Although quite effective, neither a partial nor a total penectomy is a desirable therapeutic choice. Because of the rarity of penile cancer, no randomized trials comparing various treatment options have been or are likely to be completed. High success rates achievable with surgical salvage have made possible attempts at organ preservation with the use of radiation alone. Similarly, the relative paucity of cases has precluded the development of standardized radiotherapy management; a minimum dose of 60–65 Gy is needed for control of the primary tumor. Careful patient selection is important, with optimal candidates described as having distal, well to moderately well-differentiated tumors that are ≤4 cm in diameter.

Both EBRT and interstitial brachytherapy may be used to treat penile lesions. If not previously performed, circumcision is required prior to radiotherapy treatment. Small superficial lesions of the glans and distal shaft may be treated with orthovoltage or low-energy electron beams. Larger, invasive lesions of the penis, penile urethra, and bulbous urethra are managed with EBRT alone or EBRT followed by an interstitial brachytherapy boost. Brachytherapy is required to achieve doses to the primary >65 Gy. Prophylactic irradiation of the bilateral inguinal and pelvic lymph nodes to 45–50 Gy accompanies management of all but the most superficial primary lesions. Palpable and clinically suspicious nodes receive 65–70 Gy. Local failure rates of 15–37% have been reported (Sarin et al, 1997; Rozan, 1995). Lesions of the prostatic urethra are managed in a manner similar to that of prostate cancer.

The high risk of systemic failure associated with urethral tumors results in 5-year survival rates of 55% and
15% for distal and proximal urethral tumors, respectively (Heysek et al, 1985). Neoadjuvant chemotherapy and radiotherapy, with surgery reserved for salvage, is one strategy for more advanced lesions (Husein, Benedetto and Sridhar, 1990; Eisenberger, 1992). Regarding treatment toxicity, acute, transient sequelae include brisk, moist desquamation of the skin of the penis, urinary frequency, urgency, dysuria, nocturia, and intermittent diarrhea. The risk of soft tissue necrosis, fibrosis, and phimosis are dose-limiting sequelae for penile irradiation. Urethral or meatal stricture or both are the most common complication of penile and urethral irradiation and are dose and technique dependent. Symptomatic strictures are managed with urethral dilation, with urethrotomy reserved for severe cases. Although sexual activity is almost uniformly interrupted during the course of treatment, most patients maintain full or slightly diminished potency after radiotherapy (Opjordsmoen and Fossa, 1994).

**TESTICLE TUMORS**

Testicular cancer is the most common malignancy in men aged 15–34 years. For unknown reasons, the incidence of testicular carcinoma has been increasing worldwide, with the greatest increases seen in seminoma. Testicular cancer remains one of the most curable cancers with 5-year relative survival rates of 96–99% for non-metastatic cases. EBRT has played a primary role in the management of pure seminoma germ cell tumors (GCTs) of the testicles, but predominantly an adjuvant or palliative role in management of nonseminomatous GCT (NSGCT). Furthermore, EBRT has a limited role as palliative therapy in disseminated NSGCT, since patients with brain metastases may still be cured with chemotherapy.

**Germ Cell Tumors**

Seminoma accounts for 40% of testicular GCTs and occurs in slightly older men (median age, 33 years) than does nonseminoma testicular GCT. Nonseminoma GCTs (embryonal carcinoma, teratoma, choriocarcinoma, yolk sac tumor, and mixed GCT) comprise the remaining 60% of testicular tumors, peak at a slightly younger age (median age, 27 years), and are associated with elevations in β-hCG or alpha fetoprotein or both in 80% of cases. Radical inguinal orchiectomy with high ligation of the spermatic cord remains the therapeutic mainstay for pure seminomatous GCT, followed by surveillance, radiotherapy, or chemotherapy.

Adjuvant radiotherapy has been used to reduce the risk of local and regional recurrence in ipsilateral pelvic and para-aortic lymph nodes. Figure 25–5 shows the incidence

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**Figure 25–5.** A: Incidence and location of lymph nodes at risk for an early-stage left-sided testicular seminoma. B: Incidence and location of lymph nodes at risk for an early-stage right-sided testicular seminoma. (Adapted from Donohue JP et al: Distribution of nodal nets in nonseminomatous testis cancer. J Urol 1982;126:315.)
and location of nodal metastases in early-stage left- and right-sided testicular tumors. The classic prophylactic post-orchiectomy EBRT portal used to reduce the incidence of ipsilateral pelvic and para-aortic nodal recurrences is shown in Figure 25–6B. The exquisite radiosensitivity of pure seminoma has permitted progressive reduction in the radiotherapy dose used to treat seminoma, without a reduction relapse-free survival rates (Jones et al, 2005; Gurkaynak et al, 2003; Niewald et al, 1995). The role for EBRT in the management of pure seminoma has changed greatly over the past 3–5 years. Since equivalent disease-specific and overall survival rates may be achieved with adjuvant EBRT or surveillance for stage I seminoma (pT1-3, N0, M0, S0), in addition to the absence of treatment-related sequelae and the risk of secondary malignancy (Huyge, 2004; Zagars, 2004; Travis, 1997; Chao, 1995), surveillance has become a preferred approach for many centers (Choo et al, 2005; Warde, 1995, 2002, 1997; Miki et al, 1998).

Investigators at Princess Margaret Hospital (PMH) noted 80–85%, 5-year relapse-free rate (RFR) for stage I patients participating in an active surveillance protocol, compared with those undergoing adjuvant EBRT (95–99%) (Warde, 2005). Comparable 5-year cause-specific survival rates were achieved with EBRT salvage following surveillance and post-orchiectomy EBRT, 99.8% and 100%, respectively. The 10-year actuarial risk of requiring chemotherapy salvage was not significantly increased with surveillance.

The primary site of relapse in the surveillance population was isolated para-aortic lymph nodes (89%). Only 10% of relapses involved pelvic nodal regions. Seventy percent of relapses following prophylactic para-aortic and pelvic nodal EBRT were also in supra-diaphragmatic locations. These findings prompted their conclusion that surveillance should be the standard of care in stage I seminoma (Warde, 2005). However, until recently, surveillance had not been widely accepted as a management standard. A pattern of care study evaluating treatment practices at hospital centers in Canada and the United States showed significant variations in practice patterns (Choo et al, 2002). In addition, not all patients choose post-orchiectomy observation, nor are all patients suitable candidates for surveillance protocols. Patients who are not compliant and patients with pathologic features, which are associated with significantly higher recurrence rates including tumors >4 cm, rete testis invasion, lymphovascular space invasion, and age ≤33 years may not be ideal candidates for surveillance (Richie, 2003; Parker et al, 2002; Warde, 1997, 2002). Additional considerations are the long-term side effects of frequent radiographic examinations over several years and the cost associated with surveillance. A summary of surveillance guidelines for stage I testicular seminoma is shown in Table 25–7.

The pattern of recurrence following surveillance and prophylactic EBRT spurred investigations of use of smaller treatment fields, targeting the para-aortic nodes and omitting treatment of the pelvic nodes (limited-
field irradiation; Figure 25–6A), including a randomized trial by the Medical Research Council (MRC). Relapse-free survival rates using limited-field EBRT were equivalent to those seen with classic “dog-leg” irradiation, and hematologic, gastrointestinal, and gonadal toxicity were reduced. Pelvic nodal recurrences, however, were higher with limited-field EBRT (Niazi et al, 2005; Rowland et al, 2005; Fossa et al, 1999; Kiricuta, Sauer, and Bohndort, 1996; Logue et al, 2003; Sultanem, 1998). Patients who elect limited-field EBRT should have pelvic surveillance for a minimum of 10 years as a component of their treatment plan. Pelvic nodal recurrences are readily salvaged with EBRT or chemotherapy (Power et al, 2005). In addition to the benefit of reduced treatment portals, it is reasonable to assume that increased use of IMRT will lead to further reductions in EBRT-related toxicity.

The greatest change in the management of stage I seminoma has been the use single agent carboplatin. Five Phase II trials conducted in Europe assessed relapse-free survival rates with two cycles of post-orchiectomy, adjuvant carboplatin. With follow-up ranging from 14–74 months, relapse rates were <1% and there was <5% grade three to four hematologic toxicity. The MRC conducted a prospective Phase III trial that randomized men with stage I seminoma to 20–30 Gy limited-field (para-aortic nodal) or large-field (pelvis and para-aortic nodal irradiation) EBRT or one cycle of adjuvant carboplatin (AUC7) (Oliver et al, 2004; Oliver et al, 2005; Reiter et al, 2001; Dieckmann et al, 2000; Krege et al, 1997; Germa-Lluch et al, 2002). At 4 years median follow-up, the results indicated that single-cycle carboplatin was non-inferior to prophylactic nodal irradiation; relapse-free survival rates were equivalent, patients receiving carboplatin were less likely to take time off from work and significantly fewer secondary tumors were reported.

With an absence of consensus regarding treatment for stage I seminoma, the Spanish Germ Cell Cooperative group developed a risk-adapted management strategy (Figure 25–7) (Aparicio et al, 2005). In January 2007, the NCCN published new practice guidelines for the management of testicular seminoma (http://www.nccn.org) incorporating the data from the contemporary seminoma trials. The impetus for these treatment recommendations appears to be the judicious use of adjuvant therapy in patients with significant risk of disease relapse. In so doing the late toxicities associated with chemotherapy and radiotherapy may be reduced or avoided.

The use for adjuvant post-orchiectomy EBRT in stage II seminoma (pT1-3, N1-3, M0, S0-1) is determined by the bulk of the retroperitoneal lymphadenopathy. Stage IIA (single or multiple nodes all ≤ 2 cm) and IIB (single or multiple nodes 2–5 cm) patients are successfully treated with EBRT directed to the para-aortic and ipsilateral pelvic lymph nodes, using the so called “hockey stick” or “dog-leg” fields (Figure 25–6B). Doses of 20–30 Gy to the

**Table 25–7. Surveillance Guidelines for Stage I Testicular Seminoma.**

<table>
<thead>
<tr>
<th>Frequency</th>
<th>Test</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Every 3–4 months, years 1–3**</td>
<td>History and physical</td>
<td>Serum AFP, beta-hCG, LDH</td>
</tr>
<tr>
<td>Every 3–4 months, years 1–10</td>
<td>Abdominal/pelvic CT scan</td>
<td></td>
</tr>
<tr>
<td>Every 6 months years 1–10</td>
<td>Chest x-ray</td>
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<tr>
<td><strong>After year 3 the frequency changes to every 6 months until year 7 and then annually.</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>AFP, alpha fetoprotein; hCG, human chorionic gonadotrophin; LDH, lactate dehydrogenase.</strong></td>
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**Figure 25–7. Risk-adapted strategy for the management of stage I seminoma.**
pelvis and para-aortic nodes followed by a 5–10-Gy boost to bulky nodes yield 5-year and 10-year relapse-free, cause-specific, and overall survival rates of 85%, 94%, and 93%, respectively (Rowland et al, 2005; Chung et al, 2004; Classen et al, 2003; Warde, 1998). Heterogeneity within clinical stages IIA and IIB seminoma due to the presence of one versus multiple nodal masses has led to the suggested use of single agent carboplatin with EBRT, particularly for stage IIB (Patterson, et al. 2001). Patients with bulky retroperitoneal nodal disease (≥ 25 cm, IIC) have high rates of distant relapse. Standard of care is three cycles of cisplatin-etoposide and bleomycin (PEB) or four cycles of etoposide and cisplatinum, followed by surveillance. Residual disease < 3 cm may be closely observed, treated with radiotherapy, or surgically resected. Masses ≥ 3 cm should be resected. Prophylactic mediastinal irradiation for stage II seminoma has long been abandoned. Supradiaphragmatic recurrence rates approximate 3% and may be successfully salvaged with multiagent chemotherapy. EBRT has no role in the primary management of stages IIC, III, and IV seminoma.

A. TOXICITY FROM RADIOTHERAPY TREATMENT OF TESTICULAR SEMINOMA

Treatment-related toxicity associated with pelvic and para-aortic EBRT are primarily gastrointestinal in nature and can be easily managed. The most common sequelae may include nausea, diarrhea, rectal urgency, peptic ulcer disease, GERD, and transient reduction in spermatogenesis (Garcia-Serra et al, 2005; Joos et al, 1997). It has been estimated that 50% or more of the men diagnosed with testicular cancer have impaired spermatogenesis at the time of diagnosis, complicating precise characterization of EBRT effects on fertility (Fossa et al, 1986; Gordon et al, 1997; Pasquolotto, 2003). The severity and duration of oligospermia appears to be dose related and testicular scatter doses are thought to be responsible for fertility disturbances. Radiation-related disturbances in spermatogenesis can be minimized by tailoring scatter doses to ≤ 20 cGy and the use of smaller treatment fields (Joos et al, 1997; Melchor et al, 2001). This is readily achieved with contemporary treatment approaches. Patients treated with post-orchiectomy EBRT have >50% chance of regaining normal spermatogenesis and all patients regain at least some spermatogenesis, usually within 1–2 years after treatment completion (Gordon et al, 1997; Nalesnik et al, 2004). The existence of severe disturbances in spermatogenesis late in the post-EBRT course probably result from impaired pretreatment sperm cell production and less so the impact of scatter radiation on the remaining testicle.

A small but measurable increase in the risk of second malignancies after a 10–20-year latent period has been reported (Travis, 1997; Zagars, 2004; Hughes et al, 2003). In some reports however, the overall observed incidence of second nonseminomatous malignancy was not significantly increased when compared with the expected incidence (Chao, 1995; Travis, 1997).

SUMMARY

Therapeutic radiation has an extensive history in the management of genitourinary (GU) malignancies. Major advances have been made in its use, particularly for prostate cancer, muscle-invasive TCC of the bladder and testicular seminoma, and a continued role for radiation therapy in the multidisciplinary management of genitourinary malignancies seems certain. Overall treatment outcomes continue to improve, accompanied by diminishing rates of toxicity. With continued technologic development, including the use of adaptive radiotherapy, the discovery and application of novel treatment agents, and the combined efforts of each discipline within urologic oncology, patients with GU tumors have greater opportunities to become and remain disease free.

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General, Radiobiology, & Prostate Cancer


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Introduction


Bladder Cancer


Kidney, Renal Pelvis, and Ureter


Cancer of the Female Urethra


Penis and Male Urethra


Testicular Cancer


INTRODUCTION

The bladder, in concert with the urethra and the pelvic floor, is responsible for storage and periodic expulsion of urine. The integrated function of these components of the lower urinary tract (LUT) is dependent on a complex control system in the brain, spinal cord and peripheral ganglia, and on local regulatory factors (de Groat, 2006). Dysfunction of the central nervous control systems or of the components of the LUT can either produce insufficient voiding and retention of urine, or different types of urinary incontinence (mainly urgency and stress incontinence), or the symptom complex of the “overactive bladder” (OAB), characterized by urgency, frequency with or without urgency incontinence, often with nocturia (Abrams et al, 2002).

Pharmacologic treatment of urinary incontinence is the main option, and several drugs with different modes and sites of action have been tried (Andersson and Wein, 2004; Ouslander, 2004; Zinner, Koke, and Viktrup, 2004). However, to be able to optimize treatment, knowledge about the mechanisms of micturition and of the targets for treatment is necessary. Theoretically, failure to store urine can be improved by agents that decrease detrusor activity and increase bladder capacity, and/or increase outlet resistance.

In this chapter, a brief review is given of the normal nervous control of the LUT and of some therapeutic principles used in the treatment of urinary incontinence.

NEURAL CIRCUITS CONTROLLING STORAGE AND EXPULSION OF URINE

Normal micturition occurs in response to afferent signals from the LUT (Shefchyk, 2002; Holstege, 2005; Sugaya et al, 2005; de Groat, 2006). Both bladder filling and voiding are controlled by neural circuits in the brain, spinal cord, and peripheral ganglia. These circuits coordinate the activity of the smooth muscle in the detrusor and urethra with that of the striated muscles in the urethral sphincter and pelvic floor. Suprapontine influences are believed to act as on-off switches to shift the LUT between the 2 modes of operation: storage and elimination. In adults, urine storage and voiding are under voluntary control and depend upon learned behavior. In infants, however, these switching mechanisms function in a reflex manner to produce involuntary voiding. In adults, injuries or diseases of the central nervous system (CNS) can disrupt the voluntary control of micturition and cause the reemergence of reflex micturition, resulting in OAB and detrusor overactivity (DO). Because of the complexity of the CNS control of the LUT, OAB and DO can occur as a result of a variety of neurological disorders as well as changes in the peripheral innervation and smooth and skeletal muscle components (Andersson and Wein, 2004; Andersson and Arner 2004).

Filling of the bladder and voiding involve a complex pattern of afferent and efferent signaling in parasympathetic (pelvic nerves), sympathetic (hypogastric nerves), and somatic (pudendal nerves) pathways. These pathways constitute reflexes, which either keep the bladder in a relaxed state, enabling urine storage at low intravesical pressure, or which initiate bladder emptying by relaxing the outflow region and contracting detrusor. Integration of the autonomic and somatic efferents result in that contraction of the detrusor muscle is preceded by a relaxation of the outlet region, thereby facilitating bladder emptying. On the contrary, during the storage phase, the detrusor muscle is relaxed and the outlet region is contracted to maintain continence.

PARASYMPATHETIC PATHWAYS

The sacral parasympathetic pathways mediate contraction of the detrusor smooth muscle and relaxation of the outflow region. The preganglionic parasympathetic neurones are located to the sacral parasympathetic nucleus (SPN) in the spinal cord at the level of S2–S4. The axons pass
through the pelvic nerves and synapse with the postganglionic nerves in either the pelvic plexus, in ganglia on the surface of the bladder (vesical ganglia), or within the walls of the bladder and urethra (intramural ganglia). The ganglionic neurotransmission is predominantly mediated by acetylcholine acting on nicotinic receptors, although the transmission can be modulated by adrenergic, muscarinic, purinergic, and peptidergic presynaptic receptors. The postganglionic neurons in the pelvic nerve mediate the excitatory input to the normal human detrusor smooth muscle by releasing acetylcholine acting on muscarinic receptors (see below). However, an atropine-resistant (nonadrenergic, noncholinergic: NANC) contractile component is regularly found in the bladders of most animal species. Such a component can also be demonstrated in functionally and morphologically altered human bladder tissue (O’Reilly et al, 2002), but contributes only to a few percent to normal detrusor contraction (Andersson and Wein, 2004). Adenosine triphosphate (ATP) is the most important mediator of the NANC contraction, although the involvement of other transmitters cannot be ruled out (Andersson and Wein, 2004). The pelvic nerve also conveys parasympathetic nerves to the outflow region and the urethra. These nerves exert an inhibitory effect on the smooth muscle, by releasing nitric oxide and other transmitters (Andersson and Wein, 2004).

**SYMPATHETIC PATHWAYS**

The sympathetic innervation of the bladder and urethra originates from the intermediolateral nuclei in the thoracolumbar region (T10–L2) of the spinal cord. The axons leave the spinal cord via the splanchnic nerves and travel either through the inferior mesenteric ganglia (IMF) and the hypogastric nerve, or pass through the paravertebral chain to the lumbosacral sympathetic chain ganglia and enter the pelvic nerve. Thus, sympathetic signals are conveyed in both the hypogastric nerve and the pelvic nerve. The ganglionic sympathetic transmission is, like the parasympathetic preganglionic transmission, predominantly mediated by acetylcholine acting on nicotinic receptors. Some preganglionic terminals synapse with the postganglionic cells in the paravertebral ganglia or in the IMF, while other synapse closer to the pelvic organs, and short postganglionic neurones innervate the target organs. Thus, the hypogastric and pelvic nerves contain both pre- and postganglionic fibre. The predominant effect of the sympathetic innervation is to contract the bladder base and the urethra. In addition, the sympathetic innervation inhibits the parasym pathetic pathways at spinal and ganglionic levels. In the human bladder, noradrenaline is released in response to electrical stimulation in vitro, and the normal detrusor response to released noradrenaline is relaxation. However, the importance of the sympathetic innervation for relaxation of the human detrusor has never been established. In contrast, in several animal species the adrenergic innervation has been demonstrated to mediate relaxation of the detrusor during filling (Andersson and Arner, 2004).

**SOMATIC PATHWAYS**

The somatic innervation of the urethral rhabdosphincter and of some perineal muscles (eg, compressor urethrae and urethrovaginal sphincter) is provided by the pudendal nerve. These fibers originate from sphincter motor neurons located in the ventral horn of the sacral spinal cord (levels S2–S4) in a region called Onuf’s (Onufrowicz’s) nucleus (Thor and Donatucci, 2004).

**AFFERENT PATHWAYS**

The afferent nerves to the bladder and urethra originate in the dorsal root ganglia at the lumbosacral level of the spinal cord and travel via the pelvic nerve to the periphery. Some afferents originate in dorsal root ganglia at the thoracolumbar level and travel peripherally in the hypogastric nerve. The afferent nerves to the striated muscle of the external urethral sphincter travel in the pudendal nerve to the sacral region of the spinal cord. The most important afferents for the micturition process are myelinated Aδ-fibers and unmyelinated C-fibers traveling in the pelvic nerve to the sacral spinal cord, conveying information from receptors in the bladder wall. The Aδ-fibers respond to passive distension and active contraction, thus conveying information about bladder filling. The activation threshold for Aδ-fibers is 5–15 mm H₂O. This is the intravesical pressure at which humans report the first sensation of bladder filling. C-fibers have a high mechanical threshold and respond primarily to chemical irritation of the bladder urothelium/suburothelium or to cold. Following chemical irritation, the C-fiber afferents exhibit spontaneous firing when the bladder is empty and increased firing during bladder distension. These fibers are normally inactive and are therefore termed “silent fibers.” Afferent information about the amount of urine in the bladder is continuously conveyed to the mesencephalic periaqueductal gray (PAG), and from there to the pontine micturition center (PMC), also called Barrington’s nucleus (Holstege, 2005; Kuipers, Mouton, and Holstege, 2006).

**AFFERENT SIGNALING FROM THE UROTHELIUM/SUBUROTHELIUM**

Recent evidence suggests that the urothelium/suburothelium may serve not only as a passive barrier, but also as a specialized sensory and signaling unit, which, by producing nitric oxide, ATP, and other mediators, can control the activity in afferent nerves, and thereby the initiation of the micturition reflex (Andersson, 2002; de Groat, 2004: Birder and de Groat, 2007). The urothelium has been shown to express for example, nicotinic, muscarinic, tachykinin, adrenergic, bradykinin, and transient receptor potential (TRP)
receptors (de Groat, 2004; Birder et al, 2002; Birder et al, 2001). Low pH, high K⁺, increased osmolality, and low temperatures can all influence afferent nerves, possibly via effects on the vanilloid receptor (capsaicin- [CAP] gated ion channel, TRPV1), which is expressed both in afferent nerve terminals and in the urothelial cells (Birder et al, 2002; Birder et al, 2001). A network of interstitial cells, extensively linked by Cx43-containing gap junctions, was found to be located beneath the urothelium in the human bladder (Sui et al, 2002; Sui, Wu, and Fry, 2004; Brading and McCloskey, 2005). This interstitial cellular network was suggested to operate as a functional syncytium, integrating signals and responses in the bladder wall. The firing of suburothelial afferent nerves, conveying sensations and regulating the threshold for bladder activation, may be modified by both inhibitory (eg, nitric oxide) and stimulatory (eg, ATP, tachykinins, prostanoids) mediators. ATP, generated by the urothelium, has been suggested as an important mediator of urothelial signaling (Andersson, 2002). Supporting such a view, intravesical ATP induces DO in conscious rats (Pandita and Andersson, 2002). Furthermore, mice lacking the P2X₃ receptor were shown to have hypoactive bladders (Cockayne et al, 2000; Vlaskovska et al, 2001). Interstitial cells can also be demonstrated within the detrusor muscle (Brading and McCloskey, 2005). They may be involved in impulse transmission, but their role has not been clarified.

There seem to be other, thus far unidentified, factors in the urothelium that could influence bladder function (Andersson and Wein, 2004). Even if these mechanisms can be involved in, for example, the pathophysiology of OAB, their functional importance remains to be established.

**NEURAL CONTROL OF BLADDER FILLING**

During the storage phase the bladder has to relax in order to maintain a low intravesical pressure. Urine storage is regulated by 2 separate storage reflexes, of which one is sympathetic (autonomic) and the other is somatic (Thor and Donatucci, 2004). The sympathetic storage reflex (pelvic-to-hypogastric reflex) is initiated as the bladder distends (myelinated Aδ-fibers) and the generated afferent activity travels in the pelvic nerves to the spinal cord. Within the spinal cord, sympathetic firing from the lumbar region (L1–L3) is initiated, which, by effects at the ganglionic level, decreases excitatory parasympathetic inputs to the bladder. Postganglionic neurons release noradrenaline, which facilitates urine storage by stimulating β₂-adrenoceptors (ARs) in the detrusor smooth muscle (see below). As mentioned previously, there is little evidence for a functionally important sympathetic innervation of the human bladder, which is in contrast to what has been found in several animal species. The sympathetic innervation of the human bladder is found mainly in the outlet region, where it mediates contraction. During micturition, this sympathetic reflex pathway is markedly inhibited via supraspinal mechanisms to allow the bladder to contract and the urethra to relax. Thus, the Aδ afferents and the sympathetic efferent fibers constitute a vesico-spinal-vesical storage reflex, which maintains the bladder in a relaxed mode while the proximal urethra and bladder neck are contracted.

In response to a sudden increase in intra-abdominal pressure, such as during a cough, laugh, or sneeze, a more rapid somatic storage reflex (pelvic-to-pudendal reflex), also called the guarding or continence reflex, is initiated. The evoked afferent activity travels along myelinated Aδ afferent nerve fibers in the pelvic nerve to the sacral spinal cord, where efferent somatic urethral motor neurons, located in the nucleus of Onuf, are activated. Afferent information is also conveyed to the PAG and from there to the PMC (the L region). From this center impulses are conveyed to the motor neurons in the nucleus of Onuf. Axons from these neurons travel in the pudendal nerve and release acetylcholine, which activates nicotinic cholinergic receptors on the rhabdosphincter, which contracts. This pathway is tonically active during urine storage. During sudden abdominal pressure increases, however, it becomes dynamically active to contract the rhabdosphincter. During micturition, this reflex is strongly inhibited via spinal and supraspinal mechanisms to allow the rhabdosphincter to relax and permit urine passage through the urethra. In addition to this spinal somatic storage reflex, there is also supraspinal input from the pons, which projects directly to the nucleus of Onuf and is of importance for voluntary control of the rhabdosphincter (Holstege, 2005; Sugaya et al, 2005; Blok, de Weerd, and Holstege, 1997).

**NEURAL CONTROL OF BLADDER EMPTYING**

**Vesico-Bulbo-Vesical Micturition Reflex**

Electrophysiological experiments in cats and rats provide evidence for a voiding reflex mediated through a vesicobulbo-vesical pathway involving neural circuits in the pons, which constitute the PMC. Other regions in the brain, important for micturition, include the hypothalamus and cerebral cortex (Holstege, 2005; Griffiths, 2004; Griffiths et al, 2005). Bladder filling leads to increased activation of tension receptors within the bladder wall and thus to increased afferent activity in Aδ-fibers. These fibers project on spinal tract neurones mediating increased sympathetic firing to maintain continence as discussed above (storage reflex). In addition, the spinal tract neurones convey the afferent activity to more rostral areas of the spinal cord and the brain. As mentioned previously, one important receiver of the afferent information from the bladder is the PAG in the rostral brain stem (Holstege, 2005; Kuipers, Mouton, and Holstege, 2006). The PAG receives information from both afferent neurones in the bladder and from more rostral areas in the brain, that is, cerebral cortex and hypothalamus. This information is integrated
in the PAG and the medial part of the PMC (the M region), which also control the descending pathways in the micturition reflex. Thus, PMC can be seen as a switch in the micturition reflex, inhibiting parasympathetic activity in the descending pathways when there is low activity in the afferent fibers, and activating the parasympathetic pathways when the afferent activity reaches a certain threshold. The threshold is believed to be set by the inputs from more rostral regions in the brain. In cats, lesioning of regions above the inferior colliculus usually facilitates micturition by elimination of inhibitory inputs from more rostral areas of the brain. On the other hand, transections at a lower level inhibit micturition. Thus, the PMC seems to be under a tonic inhibitory control. A variation of the inhibitory input to PMC results in a variation of bladder capacity. Experiments on rats have shown that the micturition threshold is regulated by, for example, gamma-amino butyric acid (GABA)-ergic inhibitory mechanisms in the PMC neurones.

**Vesico-Spinal-Vesical Micturition Reflex**

Spinal lesions rostral to the lumbo-sacral level interrupt the vesico-bulbo-vesical pathway and abolish the supra spinal and voluntary control of micturition (Anderson and Wein, 2004). This results initially in an areflexic bladder accompanied by urinary retention. An automatic vesico-spi- nal vesical micturition reflex develops slowly, although voiding is generally insufficient due to bladder-sphincter dyssynergia, that is, simultaneous contraction of bladder and urethra. It has been demonstrated in chronic spinal cats that the afferent limb of this reflex is conveyed through unmyelinated C-fibers, which usually do not respond to bladder distension, suggesting changed properties of the afferent receptors in the bladder. Accordingly, the micturition reflex in chronic spinal cats is blocked by CAP, which blocks C-fiber-mediated neurotransmission.

**TARGETS FOR PHARMACOLOGIC INTERVENTION**

**CNS TARGETS**

Anatomically, several CNS regions may be involved in micturition control: supraspinal structures, such as the cortex and diencephalon, midbrain, and medulla, but also spinal structures (Holstege, 2005; Sugaya et al, 2005; Griffths, 2004; Griffths et al, 2005). Several transmitters are involved in the micturition reflex pathways described above and may be targets for drugs aimed for control of micturition (de Groat and Yoshimura, 2001). However, few drugs with a CNS site of action have been developed (Andersson and Pehrson, 2003).

**Opioid Receptors**

Endogenous opioid peptides and corresponding receptors are widely distributed in many regions in the CNS of importance for micturition control (de Groat and Yoshimura, 2001). It has been well established that morphine, given by various routes of administration to animals and humans, can increase bladder capacity or block bladder contractions. Furthermore, given intrathecally to anesthetized rats and intravenously to humans, the mu-opioid receptor antagonist, naloxone, has been shown to stimulate micturition, suggesting that a tonic activation of mu-opioid receptors has a depressant effect on the micturition reflex. However, intrathecal naloxone was not effective in stimulating micturition in conscious rats at doses blocking the effects of intrathecal morphine (Andersson and Wein, 2004).

Morphine given intrathecally was effective in patients with DO due to spinal cord lesions, but was associated with side effects, such as nausea and pruritus. Further side effects of opioid receptor agonists comprise respiratory depression, constipation, and abuse (Andersson and Wein, 2004). Attempts have been made to reduce these side effects by increasing selectivity toward one of the different opioid receptor types. At least 3 different opioid receptors: µ, δ, and κ bind stereospecifically with morphine, and have been shown to interfere with voiding mechanisms. Theoretically, selective receptor actions, or modifications of effects mediated by specific opioid receptors, may have useful therapeutic effects for micturition control.

Tramadol is a well-known analgesic drug. By itself, it is a weak mu-receptor agonist, but it is metabolized to several different compounds, some of them almost as effective as morphine at the mu-receptor. However, the drug also inhibits serotonin (5-HT) and noradrenaline reuptake (Raffa and Friderichs, 1996). This profile is of particular interest, since both mu-receptor agonism and amine reuptake inhibition may be useful principles for treatment of DO/OAB.

When tramadol is given to a normal, awake rat, the most conspicuous changes in the cystometrogram are increases in threshold pressure and bladder capacity. Naloxone can more or less completely inhibit these effects (Pandita, Pehrson, and Christoph, 2003). However, there are differences between the effects of tramadol and morphine. Morphine has a very narrow range between the doses causing inhibition of micturition and those increasing bladder capacity and evoking urinary retention. Tramadol has effects over a much wider range of doses, which means that it could be therapeutically more useful for micturition control. It may be speculated that the difference is dependent on the simultaneous influence of the 5-HT and...
Serotonin (5-HT) Mechanisms

Lumbosacral autonomic, as well as somatic, motor nuclei (Onuf’s nuclei), receive a dense serotonergic input from the raphe nuclei, and multiple 5-HT receptors have been found at sites where afferent and efferent impulses from and to the LUT are processed (Ramage, 2006). The main receptors shown to be implicated in the control of micturition are the 5-HT\(_{1A}\), 5-HT\(_{2}\), and 5-HT\(_{3}\) receptors (Ramage, 2006). There is some evidence in the rats for serotonergic facilitation of voiding; however, the descending pathway is essentially an inhibitory circuit, with 5-HT as a key neurotransmitter (de Groat, 2002; Tai et al, 2006). Thus, electrical stimulation of 5-HT-containing neurons in the caudal raphe nucleus causes inhibition of bladder contractions (Sugaya et al, 1998). Most experiments in rats and cats indicate that activation of the central serotonergic system by 5-HT reuptake inhibitors, as well as by 5-HT\(_{1A}\) and 5-HT\(_{2}\) receptor agonists, depresses reflex bladder contractions and increases the bladder volume threshold for inducing micturition (Ramage, 2006; de Groat 2002; Tai et al, 2006). 5-HT\(_{1A}\) receptors are involved in multiple inhibitory mechanisms controlling the spinobulbospinal micturition reflex pathway. The regulation of the frequency of bladder reflexes is presumably mediated by a suppression of afferent input via an action on interneuronal pathways in the spinal cord (Tai et al, 2006), and suppression of the micturition switching circuitry in the pons. The regulation of bladder contraction amplitude may be related to an inhibition of the output from the pons to the parasympathetic nuclei in the spinal cord. Even if blockade of 5-HT\(_{1A}\) receptors seem to have effects that would be beneficial for DO treatment, there is evidence showing that blockade of these receptors lead to rapid tolerance (Ramage, 2006).

It has been speculated that selective serotonin reuptake inhibitors (SSRIs) may be useful for treatment of DO/OAB. On the other hand, there are reports suggesting that the SSRIs in patients without incontinence actually can cause incontinence, particularly in the elderly, and one of the drugs (sertraline) seemed to be more prone to produce urinary incontinence than the others (Movig et al, 2002). Patients exposed to serotonin uptake inhibitors had an increased risk (15 out of 1000 patients) for developing urinary incontinence. So far, there are no RCTs demonstrating the value of SSRIs in the treatment of DO/OAB.

GABA Mechanisms

Both in the brain and the spinal cord GABA has been identified as a main inhibitory transmitter (de Groat and Yoshimura, 2001). GABA functions appear to be triggered by binding of GABA to its inotropic receptors, GABA\(_{\alpha}\) and GABA\(_{\beta}\), which are ligand-gated chloride channels, and its metabotropic receptor, GABA\(_{\gamma}\) (Chebib and Johnston, 1999). Since blockade of GABA\(_{\alpha}\) and GABA\(_{\beta}\) receptors in the spinal cord and brain (Pehrson and Andersson, 2002) stimulated rat micturition, an endogenous activation of GABA\(_{\alpha,\beta}\) receptors may be responsible for continuous inhibition of the micturition reflex within the CNS. In the spinal cord, GABA\(_{\alpha}\) receptors are more numerous than GABA\(_{\gamma}\) receptors, except for the dorsal horn where GABA\(_{\beta}\) receptors predominate.

GABA transporters, present on neuronal and glial cells in the brain, brain stem, and spinal cord, are presumed to provide an inactivation mechanism. Four different GABA transporters (GATs) have been described. Tiagabine is a selective inhibitor of one of these GATs, GAT1, is able to increase extracellular levels of GABA, and has inhibitory effects on rat micturition (Pehrson and Andersson, 2002). Intravenous administration of tiagabine decreased micturition pressure and decreased voided volume. Tiagabine given intrathecally reduced micturition pressure and increased bladder capacity (Pehrson and Andersson, 2002), suggesting that increasing endogenous levels of GABA in the CNS may improve micturition control.

Experiments using conscious and anesthetized rats demonstrated that exogenous GABA, muscimol (GABA\(_{\alpha}\) receptor agonist), and baclofen (GABA\(_{\beta}\) receptor agonist) given intravenously, intrathecally, or intracerebroventricularly inhibit micturition (Pehrson, Lehmann, and Andersson, 2002). Baclofen given intrathecally attenuated oxyhemoglobin-induced DO, suggesting that the inhibitory actions of GABA\(_{\beta}\) receptor agonists in the spinal cord may be useful for controlling micturition disorders caused by C-fiber activation in the urothelium and/or suburothelium (Pehrson, Lehmann, and Andersson, 2002).

Stimulation of the PMC results in an immediate relaxation of the external striated sphincter and a contraction of the detrusor muscle of the bladder demonstrated in cats a direct pathway from the PMC to the dorsal gray commissure of the sacral cord (Blok, de Weerd, and Holstege, 1997). It was suggested that the pathway produced relaxation of the external striated sphincter during micturition via inhibitory modulation by GABA neurons of the motoneurons in the sphincter of Onuf (Blok, de Weerd, and Holstege, 1997). In rats, intrathecal baclofen and muscimol ultimately produced dribbling urinary incontinence (Pehrson, Lehmann, and Andersson, 2002).

Thus, normal relaxation of the striated urethral sphincter is probably mediated via GABA\(_{\alpha}\) receptors (Pehrson and Andersson, 2002; Pehrson, Lehmann, and Andersson, 2002).
Noradrenaline Mechanisms

Noradrenergic neurons in the brainstem project to the sympathetic, parasympathetic, and somatic nuclei in the lumbarosacral spinal. Bladder activation through these bulbo-sacral noradrenergic pathways may involve excitatory lumbosacral spinal. Bladder activation through these bulbo-sacral noradrenergic neurons in the brainstem project to the sacral parasympathetic, parasympathetic, and somatic nuclei in the brainstem. Gabapentin is also widely used not only for seizures and neuropathic pain, but for many other indications, such as anxiety and sleep disorders, because of its apparent lack of toxicity.

In a pilot study, Carbone et al (2003) reported on the effect of gabapentin on neurogenic DO. These investigators found a positive effect on symptoms and significant improvement in urodynamic parameters after treatment with gabapentin, and suggested that the effects of the drug should be explored in further controlled studies in both neurogenic and non-neurogenic DO. Kim et al (2004) studied the effects of gabapentin in patients with OAB and nocturia not responding to antimuscarinics. They found that 14 out of 31 patients improved with oral gabapentin. The drug was generally well tolerated, and the authors suggested that it can be considered in selective patients when conventional modalities have failed. It is possible that gabapentin and other \( \alpha_2 \delta \) ligands (eg, pregabalin and analogs) will offer new therapeutic alternatives.

Dopamine Mechanisms

Patients with Parkinson’s disease may have neurogenic DO, possibly as a consequence of nigrostriatal dopamine depletion and failure to activate inhibitory D1 receptors (Andersson, 2004). However, other dopaminergic systems may activate D2 receptors, facilitating the micturition reflex. Apomorphine, which activates both D1 and D2 receptors, induced bladder overactivity in anesthetized rats via stimulation of central dopaminergic receptors. The effects were abolished by intracollicular transection of the brain, and by prior intraperitoneal administration of the centrally acting dopamine receptor blocker, spiroperidol. It has been shown that the DO induced by apomorphine in anesthetized rats resulted from synchronous stimulation of the micturition centers in the brainstem and spinal cord, and that the response was elicited by stimulation of both dopamine D1 and D2 receptors. Blockade of central dopamine receptors may be expected to influence voiding; however, the therapeutic potential of drugs having this action has not been established (Andersson and Wein, 2004).

PERIPHERAL TARGETS

There are many possible peripheral targets for pharmacologic control of bladder function (Andersson and Arner, 2004). Although many effective drugs are available targeting these systems, most of them are less useful in the clinical situation due to the lack of selectivity for LUT, which may result in intolerable side effects.

Muscarinic Receptors

Muscarinic receptors comprise five subtypes, \( \text{M}_1 - \text{M}_5 \), encoded by 5 distinct genes, and in both animal and human bladders, the mRNAs for all muscarinic receptor subtypes have been demonstrated, with a predominance of mRNAs encoding \( \text{M}_3 \) and \( \text{M}_4 \) receptors. These receptors are also functionally coupled to G proteins, but the signal transduction systems vary (Andersson and Arner, 2004).

Detrusor smooth muscle contains muscarinic receptors mainly of the \( \text{M}_3 \) and \( \text{M}_4 \) subtypes. The \( \text{M}_3 \) receptors in the human bladder are the most important for detrusor contraction (Andersson and Wein, 2004). In the human detrusor, Schneider et al (2004) confirmed that the muscarinic receptor subtype mediating carbachol-induced contraction was the \( \text{M}_3 \) receptor, and they also demonstrated that the L-type calcium channel blocker, nifedipine, almost completely inhibited carbachol-induced detrusor contraction, whereas an inhibitor of store-operated \( \text{Ca}^{2+} \) channels caused little inhibition. The Rho-kinase inhibitor, Y 27,632, produced a concentration-dependent attenuation of the carbachol-induced contractile responses. Schneider et al (2004) concluded that carbachol-induced contraction of human detrusor is mediated via \( \text{M}_3 \) recep-
tors, and furthermore, largely depends on transmembrane Ca\(^{2+}\)-flux through nifedipine-sensitive calcium channels as well as activation of the Rho-kinase pathway. These conclusions were supported by Takahashi et al (2004) who found that in human detrusor muscle carbachol induces contraction, not only by increasing [Ca\(^{2+}\)]\(_i\) but also by increasing the Ca\(^{2+}\) sensitivity of the contractile apparatus in a Rho-kinase and protein kinase C-dependent manner.

It has been suggested that M\(_3\) receptors may oppose sympathetically mediated smooth muscle relaxation, mediated by β-ARs (Hegde, 1997). M\(_2\) receptor stimulation may also activate nonspecific cation channels and inhibit K\(_{\text{ATP}}\) channels through activation of protein kinase C. However, the functional role for the M\(_2\) receptors in the normal bladder has not been clarified, but in certain disease states, M\(_2\) receptors may contribute to contraction of the bladder. Thus, in the denervated rat bladder, M\(_2\) receptors, or a combination of M\(_2\) and M\(_3\) mediated contractile responses and the two types of receptor seemed to act in a facilitatory manner to mediate contraction (Braverman, Tallarida, and Ruggieri, 2002). In obstructed, hypertrophied rat bladders, there was an increase in total and M\(_2\) receptor density, whereas there was a reduction in M\(_3\) receptor density (Braverman and Ruggieri, 2003). The functional significance of this change for voiding function has not been established. Pontari et al (2004) analyzed bladder muscle specimens from patients with neurogenic bladder dysfunction to determine whether the muscarinic receptor subtype mediating contraction shifts from M\(_3\) to the M\(_2\) receptor subtype, as found in the denervated, hypertrophied rat bladder. They concluded that whereas normal detrusor contractions are mediated by the M\(_3\) receptor subtype, in patients with neurogenic bladder dysfunction contractions can be mediated by the M\(_2\) receptors.

Muscarinic receptors may also be located on the presynaptic nerve terminals and participate in the regulation of transmitter release. The inhibitory prejunctional muscarinic receptors are classified as M\(_1\), in the rabbit and rat, and M\(_4\) in the guinea pig, rat, and human bladder. Prejunctional facilitatory muscarinic receptors appear to be of the M\(_3\) subtype in the rat and rabbit urinary bladder (Anderson and Arner, 2004). Prejunctional muscarinic facilitation has also been detected in human bladders. The muscarinic facilitatory mechanism seems to be upregulated in overactive bladders from chronic spinal cord-transected rats. The facilitation in these preparations is primarily mediated by M\(_1\) muscarinic receptors (Somogyi et al, 2003).

Muscarinic receptors have also been demonstrated on the urothelium and in the suburothelium (Chess-Williams, 2002; Mansfield et al, 2005; Schleipfer et al, 2007), but their functional importance has not been clarified. It has been suggested that they may be involved in the release of an unknown inhibitory factor (Chess-Williams, 2002), or they may be directly involved in afferent signaling, and thus a target for antimuscarinic agents, explaining part of the efficacy of these drugs in DO/OAB (Andersson and Yoshida, 2003; Andersson, 2004; Kim et al, 2005; Yokoyama et al, 2005).

### Adrenergic Receptors

#### A. Alpha-ARs

Most investigators agree on that there is a low expression of α-ARs in the human detrusor (Michel, 2006). Malloy et al (1998) found that two-thirds of the α-AR mRNA expressed was α\(_{1D}\), and one-third was α\(_{1A}\) (there was no α\(_{1B}\)). It has been suggested that a change of subtype distribution may be produced by outflow obstruction. Hampel et al (2002) reported that in the rat there was a change in the obstructed bladder from α\(_{1A}\)-AR to α\(_{1D}\)-AR mRNA predominance. In humans, there is an α\(_{1D}\)-AR predominance already in the normal bladder, which means that a change in a similar direction as in the rat would be of minor importance provided that the number or function of the receptors does not increase. Nomiyama and Yamaguchi (2003) confirmed the low expression of α-AR mRNA in normal human detrusor, and further demonstrated that there was no upregulation of any of the adrenergic receptors with obstruction. In addition, in functional experiments they found a small response to phenylephrine at high drug concentrations with no difference between normal and obstructed bladders. Overall, in the obstructed human bladder, there seems to be no evidence for α-AR upregulation or change in subtype, although this finding was challenged by Bouchelouche et al (2005), who found an increased response to α\(_{1A}\)-AR stimulation in obstructed bladders. Whether or not this would mean that the α\(_{1D}\)-ARs in the detrusor muscle are responsible for DO or OAB is unclear. Chen et al (2005) studied micturition in α\(_{1D}\)-AR KO mice and clearly showed that these mice have a larger bladder capacity and voided volumes than their wild type controls, supporting an important role for the α\(_{1D}\)-AR in the control of voiding. However, it was not possible to draw any conclusions from their data about the location of the α\(_{1D}\)-ARs receptors involved in micturition control. As discussed by Chen et al (2005), the α\(_{1D}\)-ARs in the detrusor and their levels of expression may not always be relevant for the functional importance of this receptor subtype.

Sugaya et al (2002) investigated the effects of intrathecal tamsulosin (blocking α\(_{1D}\)-ARs) and naftopidil (blocking preferably on α\(_{1A}\)-ARs) on isovolumetric bladder contractions in rats. Intrathecal injection of tamsulosin or naftopidil transiently abolished these contractions. The amplitude of contraction was decreased by naftopidil, but not by tamsulosin. It was speculated that in addition to the antagonistic action of these agents on the α\(_{1A}\)-ARs of prostatic smooth muscle, both agents (especially naftopidil) may also act on the lumbosacral cord (α\(_{1D}\)-ARs). This observation is
of particular interest considering the findings that in the human spinal cord, $\alpha_{1D}$-AR mRNA predominated overall (Smith et al., 1999). Ikemoto et al. (2003) gave tamsulosin and naftopidil to 96 patients with BPH for 8 weeks in a crossover study. Whereas naftopidil monotherapy decreased the I-PSS for storage symptoms, tamsulosin monotherapy decreased the I-PSS for voiding symptoms. However, this difference (which was suggested to depend on differences in affinity for $\alpha_1$-AR subtypes between the drugs) could not be reproduced in a randomized head to head comparison between the drugs (Gotoh et al., 2005). Overall, therefore, based on available evidence, it cannot be concluded that the $\alpha_{1D}$-ARs on the detrusor muscle should be an important target. This does not exclude that $\alpha_{1D}$-ARs located elsewhere in the bladder (ie, the vasculature; 72) or in other structures, would be of importance. All subtypes of $\alpha$-ARs can be found in different parts of the human vascular tree, and they all mediate contraction. The expression varies with vessel bed and increases with age. In the bladder, the function of the detrusor muscle is dependent on the vasculature and the perfusion. Hypoxia induced by partial outlet obstruction is believed to play a major role in both the hypertrophic and degenerative effects of partial outlet obstruction. Das et al. (2002) investigated in rats whether doxazosin affected blood flow to the bladder and reduced the level of bladder dysfunction induced by partial outlet obstruction. They found that 4 weeks’ treatment with doxazosin increased bladder blood flow in both control and obstructed rats. Furthermore, doxazosin treatment reduced the severity of the detrusor response to partial outlet obstruction. Thus, doxazosin could reduce the increase in bladder weight in obstructed animals, which could be one of the mechanisms that contributed to a positive effect on DO caused by the obstruction.

**B. Beta-ARs**

In the human detrusor, it is now generally accepted that the most important $\beta$-AR for bladder relaxation is the $\beta_3$-AR (Yamaguchi, 2002). This can partly explain why the clinical effects of selective $\beta_3$-AR agonists in DO have been controversial and largely inconclusive. On the other hand, the $\beta_2$-AR agonist, clenbuterol, inhibited electrically evoked contractions in human “unstable,” but not normal, bladder, which is in agreement with previous experiences in humans, suggesting that clenbuterol and also other $\beta_2$-AR agonists like terbutaline, may inhibit DO (Andersson and Wein, 2004).

The in vivo effects of $\beta_3$-AR agonists on bladder function have been studied in animal models. It has been shown that when compared with other agents (including antimuscarinic agents), $\beta_3$-AR agonists increase bladder capacity with no change in micturition pressure and the residual volume (Andersson and Wein, 2004). The $\beta_3$-AR seems to be an interesting target for drugs aimed for treatment of DO/OAB. However, proof of concept studies in man are required to show that this is an effective principle to treat DO/OAB.

**Ion Channels**

**C. Calcium Channels**

There is no doubt that an increase in $[Ca^{2+}]_{i}$ is a key process required for the activation of contraction in the detrusor myocyte. However, it is still uncertain whether this increase is due to influx from the extracellular space and/or release from intracellular stores. Furthermore, the importance of each mechanism in different species, and also with respect to the particular transmitter studied, has not been firmly established (Kajioka et al., 2002).

A decrease of the membrane potential (depolarization) increases the open probability for calcium channels, thereby increasing the calcium influx. Thus, the channels dependent on the membrane potential are termed voltage-operated calcium channels (VOCC). Elevated intracellular calcium levels are also believed to initiate release of calcium from intracellular stores (calcium-induced calcium release). Thus, regulation of the intracellular calcium concentration in smooth muscle cells is one conceivable way to modulate bladder contraction. Dihydropyridines, for example, nifedipine, have a potent inhibitory effect on isolated detrusor muscle and clinically in patients with DO (Andersson and Arner, 2004).

Theoretically, inhibition of calcium influx by means of calcium antagonists would be an attractive way of inhibiting DO/OAB. However, there have been few clinical studies of the effects of calcium antagonists in patients with DO. Naglie et al. (2002) evaluated the efficacy of nifedipine for geriatric urgency incontinence in a randomized, double-blind, placebo-controlled crossover trial, and concluded that this treatment was unsuccessful.

Thus, available information does not suggest that systemic therapy with calcium antagonists is an effective way to treat DO/OAB (Andersson and Wein, 2004).

**D. Potassium Channels**

Potassium channels represent another mechanism to modulate the excitability of the smooth muscle cells. There are several different types of $K^+$-channels and at least 2 subtypes have been found in the human detrusor: ATP-sensitive $K^+$-channels ($K_{ATP}$) and large conductance calcium-activated $K^+$-channels ($BK_{Ca}$). Studies on isolated human detrusor muscle and on bladder tissue from several animal species have demonstrated that $K^+$-channel openers reduce spontaneous contractions as well as contractions induced by carbachol and electrical stimulus. However, the lack of selectivity of presently available $K^+$-channel blockers for the bladder versus the vasculature has thus far limited the use of these drugs. The first generation of $K_{ATP}$ channel openers, such as cromakalim and pinacidil, were found to be more potent as inhibitors of vascular smooth muscle than of detrusor mus-
cle (Andersson and Arner, 2004). No effects of cromakalim or pinacidil on the bladder were found in studies on patients with spinal cord lesions or detrusor instability secondary to outflow obstruction. Also with more recently developed K<sub>ATP</sub> channel openers, claimed to have selectivity toward the bladder, negative results have been obtained in an RTC on patients with idiopathic DO (Chapple, Patroneva, and Raines, 2006).

Thus, at present there is no clinical evidence to suggest that K<sup>+</sup> channel openers represent a treatment alternative for DO/OAB (Andersson and Wein, 2004).

**Vanilloid Receptors**

Appropriate bladder function is dependent on an intact afferent signaling from the bladder to the CNS. This signaling conveys information about bladder filling and the status of the tissue, for example, presence of infectious agents and so on. The afferent nerves consist of small slowly conducting myelinated Aδ-fibers and slowly conducting unmyelinated C-fibers. The former are excited by mechanoreceptors and convey information about bladder filling, while C-fibers mediate painful sensations recognized by chemoreceptors (Candenas, Lecci, and Pinto, 2005).

By means of CAP, a subpopulation of primary afferent neurons innervating the bladder and urethra, the “CAP-sensitive nerves,” has been identified. It is believed that CAP exerts its effects by acting on specific “vanilloid” receptors (TPVR1), on these nerves. CAP exerts a biphasic effect: initial excitation is followed by a long-lasting blockade, which renders sensitive primary afferents (C-fibers) resistant to activation by natural stimuli. In sufficiently high concentrations, CAP is believed to cause “desensitization” initially by releasing and emptying the stores of neuropeptides, and then by blocking further release. Resiniferatoxin (RTX) is an analogue of CAP, approximately 1000 times more potent for desensitization than CAP, but only a few hundred times more potent for excitation. Possibly, both CAP and RTX can have effects on Aδ-fibers. It is also possible that CAP at high concentrations (mM) has additional nonspecific effects.

The rationale for intravesical instillations of vanilloids is based on the involvement of C-fibers in the pathophysiology of conditions such as bladder hypersensitivity and neurogenic DO. In the healthy human bladder, C-fibers carry the response to noxious stimuli, but they are not implicated in the normal voiding reflex. After spinal cord injury, major neuroplasticity appears within bladder afferents in several mammalian species, including man (de Groat and Yoshimura, 2006). C-fiber bladder afferents proliferate within the suburothelium and become sensitive to bladder distention. Those changes lead to the emergence of a new C-fiber-mediated voiding reflex, which is strongly involved in spinal neurogenic DO. Improvement of this condition by defunctionalization of C-fiber bladder afferents with intravesical vanilloids has been widely demonstrated in humans and animals.

Cystometric evidence that CAP-sensitive nerves may modulate the afferent branch of the micturition reflex in humans was originally presented by Maggi et al (1989), who instilled CAP (0.1–10 µM) intravesically in 5 patients with hypersensitivity disorders with attenuation of their symptoms a few days after administration. Intravesical CAP, given in considerably higher concentrations (1–2 mM) than those administered by Maggi et al (1989), has since been used with success in neurological disorders such as multiple sclerosis or traumatic chronic spinal lesions. Side effects of intravesical CAP include discomfort and a burning sensation at the pubic/urethral level during instillation, an effect that can be overcome by prior instillation of lidocaine, which does not interfere with the beneficial effects of CAP. No premalignant or malignant changes in the bladder have been found in biopsies of patients who had repeated CAP instillations for up to 5 years.

The beneficial effect of CAP and RTX has been demonstrated in several studies including RCTs (Lazzeri et al, 2004; Reitz and Schurch, 2004; Silva et al, 2005).

Available information (including data from randomized controlled trials) suggests that both CAP and RTX may have useful effects in the treatment of neurogenic DO. There may be beneficial effects also in non-neurogenic DO in selected cases refractory to antimuscarinic treatment. RTX is an interesting alternative to CAP, but the clinical development of the drug is hampered by formulation problems.

**Botulinum Toxin-Sensitive Mechanisms**

Seven immunologically distinct antigenic subtypes of botulinum toxin (BTX) have been identified: A, B, C1, D, E, F, and G. Types A and B are in clinical use in urology, but most studies have been performed with BTX A type. BTX is believed to act mainly by inhibiting acetylcholine release from cholinergic nerve terminals interacting with the protein complex necessary for docking acetylcholine vesicles, but the mechanism of action may be more complex (Yokoyama et al, 2002; Smith et al, 2003; Simpson, 2004; Apostolidis, Dasgupta, and Fowler, 2006). Apostolidis et al (2006) proposed that a primary peripheral effect of BTX is “the inhibition of release of acetylcholine, ATP, substance P, and reduction in the axonal expression of the CAP and purinergic receptors. This may be followed by central desensitization through a decrease in central uptake of substance P and neurotrophic factors.”

The BTX-produced chemical denervation is a reversible process and axons are regenerated in about 3–6 months. Given in adequate amounts BTX inhibits release not only of acetylcholine, but also of several other transmitters. The BTX molecule cannot cross the blood-brain barrier and therefore has no CNS effects.
BTX injected into the external urethral sphincter was initially used to treat spinal cord injured patients with detrusor-external sphincter dyssynergia (Yokoyama et al, 2002; Smith et al, 2003). The use of BTX has increased rapidly, and successful treatment of neurogenic DO by intravesical BTX injections has now been reported by several groups (Leippold, Reitz, and Schurch, 2003; Cruz and Silva, 2004; Sahai et al, 2005). BTX may also be an alternative to surgery in children with intractable OAB (Schurch and Corcos, 2005). However, toxin injections may also be effective in refractory idiopathic DO (Rapp et al, 2004). Adverse effects, for example, generalized muscle weakness, have been reported (De Laet and Wyndale, 2005), but seem to be rare.

**CONCLUSIONS**

To effectively control bladder activity, and to treat urinary incontinence caused by DO/OAB, identification of suitable targets for pharmacological intervention is necessary. Such targets may be found in the CNS or peripherally. Drugs specifically directed for control of bladder activity are under development and will hopefully lead to improvement of the present treatment of urinary incontinence.

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The urinary bladder is probably the only visceral smooth-muscle organ that is under complete voluntary control from the cerebral cortex. Normal bladder function requires coordinated interaction of sensory and motor components of both the somatic and autonomic nervous systems. Because many levels of the nervous system are involved in the regulation of voiding function, neurologic diseases often cause changes in bladder function. Examples are multiple sclerosis, spinal cord injury, cerebrovascular disease, Parkinson disease, diabetes mellitus, meningomyelocele, and amyotrophic lateral sclerosis. Injury to the sacral roots or pelvic plexus from spinal surgery, herniation of an intervertebral disk, or pelvic surgery (hysterectomy, abdominoperineal resection) can also cause neuropathic bladder.

Significant bladder dysfunction may occur as a result of poor voiding habits in childhood or of degenerative changes in bladder muscle and nerve endings caused by aging, inflammation, or anxiety disorders. All the above conditions can disrupt efficient reflex coordination between sphincter and bladder, and with time, this leads to symptomatic dysfunction.

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**NORMAL VESICAL FUNCTION**

**ANATOMY & PHYSIOLOGY**

**The Bladder Unit**

The bladder wall is composed of a syncytium of smooth-muscle fibers that run in various directions; near the internal meatus, however, 3 layers are distinguishable: a middle circular layer and inner and outer longitudinal layers. In females the outer layer extends down the entire length of the urethra, while in males it ends at the apex of the prostate. The muscle fibers become circular and spirally oriented around the urethra junction and thus function as part of the smooth-muscle sphincter. The middle circular layer ends at the internal meatus of the bladder and is most developed anteriorly. The inner layer remains longitudinally oriented and reaches the distal end of the urethra in females and the apex of the prostate in males. The convergence of these muscle fibers forms a thickened bladder neck, which functions as the internal sphincter.

The normal bladder is able to distend gradually to a capacity of 400–500 mL without appreciable increase in intravesical pressure. When the sensation of fullness is transmitted to the sacral cord, the motor arc of the reflex causes a powerful and sustained detrusor contraction and urination if voluntary control is lacking (as in infants). As myelination of the central nervous system progresses, the young child is able to suppress the sacral reflex so that he or she can urinate when it is appropriate.

The functional features of the bladder include (1) a normal capacity of 400–500 mL, (2) a sensation of fullness, (3) the ability to accommodate various volumes without a change in intraluminal pressure, (4) the ability to initiate and sustain a contraction until the bladder is empty, and (5) voluntary initiation or inhibition of voiding despite the involuntary nature of the organ.

**The Sphincteric Unit**

In both males and females, there are 2 sphincteric elements: (1) an internal involuntary smooth-muscle sphincter at the bladder neck, and (2) an external voluntary striated-muscle sphincter from the prostate to the membranous urethra in males and at the mid urethra in females.

The bladder neck sphincter is a condensation of smooth muscle of the detrusor. This area is rich in sympathetic innervation. Active contraction of the bladder neck region occurs simultaneously with seminal emission, just before ejaculation. In the filling phase, the bladder neck remains closed to provide continence. It opens during both spontaneous contraction and contraction induced by stimulation of the pelvic nerve, indicating that it can be pulled open by contraction of the longitudinal muscles.

The external sphincter is composed of slow-twitch, small striated muscle fibers. This voluntary sphincter maintains a constant tonus and is the primary continence mechanism. While the resting tone is maintained by the slow-twitch intrinsic muscle, it can be voluntarily increased by contraction of the striated muscles of the pelvic floor (eg, levator ani), which contain both fast- and slow-twitch fibers. The levator muscles also contribute indirectly to continence through support of the bladder base. Weakness of the pelvic floor may impair the efficiency of the closure
mechanism of the otherwise normal bladder and sphincter. During abdominal straining, the diaphragm and abdominal muscles contract, and the increased abdominal pressure is transmitted to add to the intravesical pressure. A reflex contraction of the pelvic musculature together with transmitted abdominal pressure helps to close the urethra and prevent stress incontinence.

Relaxation of the sphincter is mostly a voluntary act without which voiding is normally inhibited. Failure to initiate sphincteric relaxation is a mechanism of urinary retention often seen in children with dyssynergic voiding. In infancy, the detrusor behaves in an uninhibited fashion. As the central nervous system matures, children learn to suppress or enhance the micturition reflex through voluntary contraction or relaxation of the pelvic musculature.

The Ureterovesical Junction

The function of the ureterovesical junction is to prevent backflow of urine from the bladder to the upper urinary tract. Longitudinal muscle from the ureter contributes to the makeup of the trigone. Stretching of the trigone has an occlusive effect on the ureteral openings. During normal detrusor contraction, the increased pull on the ureters prevents reflux of urine. Conversely, the combination of detrusor hypertrophy and trigonal stretch owing to residual urine can significantly obstruct the flow of urine from the ureters into the bladder.

INNERVATION & NEUROPHYSIOLOGY

Nerve Supply

The lower urinary tract receives afferent and efferent innervation from both the autonomic and somatic nervous systems. The parasympathetic innervation originates in the second to fourth sacral segments. The cholinergic postganglionic fibers supply both the bladder and smooth-muscle sphincter. The sympathetic nerves originate at T10–L2. The noradrenergic postganglionic fibers innervate the smooth muscles of the bladder base, internal sphincter, and proximal urethra. Somatic motor innervation originates in S2–3 and travels to the striated urethral sphincter via the pudendal nerve. Some motor neurons to the tonic small muscle fibers of the striated sphincter may also project through the pelvic nerve (Gosling and Dixon, 1990).

There are both somatic and visceral afferents from the bladder and urethra. The somatic afferent is carried by the pudendal nerve, while the visceral afferent projects through the sympathetic and parasympathetic nerves to their respective spinal areas.

The normal afferent pathway is mediated largely by Aδ-fibers, which send information about the state of bladder fullness to the pontine micturition center. After spinal disruption, a different type of afferent pathway emerges, mediated by capsaicin-sensitive C-fibers that drive a spinal segmental reflex pathway, causing neurogenic detrusor overactivity. The common sources of afferent information for either pathway are likely to be afferents from the urothelium, lamina propria, and afferents that originate in the bladder wall (Fowler, 2002). On the other hand, the thoracolumbar visceral afferents may transmit discomfort and pain (Janig and Koltzenburg, 1993).

The Micturition Reflex

Intact reflex pathways via the spinal cord and the pons are required for normal micturition. Afferents from the bladder are essential for the activation of the sacral center, which then causes detrusor contraction, bladder neck opening, and sphincteric relaxation. The pontine center, through its connection with the sacral center, may send either excitatory or inhibitory impulses to regulate the micturition reflex. Electrical or chemical stimulation of the neurons in the medial pontine micturition center generates contraction of the detrusor and relaxation of the external sphincter. Disruption of pontine control, as in upper spinal cord injury, leads to contraction of the detrusor without sphincteric relaxation (detrusor-sphincter dyssynergia).

In pathologic conditions affecting the urethra (eg, urethritis or prostatitis) or the bladder (eg, cystitis or obstructive hypertrophy), uninhibited detrusor contraction may occur because of facilitation of the micturition reflex (Figure 27–1).

The Storage Function

The external sphincter plays an important role in urine storage. The afferents from pelvic and pudendal nerves activate both the sacral and lateral pontine nerves; this enhances sphincteric contraction while suppressing the parasympathetic impulse to the detrusor. Voluntary tightening of the sphincter can also inhibit the urge to urinate. In addition, activation of sympathetic nerves increases urethral resistance and facilitates bladder storage (Figure 27–2).

Cerebral (Suprapontine) Control

Although micturition and urine storage are primarily functions of the autonomic nervous system, these are under voluntary control from suprapontine cerebral centers, so that other groups of muscles (arm, leg, hand, bulbocavernosus) can be integrated to assist in urination at the appropriate time and place. Cerebral lesions (eg, from tumor, Parkinson’s disease, vascular accident) are known to affect the perception of bladder sensation and result in voiding dysfunction.

Neurotransmitters & Receptors

In parasympathetic innervation, acetylcholine and nicotinic receptors mediate pre- to postganglionic transmis-
sion, while acetylcholine and M2 muscarinic receptors mediate the postganglionic neuron–smooth muscle transmission. In some species, adenosine triphosphate (ATP) is released with acetylcholine and acts on purinoceptors (P2) in the smooth-muscle cell. In sympathetic nerves, noradrenaline can act on the beta-2-adrenoreceptors to relax the detrusor or the alpha-1-receptors to contract the bladder neck and the external sphincter.

In addition, many neuropeptides, which usually colocalize with the classic transmitters, are also found in the genitourinary tract. Neuropeptide Y, encephalin, and vasoactive intestinal polypeptide (VIP) are found in cholinergic postganglionic neurons, while calcitonin gene-related peptide (CGRP), VIP, substance P, cholecystokinin, and encephalins are distributed in sacral visceral afferent fibers. These peptides are thought to be involved in modulation of efferent and afferent neurotransmissions.

**URODYNAMIC STUDIES (SEE ALSO CHAPTER 28)**

**Micturition**

Urodynamic studies are techniques used to obtain graphic recordings of activity in the urinary bladder, urethral sphincter, and pelvic musculature. The 3 current methods involve use of gas or water to transfer pressure to a transducer housed near a polygraph or use of a transducer-tipped catheter to transfer pressure recordings directly to a polygraph. All the techniques have limitations, with gas being the least reliable of the three. Pressure recordings may be complemented by elec-
tromyography of the perineal musculature, ultrasound, or radiography (Figure 27–3).

**Uroflowmetry**

Uroflowmetry is the study of the flow of urine from the urethra. Uroflowmetry is best performed separately from all other tests and, whenever possible, as a standard office screening or monitoring procedure. The normal peak flow rate for males is 20–25 mL/s and for females 20–30 mL/s. Lower flow rates suggest outlet obstruction or a weak detrusor; higher flow rates suggest bladder spasticity or excessive use of abdominal muscles to assist voiding. Intermittent flow patterns generally reflect spasticity of the sphincter or straining to overcome resistance in the urethra.

**Cystometry**

Cystometry is the urodynamic evaluation of the reservoir function of the bladder. Cystometry is most informative when combined with studies of the external urethral sphincter and pelvic floor.

Normal bladder capacity is 400–500 mL. Bladder pressure during filling should remain low up to the point of voiding. The first desire to void is generally felt when the volume reaches 150–250 mL, but detrusor filling pressure should remain unchanged until there is a definite sense of fullness at 350–450 mL, the true capacity of the bladder. Detrusor contractions before this point are considered abnormal and the result of hyperreflexic or uninhibited behavior. Normal voiding pressures in the bladder should not rise above 30 cm of water pressure. With normal voiding there should not be any residual urine, and voiding should be accomplished without straining.

**Urethral Pressure Recordings**

Normal voiding requires a synergic action of the bladder (contraction) and urethra (relaxation). High pressures in the bladder during voiding reflect abnormal resistance in the urethral outlet. Increased outlet resistance can result from benign prostatic hypertrophy, urethral stricture, bladder neck contracture, or spasm of the external urethral sphincter. Low resistance in the urethral outlet generally reflects compromised function of the sphincter mechanism. Recording of urethral pressures with the bladder at rest as well as during contraction helps determine the presence of functional or anatomic disorders.

**Electromyography**

With electromyography, the activity of the striated urethral muscles can be monitored without obstructing the urethral lumen. In the normal urethra, activity increases slightly as the bladder fills and falls precipitously just before voiding begins. Denervation results in an overall decrease...
in activity as well as production of denervation potentials. An overall increase in activity reflects a state of hyperreflexia. The technique provides a sensitive assessment of urethral and pelvic muscle behavior. The disadvantages of the technique are dependence on accurate needle position and a tendency to record artifacts.

**ABNORMAL VESICAL FUNCTION**

**CLASSIFICATION OF NEUROPATHIC BLADDER**

The traditional classification was according to neurologic deficit. Thus, the terms motor, spastic, upper motor neuron, reflexic, and uninhibited were used to describe dysfunction found with injury above the spinal cord micturition center. Coordination between the bladder and the sphincter was considered either balanced or unbalanced. The terms flaccid, atonic, areflexic, and sensory were used to describe loss of ability of the bladder to contract owing to injury of the pelvic nerves or the spinal micturition center. Dysfunction with both types of features was described as mixed.

Descriptions of neuromuscular dysfunction of the lower urinary tract should be individualized because no 2 neural injuries (no matter how similar) result in the same type of dysfunction. The standardization committee for the International Continence Society has attempted to create a functional classification that is easy to understand and provides a simple basis for therapy (Table 27–1).

### 1. Neuropathic Bladder Due to Lesions Above the Sacral Micturition Center

Most lesions above the level of the cord where the micturition center is located will cause bladder spasticity. Sacral reflex arcs remain intact, but loss of inhibition from higher centers results in spastic bladder and sphincter behavior on the segmental level. The degree of spasticity varies between the bladder and sphincter, from lesion to lesion, and from patient to patient with similar lesions. Common lesions found above the brain stem that affect voiding include dementia, vascular accidents, multiple sclerosis, tumors, and inflammatory disorders such as encephalitis or meningitis. These lesions can produce a wide range of functional changes, including precipitate urge, frequency, residual urine, retention of urine, recurrent urinary tract infections, and gross incontinence. Symptoms range from mild to disabling. Obviously, incontinence is especially troublesome. If the lesion is above the pontine micturition center, detrusor–striated sphincter dyssynergia usually does not occur. However, leakage may occur because the need to void cannot be felt or because the sphincter becomes more relaxed and can no longer inhibit spontaneous voiding. Lesions of the internal capsule include vascular accidents and Parkinson’s disease. Both spastic and semiflaccid voiding disorders are found with these lesions.

Spinal cord injury can be the result of trauma, herniated intervertebral disk, vascular lesions, multiple sclerosis, tumor, syringomyelia, or myelitis, or it may be iatrogenic. Traumatic spinal cord lesions are of greatest clinical concern. Partial or complete injuries may cause equally severe genitourinary dysfunction. Sphincter spasticity and voiding dyssynergia can lead to detrusor hypertrophy, high voiding pressures, ureteral reflux, or ureteral obstruction. With time, renal function may become compromised. If infection is combined with back pressure on the kidney, loss of renal function can be particularly rapid.

Spinal cord injuries at the cervical level are often associated with a condition known as autonomic dysreflexia.

### Table 27–1. Various Classifications of Neuropathic Bladder.

<table>
<thead>
<tr>
<th><strong>International Continence Society</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Detrusor: Normal (N), hyperreflexic (+), hyporeflexic (–)</td>
</tr>
<tr>
<td>Striated sphincter: Normal (N), hyperactive (+), incompetent (–)</td>
</tr>
<tr>
<td>Sensation: Normal (N), hypersensitive (+), hyposensitive (–)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Bors and Comarr</strong></th>
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<tbody>
<tr>
<td>Sensory neuron lesion</td>
</tr>
<tr>
<td>Motor neuron lesion (balanced or unbalanced)</td>
</tr>
<tr>
<td>Sensory motor neuron lesion</td>
</tr>
<tr>
<td>Upper motor neuron lesion</td>
</tr>
<tr>
<td>Lower motor neuron lesion</td>
</tr>
<tr>
<td>Mixed upper and lower motor neuron lesion</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Nesbit, Lapides, and Baum</strong></th>
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<tbody>
<tr>
<td>Sensory neuron lesion</td>
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<tr>
<td>Motor neuron lesion</td>
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<tr>
<td>Uninhibited bladder</td>
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<tr>
<td>Reflex bladder</td>
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<tr>
<td>Autonomous bladder</td>
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<table>
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<tr>
<th><strong>Krane</strong></th>
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<tbody>
<tr>
<td>Detrusor hyperreflexia</td>
</tr>
<tr>
<td>Coordinated sphincters</td>
</tr>
<tr>
<td>Striated muscle dyssynergia</td>
</tr>
<tr>
<td>Smooth muscle dyssynergia</td>
</tr>
<tr>
<td>Detrusor areflexia</td>
</tr>
<tr>
<td>Coordinated sphincters</td>
</tr>
<tr>
<td>Nonrelaxing striated sphincter</td>
</tr>
<tr>
<td>Denervated striated sphincter</td>
</tr>
<tr>
<td>Nonrelaxing smooth sphincter</td>
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<table>
<thead>
<tr>
<th><strong>Wain, Benson, and Raezer</strong></th>
</tr>
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<tbody>
<tr>
<td>Failure to empty</td>
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<tr>
<td>Failure to store</td>
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</table>
Because the lesions occur above the sympathetic outflow from the cord, hypertensive blood pressure fluctuations, bradycardia, and sweating can be triggered by insertion of a catheter, mild overdistention of the bladder with filling, or dysynergic voiding.

In summary, the spastic neuropathic bladder is typified by (1) reduced capacity, (2) involuntary detrusor contractions, (3) high intravesical voiding pressures, (4) marked hypertrophy of the bladder wall, (5) spasticity of the pelvic-striated muscle, and (6) autonomic dysreflexia in cervical cord lesions.

2. Neuropathic Bladder Due to Lesions at or Below the Sacral Micturition Center

Injury to the Detrusor Motor Nucleus

The most common cause of flaccid neuropathic bladder is injury to the spinal cord at the micturition center, S2–4. Other causes of anterior horn cell damage include infection due to poliovirus or herpes zoster and iatrogenic factors such as radiation or surgery. Herniated disks can injure the micturition center but more commonly affect the cauda equina or sacral nerve roots. Myelodysplasias could also be grouped here, but the mechanism is actually failure in the development or organization of the anterior horn cells. Lesions in this region of the cord are often incomplete, with the result commonly being a mixture of spastic behavior with weakened muscle contractility. Mild trabeculation of the bladder may occur. External sphincter and perineal muscle tone are diminished. Urinary incontinence usually does not occur in these cases because of the compensatory increase in bladder storage. Because pressure in the bladder is low, little outlet resistance is needed to provide continence. Evacuation of the bladder may be accomplished by straining, but with variable success.

Injury to the Afferent Feedback Pathways

Flaccid neuropathic bladder also results from a variety of neuropathies, including diabetes mellitus, tabes dorsalis, pernicious anemia, and posterior spinal cord lesions. Here the mechanism is not injury of the detrusor motor nucleus but a loss of sensory input to the detrusor nucleus or a change in motor behavior due to loss of neurotransmission in the dorsal horns of the cord. The end result is the same. Loss of perception of bladder filling permits overstretching of the detrusor. Atony of the detrusor results in weak, inefficient contractility. Capacity is increased and residual urine significant.

In summary, the flaccid neuropathic bladder is typified by (1) large capacity, (2) lack of voluntary detrusor contractions, (3) low intravesical pressure, (4) mild trabeculation (hypertrophy) of the bladder wall, and (5) decreased tone of the external sphincter.

3. Injury Causing Poor Detrusor Distensibility

Another cause of atonic neuropathic bladder is peripheral nerve injury. This category includes injury caused by radical surgical procedures such as low anterior resection of the colon or Wertheim hysterectomy. This type of dysfunction has been referred to as autonomous because the smooth muscle remains active but there is no central reflex to organize muscle activity. The end result is a bladder that stores poorly owing to failure to accommodate with filling. There is a rather steep pressure rise in the bladder with filling due to hypertonicity in the detrusor wall.

Radiation therapy can result in denervation of the detrusor or the sphincter. More commonly, it damages the detrusor, resulting in fibrosis and loss of distensibility. Other inflammatory causes of injury to the detrusor include chronic infection, interstitial cystitis, and carcinoma in situ. These lesions produce a fibrotic bladder wall that, for obvious reasons, distends poorly.

Selective Injury to the External Sphincter

Pelvic fracture often tears the nerves to the external sphincter. Selective denervation of the external sphincter muscle, with incontinence, can follow if the bladder neck is not sufficiently competent. Radical surgery in the perineum is highly unlikely to damage the motor innervation to the urethra, although sensory innervation of the external sphincter can be affected.

4. Injury to the Spinal Cord

Spinal shock may last from a few weeks to 6 months (usually 2–3 months). Reflex response in striated muscle is usually present from the time of injury but is suppressed. With time, the reflex excitability of striated muscle progresses until a spastic state is achieved. Smooth muscle is much slower to develop this hyperreflex activity and, unlike striated muscle, loses spontaneous response after the injury. Urinary retention is the rule, therefore, in the early months following injury.

Urodynamic studies are indicated periodically to monitor the progressive return of reflex behavior. In the early recovery stages, a few weak contractions of the bladder may be found. Later, in injuries above the micturition center, more significant reflex activity will be found. Low-pressure storage can be managed via intermittent catheter-
1. Spastic Neuropathic Bladder

Spastic neuropathic bladder results from partial or extensive neural damage above the conus medullaris (T12). The bladder functions on the level of segmental reflexes, without efficient regulation from higher brain centers.
NEUROPATHIC BLADDER DISORDERS

D. X-RAY FINDINGS

Periodic excretory urograms and retrograde cystograms are essential because complications are common. A trabeculated bladder of small capacity is typical of this type of neuropathic dysfunction. The bladder neck may be dilated. The kidneys may show evidence of pyelonephritic scarring, hydronephrosis, or stone disease. The ureters may be dilated from obstruction or reflux. A voiding film often clearly outlines a narrowed zone created by the spastic sphincter but may also identify a strictured segment of the urethra. Most, if not all, of these features can be detected with ultrasound. Magnetic resonance imaging is especially useful for the sagittal view it offers of the bladder neck and posterior urethral zones.

E. INSTRUMENTAL EXAMINATION

Cystoscopy and panendoscopy help assess the integrity of the urethra and identify stricture sites. The bladder shows variable degrees of trabeculation, occasionally with diverticula. Bladder capacity, stones, competency of the ureteral orifices, changes secondary to chronic infection or indwelling catheters, and the integrity of the bladder neck and external urethral sphincter can be assessed.

F. URODYNAMIC STUDIES

Combined recording of bladder and urethral sphincter activity during filling will reveal a low-volume bladder with spastic dyssynergy of the external sphincter (Figure 27–4). High voiding pressures in the bladder are not unusual. Ureretal reflux or obstruction is more likely if voiding pressures exceed 40 cm of water. A high resting pressure is noted in the external sphincter on the urethral pressure profile, and labile spastic behavior is noted during filling and voiding. Various auras replace a true sense of bladder filling, for example, sweating, vague abdominal discomfort, spasm of the lower extremities. Movement of a catheter in the urethra can trigger detrusor contraction and voiding.

2. Mildly Spastic Neuromuscular Dysfunction

Incomplete lesions of the cerebral cortex, pyramidal tracts, or spinal cord may weaken, but not abolish, cerebral restraint. The patient may have frequency and nocturia or urinary incontinence due to precipitous urge or voiding. Common causes include brain tumors, Parkinson’s disease, multiple sclerosis, dementias, cerebrovascular accidents, prolapsed disks, or partial spinal injury.

In many cases, the cause is unclear. The hyperreflexic behavior often seems to be associated with a peripheral abnormality (eg, prostatitis, benign prostatic hypertrophy, urethritis) or follows pelvic surgery (eg, anterior colporrhaphy, anteroposterior tumor resection). Symptoms are commonly associated with psychological factors.

Clinical Findings

A. SYMPTOMS

Frequency, nocturia, and urgency are the principal symptoms. Hesitancy, intermittency, double voiding, and residual urine are also common. Incontinence may vary from pre- or postvoiding dribbling to complete voiding that the patient does not recognize or cannot inhibit once started.

B. SIGNS

The degree of voiding dysfunction does not parallel neurologic deficits. Slight physical disabilities can be associated with gross disturbances in bladder function, and the reverse is also true. However, it is always important to check lower extremity and perineal reflexes for evidence of hyperreflexia. Sensory or motor deficits may also be detected in the lumbar or sacral segments.

C. X-RAY FINDINGS

In the early stages, radiologically evident change is minimal for the most part in both the lower and upper urinary tracts. Low bladder volume and mild trabeculation of the bladder are usually evident.

D. INSTRUMENTAL EXAMINATION

Cystoscopy and urethroscopy are generally unremarkable. Mild detrusor and sphincter irritability and diminished capacity can be demonstrated.

E. URODYNAMIC STUDIES

The behavior patterns of the sphincter and bladder are similar to those of the previous group but on a milder scale. Uninhibited detrusor activity, evident urodynamically, may not be associated with the same symptom pattern on the clinical level. The patient occasionally perceives a sense of urgency and the need to void. However, these sensations may not be present, and the patient may complain of the actual leakage as the main inconvenience.
Morphologic changes in the bladder are slight, with changes in the upper urinary tract occurring rarely and late because of lower pressures in the bladder.

### 3. Flaccid (Atonic) Bladder

Direct injury to the peripheral innervation of the bladder or sacral cord segments S2–4 results in flaccid paralysis of the urinary bladder. Characteristically, the capacity is large, intravesical pressure low, and involuntary contractions absent. Because smooth muscle is intrinsically active, fine trabeculations in the bladder may be seen. Common causes of this type of bladder behavior are trauma, tumors, tabes dorsalis, and congenital anomalies (eg, spina bifida, meningomyelecele).

### Clinical Findings

#### A. Symptoms

The patient experiences flaccid paralysis and loss of sensation affecting the muscles and dermatomes below the level of injury. The principal urinary symptom is retention with overflow incontinence. Male patients lose their erections. Surprisingly, despite weakness in the striated sphincter, neither bowel nor urinary incontinence is a major factor. Storage pressures within the bladder remain below the outlet resistance.

#### B. Signs

Neurologic changes are typically lower motor neuron. Extremity reflexes are hypoactive or absent. Sensation is diminished or absent. It is important to check sensation over the penis (S2) and perianal region (S2–3) for evidence of a mixed or partial injury. Anal tone (S2) should be compared with levator tone (S3–4), again for evidence of a mixed injury. Similarly, sensation over the outside of the foot (S2), sole of the foot (S2–3), and large toe (S3) should be compared for evidence of mixed injury. Occasionally, extremity findings do not parallel those of the perineum, with the pattern being absent sensation and tone in the feet but partial tone or sensation in the perineum. This is especially true in patients who have spina bifida or meningomyelecele.

#### C. Laboratory Findings

Repeated urination at regular intervals is no less important in this group than in others. Infection with white blood cells (leukocytes) and bacteria may occur because of the need for bladder catheterization. Advanced renal change is unusual because bladder storage is under low pressure, but chronic renal failure secondary to pyelonephritis, hydronephrosis, or calculus formation is still possible.

#### D. X-Ray Findings

A plain film of the abdomen may reveal fracture of the lumbar spine or extensive spina bifida. Calcific shadows compatible with urinary stone may be seen. Excretory urograms should be performed initially to check for calculus, hydronephrosis, pyelonephritic scarring, or ureteral obstruction secondary to an overdistended bladder. A cystogram may detect morphologic changes in the detrusor (it is usually large and smooth walled); vesicoureteral reflux may be present. Checks on the integrity of both the lower and upper urinary tracts can subsequently be made using ultrasound.

### E. Instrumental Examination

Visual inspection is performed to rule out pathologic changes (eg, bladder stones, urethral stricture, or ureteral reflux or obstruction).

Cystoscopy and urethroscopy performed some months or weeks after the injury will confirm the laxity and areflexia of the sphincter and pelvic floor; the bladder neck is usually funneled and open and the bladder should be large and smooth walled. The integrity of the ureteral orifices should be normal. Fine trabeculation may be evident.

### F. Urodynamic Studies

The urethral pressure profile reflects low smooth and striated sphincter tone. Bladder filling pressures are low; detrusor contractions are weak or absent; voiding is accomplished by straining or by the Credé maneuver, if at all; and there is a large volume of residual urine. Awareness of filling is markedly diminished and usually results from stretch on the peritoneum or abdominal distention.

### G. Denervation Hypersensitivity

This test is classically performed by giving bethanechol chloride (Urecholine) 15 mg subcutaneously. A cystometrogram is performed after 20 minutes, and the results are compared with the findings obtained before the bethanechol was given. If the results are positive, a rise in filling pressure of more than 15 cm of water is noted, with a shift in the filling curve to the left; the same behavior in the bladder is noted only at a lower filling volume and slightly higher pressure. A finding of no change on filling reflects myogenic damage to the detrusor. A more physiologic way to perform the test is to fill the bladder to about half its capacity, administer urecholine, and monitor for change in storage pressure. The ice water test also checks for detrusor hypersensitivity.

Bethanechol does not facilitate a detrusor contraction; it can only increase tone in the detrusor wall, which in turn might trigger the voiding reflex. The test is not a check on the integrity of the voiding reflex but demonstrates denervation hypersensitivity in flaccid bladders and differentiates this condition from myogenic damage.

The test is not applicable in patients with reduced bladder capacity, decreased compliance (ie, sharp rise in detrusor-filling pressure), or forceful contractions of the detrusor.
DIFFERENTIAL DIAGNOSIS OF NEUROPATHIC BLADDER

The diagnosis of neuropathic bladder is usually obvious from the history and physical examination. Neural impairment is evidenced by abnormal sacral reflex activity and decreased perineal sensation. Some disorders with which neuropathic bladder may be confused are cystitis, chronic urethritis, vesical irritation secondary to psychic disturbance, myogenic damage, interstitial cystitis, cystocele, and infravesical obstruction.

Cystitis

Inflammation of the bladder, both nonspecific and tuberculous, causes frequency of urination and urgency, even to the point of incontinence. Infections secondary to residual urine caused by neuropathic behavioral disturbance should be ruled out.

The urodynamics of the inflamed bladder are similar to those of the uninhibited neuropathic bladder. However, with inflammation, symptoms disappear after definitive antibiotic therapy, and the urodynamic behavior reverts to normal. If symptoms persist or infections return repeatedly, a neuropathic behavioral abnormality should be considered (e.g., multiple sclerosis or even idiopathic detrusor-sphincter dysfunction).

Chronic Urethritis

Symptoms of frequency, nocturia, and burning on urination may be due to chronic inflammation of the urethra not necessarily associated with infection. The urodynamics will show an irritable urethral sphincter zone with labile, spastic tendencies. The cause is unknown.

Vesical Irritation Secondary to Psychic Disturbance

Anxious, tense individuals or those with pathologic psychological fixation on the perineum may present a long history of periodic bouts of urinary frequency or perineal or pelvic pain. The clinical picture and urodynamic findings are similar to those described previously for chronic urethritis. Often, however, if the patient’s anxiety can be allayed, the symptoms subside. The underlying problem is one of excessive pelvic muscle tension and inefficient sphincter behavior. Some of the symptoms may improve with manual therapy of pelvic floor myofascial trigger points (Weiss, 2001).

Interstitial Cystitis

Interstitial cystitis is poorly understood chronic inflammation of the bladder. The typical patient is a woman over 40 years of age, with symptoms of frequency, nocturia, urgency, and suprapubic pain. The symptoms are brought on by bladder distention. Capacity is limited (often <100 mL in the most symptomatic and disabled patients). Urinalysis is normal, and there is no residual urine. Urodynamic studies show a hypertonic, poorly compliant bladder. Distention of the bladder with cystoscopy produces bleeding from petechial hemorrhages and fissuring in the mucosa. The condition represents an end-stage inflammatory process of unknown cause in the detrusor.

Cystocele

Relaxation of the pelvic floor following childbirth may cause some frequency, nocturia, and stress incontinence. Residual urine may be present and predispose to infection. Loss of urine occurs with lifting, standing, or coughing. Pelvic examination usually reveals relaxation of the anterior vaginal wall and descent of the urethra and bladder when the patient strains to void.

Bladder Outlet Obstruction

Urethral strictures, benign or malignant enlargement of the prostate gland, and congenital urethral valves all can produce significant obstruction of the urinary outlet. Hypertrophy (i.e., trabeculation) of the detrusor develops, and residual urine can accumulate. Uninhibited detrusor activity is often found at this stage and resembles that of the spastic neuropathic bladder.

If decompensation occurs, the vesical wall becomes attenuated and atonic, and capacity may be markedly increased. Overflow incontinence may develop. The behavior of the bladder is similar to that of the flaccid neuropathic bladder.

If the difficulty is nonneuropathic, the anal sphincter tone is normal and the bulbocavernous reflex intact. Peripheral sensation, voluntary muscle contraction, and limb reflexes should also be normal. Cystoscopy and urethroscopy reveal the local lesion causing obstruction. Once the obstruction is relieved, bladder function improves but may never return to normal.

TREATMENT OF NEUROPATHIC BLADDER

The treatment of any form of neuropathic bladder is guided by the need to restore low-pressure activity to the bladder. In doing so, renal function is preserved, continence restored, and infection more readily controlled. Reflex evacuation may develop if detrusor integrity is protected and trigger techniques are practiced.

1. Spinal Shock

Following severe injury to the spinal cord, the bladder becomes atonic. With suprasegmental spinal injuries, the
bladder gradually recovers its contractile capabilities within months. A spastic state evolves, the degree of which varies from patient to patient according to level of injury. Injuries to the sacral cord, if complete enough, may leave the bladder permanently flaccid. More often, however, these lesions are partial, and a mixed degree of detrusor-sphincter spasticity is found along with a variable degree of weakness.

During the spinal shock stage, some type of bladder drainage must be instituted immediately and maintained. Chronic overdistention can damage the detrusor smooth muscle and limit functional recovery of the bladder. Intermittent catheterization using strict aseptic technique has proved to be the best form of management. This avoids urinary tract infection as well as the complications of an indwelling catheter (eg, urethral stricture, abscess, erosions, stones).

If a Foley catheter becomes necessary, a few principles need to be followed. The catheter should not be larger than 16F and preferably should be made of silicon, and it should be taped to the abdomen. Taping the catheter to the leg puts unnecessary stress on the penoscrotal junction and bulbous urethra (ie, the curves in the urethra), and this can lead to stricture formation. The catheter should be changed with sterile procedure every 2–3 weeks.

Some urologists advocate the use of suprapubic cystostomy rather than a urethral catheter to avoid the risks associated with permanent indwelling catheters. Certainly, whenever catheter-related complications occur, the physician should not hesitate in resorting to cystostomy drainage.

Irrigation of the bladder with antibiotic solutions, use of systemic antibiotics, or covering the tip of the meatus with antibiotic creams does not significantly lower the long-term risk of bladder infection. Keeping the meatus lubricated does help avoid meatal stricturing, however.

As peripheral reflex excitability gradually returns, urodynamic evaluation should be performed. A cystogram is needed to rule out reflux. The urodynamic study should be repeated every 3 months as long as spasticity is improving and then annually to check for complications of the upper urinary tract.

To control infection, a fluid intake of at least 2–3 L/day should be maintained (100–200 mL/h) if at all possible. This reduces stasis and decreases the concentration of calcium in the urine. Renal and ureteral drainage are enhanced by moving the patient frequently, with ambulation in a wheelchair as soon as possible, and even by raising the head of the bed. These measures improve ureteral transport of urine, reduce stasis, and lower the risk of infection. Additional measures aid in prophylaxis for calculus formation (eg, reduction of intake of calcium and oxalate and elimination of vitamin D in the diet).

2. Specific Types of Neuropathic Bladder

Once a neuropathic voiding disorder is established, regardless of cause, the following steps should be taken to attain optimum function.

Spastic Neuropathic Bladder

A. PATIENT WITH REASONABLE BLADDER CAPACITY

To consider a bladder rehabilitated to a functional state, a patient should be able to go 2–3 hours between voiding and not be incontinent during this interval. Voiding is initiated using trigger techniques—tapping the abdomen suprapublically, rugging on the pubic hair, squeezing the penis, or scratching the skin of the lower abdomen, genitalia, or thighs. Patients can accomplish this on their own unless they are high quadriplegics with no upper limb function.

Some patients in this category can empty the bladder completely but are incontinent due to inconvenient triggering of the voiding reflex. They may be helped by low-dose anticholinergic medication or by neural stimulation.

B. PATIENT WITH MARKEDLY DIMINISHED FUNCTIONAL VESICAL CAPACITY

If the functional capacity of the bladder is under 100 mL, involuntary voiding can occur as often as every 15 minutes. Satisfactory training of the bladder cannot be achieved, and alternative measures must be taken. First, the possibility that reduced functional bladder capacity is due to a large residual volume of urine must be ruled out. One of the following treatment regimens can then be administered.

1. A permanent indwelling catheter with or without anticholinergic medication.

2. A condom catheter and a leg bag in males if residual urine volumes are small and the patient does not have bladder pressures above 40 cm of water on urodynamic evaluation. If either of these parameters is found, the upper urinary tract is considered at risk from obstruction or reflux.

3. Performance of a sphincterotomy in males. It is possible to turn the bladder into a urinary conduit by surgically eliminating all outlet resistance from the bladder. This option should be used only when other options have failed, as it is irreversible. Patients having this procedure usually have more serious sequelae of a highly spastic bladder (ie, upper urinary tract dilatation, recurrent urinary tract infections, or marked autonomic dysreflexia).

4. Conversion of the spastic bladder to a flaccid bladder through sacral rhizotomy. Complete surgical section or percutaneous heat fulguration of the S3 and S4 roots is necessary. Chemical rhizotomy is
unreliable, as spasticity usually returns after 6–9 months. These procedures may cause loss of reflex erections, and the decision to perform them should be weighed accordingly. They can relieve spasticity, lower intravesical pressures, increase bladder storage, and decrease the risk of damage to the upper urinary tract. The bladder would then be managed as a flaccid bladder (see below).

5. Neurostimulation of the sacral nerve roots to accomplish bladder evacuation (see section following).

6. Urinary diversion for irreversible, progressive upper urinary tract deterioration. A variety of procedures are available, including the standard ileal conduit, cutaneous ureterostomies, transureteroureterostomy, or non-refluxing urinary reservoir (eg, Mainz pouch, Koch pouch, or one of several other continent diversions designed to protect the upper urinary tract and kidneys).

7. In females with a spastic bladder, one does not have the option of performing a sphincterotomy. If pharmacologic methods are unsuccessful, surgical conversion to a flaccid, low-pressure system or a urinary diversion should be considered.

C. PARASYMPATHOLYTIC DRUGS

Because of the chronic nature of the neuropathic bladder, patients are not always willing to tolerate the side effects of parasympatholytic drugs. Several drugs in this category can be alternated to reduce the side effects of either drug. They also may be useful when given with skeletal muscle relaxants. Dosages must be individualized. Commonly used drugs and dosages are as follows: oxybutynin chloride (Ditropan), 5 mg 2–3 times daily; Ditropan XL, once daily; dicyclomine hydrochloride (Bentyl), 80 mg in 4 equally divided doses daily; methantheline bromide (Banthine), 50–100 mg every 6 hours; and propantheline bromide (Pro-Banthine), 15 mg 30 minutes before meals and 30 mg at bedtime; and Tolterodine (Detrol), 2 mg 2 times daily. These drugs may not be effective if incontinence is the result of uninhibited sphincter relaxation or compliance changes in the bladder wall.

D. BOTULINUM-A TOXIN

Several centers have investigated injection of 85–300 Units of botulinum-A toxin into 30–40 sites in the detrusor muscle in both children and adults who have detrusor hyperreflexia. The early results are promising, as shown by a significant increase in bladder capacity and compliance as well as symptomatic improvement for several weeks after injection (Schulte-Baukloh, 2002).

E. INTRAVESICAL INSTILLATION OF CAPSAICIN OR RESINIFERATOXIN

Capsaicin and resiniferatoxin are specific C-fiber afferent neurotoxins. After spinal cord injury, C-fiber afferents proliferate in the bladder mucosa and are involved in detrusor hyperreflexia. In a study of 24 spinal cord–injured patients with refractory detrusor hyperreflexia treated with a single dose of 2 mM capsaicin in 30 mL ethanol plus 70 mL of normal saline or 100 nM resiniferatoxin in 100 mL of normal saline, Cruz (1998) found no significant urodynamic or clinical improvement in the capsaicin arm at 30 and 60 days of follow-up. In the resiniferatoxin arm, the mean uninhibited detrusor contraction threshold plus or minus standard deviation increased from 176 ± 54 to 250 ± 107 mL at 30 days (P < .05) and to 275 ± 98 mL at 60 days (P < .01). Mean maximum bladder capacity increased from 196 ± 75 to 365 ± 113 mL at 30 days (P < .001) and to 357 ± 101 mL at 60 days (P < .001). Daily catheterizations and incontinent episodes were significantly decreased at 30 and 60 days of follow-up. Autonomic dysreflexia, limb spasms, suprapubic discomfort, and hematuria developed in most patients who received capsaicin but in none who received resiniferatoxin (Giannantoni et al, 2002).

F. NEUROSTIMULATION (BLADDER PACEMAKER)

Neuroprosthetics are becoming an established alternative to managing selective neuropathic bladder disorders. Patients are evaluated for a bladder pacemaker primarily by urodynamic monitoring of bladder and sphincter responses to trial stimulation of the various sacral nerve roots. Selective blocks are then prepared to the right and left pudendal nerves. If voiding is produced, patients are considered suitable for a neuroprosthesis. Other factors such as detrusor storage capability, sphincter competence, age, kidney function, and overall neurologic and psychological status are also taken into consideration.

Electrodes are implanted on the motor (ventral) nerve roots of those sacral nerve roots that will produce detrusor contraction on stimulation (always S3, occasionally S4). Steps are then taken to reduce sphincter hyperreflexia by selectively dividing the sensory (dorsal) component of these same sacral nerve roots and selective branches of the pudendal nerves. The electrodes are connected to a subcutaneous receiver that can be controlled from outside the body. Bladder or bowel evacuation or continence can then be controlled selectively by the external transmitter.

The first 2 are accomplished by reducing intravesical pressures. This step protects the integrity of the upper urinary tract and restores continence by increasing storage capacity. Both can be achieved by combining neurostimulation of the sphincter with selective sacral neurotomies. This approach preserves sphincter integrity and avoids the need for drugs. Other options include complete bladder denervation or bladder augmentation.

The third goal, restoration of controlled evacuation, eliminates the need for catheters and associated risk of infection. This is the most difficult goal to achieve, and patients need to be carefully evaluated for their suitability.
Flaccid Neuropathic Bladder

If the neurologic lesion completely destroys the micturition center, volitional voiding cannot be accomplished without manual suprapubic pressure, that is, the Credé maneuver. Bladder evacuation can be accomplished by straining, using the abdominal and diaphragmatic muscles to raise intra-abdominal pressures. Partial injuries to the lower spinal cord (T10–11) result in a spastic bladder and a weak or weakly spastic sphincter. Incontinence can then result from spontaneous detrusor contraction.

A. BLADDER TRAINING AND CARE

In partial lower motor neuron injury, voiding should be tried every 2 hours by the clock to avoid embarrassing leakage. This helps protect the bladder from overdistention due to a buildup of residual urine.

B. INTERMITTENT CATHETERIZATION

Any patient with adequate bladder capacity can benefit from regular intermittent catheter drainage every 3–6 hours. This technique eliminates residual urine, helps prevent infection, avoids incontinence, and protects against damage to the upper urinary tract. It simulates normal voiding and is easily learned and adapted by patients. It is an extremely satisfactory solution to the problems of the flaccid neuropathic bladder. A clean technique is used rather than the inconvenient, expensive, sterile technique. Urinary tract infections are infrequent, but if they occur, a prophylactic antibiotic can be given once daily. The method is contraindicated if ureteral reflux is present, unless the reflux is mild and the bladder emptied frequently.

C. SURGERY

Transurethral resection is indicated for hypertrophy of the bladder neck or an enlarged prostate, either of which may cause obstruction of the bladder outlet and retention of residual urine. It also may be performed in some male patients to weaken the outlet resistance of the bladder to prevent leakage. This helps protect the bladder from overdistention due to a buildup of residual urine.

Complete urinary incontinence due to sphincter incompetence can be managed by implanting an artificial sphincter. Bladder pressure should be low, however, for this to be successful. Bladder neck resection also may be considered as a way to increase outlet resistance. Incontinence in this group of patients can be treated with drugs or neurostimulation if it results from mild bladder spasticity.

D. PARASYMPATHOMIMETIC DRUGS

The stable derivatives of acetylcholine are at times of value in assisting the evacuation of the bladder. Although they do not initiate or effect bladder contraction, they do provide increased bladder tonus. They may be helpful in symptomatic treatment of the milder types of flaccid neuropathic bladder. Drugs may be tried empirically, but usefulness is best gauged during urodynamic evaluation. If filling pressure or resting tonus is increased after bethanechol chloride (Urecholine) is administered, evacuation of the bladder through trigger reflexes or straining should be more effective. The drug then should be clinically helpful.

Bethanechol chloride is the drug of choice. It is given orally, 25–50 mg every 6–8 hours. In special situations (eg, urodynamic study or immediately following operation), it may be given subcutaneously, 5–10 mg every 6–8 hours.

Neuropathic Bladder Associated with Spina Bifida

Spina bifida is incomplete formation of the neural arch at various levels of the spine. The defect is recognized at birth and closed immediately to prevent infection. The scarring that results can entrap and tether nerves in the cauda equina. With failure of the neural arch to close, there is failure of anterior horn cell development and organization. The end result is a mixed type of neuropathic defect. Roughly two-thirds of patients have a spastic bladder with weakness in the feet and toes. About one-third have a flaccid bladder. Often, there is a greater degree of flaccidity in the pelvic floor than in the detrusor. The goals of therapy are to control incontinence and preserve renal function.

A. CONSERVATIVE TREATMENT

Clean intermittent catheterization is the best management. Parents can be taught to do this for the child, and eventually the child can take over this function. Frequency should be determined by the storage capacity of the bladder and the fluid intake, usually every 3–6 hours. An anticholinergic drug may be required to mediate bladder spasticity and improve storage function in order to control incontinence.

1. Mild symptoms—If there is occasional dribbling or some residual urine associated with lack of desire to void, the patient should try to void every 2 hours when awake. Manual suprapubic pressure enhances the efficiency of emptying. An external condom catheter or a small pad can be worn to protect against small-volume losses of urine.

2. More severe symptoms—If urinary incontinence is associated with residual urine or if ureteral reflux is found, the following steps should be taken:

   a. Hypotonic bladder—If reflux has been demonstrated, intermittent self-catheterization 4–6 times a day may protect the upper urinary tract from deterioration and the consequences of pyelonephritis. Ureteral reimplantation can be considered for bilateral reflux or a transureteroureterostomy for single-sided reflux if all other considerations are favorable. Intermittent catheterization should then be reinstituted.
b. Hypertonic bladder—The problem with patients in this category is more serious because the bladder is spastic with reduced capacity and the sphincter is hypotonic. Virtually constant dribbling can result. The cystogram will reveal heavy trabeculation of the bladder, often with reflux and advanced hydronephrostonephrosis. Anticholinergic medication should be given, and an indwelling catheter should be inserted for several months. Once upper urinary tract dilatation has improved and the bladder has been restored to a more spheric shape, intermittent catheterization may be reinstituted. With time and care, many of these children develop a more balanced type of bladder behavior. Continence may be gained without compromising the upper urinary tract.

Most of these patients will not require urinary diversion if they are carefully followed up and if the parents actively participate in their care.

**B. Surgical Treatment**

If the bladder is of the spastic type with diminished capacity, there are several surgical options short of actual urinary diversion. Sacral nerve block during urodynamic evaluation helps in determining whether sacral nerve root section would be beneficial. This helps in cases of spastic bladder but not in cases of poorly compliant, fibrotic bladder. Sectioning the S3 nerves reduces intravesical pressures, improves storage, and reduces the risk of reflux or obstruction of the ureters.

For the patient with a mildly spastic bladder and reasonable storage capacity (>200 mL), urinary incontinence might be controlled via electrostimulation of the pelvic floor. Many of these patients have intact nerves to the sphincter. These can be stimulated to enhance sphincter tone and inhibit voiding. If the bladder has a limited capacity with poor compliance and poor contractility, augmentation cystoplasty followed by intermittent self-catheterization is the treatment of choice.

If the refluxing patient has recurrent fever (equivalent to pyelonephritis) despite the presence of an indwelling catheter or if incontinence cannot be controlled because of poor detrusor compliance, urinary diversion must be considered. Nonrefluxing continent reservoirs offer the most favorable long-term outlook for preservation of the upper urinary tract.

**3. Control of Urinary Incontinence**

**In the Hospital**

Urinary incontinence is one of the most distressing aspects of neurovesical dysfunction, especially when the bladder has otherwise adequate function. The problem is minimized in men who are hospitalized because supervision is available, bathrooms are nearby, and a bedside urinal is always available. Women have a greater problem because they must use a bedpan or may require an indwelling catheter. Catheters have associated risks and do not always control leakage associated with spastic bladder. No simple, satisfactory solution to this problem has been devised for women.

**After Discharge**

After discharge from the hospital, most men with spastic bladders rely on a condom catheter for protection against leakage and for practical urine collection. The only exception is patients who are predictably dry between catheterizations. The condom catheter attaches to the penis without pressure and has a conduit to a leg bag. The adhesives are nonirritating and long lasting. Problems involved in keeping these catheters in place are limited to noncircumcised patients and those with large suprapubic fat pads that shorten the length of the shaft of the penis. Circumcision or placement of a penile prosthesis will correct for these limitations.

Urethral compression by means of a Cunningham clamp is occasionally preferred by patients. This protects only against low-pressure leakage, however, and if it is applied too tightly, a urethral diverticulum may develop.

Other types of external collection devices are available (McGuire urinal, Texas catheter), but with advancements in adhesive glues for condom catheters and use of penile prostheses, the other methods are being used less frequently.

**Neurostimulation**

Extensive research continues to be conducted on methods of restoring complete voluntary control over the storage and evacuation functions of the bladder. Sacral and pudendal nerve anatomy has been determined so that surgical exposure of these nerves and their branches is possible. An electrode can be placed for selective stimulation of the bladder, levator, and urethral or anal sphincters. A number of possibilities exist for neurostimulation or rhizotomy, but only a few are practical. Urodynamic evaluation of bladder function following a nerve block or during neurostimulation can help determine the therapeutic value of these treatments.

Single or multiple electrodes can be placed on selected nerves and coupled to a subcutaneous receiver. The desired function (continence or evacuation) can be selected. Usually, one or the other is needed in any one patient. Much will change in this approach as technologic advances become adapted to the increased understanding of bladder physiology. Striking successes are also being seen with electroevacuation in highly selected patients.

**Complications of Neuropathic Bladder**

The principal complications of the neuropathic bladder are recurrent urinary tract infection, hydronephrosis secondary to ureteral reflux or obstruction, and stone formation.
The primary factors contributing to these complications are the presence of residual urine, sustained high intravesical pressures, and immobilization, respectively.

Incontinence in neuropathic disorders may be passive, as in flaccid lesions when outlet resistance is compromised, or may be the result of uninhibited detrusor contractions, as in spastic lesions.

Infection

Infection is virtually inevitable with the neuropathic bladder state. During the stage of spinal shock that follows cord injury, the bladder must be emptied by catheterization. Sterile intermittent catheterization is recommended at this stage, but for practical purposes or for the sake of convenience, a Foley catheter is often left indwelling. Chronic catheter drainage guarantees infection regardless of any preventive measures taken. Nevertheless, a recent clinical trial of colonization of the bladder with nonpathogenic *Escherichia coli* showed some promise; it significantly reduced the episodes of infection in a group of spinal cord injury patients with neurogenic bladder (Darouiche et al, 2001).

The upper urinary tract is usually protected from infection by the integrity of the ureterovesical junction. If this becomes incompetent, infected urine will reflux up to the kidneys. Decompensation of the ureterovesical junction results from the high intravesical pressures generated by the spastic bladder. It is most important that these cases be treated aggressively with an intensive program of self-catheterization and anticholinergic medication. The Credé maneuver should not be used.

A number of infective complications can result from the presence of a chronically indwelling Foley catheter. These include cystitis and periurethritis resulting from mechanical irritation. A periurethral abscess may follow, with formation of a fistula via eventual rupture of the abscess through the perineal skin. Drainage may also take place through the urethra, with the end result being a urethral diverticulum. Infection may travel up into the prostatic ducts (prostatitis) or seminal vesicles (seminal vesiculitis) and along the vas into the epididymis (epididymitis).

**A. Treatment of Pyelonephritis**

Episodic renal infection should be treated aggressively with appropriate antibiotics to prevent renal loss. The source and cause of infection should be eliminated if possible.

**B. Treatment of Epididymitis**

This condition is a complication of either dyssynergic voiding or an indwelling catheter. Treatment consists of appropriate antibiotics, bed rest, and scrotal elevation. The indwelling catheter should be removed or replaced with a suprapubic catheter. Preferred long-term management is to place the patient on an intermittent self-catheterization program. Rarely, ligation of the vas is required.

**Hydronephrosis**

Two mechanisms lead to back pressure on the kidney. Early, the effect of trigonal stretch secondary to residual urine and detrusor hypertonicity becomes compounded by evolving trigonal hypertrophy. The combination causes abnormal pull on the ureterovesical junction, with increased resistance to the passage of urine. A “functional” obstruction results, which leads to progressive ureteral dilatation and back pressure on the kidney. At this stage, this condition can be relieved by continuous catheter drainage or by combined intermittent catheter drainage and use of anticholinergics.

A delayed consequence of trigonal hypertrophy and detrusor spasticity is reflux due to decompensation of the ureterovesical junction. The causative factor appears to be a combination of high intravesical pressure and trabeculation of the bladder wall. The increased stiffness of the ureterovesical junction weakens its valve-like function, slowly eroding its ability to prevent reflux of urine during forceful bladder contractions.

When ureteral reflux is detected by cystography, previous methods of bladder care must be radically adjusted. An indwelling catheter may manage the problem temporarily. However, if the reflux persists after a reasonable period of drainage, antireflux surgery must be considered. In addition, measures to reduce high intravesical pressure are needed (bladder augmentation, sacral rhizotomy, transurethral resection of the bladder outlet, or sphincterotomy). Progressive hydronephrosis may require nephrectomy. Urinary diversion is a last resort, which should be avoidable if the patient is followed up regularly.

**Calculus**

A number of factors contribute to stone formation in the bladder and kidneys. Bed rest and inactivity cause demineralization of the skeleton, mobilization of calcium, and subsequent hypercalciuria. Recumbency and inadequate fluid intake both contribute to urinary stasis, possibly with increased concentration of urinary calcium. Catheterization of the neurogenic bladder may introduce bacteria. Subsequent infection is usually due to a urea-splitting organism, which causes the urine to become alkaline, with reduced solubility of calcium and phosphate.

**A. Bladder Stones**

Because these stones are usually soft, they can be crushed and will wash out through a cystoscope sheath. Occasionally, they are large and need to be removed via a suprapubic cystotomy.

**B. Ureteral Stones**

Virtually all ureteral stones can now be removed by antegrade or retrograde retrieval methods or by extracorporeal shock wave lithotripsy (ESWL).
C. RENAL STONES

In a patient with neurogenic bladder, kidney stones generally are the result of infection; if the infection is untreated, the stones become the source of persistent renal infection and eventual renal loss. Most of the stones in the renal pelvis can be removed by either a percutaneous endoscopic procedure or ESWL. Occasionally, a large staghorn stone may require open surgery.

Renal Amyloidosis

Secondary amyloidosis of the kidney is a common cause of death in patients with neuropathic bladder. It is a result of chronic debilitation in patients with difficult decubitus ulcers and poorly controlled infection. Fortunately, due to better medical care, this is an uncommon finding today.

Sexual Dysfunction

Men who have had traumatic cord or cauda equina lesions experience varying degrees of sexual dysfunction. Those with upper motor lesions fare well, with the majority having reflexogenic erectile capability. Dangerous elevations in blood pressure can occur with erections in patients with high thoracic or cervical lesions. Problems of quality of erection or premature detumescence are found with all levels of injury. Patients with lower motor lesions are, as a rule, impotent, unless the lesion is incomplete. There is a high degree of variability in the sexual capabilities of patients with all levels of spinal injury. Fortunately, sexual function can be restored to most patients by oral sildenafil, transurethral medications, a vacuum erection device, intracavernous injection, or a penile prosthesis.

Often, patients with spinal injury lose the ability to ejaculate even with preservation of functional erections. This is a result of lost coordination between reflexes normally synchronized through higher center regulation. Patients may have the capability to ejaculate after an erection, but are either unable to trigger this sexual event or are unable to trigger it in proper sequence. Techniques using vibratory stimulation of the penis or transrectal electrical stimulation have been developed to accomplish semen collection in patients with “functional infertility.”

Autonomic Dysreflexia

Autonomic dysreflexia is sympathetically mediated reflex behavior triggered by sacral afferent feedback to the spinal cord. The phenomenon is seen in patients with cord lesions above the sympathetic outflow from the cord. As a rule, it occurs in rather spastic lesions above T1 but on occasion in lesions of mild spasticity or those as low as T5. Symptoms include dramatic elevations in systolic or diastolic blood pressure (or both), increased pulse pressure, sweating, bradycardia, headache, and piloerection. Symptoms are brought on by overdistention of the bladder. Immediate catheterization is indicated and usually brings about prompt lowering of blood pressure. Oral nifedipine (20 mg) has been shown to alleviate this syndrome when given 30 minutes before cystoscopy (Dykstra, Sidi, and Anderson, 1987) or electroejaculation (Steinberger et al, 1990). The acute hemodynamic effect can be managed with a parenteral ganglionic blocking agent or alpha-adrenergic blockers (Barrett and Wein, 1987). Sphincterotomy and peripheral rhizotomy have been used by some to prevent recurring autonomic dysreflexia.

PROGNOSIS

The greater threat to the patient with a neuropathic bladder is progressive renal damage (pyelonephritis, calculus, and hydronephrosis). Advances in the management of the neuropathic bladder, together with better follow-up of patients at regular intervals, have substantially improved the outlook for long-term survival.

REFERENCES

Urodynamic study is an important part of the evaluation of patients with voiding dysfunctions—dysuria, urinary incontinence, neuropathic disorders, and so on. Formerly, the examiner simply observed the act of voiding, noting the strength of the urinary stream, and drawing inferences about the possibility of obstruction of the bladder outlet. In the 1950s, it became possible to observe the lower urinary tract by fluoroscopy during the act of voiding; and in the 1960s, the principles of hydrodynamics were applied to lower urinary tract physiology. The field of urodynamics now has clinical applications in evaluating voiding problems resulting from lower urinary tract disease.

The nomenclature of the tests used in urodynamic studies is not yet settled, and the meanings of urodynamic terms are sometimes overlapping or confusing. In spite of these difficulties, urodynamic tests are extremely valuable. Symptoms elicited by the history or by physical, endoscopic, or even radiographic examination often must be investigated further by urodynamic tests so that therapy can be devised that is based on an understanding of the altered physiology of the lower urinary tract.

As is true of many high-technology testing procedures (eg, electrocardiography, electroencephalography), urodynamic tests have the greatest clinical validity when their interpretation is left to the treating physician, who should either supervise the study or be responsible for correlating all of the findings with personal clinical observations.

FUNCTIONS RELEVANT TO URODYNAMICS & TESTS APPLICABLE TO EACH

Urodynamic study of the lower urinary tract can provide useful clinical information about the function of the urinary bladder, the sphincteric mechanism, and the voiding pattern itself.

**Bladder function** has been classically studied by cystography and fluoroscopy. Urodynamic studies use cystometry. Conventional radiographic studies and urodynamic studies can, of course, be usefully combined.

**Sphincteric function** depends on 2 elements: the smooth muscle sphincter and the voluntary sphincter. The activity of both elements can be recorded urodynamically by pressure measurements; the activity of the voluntary sphincter also can be recorded by electromyography.

The act of voiding is a function of the interaction between bladder and sphincter, and the result is the **flow rate**. The flow rate is one major aspect of the total function of the lower urinary tract. It is generally recorded in milliliters per second as well as by total urine volume voided. The simultaneous recording of bladder activity (by intraluminal pressure measurements), sphincteric activity (by electromyography or pressure measurements), and flow rate reveals interrelationships among the 3 elements. Each measurement may give useful information about the normality or abnormality of one specific aspect of lower urinary tract function. A more complete picture is provided by integrating all 3 lower tract elements in a simultaneously recorded comparative manner. This comprehensive approach may involve synchronous recordings of variable pressures, flow rate, volume voided, and electrical activity of skeletal musculature around the urinary sphincter (electromyography), along with fluoroscopic imaging of the lower urinary tract. The multiple pressures to be recorded are quite variable and usually include intravesical pressure, intraurethral pressure at several levels, intra-abdominal pressure, and anal sphincteric pressure as a function of muscular activity of the pelvic floor.

The techniques of urodynamic study must be tailored to the needs of specific patients. Each method has advantages and limitations depending on the requirements of the study. In one patient, results of a single test might be sufficient to establish the diagnosis and suggest appropriate therapy; in another, many more studies might be necessary.

**PHYSIOLOGIC & HYDРОDYNAMIC CONSIDERATIONS**

**URINARY FLOW RATE**

Because urinary flow rate is the product of detrusor action against outlet resistance, a variation from the normal flow rate might reflect dysfunction of either. The normal flow rate from a full bladder is about 20–25 mL/s in men and 25–30 mL/s in women. These variations are directly
related to the volume voided and the person’s age. Obstruction should be suspected in any adult voiding with a full bladder at a rate of less than 15 mL/s. A flow rate less than 10 mL/s is considered definite evidence of obstruction. Occasionally, one encounters “supervoiders” with flow rates far above the normal range. This may signify low outlet resistance but is of less concern clinically than obstruction.

**Outlet Resistance**

Outlet resistance is the primary determinant of flow rate and varies according to mechanical or functional factors. Functionally, outlet resistance is primarily related to sphincteric activity, which is controlled by both the smooth sphincter and the voluntary sphincter. The smooth sphincter is rarely overactive in women; we have never seen an example of it in any of our urodynamic evaluations. Overactivity of the smooth sphincter is rarely seen in men also but it may occur in association with hypertrophy of the bladder neck due to neurogenic dysfunction or distal obstruction. However, such cases must be critically evaluated before this conclusion is reached.

Increased voluntary sphincteric activity is not uncommon. It is often neglected as a primary underlying cause of increased sphincteric resistance. It is manifested either as lack of relaxation or as actual overactivity during voiding. The normal voluntary sphincter provides adequate resistance, along with the smooth sphincter, to prevent escape of urine from the bladder; if the voluntary sphincter does not relax during detrusor contraction, partial functional obstruction occurs. Overactivity of the sphincter, resulting in increased outlet resistance, is usually a neuropathic phenomenon. However, it can also be functional, resulting from irritative phenomena such as infection or other factors—chemical, bacterial, hormonal, or, even more commonly and often not appreciated, psychological.

**Mechanical Factors**

Mechanical factors resulting in obstruction to urine flow are the easiest to identify by conventional methods. In women, they may take the form of cystoceles, urethral kinks, or, most commonly, iatrogenic scarring, fibrosis, and compression from previous vaginal or periurethral operative procedures. Mechanical factors in men are well known to all urologists; the classic form is benign prostatic hypertrophy. Urethral stricture from various causes and posterior urethral valves are other common causes of urinary obstruction in men, and there are many others.

Normal voiding with a normal flow rate is the product of both detrusor activity and outlet resistance. A high intravesical pressure resulting from detrusor contraction is not necessary to initiate voiding, because outlet resistance has usually dropped to a minimum. Sphincteric relaxation usually precedes detrusor contraction by a few seconds, and when relaxation is maximal, detrusor activity starts and is sustained until the bladder is empty.

**Variations in Normal Flow Rate**

The sequence just described is not essential for normal flow rates. The flow rate may be normal in the absence of any detrusor contraction if sphincteric relaxation is assisted by increased intra-abdominal pressure from straining. Persons with weak outlet resistance and weak sphincteric control can achieve a normal flow rate by complete voluntary sphincteric relaxation without detrusor contraction or straining. A normal flow rate can be achieved in spite of increased sphincteric activity or lack of complete relaxation if detrusor contraction is increased to overcome outlet resistance.

Because a normal flow rate can be achieved in spite of abnormalities of one or more of the mechanisms involved, recording the flow rate alone does not provide insight into the precise mechanisms by which it occurs. Distinction between patterns of flow can be difficult. For practical purposes, if the flow rate is adequate and the recorded pattern and configuration of the flow curve are normal, these variations may not be clinically significant except in rare cases.

**Nomenclature**

The study of urinary flow rate itself is usually called uroflowmetry. The flow rate is generally identified as maximum flow rate, average flow rate, flow time, maximum flow time (the time elapsed before maximum flow rate is reached), and total flow time (the aggregate of flow time if the flow has been interrupted by periods of no voiding) (Figure 28–1). The flow rate pattern is characterized as continuous or intermittent, etc.

**Pattern Measurement of Flow Rate**

A normal flow pattern is represented by a bell-shaped curve (Figure 28–1). However, the curve is rarely completely smooth; it may vary within limits and still be normal. Flow rate can be determined by measuring a 5 seconds’ collection at the peak of flow and dividing the amount obtained by 5 to arrive at the average rate per second. This rough estimate is useful, especially if the flow rate is normal and the values are above 20 mL/s.

In modern practice, the flow rate is more often recorded electronically: The patient voids into a container on top of a measuring device that is connected to a transducer, the weight being converted to volume and recorded on a chart in milliliters per second. Figure 28–2 is an example of such a recording from a normal man. The general bell-shaped curve is quite clear, and the tracing shows all of the values discussed previously: total flow time, maximum flow time, maximum flow rate, average flow rate, and total volume voided. Occasional supervoiders can
URODYNAMIC STUDIES

exceed the limits of the chart, but this is usually not of clinical concern (Figure 28–3). A possible variation in the bell appearance is seen in Figure 28–4.

The overall appearance of the flow curve may disclose unsuspected abnormalities. In Figure 28–5, for example, flow time is greatly prolonged. Maximum flow rate may not be low, but the average flow rate is very low—though the maximum flow rate is at one point within the normal range. Such fluctuation in flow rate is most commonly related to variations in voluntary sphincteric activity. In Figure 28–6, this pattern is extreme: Maximum flow rate never exceeds 15 mL/s, and average flow rate is about 10 mL/s, which is indicative of obstruction. (Again, this fluctuation in pattern probably reflects sphincteric hyperactivity.)

The flow rate pattern reveals a great deal about the forces involved. For example, if the patient is voiding without the aid of detrusor contractions—primarily by straining—this can be easily deduced from the pattern of the flow rate. Figure 28–7 shows an example of intermittent voiding, primarily by straining, with no detrusor activity, and at a rate that sometimes does not reach the usual peaks. With experience, one becomes expert at detecting the mechanisms underlying abnormalities in flow rate. For example, in Figure 28–5, the maximum flow rate is in the normal range, the average flow rate is slightly low, and the curve has a general bell pattern, yet brief partial intermittent obstructions to flow can be readily interpreted as due to overactivity of the voluntary sphincter, a mild form of detrusor/sphincter dyssynergia (see discussion following).

Flow rates in mechanical obstruction are totally different, classically in the range of 5–6 mL/s; flow time is greatly prolonged, and there is sustained low flow with minimal variation (Figure 28–8). Figure 28–9 is a striking example of a curve for a patient with benign prostatic hypertrophy. No simultaneous studies are needed with such a pattern, since the pattern is obviously one of mechanical obstruction.

Figure 28–1. Uroflowmetry. Basic elements of maximum flow, average flow, total flow time, and total volume voided.

Figure 28–2. Classic normal flow rate, with peak of about 30 mL/s and average of about 20 mL/s. On the horizontal scale, one large square equals 5 s.
Figure 28–3. Flow rate of “supervoider.” Maximum flow rate exceeds limits of chart. Tracing shows fast buildup and complete bladder emptying of large volume of urine in a very short period. On the horizontal scale, one large square equals 5 s.

Figure 28–4. Normal flow rate with some variation in appearance of curve. Note rapid pressure rise but progressive increase to maximum, followed by a sharp drop. There is also fluctuation in ascending limb of tracing. On the horizontal scale, one large square equals 5 s.
**Figure 28–5.** Rather low flow rate (not exceeding 10 mL/s), yet at one point the peak reaches 27–32 mL/s. Note again fluctuation in flow. On the horizontal scale, one large square equals 5 s.

**Figure 28–6.** Very low flow rate of short duration and small volume. Note that maximum flow is not above 15 mL/s; however, flow average is less than 10 mL/s, and flow is almost completely interrupted in the middle. On the horizontal scale, one large square equals 5 s.

**Figure 28–7.** Classic flow rate due to abdominal straining with no detrusor activity. See effect of spurts of urine with complete interruption between them; patient cannot sustain increased intra-abdominal pressure. On the horizontal scale, one large square equals 5 s.

**Figure 28–8.** Flow rate in a case of urinary obstruction showing very low average flow rate (not above 5 or 6 mL/s). Prolonged duration of flow is associated with incomplete emptying. On the horizontal scale, one large square equals 5 s.

**Figure 28–9.** Classic low flow rate of bladder outlet obstruction (benign prostatic hypertrophy), markedly prolonged flow time, and fluctuation due to attempt at improving flow by increasing intra-abdominal pressure. On the horizontal scale, one large square equals 5 s.
Reduced flow rate in the absence of mechanical obstruction is due to some impairment of sphincteric or detrusor activity. This is seen in a variety of conditions, for example, normal detrusor contraction with no associated sphincteric relaxation and normal detrusor contraction with sphincteric overactivity, which is more serious. These 2 entities are commonly referred to as detrusor/sphincter dyssynergia. If with detrusor contraction the sphincter does not relax and open up or (worse) if it becomes overactive, urine flow is obstructed (ie, flow rate is reduced and of abnormal pattern). Reduced flow rate may occur even with increased detrusor activity if the latter is not adequate to overcome sphincteric resistance.

So many variations are possible in the shape of the flow curve—no matter how accurately the flow is recorded or how often the study is repeated to confirm abnormal findings—that it is beneficial to relate it to simultaneous recordings, such as of bladder pressure, pelvic floor electromyography, urethral pressure profile, or simply cinefluoroscopy. Nevertheless, by itself it can be one of the most valuable urodynamic studies undertaken to evaluate a specific type of voiding dysfunction. Flowmetry not only is of diagnostic value but also is valuable in follow-up studies and in deciding on treatment. In some cases, however, flowmetry alone does not provide enough data about the abnormality in the voiding mechanism. More information must then be obtained by evaluation of bladder function.

BLADDER FUNCTION

The basic factors of normal bladder function are bladder capacity, accommodation, sensation, contractility, voluntary control, and response to drugs. All of them can be evaluated by cystometry. If all are within the normal range, bladder physiology can be assumed to be normal. Evaluation of every factor has its own implication and, before a definitive conclusion is reached, must be examined in the light of associated manifestations and findings.

**Capacity, Accommodation, & Sensation**

Cystometry can be done by either of 2 basic methods: (1) allowing physiologic filling of the bladder with secreted urine and continuously recording the intravesical pressure throughout a voiding cycle (starting the recording when the patient’s bladder is empty and continuing it until the bladder has been filled—at which time the patient is asked to urinate—and voiding begins) or (2) by filling the bladder with water and recording the intravesical pressure against the volume of water introduced into the bladder. With the first (physiologic filling) method, the assessment of bladder function is based on voided volume (assuming that the presence of residual urine has been ruled out). The second method permits accurate determination of the volume distending the bladder and of the pressures at each level of filling, yet it has inherent defects: fluid is introduced rather than naturally secreted, and bladder filling occurs more rapidly than it normally does.

The cystometrogram (Figure 28–10) is obtained during the phase of bladder filling; the volume of fluid in the bladder is plotted against the intravesical pressure to show bladder wall compliance to filling. The normal cystometric curve shows a fairly constant low intravesical pressure until the bladder nears capacity, then a moderate rise until capacity is reached, and then a sharp rise as voiding is initiated. Normally, the sensation of fullness is first perceived when the bladder contains 100–200 mL of fluid and strongly felt as the bladder nears capacity; the desire to void occurs when the bladder is full (normal capacity, 400–500 mL). However, the bladder has a power of accommodation, that is, it can maintain an almost constant intraluminal pressure throughout its filling phase regardless of the volume of fluid present, and this directly influences compliance. As the bladder progressively accommodates larger volumes with no change in intraluminal pressure, the compliance values become higher (Compliance = Volume/Pressure) (Figure 28–10).

![Figure 28–10. Cystometrogram of patient with normal bladder capacity. Note stable intravesical pressure during filling phase; slight rise at end of filling phase, indicating bladder capacity perceived as sense of fullness; and sharp rise at end (voiding contraction).](image-url)
Contractility & Voluntary Control

The bladder normally shows no evidence of contractility or activity during the filling phase. However, once it is filled to capacity and the patient perceives the desire to urinate and consciously allows urination to proceed, strong bladder contractions occur and are sustained until the bladder is empty. The patient can of course consciously inhibit detrusor contraction. Both of these aspects of voluntary detrusor control must be assessed during cystometric study in order to rule out uninhibited bladder activity and to determine whether the patient can inhibit urination with a full bladder and initiate urination when asked to do so. The latter is occasionally difficult to verify clinically because of conscious inhibition by a patient who may be embarrassed by the unnatural circumstances.

Responses to Drugs

Drugs are being used with increasing frequency in the evaluation of detrusor function. They can help to diagnose underlying neuropathy and to determine whether drug treatment might be of value in individual cases. Study of the relationship of bladder capacity to intravesical pressure and bladder contractility gives a rough evaluation of the patient’s bladder function. Low intravesical pressure with normal bladder capacity might not be significant, whereas low pressure with a very large capacity might imply sensory loss or a flaccid lower motor neuron lesion, a chronically distended bladder, or a large bladder due to myogenic damage. High pressure (usually associated with reduced capacity) that rises rapidly with bladder filling is most commonly due to inflammation, enuresis, or reduced bladder capacity. However, uninhibited bladder activity during this high-pressure filling phase might indicate neuropathic bladder or an upper motor neuron lesion.

The parasympathetic drug bethanechol chloride (Urecholine) is often used to assess bladder muscle function in patients with low bladder pressure associated with lack of detrusor contraction. No response to bethanechol suggests myogenic damage; a normal response indicates a bladder of large capacity with normal musculature; and an exaggerated response indicates a lower motor neuron lesion. The test has so many variables that it must be done meticulously to give reliable results.

Testing with anticholinergic drugs or muscle depressants may be helpful in the evaluation of uninhibited detrusor contraction or increased bladder tone and low compliance. The information thus obtained can be useful in choosing drugs for treatment.

Recording of Intravesical Pressure

Intravesical pressure can be measured directly from the vesical cavity, either by a suprapubic approach or via a transurethral catheter. The pressure inside the bladder is actually a function of both intra-abdominal and intravesical pressure. Thus, true detrusor pressure is the pressure recorded from the bladder cavity (intravesical pressure) minus intra-abdominal pressure. This point is important because variations in intra-abdominal pressure may alter the recorded intravesical pressure, and if the recorded intravesical pressure is mistakenly considered to reflect only detrusor pressure and not increased intra-abdominal pressure due to straining as well, erroneous conclusions may be reached.

Whenever possible, intra-abdominal pressure should be recorded simultaneously with intravesical pressure, since there is no other way to determine the true detrusor pressure. Intra-abdominal pressure is usually recorded by a small balloon catheter inserted high in the rectum and connected to a separate transducer.

The most valuable part of the cystometric study is the determination of voiding activity or voiding contraction. The characteristics of intravesical pressure can be quite significant. Normally, voiding contractions are not high (20–40 cm of water); this magnitude of intravesical pressure is generally adequate to deliver a normal flow rate of 20–30 mL/s and completely empty the bladder if it is well sustained. A higher voiding pressure is indicative of possible increase in outlet resistance yet denotes an overactive, healthy detrusor musculature. Figure 28–11 shows a normal flow rate associated with normal detrusor contraction at a magnitude of 20 cm of water that is well sustained and of short duration and results in complete bladder emptying.

The quality of bladder pressure can also be informative, even without simultaneous recording of flow rate. In such cases, however, it is preferable to record flow rate under normal circumstances. A well-sustained detrusor contraction, high at initiation and sustained at normal values, is seen in Figure 28–12. In Figure 28–13, the voiding pressure is too high—there is an element of sphincteric dysynergia triggering variations in voiding pressures and flow rate. Simultaneous recording of bladder and intra-abdominal pressures would provide more information. As suggested previously, recording the intravesical pressure alone does not give as much information as may be required, and increased intra-abdominal pressure might be mistaken for detrusor action. This situation is illustrated in Figure 28–14. The bladder pressure appears to indicate good detrusor function; nevertheless, simultaneous recording of intra-abdominal pressure makes it clear that all of the apparent changes in vesical intraluminal pressure in fact represent variations in intra-abdominal pressure.

Figure 28–15 shows the 2 pressures recorded on the same chart, on the same channel, by having the writing pen share the time between 2 transducers—one recording intra-abdominal pressure; the other, intravesical pressure.

A. Pathologic Changes in Bladder Capacity

The bladder capacity is normally 400–500 mL, but it can be reduced or increased in a variety of disorders and lesions
Some common causes of reduced bladder capacity are enuresis, urinary tract infection, contracted bladder, upper motor neuron lesion, and defunctionalized bladder. Reduced capacity also may occur in association with incontinence and in postsurgical bladder. Increased bladder capacity is not uncommon in women who have trained themselves to retain large volumes of urine. Bladder capacity is increased also in sensory neuropathic disorders, lower motor neuron lesions, and chronic obstruction from myogenic damage. It is important to relate bladder capacity to the intravesical pressure (Table 28–2). Slight variations in bladder capacity with no change in bladder pressure might be of less significance than the reverse. What is usually of greatest significance is the bladder with reduced capacity associated with normal pressure or, more important, with increased pressure, or the bladder with large capacity associated with decreased pressure.

**B. PATHOLOGIC CHANGES IN ACCOMMODATION**

Accommodation reflects intravesical pressure in response to filling. In a bladder with normal power of accommodation—in which case the micturition center of the spinal cord is controlled by the central nervous system—intravesical pressure does not vary with progressive bladder filling until capacity is reached; in other words, when compliance is reduced, there will be a progressive increase in intravesical pressure and loss of accommodation. This usually occurs at smaller volumes and with reduced capacity. The patient being studied by cystometry can always note the presence or absence of a sensation of fullness. One normally does not sense volumes in the bladder but only changes in pressure.

**C. PATHOLOGIC CHANGES IN SENSATION**

A slight rise in intravesical pressure on cystometry signifies that the bladder is full to normal capacity and that the patient is perceiving it. This sign is usually absent in pure sensory neuropathy and in mixed sensory and motor loss. (Other sensations can be tested for in different ways; see Chapter 26.)

**D. PATHOLOGIC CHANGES IN CONTRACTILITY**

The bladder is normally capable of sustaining contraction until it is empty. Absence of residual urine after voiding usually denotes well-sustained contractions. Neuropathic
dysfunction is usually associated with residual urine of variable amount depending on the type of dysfunction. Significant outlet resistance—mechanical or functional—is also a cause of residual urine.

Cystometric study may disclose complete absence of detrusor contractility due to motor or sensory deficits or conscious inhibition of detrusor activity (Table 28–3). Detrusor hyperactivity is shown as uninhibited activity, usually due to interruption of the neural connection between spinal cord centers and the higher midbrain and cortical centers.

An integrated picture of bladder capacity, intravesical pressure, and contractility is useful for general assessment of the basic physiologic mechanisms of the bladder. Low intravesical pressure in a patient with normal bladder capacity may have no clinical significance, whereas low pressure with a very large capacity may signify sensory loss or a flaccid lower motor neuron lesion, a chronically distended bladder, or a large bladder due to myogenic damage. High pressure (usually associated with reduced capacity) that rises rapidly with bladder filling is most commonly associated with inflammation, enuresis, or reduced bladder capacity. However, uninhibited activity during the interval of rising pressure that occurs with bladder filling may indicate a neurogenic bladder or an upper motor neuron lesion.

![Figure 28–13. Simultaneous recording of flow rate and intra-abdominal pressure; intravesical pressure overlap in top recording. Note very high voiding pressure. However, flow rate is relatively low, with some interruption most likely due to sphincteric overactivity. On the horizontal scale, one large square equals 5 s.](image)

![Figure 28–14. Simultaneous recording of intra-abdominal and intravesical pressures. If one considers only intravesical pressure (upper recording), one might assume adequate detrusor contraction. Comparison with intra-abdominal pressure (lower recording) shows that they are almost identical and that there is no detrusor contraction at all.](image)

![Figure 28–15. Simultaneous recording of 2 measurements—intravesical pressure (top) and intra-abdominal pressure (bottom)—on a single channel. The difference between the two can be clearly seen as pure detrusor contraction.](image)

<table>
<thead>
<tr>
<th>Causes of reduced bladder capacity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Enuresis or incontinence</td>
</tr>
<tr>
<td>Bladder infections</td>
</tr>
<tr>
<td>Bladder contracture due to fibrosis (from tuberculosis, interstitial cystitis, etc)</td>
</tr>
<tr>
<td>Upper motor neuron lesions</td>
</tr>
<tr>
<td>Defunctionalized bladder</td>
</tr>
<tr>
<td>Postsurgical bladder</td>
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</tbody>
</table>

<table>
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<tr>
<th>Causes of increased bladder capacity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensory neuropathic disorders</td>
</tr>
<tr>
<td>Lower motor neuron lesions</td>
</tr>
<tr>
<td>Megacystis (congenital)</td>
</tr>
<tr>
<td>Chronic urinary tract obstruction</td>
</tr>
</tbody>
</table>

Table 28–1. Causes of Reduced or Increased Bladder Capacity.

Note: Normal capacity in adults is 400–500 mL.
CHAPTER 28

SPHINCTERIC FUNCTION

Urinary sphincteric function can be evaluated either by recording the electromyographic activity of the voluntary component of the sphincteric mechanism or by recording the activity of both smooth and voluntary components by measuring the intraurethral pressure of the sphincteric unit. The latter method is called pressure profile measurement (profilometry).

Profilometry

The urethral pressure profile is determined by recording the pressure in the urethra at every level of the sphincteric unit from the internal meatus to the end of the sphincteric segment. Water profilometry, which requires a flow rate of about 2 mL/min, gives fairly accurate results. It may be used for screening patients with incontinence or functional obstruction, but it is not very sensitive and only provides information about total urethral pressure. The membrane catheter and microtransducer techniques of profilometry described in the following sections provide much more accurate and detailed information.

A. MEMBRANE CATHETER TECHNIQUE

Membrane catheters used for recording pressure profiles usually have several channels, so that several measurements can be obtained simultaneously. One such catheter used at UCSF has 4 lumens and an outside diameter of 7F. Two of the four lumens are open at the end, one for bladder filling and the other for recording bladder pressure; the other two lumens, which are situated 7 cm and 8 cm from the catheter tip, are covered by a thin membrane with a small chamber underneath (Figure 28–16). The space under the membrane and the lumen connected to it are filled with fluid, free of any gas, and connected to a pressure transducer. The pressure under this membrane should be zero at the level of the transducer so that it can register any pressure applied to the membrane whatever its level at any time. The catheter also has radiopaque markers at 1-cm intervals starting at the tip, with a heavier marker every 5 cm; it also has a special marker showing the site of each membrane. The markers permit fluoroscopic visualization of the catheter and the membrane levels during the entire study.

B. MICROTRANSUDER TECHNIQUE

The results of microtransducer profilometry are as accurate as those obtained with the membrane catheter. Two microtransducers can be mounted on the same catheter, one at the tip for recording of bladder pressure and the other about 5–7 cm from the tip to record the urethral pressure profile as the catheter is gradually withdrawn from the bladder cavity to below the sphincteric segment.

Electromyographic Study of Sphincteric Function

Electromyography alone gives useful information about sphincteric function, but it is most valuable when done in conjunction with cystometry. There are several techniques for electromyographic studies of the urinary sphincter: either surface electrodes or needle electrodes are used. Surface electrode recordings can be obtained either from the lumen of the urethra in the region of the voluntary sphincter or, preferably, from the anal sphincter by using an anal plug electrode. Recording via needle electrodes can be obtained from the anal sphincter, from the bulk of the musculature of the pelvic floor, or from the external sphincter itself, though in the latter case the placement is difficult and the accuracy of the results is questionable.

Direct needle electromyography of the urethral sphincter provides the most accurate information. Because the technique is difficult, however, simpler approaches are generally used. The anal sphincter is readily accessible for electromyographic testing, and testing of any area of the

Table 28–2. Relationship between Intravesical Pressure and Capacity in Various Diseases.

<table>
<thead>
<tr>
<th>Low intravesical pressure</th>
<th>Normal capacity</th>
<th>Large capacity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensory deficits (diabetes mellitus, tabes dorsalis)</td>
<td>Flaccid lower motor neuron lesions</td>
<td>Large bladder (due to repeated stretching)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>High intravesical pressure</th>
<th>Rapidly rising</th>
<th>Reduced capacity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inflammation</td>
<td>Enuresis</td>
<td>Uninhibited contraction</td>
</tr>
<tr>
<td>Uninhibited neurogenic bladder</td>
<td></td>
<td>Upper motor neuron lesions</td>
</tr>
</tbody>
</table>

Table 28–3. Variations in Detrusor Contractility in Various Diseases.

<table>
<thead>
<tr>
<th>Normal contractions</th>
<th>Normal volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>Absent or weak contractions</td>
<td>Well-sustained contractions</td>
</tr>
<tr>
<td>Sensory neuropathic disorders</td>
<td>Conscious inhibition of contractions</td>
</tr>
<tr>
<td>Lower motor neuron lesions</td>
<td></td>
</tr>
<tr>
<td>Uninhibited contractions</td>
<td>Upper motor neuron lesions</td>
</tr>
<tr>
<td>Cerebrovascular lesions</td>
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</tbody>
</table>
pelvic floor musculature generally reflects the overall electrical activity of the pelvic floor, including the external sphincter. Electromyography is not simple, and the assistance of an experienced electromyographer is probably essential. Electromyographic study makes use of the electrical activity that is constantly present within the pelvic floor and external urinary sphincter at rest and that increases progressively with bladder filling. If the bladder contracts for voiding, electrical activity ceases completely, permitting free flow of urine, and is resumed at the termination of detrusor contraction to secure closure of the bladder outlet (Figure 28–17). Electromyography is important in showing this effect and, along with bladder pressure measurement, can pinpoint the exact time of detrusor contraction. Persistence of electromyographic activity during the phase of detrusor contraction for voiding—or, even worse, its overactivity during that phase—interferes with the voiding mechanism and leads to incoordination between detrusor and sphincter (detrusor/sphincter dyssynergia). During the interval of detrusor contraction, increased electromyographic activity interferes with the free flow of urine, as can be shown by simultaneous recording of flow rate.

Electromyographic recording shows only the activity of the voluntary component of the urinary sphincteric mechanism and the overall activity of the pelvic floor. More information is gained when the electromyogram is recorded simultaneously with detrusor pressure or flow rate. However, this method gives no information about the smooth component of the urinary sphincter.

### Pressure Measurement for Evaluation of Sphincteric Function

Perfusion profilometry, usually performed with the patient supine and with an empty bladder, provides a simple pressure profile that allows determination of the maximum pressure within the urethra. This is adequate for screening patients with incontinence or functional obstruction. However, in order to determine the maximum closure pressure (see section following), the bladder pressure must be recorded simultaneously with the urethral pressure profilometry.

*Figure 28–17. Simultaneous recording of bladder pressure, flow rate, and electromyography of anal sphincter. With rise in bladder pressure for voiding, start of flow rate has a smooth, continuous, bell-shaped curve. Note also complete absence of electromyographic activity of the anal sphincter throughout the voiding act. On the horizontal scale, one large square equals 5 s.*
Such simultaneous recording is not possible with perfusion profilometry. The membrane catheter and microtransducer techniques of profilometry, because they use multichannel recording, routinely provide much more detailed information; at least 4 distinct sets of measurements can be obtained from the simplest pressure profile made using the membrane catheter or microtransducer (Figure 28–18): (1) the maximum pressure exerted around the sphincteric segment, (2) the net closure pressure of the urethra, (3) the distribution of this closure pressure along the entire length of the sphincter, and (4) the exact functional length of the sphincteric unit and its relation to the anatomic length.

A. TOTAL PRESSURE

The urethral pressure profile recording shows the pressure directly recorded within the urethral lumen along the entire length of the urethra from internal to external meatus. From this measurement, the maximum pressure exerted around the sphincteric segment can be determined.

B. CLOSURE PRESSURE

The urethral closure pressure is the difference between intravesical pressure (bladder pressure) and urethral pressure, that is, the net closure pressure. The maximum closure pressure is the most important measurement in evaluating the activity of the sphincteric unit and its responses to various factors.

C. DISTRIBUTION OF CLOSURE PRESSURE

As the catheter is withdrawn down the urethra, the closure pressure at various levels along the entire length of the sphincteric segment is recorded.

D. FUNCTIONAL LENGTH OF SPHINCTERIC UNIT

The functional length of the sphincteric unit is the portion with positive closure pressure, that is, where urethral pressure is greater than bladder pressure. The distinction between anatomic length and functional length is important. Regardless of the anatomic length, the effectiveness of the urethral sphincter may be limited to a shorter segment. In women, the pressure is normally rather low at the level of the internal meatus but builds up gradually until it reaches its maximum in the midurethra, where the voluntary sphincter is concentrated; it slowly drops until it is at its lowest at the external meatus. On the basis of these measurements, it is clear that the anatomic and functional lengths of the normal urethra in women are about the same and that the maximum closure pressure is at about the center of the urethra—not at the level of the internal meatus. In men, the pressure profile is slightly different: the functional length is longer, and the maximum closure pressure builds up in the prostatic segment, reaches a peak in the membranous urethra, and drops as it reaches the level of the bulbous urethra (Figure 28–19). The entire functional length in men is about 6–7 cm; in women, it is about 4 cm.

Dynamic Changes in Pressure Profile

The usefulness of the pressure profile is enhanced if the examiner notes the sphincteric responses to various physiologic stimuli: (1) postural changes (supine, sitting, standing), (2) changes in intra-abdominal pressure (sharp increase with coughing; sustained increase with bearing down), (3) voluntary contractions of the pelvic floor musculature to assess activity of the voluntary sphincter, and (4) bladder filling. The latter test consists of making baseline recordings with both an empty bladder and a full bladder and comparing these recordings with recordings made under conditions of stress (coughing, bearing down) and during voluntary contraction with an empty bladder and a full bladder.

A simple pressure profile is informative but does not provide data that will delineate and identify specific sites of sphincteric dysfunction. The advantage of using a membrane catheter or microtransducer is that the pressure profile can be expanded by slowing the rate of withdrawal of the catheter and speeding up the motion of the recording paper. Since the catheter can be held at different levels for any length of time, other tests can be made and their effects monitored. Response to stress (particularly when standing), response to bladder distention, response to changes in position, the effects of drugs, and the effects of nerve stimulation can all be evaluated if needed. Bladder filling normally leads to increase in tonus of the sphincteric element, with some rise in closure pressure, especially when bladder filling approaches maximum capacity. Stress from coughing or straining also normally results in sustained or increased clo-
sure pressure (Figure 28–20). When the patient stands up, closure pressure is usually substantially increased (Figure 28–21). Testing for activity of the voluntary sphincter by the hold maneuver (asking the patient to actively contract the perineal muscles) produces a significant rise in urethral pressure (Figure 28–22). When the effects of all of these responses are recorded concomitantly with intravesical pressure, the data can be interrelated and the exact closure pressure at any given time can be ascertained.

The response to stress with the patient standing usually should be recorded also. Especially in cases of stress incontinence, weakness of the sphincteric mechanism may not be apparent with the patient sitting or supine but becomes clear when the patient stands up.

The effectiveness of drugs in increasing or reducing the urethral pressure profile can also be tested. For example, phenoxybenzamine (Regitine) can be administered and the urethral pressure profile recorded; a drop in pressure indicates that alpha-blockers may be an effective means of decreasing urethral resistance, with obvious implications for the management of urinary obstruction. Anticholinergic drugs can be tested for possible use as detrusor depressants. Detrusor activity can be investigated by administering bethanechol chloride (Urecholine) and simultaneously recording bladder and urethral pressures.

Figure 28–19. Normal male urethral pressure profile showing progressive rise throughout prostatic segment and peak being reached in membranous urethra. (Reproduced, with permission, from Tanagho EA: Membrane and microtransducer catheters: Their effectiveness for profilometry of the lower urinary tract. Urol Clin North Am 1979;6:110.)

Figure 28–20. Simultaneous recording of intraurethral (U) and intravesical (B) pressures and their responses to coughing and bearing down. Rise in intravesical pressure as a result of increase in intra-abdominal pressure is associated with simultaneous rise in intraurethral pressure, maintaining a constant closure pressure.

Figure 28–21. Urethral pressure profile of normal woman in sitting and standing positions. Note marked improvement in closure pressure (in both functional length and magnitude) when patient stands up. (Reproduced, with permission, from Tanagho EA: Urodynamics of female urinary incontinence with emphasis on stress incontinence. J Urol 1979;122:200.)
Characteristics of Normal Pressure Profile (Figure 28–23)

The basic features of the ideal pressure profile are not easily defined. In women, the normal urethral pressure profile has a peak of 100–120 cm of water, and the closure pressure is in the range of 90–100 cm of water. Closure pressure is lowest at the level of the internal meatus, gradually builds up in the proximal 0.5 cm, and reaches its maximum about 1 cm below the internal meatus. It is sustained for another 2 cm and then starts to drop in the distal urethra. The functional length of a normal adult female urethra is about 4 cm. The response to stress with coughing and bearing down is sustained or augmented closure pressure. Standing up also increases this pressure, with maximum rise in the midsegment.

Pressure Profile in Pathologic Conditions

A. URINARY STRESS INCONTINENCE

The classic pressure changes noted in this type of incontinence are as follows:

1. Low urethral closure pressure
2. Short urethral functional length at the expense of the proximal segment
3. Weak responses to stress
4. Loss of urethral closure pressure with bladder filling
5. Fall in closure pressure on assuming the upright position
6. Weak responses to stress in the upright position

B. URINARY URGE INCONTINENCE

The most pertinent pressure changes in urinary urge incontinence are normal or high closure pressures with normal responses to stress, normal responses to bladder filling, and normal responses when the patient stands up. Urge incontinence can result from any of the following mechanisms (Figure 28–24):

1. Detrusor overactivity, with active detrusor contractions overcoming urethral resistance and leading to urine leakage.
2. The exact reverse, that is, a constant detrusor pressure with no evidence of detrusor overactivity but with urethral instability in that urethral pressure becomes less than bladder pressure, so that urine leakage occurs without any detrusor contraction.
3. A combination of the 2 preceding mechanisms (the most common form), that is, some drop in closure pressure and some rise in bladder pressure. In such cases, the drop in urethral pressure is often the initiating factor.

C. COMBINATION OF STRESS & URGE INCONTINENCE

In this common clinical condition, profilometry is used to determine the magnitude of each component, that is, whether the incontinence is primarily urge, primarily stress, or both equally. As a guide to treatment, profilometric studies sometimes show that stress incontinence precipitates urge incontinence. The stress elements initiate urine leakage in the proximal urethra, exciting detrusor response and sphincteric relaxation and ending with complete urine leakage. Once the stress components are corrected, the urge element disappears. This combination cannot be detected clinically.

D. POSTPROSTATECTOMY INCONTINENCE

After prostatectomy, there is usually no positive pressure in the entire prostatic fossa, minimal closure pressure at the apex of the prostate, and normal or greater than nor-
mal pressure within the voluntary sphincteric segment of the membranous urethra. It is the functional length of the sphincteric segment above the genitourinary diaphragm that determines the degree of incontinence; the magnitude of closure pressure in the voluntary sphincteric segment has no bearing on the patient’s symptoms. High pressure is almost always recorded within the voluntary sphincter despite the common belief that what someone termed “iatrogenically induced incontinence” is due to damage to the voluntary sphincter—which is definitely not the case.

E. DETRUSOR/SPHINCTER DYSSYNERGIA

In this situation, findings of cystometric studies are normal at the filling phase, with possible closure pressure above average. However, the pathologic entity becomes clear when the patient attempts to void: Detrusor contraction is associated with a simultaneous increase in urethral closure pressure instead of a drop in pressure. This is a direct effect of overactivity of the voluntary component, leading to obstructive voiding or low flow rate and frequent interruption of voiding. This phenomenon is commonly seen in patients with supraspinal lesions. It can be encountered in several other conditions as well.

**Value of Simultaneous Recordings**

Measurement of each of the physiologic variables described previously gives useful clinical information. A rise in intravesical pressure has greater significance when related to intra-abdominal pressure. The urine flow rate is more significant if recorded in conjunction with the total volume voided as well as with evidence of detrusor contraction. The urethral pressure profile is more significant when related to bladder pressure and to variations in intra-abdominal pressure and voluntary muscular activity. And for greatest clinical usefulness, all data must be recorded simultaneously so that the investigator can analyze the activity involved in each sequence.

At a minimum, a proper urodynamic study should include recordings of intravesical pressure and intra-abdominal pressure (true detrusor pressure is intravesical pressure minus intra-abdominal pressure), urethral pressure or electromyography, flow rate, and, if possible,
voided volume. For a complete study, the following are necessary: intra-abdominal pressure, intravesical pressure, urethral sphincteric pressure at various (usually 2) levels, flow rate, voided volume, anal sphincteric pressure (as a function of pelvic floor activity), and electromyography of the anal or urethral striated sphincter. These physiological data are recorded with the patient quiet as well as during activity (ie, voluntary increase in intra-abdominal pressure, changes in the state of bladder filling, voluntary contraction of perineal muscles, or—more comprehensively—an entire voiding act starting from an empty bladder; continuing through complete filling of the bladder, and initiation of voiding; and ending when the bladder is empty).

The data derived from urodynamic studies are descriptive of urinary tract function. Simultaneous visualization of the lower urinary tract as multiple recordings are made gives more precise information about the pathologic changes underlying the symptoms. By means of cinerouroscopy, the examiner can observe the configuration of the bladder, bladder base, and bladder outlet during bladder filling (usually with radiopaque medium). The information obtained can then be correlated with the level of catheters, with pressure recordings, and with changes in pelvic floor support during voiding. Combined cinerouroscopy and pressure measurements thus represent the ultimate in urodynamic studies.

**REFERENCES**

**Urethra & Bladder**


Urinary incontinence is a major health issue that affects more than 200 million people worldwide. The direct cost in the United States alone is $16.3 billion, of which 75% is for the management of women with this condition. Incontinence also results in psychological and medical morbidity, significantly impacting health-related quality of life in a manner similar to other chronic medical conditions including osteoporosis, chronic obstructive pulmonary disease, and stroke. Overall prevalence of female incontinence is reported at 38%, increasing with age from 20–30% during young adult life to almost 50% in the elderly (Anger et al, 2006). Recent advances in the understanding of pathophysiology, as well as development of novel pharmacotherapy and surgical techniques for stress, mixed, and urge incontinence (UI), have redefined contemporary care of this patient group.

PATHOPHYSIOLOGY

Elderly patients frequently accept urinary incontinence as a sign of aging and fail to seek help. In fact, it is a manifestation of an underlying disease; occasionally it is transient and resolves spontaneously, but most often it is chronic and progressive. Transient incontinence may occur after childbirth or may be associated with an acute bladder infection. Chronic urinary incontinence can result from a multitude of causes and can be classified under these main headings:

- Anatomic or genuine urinary stress incontinence
- Urge incontinence
- Neuropathic incontinence
- Congenital incontinence
- False (overflow) incontinence
- Posttraumatic or iatrogenic incontinence
- Fistulous incontinence

Each entity listed has its own basic mechanism, although a combination of more than one of the varieties of incontinence is not uncommon.

A. ANATOMIC (GENUINE STRESS INCONTINENCE)

Anatomic incontinence is primarily the result of hypermobility of the vesicourethral segment owing to pelvic floor weakness. Its basic features are an essentially intact sphincteric mechanism, a weak pelvic floor support, and an anatomic abnormality. It is easily demonstrable radiologically, and restoration of the anatomy restores function.

B. TRUE UI

The basic features of true UI are detrusor instability with a normal sphincteric component, normal anatomy, and no neuropathy. Sphincteric instability is less common. Leakage occurs with either detrusor instability and spontaneous contraction or, less commonly, with sphincteric instability and relaxation.

C. NEUROPATHIC INCONTINENCE

Neuropathic incontinence varies, depending on the nerve lesion. The neuropathy is usually identifiable. The incontinence can be active (detrusor hyperreflexia) or passive (sphincteric atony) or, occasionally, a combination of the two.

D. CONGENITAL INCONTINENCE

The causes of congenital incontinence are ectopic ureters, duplicate or single system, with epispadias, extrophy, or cloacal malformation.

E. FALSE (OVERFLOW) INCONTINENCE

False incontinence is usually the result of an obstructive or neuropathic lesion. It is not true incontinence.

F. TRAUMATIC INCONTINENCE

Traumatic incontinence is associated with a fractured pelvis or with surgical damage to the sphincter during bladder neck resection or extensive internal urethrotomy; it also may result from failure of urethral diverticulectomy or repair of erosion of an artificial sphincter.

G. FISTULOUS COMMUNICATION

The fistula can be ureteral, vesical, or urethral. Most of the time, the cause is iatrogenic, from either pelvic or vaginal surgery. This chapter discusses common and significant incontinence disorders: stress, urge, mixed, overflow, and neuropathic incontinence.

Urinary incontinence is defined as any involuntary loss of urine. Normal continence in women is the end-result of coordination between the urethra, bladder, pelvic muscles,
and surrounding connective tissue elements. Under resting conditions, urethral tone is maintained by smooth and striated muscle activity, tension of the fibroelastic elements in the urethral wall, and the cushioning effect of the soft, compressible, submucosal vascular bed (Figure 29–1). The pelvic muscles support the bladder and urethra; contraction of the levator ani pulls the vagina forward toward the pubic symphysis, creating a stable backstop (Norton and Brubaker, 2006).

The major contribution to urethral resistance comes from the smooth and striated muscle components (Figures 29–2 and 29–3). In experimental animals, as well as in humans, the striated external sphincter provides about 50% of static urethral resistance, while smooth muscle is primarily responsible for proximal urethral closure pressure (Figure 29–4). The rise in pressure in the midurethra results from the combined function of the smooth musculature and the striated muscle fibers around it. To maintain continence under stress conditions, the striated urethral sphincter has to resist a raised bladder pressure owing to intra-abdominal pressure increase (Figures 29–5 and 29–6). The activity of the external sphincter helped by the pelvic floor provides for this increased urethral resistance. Involuntary loss of urine with increased intra-abdominal pressure, in the absence of detrusor contraction, is usually labeled stress incontinence. When loss of urine is associated with increased intravesical pressure owing to detrusor contraction, it is commonly referred to as UI.

Genuine stress incontinence is invariably associated with weakness of the pelvic floor support, permitting hypermobility of the vesicourethral segments, which in turn impairs the efficiency of the sphincteric musculature. The increase in intraurethral pressure observed during

**Figure 29–1.** Normal urethral pressure. Closure pressure at the level of the internal meatus is very low; the pressure rises progressively to reach its maximum at approximately the middle third of the urethra—the site of maximal condensation of striated muscle.

**Figure 29–2.** **A:** Response to pelvic nerve stimulation. Note the simultaneous, equal pressure rise in the bladder, proximal urethra (U₁), and midurethra (U₂). **B:** Vesical and sphincteric responses to an injection of the parasympathetic drug methacholine chloride. Note again the simultaneous rise in pressure at the bladder, proximal urethra (U₁), and midurethra (U₂).
coughing results mainly from contraction of the voluntary muscles with sphincteric action. Part of the rise is passive (i.e., by direct transmission), but a significant component is active (i.e., caused by reflex musculature contraction).

**URINARY STRESS INCONTINENCE**

Often seen in women after middle age (with repeated pregnancies and vaginal deliveries), urinary stress incontinence is usually a result of weakness of the pelvic floor and poor support of the vesicourethral sphincteric unit. Urethral closure pressure normally responds to bladder filling; a change in position; or stressful events such as coughing, sneezing, and bearing down. The sphincteric mechanism has its own capacity to augment urethral resistance under stress reflexively and thus to prevent leakage.

The urethral pressure profile is a good measure of the activity of the external sphincter. A static profile demon-

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**Figure 29–3.** Response of the striated component to sacral nerve stimulation. Note that bladder pressure does not change and proximal urethral pressure (U₁) rises only slightly, compared to the sharp and sustained increase in midurethral pressure (U₂).

**Figure 29–4.** A: The resistance required to force the urethra open, overcoming both voluntary and involuntary sphincteric elements. With progressively increasing pressure, the urethra opens at the critical opening pressure (in this recording, about 85 mm Hg). Once the urethra is forced open, the resistance to flow drops precipitously and becomes sustained at the level of sustained urethral resistance (in this recording, roughly 50 mm Hg). B: A similar recording obtained after administration of curare, which completely blocks voluntary sphincteric responses. Note the appreciable drop in both critical opening pressure and sustained resistance. C: Recording after administration of both curare and atropine (a combination that eliminates the activity of smooth and voluntary sphincteric elements). The critical opening pressure drops markedly and is now equal to the sustained resistance; both are very low. D: An overlap of the 3 recordings shows the contribution of each muscular element: the voluntary component contributes roughly 50% of the total resistance, while the smooth component contributes the other 50%. The minimal residual resistance is a function of the collagen elastic element of the urethral wall; this collagen element has no sphincteric significance.
strates the resting tonus of both components of the sphincteric mechanism (see Figure 29–1); a dynamic profile gives the responses of these sphincteric elements to various activities, such as an increase in bladder volume, assumption of the upright position (Figure 29–7), the prolonged stress of bearing down, or the sudden stress of coughing and sneezing (Figure 29–8). Normally, the urethral closure pressure—the net difference between the intraurethral and intravesical pressures—is maintained or augmented during stress.

Anatomy

In genuine stress incontinence, the assumption is that the intrinsic structure of the sphincter itself is intact and normal. However, it loses efficiency because of excessive mobility and loss of support. Thus, the anatomic feature of genuine stress incontinence is consistently that of hypermobility or a lowering of the position of the vesicourethral segment (or a combination of the two factors) (Figure 29–9).

The relationships among the urethra, the bladder base, and various bony points have been the object of much study. For many years the posterior vesicourethral angle has been considered a key factor indicating the presence of anatomic stress incontinence. Some authors, however, have emphasized the axis of inclination, that is, the angle between the urethral line and the vertical plane. Other investigators stress bony landmarks in the pelvis in their descriptions of the relationship of the bladder base and the vesicourethral junction to the sacrococcygeal inferior pubic point (Figure 29–10).

These descriptions illustrate that abnormal anatomic position and excessive mobility are essential elements in the diagnosis of genuine anatomic stress incontinence. To evaluate this aspect of incontinence, I recommend a simplified cystographic study (a lateral cystogram with a urethral catheter in place) to define the vesicourethral segment clearly. With the patient lying on the flat x-ray table, a lateral film is obtained, first at rest to determine the position of the vesicourethral segment in relation to the pubic bone and then with straining to ascertain its degree of mobility (Figures 29–11 and 29–12). Normally, the vesicourethral junction is opposite the lower third of the pubic bone and moves 0.5–1.5 cm with straining. It should be emphasized, however, that cystography is not the means of diagnosing stress incontinence. This demonstration of abnormal position or excessive mobility of the vesicourethral segment is helpful in confirming the cause of existing urinary incontinence. This demonstration of abnormal position or excessive mobility of the vesicourethral segment is helpful in confirming the cause of existing urinary incontinence. Some authors like to classify urinary incontinence in various stages. Stages I and II depend on the degree of hypermobility and usually relate to the amount of urinary leakage. Stage III, which most often is not associated with

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**Figure 29–5.** Urethral pressure profile. **A:** At rest. **B:** Stimulation of both the pudendal and the pelvic nerves incites the maximal response from both smooth and voluntary sphincteric elements. **C:** Pudendal stimulation alone demonstrates the contribution of the voluntary component. **D:** Pelvic nerve stimulation shows the response of the smooth-muscle component alone. **Bottom tracings:** Total maximal pressure profile obtained by stimulation of pelvic and pudendal nerves depicted by overlapping the profile of simultaneous stimulation of both nerves. The contribution and anatomic distribution of each element are clearly seen. Their summation results in the overall total responses recorded in **B** above.
hypermobility, is usually due to intrinsic sphincteric damage—most often iatrogenic.

**Urodynamic Characteristics of Stress Incontinence**

**A. PRESSURE PROFILE**

As would be expected, patients have a low urethral pressure profile with reduced closure pressure. This factor varies with the severity of the sphincteric impairment as a result of the excessive mobility: The pressure profile might be low-normal when weakness is minimal or it might be quite significant when mobility is severe. Not infrequently, however, this weakness of the pressure profile is not demonstrable when the bladder is partially full. It characteristically becomes more significant when the bladder has been distended (Figure 29–13). Also, the pressure profile may appear normal when the patient is in the resting (sitting) position; when he or she assumes the upright position in the dynamic pressure profile, the weakness becomes more apparent (Figure 29–14).

**B. FUNCTIONAL URETHRAL LENGTH**

The anatomic length of the urethra is usually maintained, yet the functional length is invariably shorter. The loss is in the proximal urethral segment (Figure 29–15). Although it might not look funneled on the cystogram, this segment has very low closure efficiency, or none at all, and its pres-
Figure 29–7. Urethral pressure profile for a patient in sitting and upright positions. An approximately 50% increase in urethral closure pressure occurs when the patient assumes the upright position. Urethral functional length is well sustained.

Figure 29–8. A: Intravesical and urethral pressure responses to the stresses of coughing, bearing down, and the hold maneuver. Note the sharp increase in intra-abdominal pressure reflected in intravesical pressure with coughing and the simultaneous greater increase in urethral pressure. The response is similar with bearing down. Closure pressure is maintained and even augmented during these periods of stress. The hold maneuver (recording membrane is in the proximal urethra) produces a minimal response in closure pressure of the proximal urethra. B: Recording comparable to that in A, but the membrane is in the midurethra. Note again the sustained closure pressure as a result of coughing and bearing down and the marked pressure increase in the midurethral segment with the hold maneuver.
Figure 29–9. Lateral cystograms in a 53-year-old woman with stress incontinence. A: Preoperative, relaxed. Note slightly low-lying vesicourethral junction. The posterior vesicourethral angle is near normal. B: With straining, excessive downward and posterior mobility of the vesicourethral segment is shown. Posterior angle almost disappears.

Figure 29–10. Diagrammatic representation of (A) the angles considered when assessing adequacy of bladder support (posterior vesicourethral angle; angle of inclination) and (B) the “SCIPP line” (sacrococcygeal inferior pubic point) and its relationship to the bladder base and the vesicourethral segment as a reference to adequate pelvic support.
sure is almost equal to intravesical pressure. The functional shortening might be minimal or it might involve more than one-half of the length of the urethra. It is important to note that the functional length, like the pressure profile, might appear normal when the bladder is partially full or the patient is in the sitting position.

C. RESPONSE TO STRESS

With the sustained stress of bearing down or the sudden stress of coughing or sneezing, the net urethral closure pressure is reduced, depending on the degree of sphincteric weakness. In severe urinary stress incontinence, any strain or increase in intravesical pressure leads to negative closure pressure and urinary leakage (Figure 29–16).

D. VOLUNTARY INCREASE IN URETHRAL CLOSURE PRESSURE

Patients with mild stress incontinence might be capable of activating their external sphincter maximally and generating a high urethral closure pressure. However, with progression of the anatomic problem and hypermobility, this voluntary increase progressively diminishes; depending on the severity of the weakness and inefficiency of the external sphincter, this weakness becomes more readily apparent.

E. RESPONSE TO BLADDER DISTENTION AND CHANGE IN POSITION

It must be emphasized that, although the features described might be normal in the resting position with minimal bladder filling, all of them can become aggravated with a full bladder or the upright position. In testing these patients urodynamically, one must ascertain the changes that occur with a full bladder and the assumption of the upright position (Figures 29–14 and 29–17).

Diagnosis

A detailed history is important, including the degree of leakage; its relation to activity, position, and state of bladder fullness; the timing of its onset; and the course of its progression. Knowledge of past surgical and obstetric history, medications taken, dietary habits, and systemic diseases (eg, diabetes) can be helpful in the diagnosis. Whether the incontinence is purely stress or purely urge or a combination of the two can be assessed, as can its degree—minimal, moderate, severe, or complete.

Physical examination is essential. The pelvic examination demonstrates laxity of pelvic support, presence of any degree of prolapse, cystocele, rectocele, and mobility of the anterior vaginal wall. A neurologic examination should be
Cystographic study for demonstration of the anatomic abnormality is important, as is urodynamic study to confirm the classic features of urinary incontinence and determine its cause. The goals of cystographic and urodynamic study are, first, to demonstrate the anatomic abnormality and its extent and, second, to assess the activity of the sphincteric mechanism and hence the potential for improvement by correcting the anatomic abnormality. In recurrent cases, repeated previous surgeries may have caused so much intrinsic damage to the sphincteric musculature that simple suspension cannot provide satisfactory results. Indirect evidence of the degree of sphincteric weakness can be obtained by measurement of what is called the leak pressure (ie, by measuring the intra-abdominal pressure through a rectal transducer during the Valsalva maneuver and noting at what pressure the

**Figure 29–12.** Lateral cystograms in two young continent women. A: Relaxed state, 28-year-old woman. B: With straining, the vesicourethral segment is displaced 0.5 cm downward and posteriorly. C: Relaxed state, 34-year-old woman. D: With straining, the vesicourethral segment is displaced 0.8 cm downward and 1 cm posteriorly.


**Figure 29–13.** Urethral pressure profile with minimally filled bladder. Bladder pressure remains constant, but urethral pressure drops progressively. Closure pressure becomes minimal at the end of bladder filling.

**Figure 29–14.** Urethral pressure profile in moderately severe stress incontinence: closure pressure with patient in the sitting position with half-distended bladder, then after the upright position is assumed. Note that closure pressure is close to 75 cm H₂O with the patient in the sitting position but decreases to approximately 35 cm H₂O with the upright position. Note also the marked shortening of functional urethral length once the upright position is assumed.

**Figure 29–15.** Urethral pressure profile in a female patient with moderate urinary stress incontinence. Note the relatively low closure pressure, the short functional urethral length, and the loss of closure pressure of the proximal 1.5 cm of urethra.

**Figure 29–16.** Urethral pressure profile in moderate stress incontinence. Note that, with the bladder relatively empty, closure pressure is close to the normal range. At the start of bladder filling, resting pressure is again normal; as filling progresses, bladder pressure remains stable and urethral closure pressure decreases progressively to a minimum with full bladder distention.
URINARY INCONTINENCE

first leakage of urine occurs). A low reading indicates a severe degree of sphincteric weakness.

Treatment

The principal treatment of urinary stress incontinence is proper suspension and support of the vesicourethral segment in a normal position. The rationalization is that, in genuine stress incontinence, the intrinsic sphincteric mechanism is intact but its efficiency is impaired because of excessive mobility in the abnormal position. Once the position is restored, the sphincteric mechanism usually regains its function.

There are numerous approaches to restoring the normal position and providing adequate support—some vaginal, others suprapubic. The suprapubic approach was popularized by the classic Marshall-Marchetti-Krantz (MMK) retropubic suspension described in 1949, in which periurethral tissue is attached to the back of the pubic symphysis. A modification was introduced by Burch in 1961, in which the anterior vaginal wall is fixed to Cooper’s ligament. Many urologic surgeons today have found that the latter technique, with modifications, provides the most lasting results (Drouin et al, 1999; Kulseng-Hanssen and Berild, 2002) (Figures 29–18 and 29–19).

With excessive sphincteric damage and intrinsic weakness, suspension alone might not be adequate and sling procedures are advised. Of the various techniques and materials, the most popular uses a strip of the anterior rectus sheath, first reported by McGuire. Raz advocates the vaginal wall sling, in which an island of the anterior vaginal wall is mobilized and used to support the vesicourethral segment. Numerous other sling materials are being used: for example, cadaveric fascia lata and various synthetic materials. Most recently, tension-free vaginal tape (TVT) has been gaining popularity. Early results for TVT with follow-up to 5 years demonstrate comparable or improved versus traditional surgical approaches (suburethral slings, urethropexy, colposuspension, or injectable bulking agents) and reported success rates of up to 80% (Ankardal et al, 2006). Potential complications include bladder injury, infection, urinary retention, hemorrhage or hematoma, erosion (vaginal or urethral), and dyspareunia. More recently, in instances of significant intrinsic sphincteric damage, local injection of bulking material, such as hyaluronic acid/detrannomer, polydimethylsiloxan (Macroplastique), and collagen, is used to increase the bladder outlet resistance for patients in whom vesicourethral mobility is not excessive and whose primary problem is intrinsic sphincteric weakness (Appell et al, 2006).

URGE INCONTINENCE

UI is defined as involuntary loss of urine, accompanied or immediately preceded by urgency. The basic feature of UI is detrusor instability and the loss of urine while attempting to inhibit micturition. Sphincteric instability is less common. The focus of this section is restricted to UI;
overactive bladder (OAB), which has replaced the term unstable bladder, and is clinically defined by symptoms of urgency, frequency, and nocturia with or without UI, is addressed only in the context of UI.

Neurogenic, myogenic, or urothelial bladder dysfunction can lead to UI or the constellation of symptoms which define OAB. OAB, with or without UI, is common in both men and women and can result from neuropathic injuries (spinal cord injury), obstruction, inflammation (interstitial cystitis), diabetes, benign prostatic hyperplasia (BPH), and so on, or be iatrogenic. Bladder hypersensitivity may originate from the urothelium or the detrusor muscle or altered neural activation at various points of the micturition cycle (e.g., persistent smooth muscle activation during filling) (Norton and Brubaker, 2006).

**Diagnosis**

Assessment of patients with symptoms of UI, or OAB, should include detailed history, including an assessment of the impact of the disorder on daily life, physical examination, urinalysis, and identification of modifiable causes such as impaired mobility. A sudden urge with uncontrolled loss of urine not associated with physical activity and leak of urine prior to reaching the bathroom are common patient complaints. Because cough-induced urinary leakage may be symptomatic of urge and SUI, this simple stress incontinence test should be performed in the office setting to rule out mixed incontinence (Dmochowski, 2006). Most women with uncomplicated urinary incontinence can be given a preliminary diagnosis at this point and treatment is initiated. Should initial management fail (usually after 8–12 week trial), or if complex conditions are present (e.g., pelvic organ prolapse, significant PVR), urodynamics or other specialized investigations are recommended.

**Treatment**

The treatment of UI often progresses from behavioral techniques (bladder training) to anticholinergic pharmacotherapy. In contrast to SUI, medical management of UI is consistently more efficacious. Options for nonresponders to drug therapy may include implanted sacral nerve stimulation (SNS). More invasive surgical procedures, including bladder reconstruction (augmentation) or urinary diversion for persistent severe UI, is rarely indicated.

Lifestyle modification includes fluid management, as large volumes can exacerbate urinary incontinence, and bladder training to correct voiding patterns, improve the ability to suppress urge, and increase bladder capacity. Maneuvers included pelvic muscle training, scheduled voiding, and relaxation techniques. The International Consultation on Continence recommends an initial voiding interval of 1 hour during waking hours, with weekly increases of 15–30 minutes until a 2–3 hour interval is achieved (Norton and Brubaker, 2006).
Anticholinergic agents, such as tolterodine, oxybutynin, and trospium, are considered first-line therapy for UI, suppressing or reducing involuntary bladder contractions, and also addressing the symptoms of idiopathic detrusor overactivity (OAB). Controlled clinical trials have demonstrated improvements in micturition frequency and episodes of incontinence, although side-effects of dry mouth and constipation can lead to discontinuation of treatment. Extended release formulations of oxybutynin (Ditropan XL) and tolterodine (Detrol LA) can be dosed once daily, increasing patient compliance. The recently introduced selective M3 receptor antagonists solifenacin (Vesicare) and darifenacin (Enablex) have also demonstrated good efficacy, safety, and patient tolerance in well-designed clinical trials for UI (Dmochowski, 2006).

Intravesical botulinum toxin A has shown significant initial promise for UI, OAB, and neurogenic detrusor overactivity, and may offer an important alternative to long-term oral pharmacotherapy or more invasive treatments. Quality of life, urgency, frequency, and bladder capacity improvements have been impressive to date. However, dosing, number and location of injection sites, and optimal treatment regimens have not been established as ongoing multicenter trials are yet to report their experience. Resiniferatoxin (an intravesical vanilloid) and oral beta-adrenergic agonists (nonspecific smooth muscle relaxants) are two other agents currently under investigation for UI and OAB treatment (Wein, 2006).

The results of randomized control trials provide evidence of SNS benefit for decreasing episodes of incontinence, pad use, voiding frequency, and improvement of bladder capacity and voided volume (Brazzelli et al, 2006). Although surgical revision was required for one-thirds of cases, no major irreversible complications were reported and benefits of SNS were reported to persist at follow-up of 3–5 years.

**MIXED URINARY INCONTINENCE**

Mixed urinary incontinence (MUI) refers to the occurrence of stress-related incontinence with symptomatic urinary urgency and UI. This disorder comprises an element of detrusor dysfunction (motor or sensory) and is associated with urethral sphincter underactivity. About one-thirds of incontinent patients have both UI linked to idiopathic overactivity and genuine SUI. Some experts now believe that MUI is now the predominant symptom grouping, with rates >50% reported in large population studies (Dmochowski, 2006). The relative incidence increases with advancing age, and occurs most commonly in women greater than 60 years old.

**Diagnosis**

The definition of MUI by the ICS emphasizes the presence of SUI and components of OAB (frequency and urgency) with or without UI, in the absence of known instigating factors. Urodynamically, detrusor overactivity is often noted. However, it should be emphasized that the underlying source of MUI may be a reflex response initiated by urine released into the proximal urethra during stress events. In this way, some individuals with SUI may mimic MUI due to a significant urge component associated with spontaneous urine loss. The diagnostics steps for MUI are the same as for SUI and are described in the “Stress Urinary Incontinence” subsection. In those individuals with equal bother (UI and SUI), or difficulty defining their symptoms, urodynamics may help define dysfunction and therapy.

**Treatment**

The presenting symptoms serve as a guide to initial therapeutic approach. The most bothersome aspect, SUI versus...
UI, is usually addressed first. If both types of incontinence are equally bothersome, treatment of the urge component is preferred in most cases.

Initial approaches include behavioral therapy, biofeedback and treatment with anticholinergics, with approximately 70% of patients experiencing symptomatic improvement with this class of medications; the notable exception is the patient with severe stress incontinence. Improvements in the total number of incontinent episodes, urinary frequency, urgency, and UI specifically, are noted (Chapple and Gormley, 2006). Once the initial treatment response is determined, further therapies can be initiated for persistent or secondary symptoms as outlined in the sections for SUI and UI, respectively.

Surgical outcomes for MUI have been reviewed for various sling techniques. Correction of a low-pressure outlet may benefit at least some patients with detrusor overactivity, although results for pure SUI remain superior. MUI symptom resolution has been demonstrated for upward of 70% of patients in some series, including a 4-year cure rate of 85% reported for the TVT approach (Dmochowski and Staskin, 2005). Current data support the use either midurethral or pubovaginal slings for MUI.

OVERFLOW INCONTINENCE

Overflow incontinence (OI) is defined as the involuntary loss of urine associated with bladder overdistension. Two primary processes are involved: urinary retention caused by bladder outlet obstruction or inadequate bladder contractions. Outflow obstruction may be secondary to BPH, bladder neck contracture or urethral stricture, or less commonly, prostate cancer in men, and due to cystocele, pelvic organ prolapse, or previous incontinence surgery in females. Impaired bladder emptying caused by decreased detrusor contractility may be the result of medications, spinal or peripheral nerve injuries, or due to long-standing overdistension. Diabetic cystopathy can result in OI as both sensory and contractile functions may be compromised. OI may also occur following transurethral prostatectomy (TURP), as urine flow is impeded by stricture, contracture, or residual adenoma.

Diagnosis

A similar approach is followed as outlined previously for the other incontinence subtypes. Reversible causes can usually be identified by patient history. Specific to OI, overdistended bladder is detected by measuring post-void residual volume with ultrasonography (preferred) or urethral catheterization immediately after the patient urinates. Normally, <50 mL of urine will remain in the bladder immediately following voiding and residual volumes of more than 200 mL indicate overdistension. Urodynamic testing and cystourethroscopy are used to determine the underlying cause, or differentiate OI from other incontinence states.

Treatment

Initial treatment of OI focuses on addressing reversible causes identified during patient evaluation such as cystocele, pelvic organ prolapse, impaired mobility, and so on.

Should such precipitating elements not be found, outlet obstruction may be treated conservatively, with adjustment of fluid intake and timed voiding. However, male patients will often require further intervention, including pharmacotherapy with alpha-adrenergic antagonists or 5α-reductase (finasteride)/dual 5α-reductase (dutasteride) inhibitors. If stricture or BPH is present, surgical intervention (TURP, incision of the bladder neck, visual internal urethrotomy) may offer definitive treatment.

For OI secondary to nonobstructive underactive detrusor, the first step is to decompress the bladder with an indwelling catheter or clean intermittent catheterization (CIC) for 7–14 days, while addressing potential reversible causes such as medications, infection, or constipation. An alpha-blocker may be initiated during this time period. Should voiding trials fail repeatedly in the patient with an contractile detrusor, CIC is the treatment of choice versus a permanent indwelling catheter (if feasible).

NEUROPATHIC INCONTINENCE

Neuropathic incontinence can be divided into 2 broad classifications: active and passive. Active neuropathic incontinence (neurogenic detrusor overactivity) is found in patients who have a spastic lesion but in whom the sphincteric mechanism, although not under voluntary control, still exerts adequate closure pressure. The presence of a hyperreflexive detrusor with uninhibited contractions increases the intravesical pressure. When intravesical pressure exceeds sphincteric pressure, there is a leakage of urine (Figure 29–20). Active incontinence is most often associated with suprasegmental, or upper motor neuron, lesions.

Passive neuropathic incontinence occurs when the sphincteric mechanism is weakened or completely lacking. Even without abnormally high intravesical pressures, any increase in intra-abdominal pressure results in urinary leakage. Passive incontinence is most often associated with lesions involving the micturition center or more distal lesions.

The more common classification of neurogenic incontinence is based on an evaluation of the functions of the lower urinary tract: incontinence owing to failure of the reservoir or to failure of retention.

A. FAILURE OF RESERVOIR FUNCTION

Loss of reservoir function in the contractile or contracted bladder can be caused by poor compliance in the detrusor muscle. Intravesical pressure rises with minimal bladder

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filling, exceeding the outlet resistance and causing urinary leakage. In contrast to the classification of active incontinence associated with suprasegmental lesions, failure of reservoir function may be found in patients who have meningomyelocele or exhibit other lower motor neuron lesions. Although these patients may have partial lesions with significant striated sphincteric activity offering some degree of resistance, early loss of bladder compliance increases intravesical pressure with minimal bladder filling and overcomes remaining outlet resistance. These patients, once recognized, must be managed aggressively because they often have a significant risk to the upper urinary tract, which can lead to possible vesicoureteral reflux, early renal deterioration, or lower ureteral obstruction.

Figure 29–20. Urodynamic recording in a patient with evidence of detrusor/sphincter dyssynergia, showing spontaneous activity in the bladder associated with a burst of activity in the external sphincter interrupting voiding. This represents a classic demonstration of upper motor neuron dysfunction leading to urinary incontinence as a result of detrusor hyperactivity or hyperreflexia.
B. FAILURE OF RETENTION FUNCTION

Complete lesions of the sacral segment or the cauda equina result in a total loss of smooth and striated sphincteric activity. The external sphincter offers minimal resistance. Most patients experiencing such failure can retain some volume, because the bladder musculature becomes atonic and intravesical pressure remains low, but any increase in intravesical pressure can cause leakage and the bladder never reaches full capacity. Consequently, the integrity of the upper urinary tract is not endangered, as in cases of reservoir failure.

Diagnosis

Diagnostic evaluation of neuropathic urinary incontinence determines whether the condition arises from detrusor or sphincteric dysfunction, or from a combination of the two. In exceptional cases, such as patients with multiple sclerosis with spinal stenosis or those with disk problems in addition to partial traumatic damage to the spinal cord, the neuropathy is less clear.

A complete urologic and neurologic history, physical examination, and urodynamics (cystometry, urethral pressure recordings, uroflowmetry) should be performed. Ultrasound evaluation, which can accurately measure renal size, and identify scarring, calculi, and hydronephrosis (obstruction or vesicoureteral reflux), is an appropriate baseline study for all patients with neuropathic lesion. This modality has largely replaced excretory urograms and retrograde cystourethrograms for screening. Other radiologic studies (voiding cystourethrogram, excretory urography, computed tomography scanning, magnetic resonance imaging) and neurologic studies (electromyography, evoked potentials) are performed as indicated. Cystourethroscopy is not recommended as part of the routine screening evaluation; if clinically indicated, it is used to assess the integrity of the urethra and identify stricture sites, diverticula, calculi, or other anatomic abnormalities.

The following are valuable in determining the underlying cause of incontinence:

1. Bladder responses to progressive filling
2. Sphincteric pressure profile and its response to progressive filling and the initiation of voiding
3. The presence of detrusor hyperreflexia and uninhibited activity (in patients with hyperreflexia)
4. Electromyographic studies of the striated urinary sphincter
5. In selected patients, response to neurostimulation of the sacral roots and pudendal nerve, with or without blocking and measurement of latencies

Differentiation among incontinence caused by uninhibited detrusor contractions, detrusor hyperreflexia, and poor bladder compliance is a straightforward process. Significant findings include sphincteric weakness, precipitous sphincteric relaxation, and decrease in pressure, as well as a lack of electromyographic activity of the pelvic floor musculature and external sphincter.

The proper diagnosis can be achieved by establishing the integrity of the sacral root reflex arc by neurostimulation of the sacral roots, with simultaneous recording of intravesical and intraurethral pressures at the levels of the internal and the external sphincter. One must be alert to the possibility of overlapping causes, and the responses of the external sphincter and pelvic floor musculature to active contraction, progressive bladder filling, and sacral root stimulation can be informative. The extent of one cause relative to another (eg, sphincteric weakness versus detrusor hyperreflexia) should be taken into account, and management should be directed toward the predominating cause of incontinence.

Treatment

Although diagnosis of the causes of incontinence is relatively easy, management can be challenging. The rehabilitation of the neuropathic condition and the alleviation of potentially damaging sequelae must be the guiding principles of management (see also Chapter 27—Neuropathic Bladder Disorders). Choices must be made according to the severity and potential progression of the lesion and the integrity of the system, with great care taken to prevent deterioration of the upper tracts. Especially in the previously mentioned cases of patients with spinal cord injury and meningomyelecele, early treatment provides the best chance of preserving the integrity of the entire urirary system.

1. Failure of Reservoir Function

Anticholinergic agents are commonly used to treat reservoir dysfunction. These medications inhibit the binding of acetylcholine to muscarinic receptors in the detrusor muscle, increasing bladder capacity and inhibiting involuntary contractions. Treatment with oxybutynin (Ditropan) 5 mg 2–3 times daily, tolteridone (Detro1) 2 mg twice daily, and single-dose extended release daily formulations (Ditropan XL and Detro1 ER), has proven to be highly effective and well tolerated (Chancellor et al, 2006). Dosages can be increased above these levels in some patients, dependent on tolerance of adverse effects and treatment response. Common side-effects include dry mouth, palpitations, constipation, nausea, or drowsiness, although newer extended-release formulations provide a reduced incidence versus immediate-release anticholinergics. Trospium chloride (Sanctura), and the selective M3 receptor antagonists solifenacin (Vesicare), darifenacin (Enablex), are newer anticholinergic options (Madersbacher and Rovner, 2006).

Propantheline bromide (Pro-Banthine) and imipramine hydrochloride (Tofranil) are considered second-line treatments for some patients, while the efficacy of flavoxate hydrochloride (Urispas) is unclear (Corcos et al, 2006).
Injection of botulinum-A toxin into the detrusor muscle for both children and adults who have failure of reservoir function is a promising new treatment that has demonstrated a significant increase in bladder capacity and compliance as well as symptomatic improvement for several weeks after cystoscopic injection (Kuo, 2006). This neurotoxin binds to the presynaptic nerve endings of cholinergic neurons, and leads to a temporary chemodenervation and the reduction of neural activity in the bladder. Further studies are required to clarify the clinical role of botulinum-A in these patients.

2. Failure of Retention Mechanism

CIC is the first line of conservative management in this group. Although such patients have very low outlet resistance, atonic bladder function may result in some retention. Most of the time, CIC every 4–6 hours avoids leakage. When CIC is not possible and the increase in intravesical pressure becomes excessive, the patient wears minimal protection against leakage. Pharmacotherapy has proven unsuccessful to date.

A. SURGICAL MANAGEMENT

1. Sphincterotomy—For spastic hyperreflexive male patients, this operation procedure can eliminate outlet resistance so that, with an external appliance or condom catheter, the bladder will remain empty. Although many consider this procedure the easiest way to preserve the upper urinary tract, it is clearly not rehabilitative and might interfere with other treatments.

2. Bladder augmentation—In patients with poor compliance owing to mural changes and chronic hypertrophy, augmentation improves reservoir function. If the sphincteric mechanism is adequate, detubularized bowel segments can be used in patients in whom the bladder will not expand even under anesthesia. Usually patients will be required to perform CIC postoperatively.

3. Artificial urinary sphincter (AUS)—For patients with severe sphincteric damage and a low-pressure, large-capacity bladder, the AUS is a useful option. In males it is applied around the bulbous urethra. When the device is deflated, the patient can void either by detrusor contraction (if some capability is preserved) or by straining and the Valsalva maneuver. A complete sphincterotomy can be performed first. CIC is hazardous in patients with an AUS and is rarely necessary.

4. Continent urinary diversion—This method should be considered only with progressively deteriorating upper urinary tract function, and even then, a simple conduit is often preferable.

5. Neurostimulation—In selected patients with detrusor hyperreflexia, stimulation of the sacral roots has effectively suppressed hyperactivity, relying on the known reflex response of the detrusor muscle to stimulate the somatic component of the sacral plexus (which aborts and inhibits detrusor contractility). If such patients with spinal cord injury, meningomyelocoele, multiple sclerosis, and other neuropahties show significant improvement after temporary testing, a permanent electrode can be placed over the most responsive root, usually S3.

6. Dorsal rhizotomy—Complete dorsal rhizotomy of S2–S4 extra- or intradurally effectively eliminates detrusor hyperreflexia and increases bladder capacity. Increases have been seen from a capacity of 150 or 200 mL to 600–800 mL. In patients with suprasegmental lesions and spastic upper motor neuron lesions, sacral root electrode implantation promotes detrusor contraction and bladder evacuation (bladder pacemaker).

REFERENCES


Chapple CR, Gormley EA: Developments in pharmacological therapy for the overactive bladder. BJU Int 2006;98(suppl 1):78.


Disorders of the adrenal glands result in classic endocrine syndromes such as Cushing’s syndrome, hyperaldosteronism, and catechol excess from pheochromocytoma (Figure 30–1). The diagnosis of these disorders requires careful endocrine evaluation and imaging with computed tomography (CT) or magnetic resonance imaging (MRI).

In addition, many adrenal lesions are discovered on cross-sectional imaging performed for other reasons. These “incidentalomas” require metabolic evaluation and assessment to determine their need for treatment.

**DISEASES OF THE ADRENAL CORTEX**

**CUSHING’S SYNDROME**

Cushing’s syndrome is the clinical disorder caused by overproduction of cortisol. Most cases (80%) are due to bilateral adrenocortical hyperplasia stimulated by overproduction of pituitary adrenocorticotropic hormone (corticotropin, ACTH), known as Cushing’s disease. About 10% of cases are due to the ectopic production of ACTH from nonpituitary tumors. Ectopic ACTH production occurs most frequently in small-cell lung carcinoma; other tumors producing ACTH include carcinoids (lung, thymic, gastrointestinal tract), islet cell tumors of the pancreas, medullary thyroid carcinoma, pheochromocytoma, and small-cell carcinoma of the prostate. Adrenal adenoma is the cause in 5% of cases and carcinoma in 5%. In children, adrenocortical carcinoma is the most common cause of Cushing’s syndrome.

**Pathophysiology**

Overproduction of cortisol by adrenocortical tissue leads to a catabolic state. This causes liberation from muscle tissue of amino acids, which are transformed into glucose and glycogen in the liver by gluconeogenesis. The resulting weakened protein structures (muscle and elastic tissue) cause a protuberant abdomen and poor wound healing, generalized muscle weakness, and marked osteoporosis, which is made worse by excessive loss of calcium in the urine.

In addition, glucose is transformed largely into fat and appears in characteristic sites such as the abdomen, supraclavicular fat pads, and cheeks. There is a tendency to diabetes, with an elevated fasting plasma glucose level in 20% of cases and diabetic glucose tolerance curve in 80%.

The cortisol excess also suppresses the immune mechanisms, which makes these patients susceptible to repeated infection. Inhibition of fibroblast function by excess cortisol further interferes with wound healing and host defenses against infection.

Hypertension is present in 90% of cases. Although the aldosterone level is not usually elevated, cortisol itself exerts a hypertensive effect when present in excessive amounts, as does 11-deoxycorticosterone. The hypertension may be accompanied by manifestation of mineralocorticoid excess (hypokalemia and alkalosis), especially in patients with the ectopic ACTH syndrome or adrenocortical carcinoma.

**Pathology**

The cells in adrenal hyperplasia resemble those of the zona fasciculata of the normal adrenal cortex. Frank adenocarcinoma reveals pleomorphism and invasion of the capsule, the vascular system, or both (Figure 30–2). Local invasion may occur, and metastases are common to the liver, lungs, bone, or brain. Histologic differentiation between adenoma and adenocarcinoma is frequently difficult.

In the presence of adenoma or malignant tumor, atrophy of the cortices of both adrenals occurs because the main secretory product of the tumor is cortisol, which inhibits the pituitary secretion of ACTH. Thus, although the tumor continues to grow, the contralateral adrenal cortex undergoes atrophy.

**Clinical Findings**

**A. SYMPTOMS AND SIGNS (FIGURES 30–3 AND 30–4)**

The presence of at least 3 of the following strongly suggests Cushing’s syndrome:

1. Obesity (with sparing of the extremities), moon face, and fat pads of the supraclavicular and dorso-cervical areas (buffalo hump).
2. Striae (red and depressed) over the abdomen and thighs.
3. Hypertension (almost always present).
4. Proximal myopathy with marked weakness, especially in the quadriceps femoris, making unaided rising from a chair difficult.
5. Emotional lability, irritability, difficulty in sleeping, and sometimes psychotic personality.
6. Osteoporosis (common), with back pain from compression fractures of the lumbar vertebrae as well as rib fractures.
7. In 80% of cases, postprandial hyperglycemia is present, and in 20% there is an elevated fasting plasma glucose level.
8. To a variable extent, there are features of adrenal androgen excess in women with Cushing’s syndrome; these are absent in the case of adenoma, most severe with carcinoma, and present to an intermediate degree with Cushing’s disease. They consist of recession of the hairline, hirsutism, small breasts, and generalized muscular overdevelopment, with deepening of the voice.

It is not possible to differentiate the cause from the clinical presentation alone.

**B. LABORATORY FINDINGS**

The leukocyte count may be elevated to the range of 12,000–20,000/µL, usually with fewer than 20% lymphocytes. Eosinophils are few in number or absent. Polycythe-
mia is present in over half the cases, with the hemoglobin ranging from 14 to 16 g/dL. Anemia, however, may occur in patients with malignant tumors ectopically secreting ACTH.

Blood chemical analyses may show an increase in serum Na⁺ and CO₂ levels and a decrease in serum K⁺ levels. Hyperglycemia may occur.

1. Specific tests for Cushing’s syndrome—The following tests are performed to determine whether the patient has Cushing’s syndrome or is an anxious individual with elevated plasma levels of cortisol.

   a. 24-hour urinary cortisol level—Urine cortisol is measured in a 24-hour urine collection (normal range, 10–50 µg/24 h). A urine cortisol value more than twofold elevated is typical of Cushing’s syndrome. False-positive elevations can occur in acute illness, depression, and alcoholism. However, obesity does not raise the level of urinary free cortisol above normal.

   b. Suppression of ACTH and plasma cortisol by dexamethasone—Dexamethasone in low doses is used to assess the feedback suppression of ACTH and cortisol production by glucocorticoids. If dexamethasone is given at 11 PM, ACTH is suppressed in normal persons but not in those with Cushing’s syndrome. Dexamethasone is useful because it has 30 times the potency of cortisol as an ACTH suppressant and it is not measured in current plasma or urine cortisol methods.

   The procedure is to give 1 mg of dexamethasone by mouth at 11 pm and to draw blood at 8–9:00 am for measurement of plasma cortisol. If the level is <5 µg/dL (normal is 5–20 µg/dL), Cushing’s syndrome can be ruled out. If the value is >10 µg/dL, Cushing’s syndrome is present. A level in the range of 5–10 µg/dL is equivocal, and the test should be repeated or the urine cortisol may be measured.

   Women taking birth control pills have high plasma cortisol levels because, as in pregnancy, the estrogen stimulates production of the cortisol-binding globulin. The pills must be withheld for at least 3 weeks before the dexamethasone suppression test. Other conditions causing false-positive responses are acute illness, depression, and alcoholism. Also, about 15% of obese patients do not suppress cortisol with this test.

2. Specific tests for differentiation of causes of Cushing’s syndrome—The various causes of Cushing’s syndrome can be determined with great accuracy (95% of cases).

Figure 30–2. Left: Histologic appearance of a typical benign adenoma of the adrenal cortex made up of a large number of identical cells from the zona fasciculata removed from a 39-year-old woman with Cushing’s syndrome. Right: Section of an adenocarcinoma removed from a 36-year-old woman with metastatic adenocarcinoma showing significant pleomorphism of the cells. Invasion of a large vein is not shown in this micrograph. Note that benign adenomas will occasionally have this appearance but without invasion of the bloodstream. (Reproduced, with permission, from Forsham PH: The adrenal cortex. In: Williams RH [editor]: Textbook of Endocrinology. 4th ed. Saunders, 1968.)
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a. Plasma ACTH level—If the diagnosis of Cushing’s syndrome has been established, this test will differentiate ACTH-dependent causes (Cushing’s disease and the ectopic ACTH syndrome) from adrenal tumors, which are ACTH-independent. The normal range is 10–50 pg/mL. Patients with Cushing’s disease have ACTH levels that range from 10 to 200 pg/mL; in the ectopic ACTH syndrome, levels are usually >200 pg/mL; and patients with adrenal tumors have suppressed ACTH levels (<5 pg/mL with the IRMA ACTH assay) and thus are easily differentiated.

b. Plasma androgen levels—In patients with adrenal adenomas, androgen levels are normal or low, and in adrenocortical carcinoma these levels are often markedly elevated.

C. X-RAY FINDINGS AND SPECIAL EXAMINATIONS

1. Localization of source of ACTH excess—When tests suggest Cushing’s disease or the ectopic ACTH syndrome and an elevated plasma level of ACTH is present, the source of ACTH must be identified. Because the great majority of these patients have Cushing’s disease and because most of the patients with ectopic ACTH secretion have an obvious malignancy, the first step is to perform pituitary MRI. These are positive in 50–60% of patients with Cushing’s disease; in the remainder, the diagnosis should be established by the sampling of ACTH levels in the venous drainage of the anterior pituitary, that is, the cavernous sinuses and inferior petrosal sinuses. If the MRI and venous sampling do not reveal a pituitary source of ACTH, CT scans of the chest and abdomen are used to localize an ectopic tumor.

2. Localization of adrenal lesions—Patients with Cushing’s syndrome with suspected adrenal tumors and suppressed ACTH levels should undergo a CT scan of the abdomen with 3-mm sections through the adrenals. Adrenal tumors causing Cushing’s syndrome are usually >3 cm in diameter (Figure 30–5) and are therefore easily visualized. Adenomas are usually 3–6 cm in diameter; carcinomas are usually >5 cm in diameter (Figure 30–6) and are frequently locally invasive or metastatic to the liver and lungs at the time of diagnosis. In patients with adrenal tumors, the contralateral adrenal is suppressed and therefore appears atrophic or normal on CT scan. The finding of bilateral adrenal enlargement is typical of Cushing’s disease or the ectopic ACTH syndrome. Ultrasound or MRI
may also be used for adrenal localization, although these techniques do not appear to offer significant advantage over CT scans.

**Complications**

Hypertension may lead to cardiac failure or stroke. Diabetes may be a problem but is usually mild. Intractable skin or systemic infections are common. Compression fractures of osteoporotic vertebrae and rib fractures may develop. Renal stones are not uncommon as a result of bone resorption. Psychosis is not uncommon; it usually subsides after successful surgery.

**Treatment**

**A. CUSHING’S DISEASE**

A pituitary microadenoma, which is the most common cause of bilateral adrenocortical hyperplasia, must be located and removed surgically. Transsphenoidal resection performed by an experienced neurosurgeon is the method of choice. Success is reported in >80% of cases, and in most instances the endocrine functions of the pituitary gland are preserved.

**B. ECTOPIC ACTH SYNDROME**

The treatment of these patients is difficult because most have an advanced malignancy and severe hypercortisolism. Removal of the primary tumor is clearly the therapy of choice; however, curative resection is limited to the few patients with benign tumors such as bronchial carcinoids. Patients with residual or metastatic tumors should be managed first with adrenal inhibitors, and if that is not successful, bilateral adrenalectomy should be considered.

**C. TOTAL BILATERAL ADRENALECTOMY**

Total bilateral adrenalectomy is indicated in patients with Cushing’s disease in whom the pituitary tumor is not resectable and in whom radiotherapy and medical therapy fail to control the cortisol excess. At present, it is best to perform bilateral adrenalectomy via a laparoscopic approach. The procedure significantly decreases morbidity and length of hospital stay. Bilateral adrenalectomy is also indicated in patients with ectopic ACTH syndrome who have life-threatening hypercortisolism that cannot be controlled by inhibitors of adrenal secretion.

1. **Preoperative preparation**—Because removal of the source of excessive cortisol will inevitably lead to temporary or permanent adrenal insufficiency, it is of the utmost importance to administer cortisol preoperatively and to continue substitution therapy after surgery to control Addison’s disease. In the postoperative period, the dose is tapered downward until oral medication provides sufficient control.

2. **Postoperative status**—The patient feels moderately well following removal of the source of excess ACTH or adrenalectomy or while receiving a high dose of hydrocortisone in excess of the usual daily output of approximately 20 mg. It is important to reduce the steroid substitution gradually over a period of several days. On the day of operation, 200 mg of cortisol is given; the dosage is then reduced gradually on successive days (150, 100, 80, 60,
and 40 mg) until a maintenance dosage of 20–30 mg cortisol combined with 0.1 mg fludrocortisone is reached.

**D. Adrenal Adenoma and Adenocarcinoma**

Virtually all adrenal adenomas and smaller adrenal carcinomas are now removed laparoscopically, again allowing decreased hospital stay and more rapid recovery from surgery. Adrenal carcinomas that are large (>8–10 cm) are likely to be metastatic or locally invasive. Thus, if there is evidence of invasion of adjacent structures or invasion of the adrenal or renal veins or the vena cava, these tumors are best approached by a traditional abdominal incision.

1. **Preoperative preparation**—Preoperative preparation is the same as that for bilateral hyperplasia, since, in this case, the remaining adrenal gland will be atrophic and thus the patient will be hypoadrenal.

2. **Postoperative treatment and follow-up**—Cortisol is administered perioperatively in the doses described above and then tapered to a replacement dose of 20–30 mg/day. Hydrocortisone is given orally in a dosage of 10 mg three times daily initially and reduced within 2–3 weeks to 10 mg daily given at 7 or 8 am. Substitution therapy may be necessary for 6 months to 2 years depending on the rate of recovery of the residual gland. Mineralocorticoid therapy is rarely necessary, since the atrophic adrenal usually produces sufficient aldosterone. Patients with adrenocortical carcinoma are usually not cured by surgery and require additional therapy.

**E. Medical Therapy**

There is no effective method of inhibiting ACTH secretion; however, adrenal hypersecretion can be controlled in many patients by inhibitors of adrenal cortisol secretion. Medical therapy is indicated in patients who either cannot undergo surgery (eg, because of debility, recent myocardial infarction) or in those who have had unsuccessful resection of their pituitary, ectopic, or adrenal tumor.

Ketoconazole is the current drug of choice; it blocks cortisol secretion by inhibiting P450c11 and P450scc. The total dose required is 800–1600 mg/day given in 2 divided doses. Side effects are adrenal insufficiency, abnormal liver function tests, and hepatotoxicity in a few patients.

Metyrapone may be used alone or may be added if ketoconazole alone is unsuccessful in normalizing cortisol levels. The usual dosage is 1–4 g daily given in 4 divided doses.

Aminoglutethimide and trilostane also inhibit adrenal secretion, but they are uncommonly used at present.

Mitotane (o,p'-DDD, Lysodren) is both an inhibitor of adrenal secretion and a cytotoxic agent that damages adrenocortical cells. It is used almost exclusively in patients with residual adrenocortical carcinoma, in whom it helps to reduce cortisol hypersecretion. The usual dosage is 6–12 g daily in 3–4 divided doses. About 70% of patients achieve a reduction in steroid secretion and 35% achieve a reduction in tumor size; however, there is no convincing evidence that the drug prolongs survival. Side effects occur in 80% of patients and include nausea, vomiting, diarrhea, depression, and somnolence.

**Prognosis**

Treatment of hypercortisolism usually leads to disappearance of symptoms and many signs within days to weeks, but osteoporosis usually persists in adults, whereas hypertension and diabetes often improve. Cushing’s disease treated by pituitary adenomectomy has an excellent early prognosis, and long-term follow-up shows a recurrence rate of about 10%. Patients with the ectopic ACTH syndrome and malignant tumors in general have a poor prognosis; these patients usually die within several months of diagnosis. Patients with benign lesions may be cured by resection of the tumor. Removal of an adrenal adenoma offers an excellent prognosis; and these patients are cured by unilateral adrenalectomy.

The outlook for patients with adrenocortical carcinoma is poor. The antineoplastic drug mitotane reduces the symptoms and signs of Cushing’s syndrome but does little to prolong survival. Radiotherapy and chemotherapy are not successful in these patients.

**ADRENAL ANDROGENIC SYNDROMES**

Adrenal androgenic syndromes are more common in females. Congenital bilateral adrenal hyperplasia and tumors, both benign and malignant, may be observed. In contrast to Cushing’s syndrome, which is protein catabolic, the androgenic syndromes are anabolic. In untreated cases, there is a marked recession of the hairline, increased beard growth, and excessive growth of pubic and sexual hair in general in both sexes. In males, there is enlargement of the penis, usually with atrophic testes; in females, enlargement of the clitoris occurs, with atrophy of the breasts and amenorrhea. Muscle mass increases and fat content decreases, leading to a powerful but trim figure. The voice becomes deeper, particularly in females; this condition is irreversible, because it is due to enlargement of the larynx. In both sexes there may be increased physical sexual aggressiveness and libido.

1. **Congenital Bilateral Adrenal Androgenic Hyperplasia**

**Pathophysiology**

A congenital defect in certain adrenal enzymes results in the production of abnormal steroids, causing pseudohermaphroditism in females and macrogenitosomia in males. The enzyme defect is associated with excess androgen production in utero. In females, the Müllerian duct
structures (eg, ovaries, uterus, and vagina) develop normally, but the excess androgen exerts a masculinizing effect on the urogenital sinus and genital tubercle, so that the vagina is connected to the urethra, which, in turn, opens at the base of the enlarged clitoris. The labia are often hypertrophied. Externally, the appearance is that of severe hypospadias with cryptorchidism.

The adrenal cortex secretes mostly anabolic and androgenic steroids, leading to various degrees of cortisol deficiency depending on the nature of the enzyme block. This increases the secretion of ACTH, which causes hyperplasia of both adrenal cortices. The cortices continue to secrete large amounts of inappropriate anabolic, androgenic, or hypertensive steroids. Absence or reduction of the usual tissue concentration of various enzymes accounts for blocks in the adrenocortical synthetic pathways.

A block at P450scc leads to the rare congenital lipoid adrenal hyperplasia with complete absence of any steroid hormone production; the infant will die at an early age unless full substitution therapy is given for life.

A block at 3β-hydroxydehydrogenase/isomerase enzyme prevents formation of progesterone, aldosterone, and cortisol. Dehydroepiandrosterone (DHEA) is produced in excess. This uncommon syndrome is characterized by adrenal insufficiency and male pseudohermaphroditism, with females showing unusual sexual development with hirsutism.

A block at P450c21, or 21-hydroxylase deficiency, which is the most common cause of congenital adrenal hyperplasia, does not allow for the transformation of 17α-hydroxyprogesterone to cortisol. This common deficiency occurs in 2 forms: the salt-losing variety, with low to absent aldosterone, and the more frequent non-salt-losing type. Infants present with adrenal insufficiency and ambiguous genitalia; older children develop pseudoprecocious puberty and accelerated growth and skeletal maturation.

A block at P450c17 with lack of 17α-hydroxylase occurs mostly in females and may not be discovered until puberty. Findings include low cortisol levels with high ACTH levels, primary amenorrhea, and sexual infantilism, as neither the glucocorticoids nor the sex steroids are produced in adequate amounts. Rarely, there is male pseudohermaphroditism. Hypertension due to excess mineralocorticoids, notably 11-deoxycorticosterone, is characterizedly present.

A block at P450c11 with lack of 11β-hydroxylase prevents formation of cortisol and corticosterone and thus leads to overproduction of adrenal androgens and 11-deoxycorticosterone. Patients usually have clinical features of mild androgen excess with hypertension and hypokalemia.

A block at P450aldo results in the inability to produce aldosterone in the zona glomerulosa; these patients present with isolated mineralocorticoid deficiency with hypotension and hyperkalemia.

Clinical Findings

A. Symptoms and Signs

In newborn girls, the appearance of the external genitalia resembles severe hypospadias with cryptorchidism. Infant boys may appear normal at birth. The earlier in intrauterine life the fetus has been exposed to excess androgen, the more marked the anomalies.

In untreated cases, hirsutism, excess muscle mass, and, eventually, amenorrhea are the rule. Breast development is poor. In males, growth of the phallus is excessive. The testes are often atrophic because of inhibition of gonadotropin secretion by the elevated androgens. On rare occasions, hyperplastic adrenocortical rests in the testes make them large and firm. In most instances, there is azoospermia after puberty.

In both males and females with androgenic hyperplasia, the growth rate is initially increased, so that they are taller than their classmates. At about age 9–10 years, premature fusion of the epiphyses caused by excess androgen causes termination of growth, so that these patients are short as adults.

B. Laboratory Findings

Urinary 17-ketosteroid levels are higher than normal for sex and age, and plasma anchostenedione, DHEA, DHEA-S, and testosterone are elevated. Plasma ACTH is also elevated, and in patients with the most common defect (ie, 21-hydroxylase deficiency), plasma 17α-hydroxyprogesterone is markedly elevated. Chromosome studies are normal.

C. X-Ray Findings

X-rays show acceleration of bone age.

D. CT Scans

Scans usually show the hypertrophied adrenals.

E. Urologic Evaluation

This is indicated to define the anatomic abnormalities.

Differential Diagnosis

Several congenital anomalies that affect the development of the external genitalia resemble adrenal androgenic syndrome. These include (1) severe hypospasias with cryptorchidism, (2) female pseudohermaphroditism of the non-adrenal type (caused by administration of androgens or gestational compounds during the pregnancy), (3) male pseudohermaphroditism, and (4) true hermaphroditism. These children show no hormonal abnormalities, and bone age and maturation are not accelerated.

Treatment

It is imperative to make the diagnosis early. Treatment of the underlying cause is medical, with the goal of suppressing
excessive ACTH secretion, thus minimizing excess androgenicity. This is accomplished by adrenal replacement with cortisol or prednisone in doses sufficient to suppress adrenal androgen production and therefore prevent virilization and rapid skeletal growth. In patients with mineralocorticoid deficiency, fludrocortisone (0.05–0.3 mg, depending on severity and age) together with good salt intake is necessary to stabilize blood pressure and body weight.

After puberty, the vagina can be surgically separated from the urethra and opened in the normal position on the perineum. Judicious administration of estrogens or birth control pills feminizes the figure in pseudohermaphrodites and improves their psyche considerably.

**Prognosis**

If the condition is recognized early and ACTH suppression is begun even before surgical repair of the genital anomaly, the outlook for normal linear growth and development is excellent. Delay in treatment inevitably results in stunted growth. In some female pseudohermaphrodites, menses begins after treatment, and conception and childbirth can occur when the anatomic abnormalities are minimal or have been surgically repaired.

**2. Adrenocortical Tumors**

Adrenocortical tumors producing androgens are most frequently carcinomas; however, a few benign adenomas have been reported. Most of the carcinomas also hypersecrete other hormones (ie, cortisol or 11-deoxycorticosterone), and thus the clinical presentation is variable. Female patients present with androgen excess, which may be severe enough to cause virilization; many of these patients also have Cushing’s syndrome and mineralocorticoid excess (hypertension and hypokalemia). In adult males excess androgens may cause no clinical manifestations, and diagnosis in these patients may be delayed until there is abdominal pain or an abdominal mass. These patients may also present with Cushing’s syndrome and mineralocorticoid excess.

The tumor can be located by CT scan, which is also used to define the extent of tumor spread. Local invasion and distant spread to the liver and lungs are common at the time of diagnosis. The primary therapy is surgical resection of the adrenal tumor, as discussed above; however, surgical cure is rare. These patients are subsequently treated with mitotane and other adrenal inhibitors, as discussed in the section on Cushing’s syndrome.

**THE HYPERTENSIVE, HYPOKALEMIC SYNDROME (PRIMARY ALDOSTERONISM)**

Excessive production of aldosterone, due mostly to aldosteronoma or to spontaneous bilateral hyperplasia of the zona glomerulosa of the adrenal cortex, leads to the combination of hypertension, hypokalemia, nocturia, and polyuria. A syndrome resembling nephrogenic diabetes insipidus may occur as a result of reversible damage to the renal collecting tubules. The alkalosis may produce tetany.

**Pathophysiology**

Excessive aldosterone, acting on most cell membranes in the body, produces typical changes in the distal renal tubule and the small bowel that lead to urinary potassium loss together with increased renal sodium reabsorption and hydrogen ion secretion. This results in potassium depletion, metabolic alkalosis, increased plasma sodium concentration, and hypervolemia. With low serum levels of potassium, the concentrating ability of the kidney is lowered and the tubules no longer respond to the administration of vasopressin by increased reabsorption of water. Finally, impairment of insulin release secondary to potassium depletion increases carbohydrate intolerance in about 50% of cases.

Plasma renin and, secondarily, plasma angiotensin are depressed by excess aldosterone as a result of blood volume expansion. Early in the course of excess aldosterone production, there may be hypertension with a normal serum potassium level. Later, the potassium level will be low as well, and this suggests the diagnosis.

**Clinical Findings**

**A. SYMPTOMS AND SIGNS**

Hypertension is usually the presenting manifestation, and the accompanying hypokalemia suggests mineralocorticoid excess. Headaches are common, nocturia is invariably present, and rare episodes of paralysis occur with very low serum potassium levels. Numbness and tingling of the extremities are related to alkalosis that may lead to tetany.

**B. LABORATORY FINDINGS**

Before the tests outlined below are done, one must ascertain that the patient is not taking oral contraceptives or other estrogen preparations, since these may increase renin and angiotensin levels and therefore aldosterone levels, thus raising the blood pressure artificially. Withdrawal of these medications for 1 week is mandatory. Diuretics must also be discontinued, since they lower blood volume and induce secondary aldosteronism and hypokalemia. Also, if the patient is following a salt-restricted diet, aldosterone is normally elevated.

In true aldosterone excess, serum sodium is slightly elevated and CO₂ increased, whereas serum potassium is very low, for example, 3 mEq/L or less. Urine and serum potassium determinations while the patient is receiving good sodium replacement provide a screening test. Potassium wasting is established if the urinary potassium level is <30
Definitive diagnosis rests on demonstration of an elevated urine or plasma aldosterone level. The initial step is to obtain simultaneous plasma aldosterone and plasma renin levels. If the aldosterone is elevated and the renin is suppressed with a ratio of <20:1, the diagnosis is established. Further confirmation can be obtained by demonstrating an elevated aldosterone level in a 24-hour urine sample.

C. LOCALIZATION

A thin-section CT scan is the initial procedure and will localize an adenoma in approximately 90% of patients (Figure 30–7). If no adenoma is visualized, adrenal vein sampling of aldosterone and cortisol will correctly differentiate adenoma from hyperplasia in virtually all cases.

Differential Diagnosis

Secondary hyperaldosteronism may accompany renovascular hypertension. This too is associated with hypokalemic alkalosis; however, the renin level is elevated rather than suppressed. Essential hypertension does not cause changes in the electrolyte pattern. Definitive tests for hyperaldosteronism show negative results.

Treatment

A. ALDOSTERONOMA

If the site of the tumor has been established, only the affected adrenal need be removed. Again, the procedure of choice is laparoscopic unilateral adrenalectomy, which is highly successful at resolving the metabolic defect.

B. BILATERAL NODULAR HYPERPLASIA

Most authorities do not recommend resection of both adrenals, since the fall in blood pressure is only temporary and electrolyte imbalance may continue. Medical treatment is recommended.

C. MEDICAL TREATMENT

If surgery must be postponed, if the hypertension is mild in an older person, or if bilateral hyperplasia is the cause, one may treat medically with spironolactone (Aldactone), 25–50 mg orally four times daily. Amiloride, a potassium-sparing diuretic, may be given in doses of up to 20–40 mg/day. Other antihypertensive agents may also be necessary.

Prognosis

Following removal of an adrenal adenoma, the hypokalemia resolves. Seventy percent of patients become normotensive and 50% show some lowering of hypertension. Bilateral nodular hyperplasia is not amenable to surgical treatment, and the results of medical treatment are only fair.

DISEASES OF THE ADRENAL MEDULLA

PHEOCHROMOCYTOMA

Pheochromocytoma, derived from the neural crest, is one of the surgically curable hypertensive syndromes. There is no sex predilection. Pheochromocytoma accounts for fewer than 1% of cases of hypertension, but it is readily diagnosed if the possibility is kept in mind. It usually occurs spontaneously, but 10% of cases occur in patients with other disorders such as neurofibromatosis or familial syndromes such as multiple endocrine neoplasia type II or von Hippel-Lindau disease. The tumor is bilateral or extra-adrenal in 10% of cases in adults and in an even greater percentage in children and is then most often familial.
Clinical Findings

A. Symptoms and Signs

Hypertension is both systolic and diastolic. Hypertension may be either sustained and indistinguishable from ordinary blood pressure elevation, or paroxysmal, coming on for variable lengths of time and then subsiding to normal levels. Such attacks are usually precipitated by trigger mechanisms of various sorts, for example, emotional upsets or straining at stool.

Headache is a frequent complaint and is commensurate in severity with the degree of hypertension. Increased sweating without appropriate causes such as exertion or environmental heat resembles the phenomenon seen during menopause and may be accompanied by flushing or blanching. Tachycardia with palpitations occurs mainly as a consequence of epinephrine rather than norepinephrine excess.

Postural hypotension is a frequent finding, as a result of diminished plasma volume. Profound weakness may occur after an attack of hypertension. Weight loss is common.

Decreased gastrointestinal motility, anxiety, and psychic instability are also common and due to excess circulating catechols.

B. Biochemical Diagnosis

The choice of biochemical test and whether to measure plasma or urine values remain controversial, but certain principles are clear: (1) Screening the hypertensive population is not recommended because of the low incidence of pheochromocytoma (about 0.1%). (2) Patients with pheochromocytoma who have sustained hypertension usually have clearly elevated catecholamines or metabolites in both urine and plasma. More than 80% of these patients have urine values two times greater than normal plasma catecholamine (Epi + Norepi) >2000 ng/L. Levels of this magnitude are unusual in patients without pheochromocytoma except in acute major illness. (3) Patients with only episodic hypertension may have normal random plasma catecholamine levels and normal 24-hour urine values. Evaluation of these patients must be directed to obtaining plasma catecholamine during an episode or having the patient collect timed urine values (eg, 2–4 hours) from the onset of an episode. (4) Suppression or stimulation tests are not recommended except in the rare instances when the diagnosis cannot be established by routine procedures.

1. Urinary measurements—Urinary measurements are the traditional diagnostic procedure. Normal values are shown in Table 30–1, and recent series in patients with pheochromocytoma are summarized in Table 30–2. These data suggest that measurements of metanephrines or catecholamine are more useful than measurement of vanillylmandelic acid, since >80% of patients have values that were elevated more than two times. Spot urinary metanephrines (MN) and normetanephrines (NMN) measured by radioimmunoassay are very simple and highly accurate.

| Table 30–1. 24-Hour Urine Measurements in Patients with Pheochromocytoma.* |
|----------------------------------|---------|--------------------|--------------------|
| **Urine**                       |         |                    |                    |
| Norepinephrine: 10–100 µg/24 h   |         |                    |                    |
| Epinephrine: Up to 20 µg/24 h    |         |                    |                    |
| Normetanephrine and metanephrine: < 1.5 mg/24 h |         |                    |                    |
| Vanillylmandelic acid (VMA): 2–9 mg/24 h |         |                    |                    |
| **Plasma**                      |         |                    |                    |
| Norepinephrine: 100–200 pg/mL    |         |                    |                    |
| Epinephrine: 30–50 pg/mL         |         |                    |                    |


At a cutpoint of 500 ng/mL creatinine for either MN or NMN, Ito et al. (1998) reported 100% sensitivity and specificity. Patients with only episodic symptoms or episodic hypertension should be studied with shorter urine collections if 24-hour studies are normal.

2. Plasma catecholamines—These values, when measured by specific methods, are elevated in most patients with pheochromocytoma; however, the frequency of false-positive values limits diagnostic utility. Thus, in patients with pheochromocytoma and sustained hypertension, 85% have plasma catecholamine values >2000 ng/L. When patients with only paroxysmal hypertension are included, however, only 75% have values >2000 ng/L. Values between 600 and 2000 ng/mL are commonly obtained in stressed or anxious patients without pheochromocytoma.

This is especially true if samples are obtained by venipuncture without prior placement of an intravenous line with the patient supine for 30 minutes. Plasma catecholamine measurements do have a role, though, because markedly elevated levels during an episode may be diagnostic; conversely, the finding of normal values during episodes of severe hypertension essentially excludes the diagnosis.

Tumor Localization

Pheochromocytomas are intra-abdominal in 98% of cases, and 90% are intra-adrenal (10% are bilateral, especially in

| Table 30–2. Catecholamines in Urine and Plasma.* |
|----------------------------------|---------|--------------------|--------------------|
|                                  | Normal  | 1–2X Elevated     | > 2X Elevated      |
|                                  | No. (%) | No. (%)            | No. (%)            |
| VMA (n = 384)                    | 41 (11) | 86 (22)            | 257 (67)           |
| MN (n = 271)                     | 12 (5)  | 33 (12)            | 226 (83)           |
| UFC (n = 319)                    | 14 (4)  | 30 (10)            | 275 (86)           |

The values listed represent the means of the normal ranges, which vary for each laboratory.

familial syndromes). Extra-adrenal pheochromocytomas are usually within the abdomen and are located along the sympathetic chain, the periaortic areas, and at the bifurcation of the aorta. The tumors may also arise from the bladder. Extra-abdominal pheochromocytomas occur in posterior mediastinum, rarely in the heart or pericardium and rarely in the neck. Tumors <2 cm in diameter are rare, and most are >3 cm (Figure 30–8). Thus, the vast majority of pheochromocytomas are larger than the lower limits of resolution of current imaging techniques.

A. CT Scans

CT is currently the initial imaging procedure of choice; with current technology, it demonstrates virtually all intra-abdominal tumors and most of those that are extra-adrenal. Small tumors in the abdomen, pelvis, and chest may be obscured by surrounding structures. CT is not useful in determining whether an adrenal mass is in fact a pheochromocytoma (ie, if the adrenal mass is found coincidentally or the catechol determinations are equivocal, the adrenal mass could be a nonfunctioning adenoma). In this case, MRI or metaiodobenzylguanidine (MIBG) techniques may be useful.

B. MRI

The accuracy of detecting pheochromocytoma with MRI is as good as that obtained with CT, but the cost is greater at most institutions. MRI has the advantage of greater diagnostic specificity in that T2-weighted images or those obtained with gadolinium enhancement show greater signal intensity of the pheochromocytoma (compared with liver) than that obtained with adrenal adenomas. Limited data suggest that MRI may be superior to CT in localizing extra-adrenal tumors.

C. MIBG Scanning

Radionuclide scanning with MIBG has assumed a prominent role in the localization of pheochromocytomas. The compound is taken up by pheochromocytomas, ganglioneuromas, neuroblastomas, and other neural crest tumors as well as some carcinoids. MIBG scans are positive in about 85–90% of patients with pheochromocytomas. The procedure is useful, however, since false-positive results are rare and a positive scan in the presence of elevated catechols is diagnostic. In addition, MIBG scans have great utility in the localization of (1) small lesions, (2) extra-adrenal lesions, (3) bilateral lesions, and (4) metastatic deposits in patients with malignant tumors.

Diagnostic Strategy

Patients in whom there is a high index of clinical suspicion and those who have a greater than twofold elevation of urine catechols should undergo an adrenal CT scan. If the CT scan reveals a unilateral tumor and the contralateral adrenal is normal, the diagnosis is established. Patients with familial syndromes and those in whom cancer is suspected should undergo MIBG scanning to determine the extent of disease. If the adrenal CT is negative, MIBG scanning or MRI of the chest and abdomen is indicated to localize the tumor. This approach localizes virtually all tumors.

If the clinical suspicion is low and urine catechols are normal, imaging procedures are not indicated. However, it is not infrequent that patients at low risk on the basis of clinical manifestations have persisting mild elevations of catecholamines. In this situation, a single negative adrenal imaging procedure should suffice to terminate the evaluation, and the patient may be followed clinically and reevaluated if appropriate.

Therapy

A. Preoperative Management

Once the diagnosis of pheochromocytoma is established, the patient should be prepared for surgery to reduce the
incidence of intraoperative complications and postoperative hypotension. The greatest experience is with the long-acting alpha-adrenergic blocker phenoxybenzamine, and its use has minimized surgical mortality and morbidity. The initial dosage is 10 mg twice daily, and patients may require hospitalization for bed rest and intravenous fluids to overcome the initially increased orthostatic hypotension that occurs in most patients. The dose may then be titrated upward every 2–3 days over several weeks until the blood pressure is <160/90 mm Hg and symptoms are abolished. Doses in the range of 100–200 mg/day are routinely used; however, no data are available that establish the superiority of these dosage levels. Higher doses of phenoxybenzamine are not associated with a higher risk of postoperative hypotension. Beta-blockers are generally unnecessary unless tachycardia and arrhythmias are present, and these occur most frequently in the minority of patients with epinephrine hypersecretion.

Metamizine (alpha-methylparatyrosine), an inhibitor of catecholamine synthesis, is also useful for preoperative management although current experience is limited. Initial dosage is 250 mg every 6 hours, and total daily dosages of 2–4 g are required. Preoperative treatment for 1–2 weeks appears to be sufficient to prevent operative complications. Metamizine can be used in conjunction with alpha-blockers.

Successful preoperative management with prazosin, calcium channel blockers, and labetalol has been reported in a few cases.

B. SURGERY

Surgery is the mainstay of therapy for pheochromocytoma; it requires adequate preoperative control of symptoms and hypertension with alpha-blockers or metamizine. Intraoperatively, hypertension is controlled with nitroprusside, and antiarrhythmics are used as needed. Adequate volume replacement is essential and in conjunction with preoperative medical therapy prevents postoperative hypotension.

If CT and MIBG show only a solitary adrenal lesion in patients with sporadic disease, a unilateral laparoscopic approach may be used. Bilateral or malignant disease may require a transabdominal approach, and even if total resection is not feasible, debulking of tumor mass facilitates subsequent medical management of catecholamine excess.

Malignant Pheochromocytoma

The incidence of cancer in pheochromocytoma has been traditionally estimated to be in the range of 10%, although recent series describe a higher incidence. Thus, all patients should undergo serial follow-up to detect early recurrences. Patients with known metastatic disease should undergo surgical debulking of accessible disease. Catecholamine excess can be controlled in most patients with alpha-blockade, metamizine, or both. Despite encouraging reports of chemotherapy or 131I-MIBG therapy, it appears that only a minority of patients have sustained remissions.

Prognosis

In general, the prognosis is good. With better understanding of the disease, surgical deaths are now rare. Blood pressure falls to normal levels in most patients with benign tumors. Patients with cancer have persisting hypertension and require the multiple therapies described above.

INCIDENTALOMA

The most common presentation of adrenal masses is incidental observation on cross-sectional imaging performed for other reasons. The differential diagnosis is quite broad (see Table 30–3) and includes benign adenoma, functional adrenal tumors as previously discussed, metastasis and benign adrenal lesions such as myelolipoma, and neurofibroma. A systematic approach is required to differentiate functional adrenal masses that deserve removal and those lesions with a significant risk of carcinoma from the more common benign nonfunctional adenoma.

Metabolic Evaluation

A careful history and physical examination with focus on obesity pattern, virilization, glucose intolerance, and hypertension is warranted. Laboratory examination with serum electrolytes including glucose and potassium should be done. If hypokalemia exists, then further tests for aldosteronoma are indicated. A 24-hour urinary free cortisol to rule out Cushing’s syndrome and urinary metanephrines and normetanephrines to rule out pheochromocytoma are recommended. Additional metabolic tests are performed when there are suspicious signs or symptoms or when screening tests are abnormal.

If the test that identified the adrenal mass was an ultrasound or CT, an MRI may be helpful to differentiate the various causes of adrenal masses. Because adrenal adenomas have abundant intracytoplasmic lipid, they can often be confirmed on CT and MRI.

Table 30–3. Differential Diagnosis of Adrenal Incidentaloma.

<table>
<thead>
<tr>
<th>Adenoma</th>
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<tbody>
<tr>
<td>Metastasis</td>
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<tr>
<td>Lymphoma</td>
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<tr>
<td>Pheochromocytoma</td>
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<td>Neuroblastoma</td>
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<tr>
<td>Adrenocortical carcinoma</td>
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<tr>
<td>Hematoma</td>
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<tr>
<td>Myelolipoma</td>
</tr>
<tr>
<td>Adrenal hyperplasia</td>
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<tr>
<td>Adrenal cyst</td>
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<tr>
<td>Granulomatous disease</td>
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<td>Hemangioma</td>
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<td>Ganglioneuroma</td>
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Imaging

Lesions that are primarily cystic on CT or MRI are typically benign and can be followed with serial imaging. Benign adrenal cysts are characterized by thin nonenhancing walls; fluid attenuation on CT and thin calcifications may be present peripherally in about 50%.

Characteristics suspicious for malignancy include solid masses that are large, hemorrhagic, or necrotic. MRI is usually heterogeneous on T1- and T2-weighted images due to internal bleeding.

Masses with gross fat on CT (Hounsfield unit [HU] <30) are myelolipomas (Figure 30–9), benign nonfunctional adrenal lesions with lipid and myeloid components. Myelolipomas are usually asymptomatic or present with pain if they bleed.

Noncontrast CT sensitivity for benign adenomas ranges from 50% to 80% depending on the HU attenuation value threshold chosen (Figure 30–5). If a threshold of <10 HU is chosen, the specificity is 84–100%. Chemical-shift imaging is an MR technique to identify intracellular lipid. In-phase and out-of-phase T1-weighted images can distinguish water and fat protons from water photons only. Sensitivities of 81–87% with specificities of 92–100% have been reported for chemical-shift MRI (Figure 30–10).

Diagnostic Algorithm

Percutaneous CT-guided biopsy may be appropriate for adrenal masses with imaging characteristics suspicious for metastasis or in patients with known malignancy. All functional adrenal masses and those >5 cm should be removed. Laparoscopic adrenalectomy is the preferred technique and
is used in most cases except very large masses suspicious for malignancy or with evidence of local extension. Nonfunctional adrenal masses <5 cm should be assessed for radiographic features concerning for malignancy and removed if they are irregular or hemorrhagic or have demonstrated growth. An individualized approach for nonsuspicious nonfunctional masses <5 cm can be taken. It may be appropriate to remove lesions in the 3–5 cm range in well-informed younger patients in order to avoid the burden of radiographic follow-up. Those <3 cm can generally be followed up with serial CTs every 6 months initially, then yearly if stable. There is growing evidence that some incidentalomas that have initial negative screening tests may be causing "subclinical" Cushing’s disease. Therefore, if a patient with an incidentaloma has symptoms or signs of Cushing’s disease, additional or repeat screening tests may be appropriate.

NEUROBLASTOMA

Neuroblastosomas are of neural crest origin and may therefore develop from any portion of the sympathetic chain. Most arise in the retroperitoneum, and 45% involve the adrenal gland. The latter offer the poorest prognosis. In childhood, neuroblastoma is the third most common neoplastic disease after leukemia and brain tumors. Most are encountered in the first 2 1/2 years of life, but a few are seen as late as the sixth decade, when they seem to be less aggressive. Abnormalities of muscle and heart and hemihypertrophy have been observed in association with neuroblastoma.

Metastases spread through both the bloodstream and lymphatics. Common sites in children include the skull and long bones, regional lymph nodes, liver, and lungs. Local invasion is common. In infants, who enjoy the best prognosis, metastases are usually limited to the liver and subcutaneous fat.

The following staging of neuroblastoma is generally accepted:

Stage I: Tumors confined to the structure of origin.
Stage II: Tumors extending in continuity beyond the organ but not crossing the midline. Ipsilateral lymph nodes may be involved.
Stage III: Tumors extending in continuity beyond the midline. Regional lymph nodes may be involved.
Stage IV: Remote disease involving skeletal organs, soft tissues, and distant lymph node groups.
Stage IV-S: Stage I or II patients with remote spread of tumor confined to one or more of the following sites: liver, skin, or bone marrow.

Clinical Findings

A. SYMPTOMS

An abdominal mass is usually noted by parents, the physician, or the patient. About 70% of patients have metastases when first seen. Symptoms relating to metastases include fever, malaise, bone pain, failure to thrive, and constipation or diarrhea.

B. SIGNS

A flank mass is usually palpable and may even be visible; it often extends across the midline. The tumor is usually nodular and fixed, since it tends to be locally invasive. Evidence of metastases may be noted: ocular proptosis from metastases to the skull, enlarged nodular liver, or a mass in bone. Hypertension is often found.

C. LABORATORY FINDINGS

Anemia is common. Urinalysis and renal function are normal. Because 70% of neuroblastomas elaborate increased levels of norepinephrine and epinephrine, urinary vanillylmandelic acid and homovanillic acid levels should be measured. Serial estimations of these substances during definitive treatment can be used as tumor markers. A return to normal levels is encouraging, while rising levels imply residual or progressive tumor. Bone marrow aspiration may reveal tumor cells.

D. X-RAY FINDINGS

Plain radiographs may show a mass and displacement of the kidneys or other organs. CT scans are used to define tumor size, vascular invasion (eg, of the vena cava), local tumor spread, and distant metastases. Further evaluation includes CT of the chest to determine whether lung metastases are present and a bone scan to define skeletal metastases. Many of these tumors take up 131I–MIB; thus this test can be used for staging.

Differential Diagnosis

Wilms tumor is also a disease of childhood. Intravenous urograms show the caliceal distortion characteristic of an intrinsic renal tumor; no such distortion is shown in neuroblastoma, which merely displaces the kidney. Hydronephrosis, polycystic renal disease, and neonatal adrenal hemorrhage may be confused with neuroblastoma. CT is very useful in differentiating the various lesions. Neuroblastomas cause the excretion of large amounts of catecholamines (eg, vanillylmandelic acid), which is not present with the other disorders.

Treatment

Surgical excision of a tumor is standard care of stage I and stage II patients. Although neuroblastoma is radiosensitive, radiation therapy is typically used as part of multimodality therapy of high-risk disease. In stage IV and high-risk stage III disease, chemotherapy is typically given followed by surgery and radiation therapy for residual disease. Useful drugs include cisplatin, cyclophosphamide, doxorubicin,
and etoposide. There is evidence that after chemotherapy and surgery/radiation for residual disease, bone marrow transplant followed by 13-cis-retinoic acid prolongs disease-specific survival in high-risk patients and is now standard care.

**Prognosis**

Patients with stage I and II disease have an 80% survival rate. Including all patients, however, long-term survival occurs in only 15% of patients. Infants have the best prognosis; their 2-year survival rate approaches 60%, and if the tumor is confined to the primary site with or without adjacent regional spread, the cure rate is about 80%. Factors that define high-risk neuroblastoma include age >1 year, metastasis, amplification of the MYCN oncogene, and particular histologic findings.

In a few infants, spontaneous maturation of neuroblastoma to ganglioneuroma has been observed. It is thought by some that x-ray treatment and chemotherapy can also accomplish this. Serial estimation of urinary catecholamines following therapy usually indicates the presence of residual tumor.

**REFERENCES**

**General**


**Cushing’s Syndrome & Adrenocortical Tumors**


**Adrenal Androgenic Syndromes**


**Hyperaldosteronism**


**Pheochromocytomas & Related Tumors**

**Incidentaloma**

**Neuroblastoma**
Disorders of the Kidneys

Jack W. McAninch, MD, FACS

CONGENITAL ANOMALIES OF THE KIDNEYS

Congenital anomalies occur more frequently in the kidney than in any other organ. Some cause no difficulty, but many (eg, hypoplasia, polycystic kidneys) cause impairment of renal function. It has been noted that children with a gross deformity of an external ear associated with ipsilateral maldevelopment of the facial bones are apt to have a congenital abnormality of the kidney (eg, ectopy, hypoplasia) on the same side as the visible deformity. Lateral displacement of the nipples has been observed in association with bilateral renal hypoplasia.

A significant incidence of renal agenesis, ectopy, malrotation, and duplication has been observed in association with congenital scoliosis and kyphosis. Unilateral agenesis, hypoplasia, and dysplasia are often seen in association with suprarelevator imperforate anus. For a better understanding of these congenital abnormalities, see the discussion of the embryology and development of the kidney in Chapter 2.

AGENESIS

Bilateral renal agenesis is extremely rare; no more than 400 cases have been reported. The children do not survive. The condition does not appear to have any predisposing factors. Prenatal suspicion of the anomaly exists when oligohydramnios is present on fetal ultrasound examination. Pulmonary hypoplasia and facial deformities (Potter facies) are usually present. Abdominal ultrasound examination usually establishes the diagnosis.

One kidney may be absent (estimated incidence: 1 in 450–1000 births). In some cases, this may be because the ureteral bud (from the Wolffian duct) failed to develop or, if it did develop, did not reach the metanephros (adult kidney). Without a drainage system, the metanephric mass undergoes atrophy. The ureter is absent on the side of the unformed kidney in 50% of cases, although a blind ureteral duct may be found. (See Chapter 2.)

Renal agenesis causes no symptoms; it is usually found by accident on abdominal or renal imaging. It is not an easy diagnosis to establish even though on inspection of the bladder the ureteral ridge is absent and no orifice is visualized, for the kidney could be present but be drained by a ureter whose opening is ectopic (into the urethra, seminal vesicle, or vagina). If definitive diagnosis seems essential, isotope studies, ultrasonography, and computed tomography (CT) should establish the diagnosis.

There appears to be an increased incidence of infection, hydronephrosis, and stones in the contralateral organ. Other congenital anomalies associated with this defect include cardiac, vertebral column, and anal anomalies as well as anomalies of the long bones, hands, and genitalia.

HYPOPLASIA

Hypoplasia implies a small kidney. The total renal mass may be divided in an unequal manner, in which case one kidney is small and the other correspondingly larger than normal. Some of these congenitally small kidneys prove, on pathologic examination, to be dysplastic. Unilateral or bilateral hypoplasia has been observed in infants with fetal alcohol syndrome, and renal anomalies have been reported in infants with in utero cocaine exposure.

Differentiation from acquired atrophy is difficult. Atrophic pyelonephritis usually reveals typical distortion of the calyces. Vesicoureteral reflux in infants may cause a dwarfed kidney even in the absence of infection. Stenosis of the renal artery leads to shrinkage of the kidney.

Such kidneys have small renal arteries and branches and are associated with hypertension, which is relieved by nephrectomy. Selective renal venography is helpful in differentiating between a congenitally absent kidney and one that is small and nonvisualized.

SUPERNUMERARY KIDNEYS

The presence of a third kidney is very rare; the presence of 4 separate kidneys in one individual has been reported only once. The anomaly must not be confused with duplication (or triplication) of the renal pelvis in one kidney, which is not uncommon.

DYSPLASIA & MULTICYSTIC KIDNEY

Renal dysplasia has protean manifestations. Multicystic kidney of the newborn is usually unilateral, nonhereditary,
and characterized by an irregularly lobulated mass of cysts; the ureter is usually absent or atretic. It may develop because of faulty union of the nephron and the collecting system. At most, only a few embryonic glomeruli and tubules are observed. The only finding is the discovery of an irregular mass in the flank. Nothing is shown on urography, but in an occasional case, some radiopaque fluid may be noted. If the cystic kidney is large, its mate is usually normal. However, when the cystic organ is small, the contralateral kidney is apt to be abnormal. The cystic nature of the lesion may be revealed by sonography, and the diagnosis can be established in utero. If the physician feels that the proper diagnosis has been made, no treatment is necessary. If there is doubt about the diagnosis, nephrectomy is considered the procedure of choice. Neoplastic changes in multicystic renal dysplasia have been noted, but this is accepted as a benign condition.

Multicystic kidney is often associated with contralateral renal and ureteral abnormalities. Contralateral ureteropelvic junction obstruction is one of the common problems noted. Diagnostic evaluation of both kidneys is required to establish the overall status of anomalous development.

Dysplasia of the renal parenchyma is also seen in association with ureteral obstruction or reflux that was probably present early in pregnancy. It is relatively common as a segmental renal lesion involving the upper pole of a duplicated kidney whose ureter is obstructed by a congenital ureterocele. It may also be found in urinary tracts severely obstructed by posterior urethral valves; in this instance, the lesion may be bilateral.

Microscopically, the renal parenchyma is “disorganized.” Tubular and glomerular cysts may be noted; these elements are fetal in type. Islands of metaplastic cartilage are often seen. The common denominator seems to be fetal obstruction.

**ADULT POLYCYSTIC KIDNEY DISEASE**

Adult polycystic kidney disease is an autosomal dominant hereditary condition and almost always bilateral (95% of cases). The disease encountered in infants is different from that seen in adults, although the literature reports a small number of infants with the adult type. The former is an autosomal recessive disease in which life expectancy is short, whereas that diagnosed in adulthood is autosomal dominant; symptoms ordinarily do not appear until after age 40. Cysts of the liver, spleen, and pancreas may be noted in association with both forms. The kidneys are larger than normal and are studded with cysts of various sizes.

**Etiology & Pathogenesis**

The evidence suggests that the cysts occur because of defects in the development of the collecting and uriniferous tubules and in the mechanism of their joining. Blind secre-
obstruction or infection, or from hemorrhage into a cyst. Gross or microscopic total hematuria is not uncommon and may be severe; the cause for this is not clear. Colic may occur if blood clots or stones are passed. The patient may notice an abdominal mass.

Infection (chills, fever, renal pain) commonly complicates polycystic disease. Symptoms of vesical irritability may be the first complaint. When renal insufficiency ensues, headache, nausea and vomiting, weakness, and loss of weight occur.

B. Signs

One or both kidneys are usually palpable. They may feel nodular. If infected, they may be tender. Hypertension is found in 60–70% of these patients. Evidence of cardiac enlargement is then noted.

Fever may be present if pyelonephritis exists or if cysts have become infected. In the stage of uremia, anemia and loss of weight may be evident. Ophthalmoscopic examination may show changes typical of moderate or severe hypertension.

C. Laboratory Findings

Anemia may be noted, caused either by chronic loss of blood or, more commonly, by the hematopoietic depression accompanying uremia. Proteinuria and microscopic (if not gross) hematuria are the rule. Pyuria and bacteriuria are common.

Progressive loss of concentrating power occurs. Renal clearance tests show varying degrees of renal impairment. About one-third of patients with polycystic kidney disease are uremic when first seen.

D. X-Ray Findings

Both renal shadows are usually enlarged on a plain film of the abdomen, even as much as 5 times normal size. Kidneys more than 16 cm in length are suspect.

The renal masses are usually enlarged and the calceal pattern is quite bizarre (spider deformity). The calyces are broadened and flattened, enlarged, and often curved, as they tend to hug the periphery of adjacent cysts. Often the changes are only slight or may even be absent on one side, leading to the erroneous diagnosis of tumor of the other kidney. If cysts are infected, perinephritis may obscure the renal and even the psoas shadows.

E. CT Scanning

CT is an excellent noninvasive technique used to establish the diagnosis of polycystic disease. The multiple thin-walled cysts filled with fluid and the large renal size make this imaging method extremely accurate (95%) for diagnosis.

F. Isotope Studies

Scans reveal multiple “cold” avascular spots in large renal shadows.

G. Ultrasonography

Sonography appears to be superior to both excretory urography and isotope scanning in diagnosis of polycystic disorders.

H. Instrumental Examination

Cystoscopy may show evidence of cystitis, in which case the urine will contain abnormal elements. Bleeding from a ureteral orifice may be noted. Ureteral catheterization and retrograde urograms are rarely indicated.

Differential Diagnosis

Bilateral hydronephrosis (on the basis of congenital or acquired ureteral obstruction) may present bilateral flank masses and signs of impairment of renal function, but ultrasonography shows changes quite different from those of the polycystic kidney.

Bilateral renal tumor is rare but may mimic polycystic kidney disease perfectly on urography. Tumors are usually localized to one portion of the kidney, whereas cysts are quite diffusely distributed. The total renal function should be normal with unilateral tumor but is usually depressed in patients with polycystic kidney disease. CT may be needed at times to differentiate between the 2 conditions.

In von Hippel-Lindau disease (angiomatous cerebellar cyst, angiomatosis of the retina, and tumors or cysts of the pancreas), multiple bilateral cysts or adenocarcinomas of both kidneys may develop. The presence of other stigmas should make the diagnosis. CT, angiography, sonography, or scintiphotography should be definitive.

Tuberous sclerosis (convulsive seizures, mental retardation, and adenoma sebaceum) is typified by hamartomaticous tumors often involving the skin, brain, retinas, bones, liver, heart, and kidneys (see Chapter 21). The renal lesions are usually multiple and bilateral and microscopically are angiomylipomas. The presence of other stigmas and use of CT or sonography should make the differentiation.

A simple cyst (see section following) is usually unilateral and single; total renal function should be normal. Urograms usually show a single lesion (Figure 31–2), whereas polycystic kidney disease is bilateral and has multiple filling defects.

Complications

For reasons that are not clear, pyelonephritis is a common complication of polycystic kidney disease. It may be asymptomatic; pus cells in the urine may be few or absent. Stained smears or quantitative cultures make the diagnosis. A gallium-67 citrate scan will definitely reveal the sites of infection, including abscess.

Infection of cysts is associated with pain and tenderness over the kidney and a febrile response. The differential diagnosis between infection of cysts and pyelonephritis
Figure 31–2. Simple cyst. **Upper left:** Large cyst displacing lower pole laterally. **Upper right:** Section of kidney showing one large and a few small cysts. **Lower left:** Excretory urogram showing soft-tissue mass in upper pole of right kidney. Elongation and distortion of upper calyces by cyst. **Lower right:** Infusion nephrotomogram showing large cyst in upper renal pole distorting upper calyces and dislocating upper portion of kidney laterally.
may be difficult, but here again a gallium scan will prove helpful. In rare instances, gross hematuria may be so brisk and persistent as to endanger life.

Treatment

Except for unusual complications, the treatment is conservative and supportive.

A. General Measures

The patient should be placed on a low-protein diet (0.5–0.75 g/kg/day of protein) and fluids forced to 3000 mL or more per day. Physical activity may be permitted within reason, but strenuous exercise is contraindicated. When the patient is in the state of absolute renal insufficiency, one should treat as for uremia from any cause. Hypertension should be controlled. Hemodialysis may be indicated.

B. Surgery

There is no evidence that excision or decompression of cysts improves renal function. If a large cyst is found to be compressing the upper ureter, causing obstruction and further embarrassing renal function, it should be resected or aspirated. When the degree of renal insufficiency becomes life-threatening, chronic dialysis or renal transplantation should be considered.

C. Treatment of Complications

Pyelonephritis must be rigorously treated to prevent further renal damage. Infection of cysts requires surgical drainage. If bleeding from one kidney is so severe that exsanguination is possible, nephrectomy or embolization of the renal or, preferably, the segmental artery must be considered as a life-saving measure. Concomitant diseases (eg, tumor, obstructing stone) may require definitive surgical treatment.

Prognosis

When the disease affects children, it has a very poor prognosis. The large group presenting clinical signs and symptoms after age 35–40 has a somewhat more favorable prognosis. Although there is wide variation, these patients usually do not live longer than 5 or 10 years after the diagnosis is made unless dialysis is made available or renal transplantation is done.

SIMPLE (SOLITARY) CYST

Simple cyst (Figures 31–2 and 31–3) of the kidney is usually unilateral and single but may be multiple and multilocular and, more rarely, bilateral. It differs from polycystic kidneys both clinically and pathologically.

Etiology & Pathogenesis

Whether simple cyst is congenital or acquired is not clear. Its origin may be similar to that of polycystic kidneys, that is, the difference may be merely one of degree. On the other hand, simple cysts have been produced in animals by causing tubular obstruction and local ischemia; this suggests that the lesion can be acquired.

As a simple cyst grows, it compresses and thereby may destroy renal parenchyma, but rarely does it destroy so much renal tissue that renal function is impaired. A solitary cyst may be placed in such a position as to compress the ureter, causing progressive hydronephrosis. Infection may complicate the picture.

Figure 31–3. Left renal cyst. Left: Computed tomography (CT) scan shows a homogeneous low-density mass (C) arising from anterior border of left kidney just posterior to tail of the pancreas. The CT attenuation value was similar to that of water, indicating a simple renal cyst. Right: After intravenous injection of contrast material, the mass did not increase in attenuation value, adding further confirmatory evidence of its benign cystic nature.
Acquired cystic disease of the kidney can arise as an effect of chronic dialysis. The spontaneous regression of cysts has occasionally been noted.

Pathology

Simple cysts usually involve the lower pole of the kidney. Those that produce symptoms average about 10 cm in diameter, but a few are large enough to fill the entire flank. They usually contain a clear amber fluid. Their walls are quite thin, and the cysts are “blue-domed” in appearance. Calcification of the sac is occasionally seen. About 5% contain hemorrhagic fluid, and possibly one-half of these have papillary cancers on their walls.

Simple cysts are usually superficial but may be deeply situated. When a cyst is situated deep in the kidney, the cyst wall is adjacent to the epithelial lining of the pelvis or calyces, from which it may be separated only with great difficulty. Cysts do not communicate with the renal pelvis (Figure 31–2). Microscopic examination of the cyst wall shows heavy fibrosis and hyalinization; areas of calcification may be seen. The adjacent renal tissue is compressed and fibrosed. A number of cases of simple cysts have been reported in children. However, large cysts are rare in children; the presence of cancer must therefore be ruled out.

Multilocular renal cysts may be confused with tumor on urography. Sonography usually makes the diagnosis. Occasionally, CT and magnetic resonance imaging (MRI) may be necessary.

The Bosniak classification of simple renal cysts is an aid to determining the chance of malignancy based on imaging criteria. Type I cysts are simple and smooth walled, with clear fluid; Type II are also benign, but may have minimal septations and a small fine rim of calcification; Type III are more complex, with more calcification, increasing septations, and a thick cyst wall; Type IV have a thickened irregular wall, often with calcifications, and a mass may be noted inside the cyst, suggesting carcinoma. Numerous variations of the findings are used as a guide in the diagnosis of renal cancer.

Clinical Findings

A. Symptoms

Pain in the flank or back, usually intermittent and dull, is not uncommon. If bleeding suddenly distends the cyst wall, pain may come on abruptly and be severe. Gastrointestinal symptoms are occasionally noted and may suggest peptic ulcer or gallbladder disease. The patient may discover a mass in the abdomen, although cysts of this size are unusual. If the cyst becomes infected, the patient usually complains of pain in the flank, malaise, and fever.

B. Signs

Physical examination is usually normal, although occasionally a mass in the region of the kidney may be palpated or percussed. Tenderness in the flank may be noted if the cyst becomes infected.

C. Laboratory Findings

Urinalysis is usually normal. Microscopic hematuria is rare. Renal function tests are normal unless the cysts are multiple and bilateral (rare). Even in the face of extensive destruction of one kidney, compensatory hypertrophy of the other kidney will maintain normal total function.

D. CT Scanning

CT appears to be the most accurate means of differentiating renal cyst and tumor (Figure 31–3). Cysts have an attenuation approximating that of water, whereas the density of tumors is similar to that of normal parenchyma. Parenchyma is made more dense with the intravenous injection of radiopaque fluid, but a cyst remains unaffected. The wall of a cyst is sharply demarcated from the renal parenchyma; a tumor is not. The wall of a cyst is thin; that of a tumor is not. CT may well supplant cyst puncture in the differentiation of cyst and tumor in many cases.

E. Renal Ultrasonography

Renal ultrasonography is a noninvasive diagnostic technique that in a high percentage of cases differentiates between a cyst and a solid mass. If findings on ultrasonography are also compatible with a cyst, a needle can be introduced into the cyst under ultrasonographic control and the cyst can be aspirated.

F. Isotope Scanning

A rectilinear scan clearly delineates the mass but does not differentiate cyst from tumor. The technetium scan, made with the camera, reveals that the mass is indeed avascular.

G. Percutaneous Cyst Aspiration with Cystography

If the studies listed leave some doubt about the differentiation between cyst and tumor, aspiration may be done. (See “Treatment,” given later.)

Differential Diagnosis

Carcinoma of the kidney also occupies space but tends to lie more deeply in the organ and therefore causes more distortion of the calyces. Hematuria is common with tumor, rare with cyst. If a solid tumor overlies the psoas muscle, the edge of the muscle is obliterated on the plain film; it can be seen through a cyst, however. Evidence of metastases (i.e., loss of weight and strength, palpable supraclavicular nodes, chest film showing metastatic nodules), erythrocy-
tosis, hypercalcemia, and increased sedimentation rate suggest cancer. It must be remembered, however, that the walls of a simple cyst may undergo cancerous degeneration. Sonography, CT scan, or MRI should be almost definitive in differential diagnosis. It is wise to assume that all space-occupying lesions of the kidneys are cancers until proved otherwise.

Polycystic kidney disease is almost always bilateral. Diffuse caliceal and pelvic distortion is the rule. Simple cyst is usually solitary and unilateral. Polycystic kidney disease is usually accompanied by impaired renal function and hypertension; simple cyst is not.

Renal cortical abscess is rare. A history of skin infection a few weeks before the onset of fever and local pain may be obtained. CT scan of the kidney will usually show signs of the abscess. The kidney may be fixed; this can be demonstrated by comparing the position of the kidney when the patient is supine and upright. Angiography demonstrates an avascular lesion. A gallium-67 scan demonstrates the inflammatory nature of the lesion, but an infected simple cyst might have a similar appearance.

Hydronephrosis may present the same symptoms and signs as simple cyst, but the urograms are quite different. Cyst causes calyceal distortion; with hydronephrosis, dilatation of the calyces and pelvis due to an obstruction is present. Acute or subacute hydronephrosis usually produces more local pain because of increased intrapelvic pressure and is more apt to be complicated by infection.

Extrarenal tumor (eg, adrenal, mixed retroperitoneal sarcoma) may displace a kidney, but rarely does it invade it and distort its calyces. If an echinococcal cyst of the kidney does not communicate with the renal pelvis, no scoleces or hooklets will be present in the urine. The wall of a hydatid cyst often reveals calcification on x-ray examination (see Figure 14–5). A skin sensitivity test (Casoni) for hydatid disease may prove helpful.

Complications (Rare)

Spontaneous infection in a simple cyst is rare, but when it occurs, it is difficult to differentiate from carbuncle. Hemorrhage into the cyst sometimes occurs. If sudden, it causes severe pain. The bleeding may come from a complicating carcinoma arising on the wall of the cyst.

Hydronephrosis may develop if a cyst of the lower pole impinges on the ureter. This in itself may cause pain from back pressure of urine in the renal pelvis. This obstruction may lead to renal infection.

Treatment

A. SPECIFIC MEASURES

(1) If renal sonography, CT, or MRI does not lead to a definitive diagnosis, renal angiography or needle aspiration of the cyst may be necessary. Should aspiration be necessary, it can be done under sonographic guidance. The recovery of clear fluid is characteristic of a benign cyst, which should be confirmed by cytologic evaluation. In some centers, contrast radiopaque fluid is injected into the cyst after aspiration for a more thorough evaluation of the cyst wall. A smooth cyst wall, free of irregularities, supports the presence of a benign cyst. If the aspirate contains blood, surgical exploration should be considered, because the chances are great that the growth is cancerous.

(2) If the diagnosis can be clearly established, one should consider leaving the cyst alone, since it is rare for a cyst to harm the kidney. Sonography is useful in follow-up of patients with cysts.

B. TREATMENT OF COMPLICATIONS

If the cyst becomes infected, intensive antimicrobial therapy should be instituted, antimicrobial drugs have been found to attain very low concentrations in the cyst fluid. Therefore, percutaneous drainage is often required. Surgical excision of the extrarenal portion of the cyst wall and drainage are curative when percutaneous drainage fails.

If hydronephrosis is present, excision of the obstructing cyst will relieve the ureteral obstruction. Pyelonephritis in the involved kidney should suggest urinary stasis secondary to impaired ureteral drainage. Removal of the cyst and consequent relief of urinary back pressure make antimicrobial therapy more effective.

Prognosis

Simple cysts can be diagnosed with great accuracy using sonography and CT. Yearly sonography is recommended as a method of following the cyst for changes in size, configuration, and internal consistency. CT may be done if changes suggest carcinoma, and aspiration may then be performed if necessary to establish a diagnosis. Most cysts cause little difficulty.

RENAL FUSION

About 1 in 1000 individuals has some type of renal fusion, the most common being the horseshoe kidney. The fused renal mass almost always contains 2 excretory systems and therefore 2 ureters. The renal tissue may be divided equally between the 2 flanks, or the entire mass may be on one side. Even in the latter case, the 2 ureters open at their proper places in the bladder.

Etiology & Pathogenesis

It appears that this fusion of the 2 metanephroi occurs early in embryologic life, when the kidneys lie low in the pelvis.
For this reason, they seldom ascend to the high position that normal kidneys assume. They may even remain in the true pelvis. Under these circumstances, such a kidney may derive its blood supply from many vessels in the area (e.g., aorta, iliacs). In patients with both ectopia and fusion, 78% have extraurologic anomalies and 65% exhibit other genitourinary defects.

Pathology (Figure 31–4)

Because the renal masses fuse early, normal rotation cannot occur; therefore, each pelvis lies on the anterior surface of its organ. Thus, the ureter must ride over the isthmus of a horseshoe kidney or traverse the anterior surface of the fused kidney. Some degree of ureteral compression may arise from this or from obstruction by one or more aberrant blood vessels. The incidence of hydronephrosis and, therefore, infection is high. Vesicoureteral reflux has frequently been noted in association with fusion.

In horseshoe kidney, the isthmus usually joins the lower poles of each kidney; each renal mass lies lower than normal. The axes of these masses are vertical, whereas the axes of normal kidneys are oblique to the spine, because they lie along the edges of the psoas muscles. On rare occasions, the 2 nephric masses are fused into one mass containing 2 pelves and 2 ureters. The mass may lie in the midline in order to open into the bladder at the proper point (crossed renal ectopy with fusion).

Clinical Findings

A. Symptoms

Most patients with fused kidneys have no symptoms. Some, however, develop ureteral obstruction. Gastrointestinal symptoms (renodigestive reflex) mimicking peptic ulcer, cholelithiasis, or appendicitis may be noted. Infection is apt to occur if ureteral obstruction and hydronephrosis or calculus develops.

B. Signs

Physical examination results are usually negative unless the abnormally placed renal mass can be felt. With horseshoe kidney, it may be possible to palpate a mass over the lower lumbar spine (the isthmus). In the case of crossed ectopy, a mass may be felt in the flank or lower abdomen.

C. Laboratory Findings

Urinalysis is normal unless there is infection. Renal function is normal unless disease coexists in each of the fused renal masses.

D. X-Ray Findings

In the case of horseshoe kidney, the axes of the 2 kidneys, if visible on a plain film, are parallel to the spine. At times the isthmus can be identified. The plain film may also reveal a large soft-tissue mass in one flank yet not show a renal shadow on the other side. Excretory urograms establish the diagnosis if the renal parenchyma has been maintained. The increased density of the renal tissue may make the position or configuration of the kidney more distinct. Urograms also visualize the pelvis and ureters.

(1) With horseshoe kidney, the renal pelves lie on the anterior surfaces of their kidney masses, whereas the normal kidney has its pelvis lying mesial to it. The most valuable clue to the diagnosis of horseshoe kidney is the presence of calyces in the region of the lower pole that point medially and lie medial to the ureter (Figure 31–4).

(2) Crossed renal ectopy with fusion shows 2 pelves and 2 ureters. One ureter must cross the midline in order to empty into the bladder at the proper point (Figure 31–4).

(3) A cake or lump kidney may lie in the pelvis (fused pelvis kidney), but again its ureters and pelves will be shown (Figure 31–4). It may compress the dome of the bladder.

CT clearly outlines the renal mass but is seldom necessary for diagnosis. With pelvic fused kidney or one lying in the flank, the plain film taken with ureteral catheters in place gives the first hint of the diagnosis. Retrograde urograms show the position of the pelves and demonstrate changes compatible with infection or obstruction. Renal scanning delineates the renal mass and its contour, as does sonography.

Differential Diagnosis

Separate kidneys that fail to undergo normal rotation may be confused with horseshoe kidney. They lie along the edges of the psoas muscles, whereas the poles of a horseshoe kidney lie parallel to the spine and the lower poles are placed on the psoas muscles. The calyces in the region of the isthmus of a horseshoe kidney point medially and lie close to the spine.

The diagnosis of fused or lump kidney may be missed on excretory urograms if one of the ureters is markedly obstructed, so that a portion of the kidney, pelvis, and ureter fails to visualize. Infusion urograms or retrograde urograms demonstrate both excretory tracts in the renal mass.

Complications

Fused kidneys are prone to ureteral obstruction because of a high incidence of aberrant renal vessels and the necessity for one or both ureters to arch around or over the renal tissue. Hydronephrosis, stone, and infection, therefore, are common. A large fused kidney occupying the concavity of the sacrum may cause dystocia.
Figure 31–4. Renal fusion. **Upper left:** Crossed renal ectopy with fusion. The renal mass lies in the left flank. The right ureter must cross over the midline. **Upper right:** Example of “sigmoid” kidney. **Lower left:** Horseshoe kidney. Pelves are anterior. Note the aberrant artery obstructing the left ureter and the low position of renal mass. **Lower right:** Pelvic kidney. Pelves are placed anteriorly. Note the aberrant blood supply.
Treatment

No treatment is necessary unless obstruction or infection is present. Drainage of a horseshoe kidney may be improved by dividing its isthmus. If one pole of a horseshoe is badly damaged, it may require surgical resection.

Prognosis

In most cases, the outlook is excellent. If ureteral obstruction and infection occur, renal drainage must be improved by surgical means so that antimicrobial therapy will be effective.

ECTOPIC KIDNEY

Congenital ectopic kidney usually causes no symptoms unless complications such as ureteral obstruction or infection develop.

Simple Ectopy

Simple congenital ectopy usually refers to a low kidney on the proper side that failed to ascend normally. It may lie over the pelvic brim or in the pelvis. Rarely, it may be found in the chest. It takes its blood supply from adjacent vessels, and its ureter is short. It is prone to ureteral obstruction and infection, which may lead to pain or fever. At times such a kidney may be palpable, leading to an erroneous presumptive diagnosis (eg, cancer of the bowel, appendiceal abscess).

Excretory urograms reveal the true position of the kidney. Hydronephrosis, if present, is evident. There is no redundancy of the ureter, as is the case with nephrophtosis or acquired ectopy (eg, displacement by large suprarenal tumor). Obstruction and infection may complicate simple ectopy and should be treated by appropriate means.

CROSSED ECTOPY WITHOUT FUSION

In crossed ectopy without fusion, the kidney lies on the opposite side of the body but is not attached to its normally placed mate. Unless 2 distinct renal shadows can be seen, it may be difficult to differentiate this condition from crossed ectopy with fusion (Figure 31–4). Sonography, angiography, or CT should make the distinction.

ABNORMAL ROTATION

Normally, when the kidney ascends to the lumbar region, the pelvis lies on its anterior surface. Later, the pelvis comes to lie mesially. Such rotation may fail to occur, although this seldom leads to renal disease. Urography demonstrates the abnormal position.

MEDULLARY SPONGE KIDNEY (CYSTIC DILATATION OF THE RENAL COLLECTING TUBULES)

Medullary sponge kidney is a congenital autosomal recessive defect characterized by widening of the distal collecting tubules. It is usually bilateral, affecting all of the papillae, but it may be unilateral. At times, only one papilla is involved. Cystic dilatation of the tubules is often present also. Infection and calculi are occasionally seen as a result of urinary stasis in the tubules. It is believed that medullary sponge kidney is related to polycystic kidney disease. Its occasional association with hemihypertrophy of the body has been noted.

The only symptoms are those arising from infection and stone formation. The diagnosis is made on the basis of excretory urograms or contrast-enhanced CT scan (Figure 31–5). The pelvis and calyces are normal, but dilated (streaked) tubules are seen just lateral to them; many of the dilated tubules contain round masses of radiopaque material (the cystic dilatation). If stones are present, a plain film will reveal small, round calculi in the pyramidal regions just beyond the calyces.

The differential diagnosis includes tuberculosis, healed papillary necrosis, and nephrocalcinosis. Tuberculosis is usually unilateral, and urography shows ulceration of calyces; tubercle bacilli are found on bacteriologic study. Papillary necrosis may be complicated by calcification in the healed stage but may be distinguished by its typical calyceal deformity, the presence of infection, and, usually, impaired renal function. The tubular and parenchymal calcification seen in nephrocalcinosis is more diffuse than that seen with sponge kidney (see Figure 16–3); the symptoms and signs of primary hyperparathyroidism or renal tubular acidosis may be found.

There is no treatment for medullary sponge kidney. Therapy is directed toward the complications (eg, pyelonephritis and renal calculi). Only a small percentage of people with sponge kidney develop complications. The overall prognosis is good. A few patients may pass small stones occasionally.

ABNORMALITIES OF RENAL VESSELS

A single renal artery is noted in 75–85% of individuals and a single renal vein in an even higher percentage. Aberrant veins and, especially, arteries occur. An aberrant artery passing to the lower pole of the kidney or crossing an infundibulum can cause obstruction and hydronephrosis. These causes of obstruction can be diagnosed on angiography or spiral CT.

ACQUIRED LESIONS OF THE KIDNEYS

ANEURYSM OF THE RENAL ARTERY

Aneurysm of the renal artery usually results from degenerative arterial disease that weakens the wall of the artery so that intravascular pressure may balloon it out. It is most
commonly caused by arteriosclerosis or polyarteritis nodosa, but it may develop secondary to trauma or syphilis. Well over 300 cases have been reported. Congenital aneurysm has been recorded. Most cases represent an incidental finding on angiography.

Aneurysmal dilatation has no deleterious effect on the kidney unless the mass compresses the renal artery, in which case some renal ischemia and, therefore, atrophy are to be expected. A true aneurysm may rupture, producing a false aneurysm. This is especially likely to occur during pregnancy. The extravasated blood in the retroperitoneal space finally becomes encapsulated by a fibrous covering as organization occurs. An aneurysm may involve a small artery within the renal parenchyma. It may rupture into the renal pelvis or a calyx.

Most aneurysms cause no symptoms unless they rupture, in which case there may be severe flank pain and even shock. If an aneurysm ruptures into the renal pelvis, marked hematuria occurs. The common cause of death is severe hemorrhage from rupture of the aneurysm. Hypertension is not usually present. A bruit should be sought over the costovertebral angle or over the renal artery anteriorly. If spontaneous or traumatic rupture has occurred, a mass may be palpated in the flank.

A plain film of the abdomen may show an intrarenal or extrarenal ringlike calcification (Figure 31–6). Urograms may be normal or reveal renal atrophy. Some impairment of renal function may be noted if compression or partial obstruction of the renal artery has developed. Aortography delineates the aneurysm. Sonography and CT scanning may prove helpful.

The differential diagnosis of rupture of an aneurysm and injury to the kidney is difficult unless a history or evidence of trauma is obtained. A hydronephrotic kidney may present a mass, but renal imaging clarifies the issue.

Because a significant number of noncalcified and large calcified aneurysms rupture spontaneously, the presence of such a lesion is an indication for operation, particularly during pregnancy. The repair of extrarenal aneurysms may be considered, but complications (eg, thrombosis) are not uncommon. If an intrarenal aneurysm is situated in one pole, heminephrectomy may be feasible. If it is in the center of the organ, however, nephrectomy is required. Therapeutic occlusion of an aneurysm by intra-arterial injection of autologous muscle tissue has been reported. Those few patients with hypertension may become normotensive following definitive surgery.
RENAL INFARCTS

Renal infarcts are caused by arterial occlusion. The major causes are subacute infective endocarditis, atrial or ventricular thrombi, arteriosclerosis, polyarteritis nodosa, and trauma. A thrombotic process in the abdominal aorta may gradually extend upward to occlude the renal artery. Renal infarcts may be unilateral or bilateral.

If smaller arteries or arterioles become obstructed, the tissue receiving blood from such a vessel will first become swollen and then undergo necrosis and fibrosis. Multiple infarcts are the rule. If the main renal artery becomes occluded, the entire kidney will react in kind. The kidney may become functionless and atrophic, therefore, as it undergoes necrosis and fibrosis.

Partial renal infarction is a silent disease, but it can result in flank pain and microscopic or gross hematuria. Sudden and complete infarction may cause renal or chest pain and at times gross or microscopic hematuria. Proteinuria and leukocytosis are found. “Epitheluria,” representing sloughing of renal tubular cells, has been noted. Tenderness over the flank may be elicited. The kidney is not significantly enlarged by arterial occlusion.

CT may fail to have contrast enhancement in a portion of the kidney with partial infarction; with complete infarction, none of the radiopaque fluid is excreted. If complete renal infarction is suspected, a radioisotope renogram should be performed. A completely infarcted kidney shows little or no radioactivity. A similar picture is seen on CT scans performed after injection of radiopaque contrast medium. Even though complete loss of measurable function has occurred, renal circulation may be restored spontaneously in rare instances. Renal angiography or CT makes the definitive diagnosis. A dynamic technetium scan will reveal no perfusion of the affected renal vasculature.

During the acute phase, infarction may mimic ureteral stone. With stone, the excretory urogram may also show lack of renal function, but even so there is usually enough medium in the tubules for a “nephrogram” to be obtained (see Figure 16–3). This will not occur with complete infarction. Evidence of a cardiac or vascular lesion is helpful in arriving at a proper diagnosis.

The complications are related to those arising from the primary cardiovascular disease, including emboli to other organs. In a few cases, hypertension may develop a few days or weeks after the infarction. It may later subside.

Although emergency surgical intervention has been done, it has become clear that anticoagulation therapy is the treatment of choice. It has been shown that an infusion of streptokinase may dissolve the embolus. Renal function returns in most cases.

THROMBOSIS OF THE RENAL VEIN

Thrombosis of the renal vein is rare in adults. It is frequently unilateral and usually associated with membranous glomerulonephritis and nephrotic syndrome. Inva-
sion of the renal vein by tumor or retroperitoneal disease can be the cause. Thrombosis of the renal vein may occur as a complication of severe dehydration and hemoconcentration in children with severe diarrhea from ileocolitis. The thrombosis may extend from the vena cava into the peripheral venules or may originate in the peripheral veins and propagate to the main renal vein. The severe passive congestion that develops causes the kidney to swell and become engorged. Degeneration of the nephrons ensues. There is usually flank pain, and hematuria may be noted. A large, tender mass is often felt in the flank. Thrombocytopenia may be noted. The urine contains albumin and red cells. In the acute stage, urograms show poor or absent secretion of the radiopaque material in a large kidney. Stretching and thinning of the calyceal infundibula may be noted. Clots in the pelvis may cause filling defects. Later, the kidney may undergo atrophy.

Ultrasonography shows the thrombus in the vena cava in 50% of cases. The involved organ is enlarged. CT scan is also a valuable diagnostic tool; visualization of the thrombus can be noted in a high percentage of cases. Recently, MRI has proved to be a very sensitive diagnostic tool. Renal angiography reveals stretching and bowing of small arterioles. In the nephrographic phase, the pyramids may become quite dense. Late films may show venous collaterals. Venacavography or, preferably, selective renal venography demonstrates the thrombus in the renal vein (Figure 31–7) and, at times, in the vena cava.

The symptoms and signs resemble obstruction from a ureteral calculus. The presence of a stone in the ureter should be obvious; some degree of dilatation of the ureter and pelvis also should be expected. Clot obstruction in the ureter must be differentiated from an obstructing calculus.

While thrombectomy and even nephrectomy have been recommended in the past, it has become increasingly clear that medical treatment is usually efficacious. The use of heparin anticoagulation in the acute phase and warfarin chronically offers satisfactory resolution of the problems in most patients. In infants and children, it is essential to correct fluid and electrolyte problems and administer anticoagulants. Fibrinolytic therapy has also been successful. Renal function is usually fully recovered.

**ARTERIOVENOUS FISTULA**

Arteriovenous fistula may be congenital (25%) or acquired. A number of these fistulas have been reported following needle biopsy of the kidney or trauma to the kidney. A few have occurred following nephrectomy secondary to suture or ligature occlusion of the pedicle. These require surgical repair. A few have been recognized in association with adenocarcinoma of the kidney.

A thrill can often be palpated and a murmur heard both anteriorly and posteriorly. In cases with a wide com-

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Figure 31–7. Thrombosis of renal vein. Selective left renal venogram showing almost complete occlusion of vein. Veins to lower pole failed to fill. Note the large size of kidney.

Figure 31–8. Arteriovenous aneurysm. Selective renal angiogram. Note the aneurysm in center of kidney, with prompt filling of the vena cava (shown by arrows).
munication, the systolic blood pressure is elevated and a widened pulse pressure is noted. Renal angiography or isotopic scan establishes the diagnosis. CT scan, sonography, and, recently, duplex ultrasound with color flow are particularly helpful. Arteriovenous fistula involving the renal artery and vein requires surgical repair or nephrectomy. Most, however, can be occluded by embolization, balloon, or steel coil. Those that develop secondary to renal biopsy tend to heal spontaneously.

ARTERIOVENOUS ANEURYSM

About 100 instances of this lesion have been reported (Figure 31–8). Most follow trauma. Hypertension is to be expected and is associated with high-output cardiac failure. A bruit is usually present. Nephrectomy is usually indicated.

RENOALIMENTARY FISTULA

Over 100 instances of renoalimentary fistula have been reported. They usually involve the stomach, duodenum, or adjacent colon, although fistula formation with the esophagus, small bowel, appendix, and rectum has been reported.

The underlying cause is usually a pyonephrotic kidney or renal cell carcinoma that becomes adherent to a portion of the alimentary tract and then ruptures spontaneously, thus creating a fistula (Figure 31–9). A few cases following trauma have been reported. The patient is apt to have symptoms and signs of acute pyelonephritis. Urography may show radiopaque material escaping into the gastrointestinal tract. Gastrointestinal series may also reveal the connection with the kidney. The treatment is nephrectomy with closure of the opening into the gut.

RENOBRONCHIAL FISTULA

Renobronchial fistulas are rare. They are caused by rupture of an infected, calculous kidney through the diaphragm.

REFERENCES

Congenital Anomalies

General


**Agenesis**

**Hypoplasia**

**Dysplasia & Multicystic Kidney**

**Polycystic Kidneys**

**Simple Cyst**

**Acquired Lesions**
**General**

**Infection-Related Renal Disorders**

**Aneurysm of the Renal Artery**

**Thrombosis of the Renal Vein**

**Arteriovenous Fistula**

**Renoalimentary Fistula**
Diagnosis of Medical Renal Diseases

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MEDICAL RENAL DISEASE—OVERVIEW

Medical renal diseases are those that involve principally the parenchyma of the kidneys. Hematuria, proteinuria, pyuria, oliguria, polyuria, pain, renal insufficiency with azotemia, acidosis, anemia, electrolyte abnormalities, and hypertension may occur in a wide variety of disorders affecting any portion of the parenchyma of the kidney, the blood vessels, or the excretory tract.

A complete medical history and physical examination, a thorough examination of the urine, and blood and urine chemistry examinations as indicated are essential initial steps in the workup of any patient.

History & Examination

A. FAMILY HISTORY

The family history may reveal disease of genetic origin, for example, tubular metabolic anomalies, polycystic kidneys, unusual types of nephritis, or vascular or coagulation defects that may be essential clues to the diagnosis.

B. PAST HISTORY

The past personal history should cover infections, injuries, and exposure to toxic agents, anticoagulants, or drugs that may produce toxic or sensitivity reactions. A history of diabetes, hypertensive disease, or autoimmune disease may be obtained. The inquiry may also elicit symptoms of uremia, debilitation, and the vascular complications of chronic renal disease, but often the patient is asymptomatic and presents with abnormal laboratory findings.

Physical Examination

Pallor, edema, hypertension, retinopathy, or stigmas of congenital and hereditary disease may be detected.

Laboratory Findings

A. URINALYSIS

Examination of the urine is the essential part of the investigation.

1. Proteinuria—Proteinuria of any significant degree (2–4+) is suggestive of medical renal disease (parenchymal involvement). Formed elements present in the urine additionally establish the diagnosis. Significant proteinuria occurs in immune-mediated glomerular diseases or disorders with glomerular involvement such as diabetes mellitus, myeloma, or amyloidosis. Interstitial nephritis, polycystic kidneys, and other tubular disorders are not associated with significant proteinuria.

2. Erythrocyte casts—Red blood cell casts point to glomerulonephritis. If red blood cell (erythrocyte) casts are not present, microscopic hematuria may or may not be of glomerular origin. Phase contrast microscope study may reveal dysmorphic changes in the erythrocytes present in the urine in patients with parenchymal renal disorders.

3. Fatty casts and oval fat bodies—Tubular cells showing fatty changes occur in degenerative diseases of the kidney (nephrosis, glomerulonephritis, autoimmune disease, amyloidosis, and damage due to toxins such as lead or mercury).

4. Granular casts—These types of casts result from degeneration of cellular casts. They are nondiagnostic of a specific renal disorder but do reflect an inflammatory condition in the kidneys.

B. OTHER FINDINGS

Abnormal urinary chemical constituents may be the only indication of a metabolic disorder involving the kidneys. These disorders include diabetes mellitus, renal glycosuria, aminoacidurias (including cystinuria), oxaluria, gout, hyperparathyroidism, hemoglobinuria, and myoglobinuria.

Examination of the Kidneys & Urinary Tract

Roentgenographic, sonographic, and radioisotopic studies provide information about the size, structure, blood supply, and function of the kidneys.

Renal Biopsy

Renal biopsy is a valuable diagnostic procedure. The technique has become well established, providing sufficient tis-
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sue for light and electron microscopy and for immunofluorescence examination. Contraindications for percutaneous kidney biopsy may include the anatomic presence of only 1 kidney, severe malfunction of one kidney even though function is adequate in the other, bleeding diathesis, and an uncooperative patient.

Clinical indications for renal biopsy, in addition to the necessity for establishing a diagnosis, include the need to determine prognosis, to follow progression of a lesion and response to treatment, to confirm the presence of a generalized disease (autoimmune disorder, amyloidosis, sarcoidosis), and to diagnose renal dysfunction in a transplanted kidney. Ultrasound or computed tomography (CT) guidance provides a more effective biopsy result. More recently, laparoscopic approach has been used by some urologists.

GLOMERULONEPHRITIS

The clinical manifestations of glomerular renal disease are apt to consist only of varying degrees of hematuria, excretion of characteristic formed elements in the urine, proteinuria, and renal insufficiency and its complications. Excluding diabetes, purported immunologic renal diseases are the most common cause of proteinuria and the nephrotic syndrome.

Alterations in glomerular architecture as observed in tissue examined by light microscopy alone can be minimal, nonspecific, and difficult to interpret. For these reasons, specific diagnoses of renal disease require targeted immune fluorescent techniques for demonstrating a variety of antigens, antibodies, and complement fractions. Electron microscopy has complemented these immunologic methods. Tissue analysis can be assisted by blood tests of immunoglobulins (Ig), complement, and other mediators of inflammation.

The 2 important humoral mechanisms leading to deposition of antibodies within the glomerulus are based on the location of the antigen, whether fixed within the kidney or present in soluble form in circulation. The fixed antigens are either a natural structural element of the glomerulus or foreign materials that have been trapped within the glomerulus for a variety of immunologic or physiochemical reasons. The best examples of the fixed natural antigens are those associated with the glomerular basement membrane (GBM). These antigens are evenly distributed in the GBM and cause characteristic linear IgG deposition, as determined in immunofluorescence studies. This process represents 5% of cases of immune-mediated glomerular disease and is referred to as anti-GBM disease. However, most patients with glomerular immune deposits have discontinuous immune aggregates caused by antibody binding to native renal cell antigens or to antigens trapped within the glomerulus. In addition, immune complexes formed in the circulation can deposit and accumulate in the GBM and the mesangium.

A group of immune-mediated nephritides characterized by necrotizing and crescentic architecture and rapid progression are referred to as pauci-immune glomerular nephritides because, while antibodies may contribute to the pathogenesis of the disease, they are rarely demonstrated within the glomeruli. These are known as antineutrophil cytoplasmic antibody (ANCA) diseases. Circulating antibodies against myeloperoxidase, MPO (P-ANCA) and proteinase 3, PR3 (C-ANCA) have been noted in microscopic angiitis and Wegener’s granulomatosis, respectively.

Cellular immune processes are likely to be stimulated and contribute in different ways in various other forms of glomerulonephritides.

The current classification of glomerulonephritis is based on the mechanism, the presence, and the localization of immune aggregates in the glomeruli.

Immunologic Mechanisms Likely

A. SUBEPITHELIAL IMMUNE DEPOSITS
1. Glomerulonephritis associated with postinfectious glomerulonephritis such as poststreptococcal glomerulonephritis
2. Membranous nephropathy idiopathic or secondary to other causes such as systemic lupus erythematosus, cancer, gold penicillamine

B. SUBENDOTHELIAL IMMUNE DEPOSITS
1. Glomerulonephritis associated with systemic lupus erythematosus, type I idiopathic membranoproliferative glomerulonephritis (MPGN), glomerulonephritis associated with hepatitis C infection, bacterial endocarditis, and shunt nephritis

C. MESANGIAL IMMUNE DEPOSITS
1. IgA nephropathy, Schönlein-Henoch purpura

D. ANTI-GBM DISEASE
1. Diffuse linear deposition of Ig

Immunologic Mechanisms Not Clearly Established

1. Minimal change nephropathy
2. Focal glomerulosclerosis
3. Hemolytic-uremic syndrome and thrombotic thrombocytopenic purpura
4. ANCA-associated disease: Wegener’s granulomatosis and small-vessel vasculitis
5. Type II MPGN (dense deposit disease)

Poststreptococcal Glomerulonephritis

A. ESSENTIALS OF DIAGNOSIS
- History of streptococcal infection
• Mild generalized edema, mild hypertension, retinal hemorrhages
• Gross hematuria; protein, erythrocyte casts, granular and hyaline casts, white blood cells (leukocytes), and renal epithelial cells in urine
• Elevated antistreptolysin O titer, hypocomplementemia

B. General Considerations

Poststreptococcal glomerulonephritis is a disease affecting both kidneys. In most cases recovery from the acute stage is complete, but progressive involvement may destroy renal tissue, leading to renal insufficiency. Acute glomerulonephritis is most common in children aged 3–10 years. By far the most common cause is an antecedent infection of the pharynx and tonsils or of the skin with group A beta-hemolytic streptococci, certain strains of which are nephritogenic. Nephritis occurs in 10–15% of children and young adults who have clinically evident infection with a nephritogenic strain. In children under age 6, pyoderma (impetigo) is the most common antecedent; in older children and young adults, pharyngitis is a common antecedent. Occasionally, nephritis may follow infection due to other agents, hence the more general term postinfectious glomerulonephritis.

The pathogenesis of the glomerular lesion has been further elucidated by the use of new immunologic techniques (immunofluorescence) and electron microscopy. A likely sequel to infection is injury to the mesangial cells in the intercapillary space. The glomerulus may then become more easily damaged by antigen-antibody complexes developing from the immune response to the infection. Complement is deposited in association with IgG or alone in a granular pattern on the epithelial side of the basement membrane.

Gross examination of the involved kidney shows only punctate hemorrhages throughout the cortex. Microscopically, the primary alteration is in the glomeruli, which show proliferation and swelling of the mesangial and endothelial cells of the capillary tuft. The proliferation of capsular epithelium occurs and around the tuft there are collections of leukocytes, erythrocytes, and exudate. Edema of the interstitial tissue and cloudy swelling of the tubular epithelium are common. When severe, typical histologic findings in glomerulitis are enlarging crescents that become hyalized and converted into scar tissue and obstruct the circulation through the glomerulus. Degenerative changes occur in the tubules, with fatty degeneration, necrosis, and ultimately scarring of the nephron.

C. Clinical Findings

1. Symptoms and signs—Often the disease is mild, and there may be no reason to suspect renal involvement unless the urine is examined. In severe cases, about 2 weeks after the acute streptococcal infection, the patient has headache, malaise, mild fever, puffiness around the eyes and face, flank pain, and oliguria. Hematuria is usually noted as “bloody” or, if the urine is acid, as “brown” or “coffee colored.” There may be moderate tachycardia, dyspnea, and moderate to marked elevation of blood pressure. Tenderness in the costovertebral angle is common.

2. Laboratory findings—The diagnosis is confirmed by examination of the urine, which may be grossly bloody or coffee colored (acid hematin) or may show only microscopic hematuria. In addition, the urine contains protein (1–3+) and casts. Hyaline and granular casts are commonly found in large numbers, but the classic sign of glomerulitis, occasionally noted, is the erythrocyte cast. The erythrocyte cast is usually of small caliber, is intensely orange or red, and may show the mosaic pattern of the packed erythrocytes held together by the clot of fibrin and plasma protein.

With the impairment of renal function (decrease in glomerular filtration rate and blood flow) and with oliguria, plasma or serum urea nitrogen and creatinine become elevated, the levels varying with the severity of the renal lesion. A mild normochromic anemia may result from fluid retention and dilution. Infection of the throat with nephritogenic streptococci is frequently followed by increasing antistreptolysin O titers in the serum, whereas high titers are usually not demonstrable following skin infections. Serum complement levels are usually low.

Confirmation of diagnosis is made by examination of the urine, although the history and clinical findings in typical cases leave little doubt. The finding of erythrocytes in a cast is proof that erythrocytes were present in the renal tubules and did not arise from elsewhere in the genitourinary tract.

3. Treatment—There is no specific treatment. Eradication of infection, prevention of overhydration and hypertension, and prompt treatment of complications such as hypertensive encephalopathy and heart failure require careful management.

4. Prognosis—Most patients with the acute disease recover completely; 5–20% show progressive renal damage. This damage may be evident only years after the immune injury. If oliguria, heart failure, or hypertensive encephalopathy is severe, death may occur during the acute attack. Even with severe acute disease, however, recovery is the rule, particularly in children.

IgA Nephropathy

Primary hematuria (idiopathic benign and recurrent hematuria, Berger’s disease) is now known to be an immune complex glomerulopathy in which deposition of IgA occurs in a granular pattern in the mesangium of the glomerulus. The associated light microscope findings are variable and range from normal to extensive crescentic glomerulonephritis.

Recurrent macroscopic and microscopic hematuria and mild proteinuria are usually the only manifestations
of renal disease. Most patients with IgA nephropathy are between the ages of 16 and 35 years at the time of diagnosis. The disease occurs much more frequently in males than females and is the most common cause of glomerulonephritis in Asians. While most patients continue to have episodes of gross hematuria or microscopic hematuria, the renal function is likely to remain stable. However, approximately 30% of patients will have progressive renal dysfunction and develop end-stage renal disease. Clinical features that indicate a poor prognosis include male sex, older age at onset of disease, the presence of nephrotic-range proteinuria, hypertension, or renal dysfunction at presentation.

There is no satisfactory therapy for IgA nephropathy. The role of immunosuppressive drugs such as steroids and cytotoxic agents is not clear, and there have been few rigorously performed controlled trials. A more intriguing approach is the use of omega-3 fatty acids (fish oils) to delay progression of the renal disease. A large prospective randomized placebo-controlled trial in patients with IgA nephropathy using 12 g of omega-3 fatty acids has shown that fish oils probably can reduce the deterioration of renal function and the number of patients in whom end-stage renal disease develops.

Rapidly Progressive Glomerulonephritis
This condition has several pulmonary-renal diseases. The patient usually gives a history of recent hemoptysis and often of malaise, anorexia, and headache. A severe acute glomerulonephritis may be accompanied by diffuse hemorrhagic inflammation of the lungs. The urine shows gross or microscopic hematuria, and laboratory findings of severely suppressed renal function are usually evident. Biopsy shows glomerular crescents, glomerular adhesions, and inflammatory infiltration interstitially. Electron microscope examination shows an increase in basement membrane material and deposition of fibrin beneath the capillary endothelium. In cases of anti-GBM disease, circulating antibody against GBM can be identified. IgG, C3, and, often, other components of the classic complement pathway can be demonstrated as linear deposits on the basement membranes of the glomeruli and the lung. This was formerly called Goodpasture’s disease.

Large doses of corticosteroids in combination with immunosuppressive therapy may be useful. Plasmapheresis to remove circulating antibody has been reported to be effective in some patients. Transplantation should be delayed until circulating antiglomerular basement antibodies have disappeared.

In contrast, some patients have crescentic glomerulonephritis but do not show any immune deposits or antibody deposition by immunofluorescence studies. They have pauci-immune idiopathic rapidly progressive glomerulonephritis (RPGN). Many of these patients are serologically ANCA-positive. High-dose prednisone pulse therapy and cytotoxic agents may result in prolonged remission.

NEPHROTIC SYNDROME
Essentials of Diagnosis and General Considerations
- Edema
- Proteinuria >3.5 g/day
- Hypoalbuminemia <3 g/dL
- Hyperlipidemia: cholesterol >300 mg/100 mL
- Lipiduria: free fat, oval fat bodies, fatty casts

Because treatment and prognosis vary with the cause of nephrotic syndrome, renal biopsy is important. Light microscopy, electron microscopy, and immunofluorescence identification of immune mechanisms diagnose most causes of nephrosis.

Glomerular diseases associated with nephrosis include the following:

Minimal Glomerular Lesions
Minimal-change nephropathy (nil disease) accounts for about 20% of cases of nephrosis in adults and 90% in children. No abnormality is visible by examination of biopsy material with the light microscope. With the electron microscope, alterations of the GBM, with effacement of foot processes of the epithelial cells, are evident. There is no evidence of immune disease by immunofluorescence studies. The response to treatment with corticosteroids is good, but for patients who have frequent relapses with steroids or are steroid resistant, a course of cyclophosphamide or chlorambucil may induce a prolonged remission. Patients who do not respond to these agents may show a favorable response with cyclosporine or tacrolimus. Renal function usually remains stable.

Focal Glomerulosclerosis
Focal glomerulosclerosis is the second most common cause of nephrotic syndrome in children and an increasing cause of the nephrotic syndrome in adults. The diagnosis is based on light microscope findings of segmental hyalinosis and sclerosis associated with effacement of the foot processes on electron microscopy. Focal glomerulosclerosis is frequently idiopathic but can be associated with human immunodeficiency virus infection and heroin use. A secondary form of focal glomerulosclerosis without the diffuse changes in foot processes may occur in patients with a solitary kidney, hyperfiltration syndromes, and reflux nephropathy. There are reports of familial variants. The response of the idiopathic form of focal glomerulosclerosis to therapy is suboptimal. Prolonged corticosteroid therapy produces remission in approximately 40% of patients.
Over a 10-year period, approximately 50% of patients will have chronic renal failure. Idiopathic focal glomerulosclerosis has a recurrence rate of 25% after transplantation.

Membranous Nephropathy

Examination of biopsy material with the light microscope shows thickening of the glomerular cells but no cellular proliferation. With the electron microscope, irregular lumpy deposits appear between the basement membrane and the epithelial cells, and new basement membrane material protrudes from the GBM as spikes or domes. Immunofluorescence studies show diffuse granular deposits of Ig (especially IgG) and complement (C3 component). As the membrane thickens, glomeruli become sclerosed and hyalinized.

The pathogenesis of most cases of membranous nephropathy in humans is unclear. Several mechanisms have been suggested. They include trapping of circulating immune complexes or binding of an antibody to scattered glomerular antigens (either present already or “planted” after a nonrenal-source antigen lodges in the glomerulus).

There is considerable controversy regarding the effectiveness of therapy with steroids or immunosuppressive agents. Therapy should be most often used in patients at high risk of progressive renal failure with the following criteria: proteinuria >5 g/day, hypertension, and elevated serum creatinine.

Membranoproliferative Glomerulonephritis—Type I and Type II

In type I MPGN, light microscopy shows thickening of glomerular capillaries, accompanied by mesangial proliferation and obliteration of glomeruli. With the electron microscope, subendothelial deposits and growth of mesangium into capillary walls are demonstrable. Immunofluorescence studies show the presence of the C3 component of complement and, rarely, the presence of Ig. The most common cause of MPGN type I is chronic hepatitis C virus infection. This condition is usually associated with high levels of IgG/IgM. Cryoimmunoglobulins may be present with normal or slightly reduced levels of complement. There is no known effective treatment.

Type II MPGN is characterized by dense deposits visible by electron microscopy and lack of findings by immunofluorescence studies. Treatment is unsatisfactory, and there is a high rate of recurrence after kidney transplantation.

Miscellaneous Diseases

Many medical illnesses that are metabolic, autoimmune, or infectious, as well as neoplastic diseases and reactions to drugs and other toxic substances can produce glomerular disease. These include diabetic glomerulopathy, systemic lupus erythematosus, ANCA-positive renal disease (including Wegener’s granulomatosis), amyloid disease, multiple myeloma, lymphomas, carcinomas, syphilis, reaction to toxins, reaction to drugs (eg, trimethadione), and exposure to heavy metals.

A rare glomerular illness has been recently described with nephrotic syndrome: fibrillary and immunotactoid glomerular nephritis. The lesions of fibrillary glomerulonephritis are characterized by randomly oriented fibril deposits 10–30 nm in diameter located within the mesangium and capillary wall. Immunotactoid glomerulopathy is characterized by deposits of microtubular structures of 18–19 nm and has been associated with lymphoproliferative disorders.

In contrast to amyloid, the deposits in both diseases are Congo red-negative. Treatment is generally unsatisfactory.

Clinical Findings in Nephrosis

A. SYMPTOMS AND SIGNS

Edema may appear insidiously and increase slowly or can appear suddenly and accumulate rapidly. Symptoms other than those related to the mechanical effects of edema are not remarkable.

On physical examination, massive peripheral edema is apparent. Signs of hydrothorax and ascites are common. Pallor is often accentuated by the edema, and striae commonly appear.

B. LABORATORY FINDINGS

The urine contains large amounts of protein, 4–10 g/24 h or more. There is a good correlation between the urine protein to creatinine ratio (from a “spot” AM urine) and the 24-hour proteinuria. For example, a ratio in excess of 3:1 in the spot urine usually correlates to a 24-hour proteinuria of 3 g. The sediment contains casts, including the characteristic fatty and waxy varieties; renal tubular cells, some of which contain fatty droplets (oval fat bodies); and variable numbers of erythrocytes. A mild normochromic anemia is common, but anemia may be more severe if renal damage is great. Nitrogen retention varies with the severity of impairment of renal function. The plasma is often lipemic, and the blood cholesterol is usually greatly elevated. Plasma protein is greatly reduced. The albumin fraction may fall to less than 2 g/dL. Serum complement is usually low in active disease. The serum electrolyte concentrations are often normal, although the serum sodium may be slightly low; total serum calcium may be low, in keeping with the degree of hypoalbuminemia and decrease in the protein-bound calcium moiety. During edema-forming periods, urinary sodium excretion is very low and urinary aldosterone excretion is elevated. If renal insufficiency (see preceding discussion) is present, the blood and urine findings are usually altered accordingly.

Renal biopsy is often essential to establish the diagnosis between the various conditions and to indicate prognosis.
C. Differential Diagnosis

The nephrotic syndrome (nephrosis) may be associated with a variety of primary renal diseases or may be secondary to a systemic process: collagen-vascular diseases (eg, disseminated lupus erythematosus, polyarteritis), diabetic nephropathy, amyloid disease, thrombosis of the renal vein, myxedema, multiple myeloma, malaria, syphilis, reaction to toxins or heavy metals, reactions to drugs, and constrictive pericarditis.

D. Treatment

An adequate diet with restricted sodium intake (0.5–1 g/d) and prompt treatment of intercurrent infection are the basis of therapy. Diuretics may be given but are often only partially effective. Salt-free albumin and other oncotic agents are of little help, and their effects are transient. The corticosteroids have been shown to be of value in treating nephrotic syndrome when the underlying disease is of minimal change—focal segmental glomerulosclerosis, systemic lupus erythematosus, or proliferative and crescentic glomerulonephritis. Steroids are often less effective in the treatment of membranous disease and membranoproliferative lesions of the glomerulus.

Alkylating agents, azathioprine, mycophenolate mofetil, cyclosporine, and tacrolimus, have been used in the treatment of nephrotic syndrome. Encouraging early results have been reported in children and adults with proliferative or membranous lesions and with systemic lupus erythematosus. It is not known what percentage of patients can be expected to benefit from these drugs.

Both corticosteroids and cytotoxic agents are commonly associated with serious side effects. At present, this form of therapy should be employed only in patients in whom the disease has proved refractory to well-established treatment regimens.

Reduction in proteinuria and improvement in nephrotic edema have been reported using low-protein diets and angiotensin-converting enzyme (ACE) inhibitors or angiotensin receptor blockers (ARBs). Most recently, studies have shown some improvements with lipid-lowering drugs.

E. Prognosis

The course and prognosis depend on the basic disease responsible for nephrotic syndrome. In most children with nephrosis (usually secondary to minimal change nephropathy), the disease appears to run a rather benign course when properly treated and to leave insignificant sequelae. Of the remaining children, most go inexorably into renal insufficiency. Adults with nephrosis fare less well. Hypertension, heavy proteinuria, and renal dysfunction are poor prognosticators.

RENAL INVOLVEMENT IN COLLAGEN DISEASES

Although it may not be accurate to classify all of these disorders as collagen diseases, disseminated lupus erythematosus, polyarteritis nodosa, microscopic angiitis scleroderma, Wegener’s granulomatosis, Henoch-Schönlein purpura and thrombotic thrombocytopenic purpura have been implicated in cases of glomerulonephritis. The urine sediment is often diagnostic, containing erythrocytes and erythrocyte casts; renal tubular cells, including some filled with fat droplets; and waxy and granular broad casts. The presence of these formed elements is indicative of active glomerular and tubular disease. The symptoms and signs of the primary disease, involving extrarenal findings (eg, pulmonary or ear, nose, or throat changes with Wegener’s granulomatosis; dermatologic abnormalities or carditis with systemic lupus erythematosus; dysphagia with scleroderma), as well as the presence of ANCA and other serologic tests, help to differentiate the form of collagen disease present. Complete renal recovery from the disease is not likely to occur, although steroid and immunosuppressive drugs (alone or in combination) may be effective for long-term amelioration.

DISEASES OF THE RENAL TUBULES & INTERSTITIUM

Interstitial Nephritis

Acute interstitial diseases are usually due to sensitivity to drugs, including antibiotics (penicillin, sulfonamides), nonsteroidal anti-inflammatory drugs, and phenytoin. The pathologic hallmark of acute interstitial nephritis is the infiltration of inflammatory cells in the interstitium. A typical presentation is a rapid deterioration in renal function associated with a recent introduction of a new drug. The finding of eosinophiluria is very suggestive of allergic interstitial nephritis. Recovery may be complete, especially if the offending drug is withdrawn. A short course with corticosteroids may hasten the recovery.

Chronic interstitial nephritis is characterized by focal or diffuse interstitial fibrosis accompanied by infiltration, with inflammatory cells ultimately associated with extensive tubular atrophy. It represents a nonspecific reaction to a variety of causes: analgesic abuse, lead and cadmium toxicity, nephrocalcinosis, urate nephropathy, radiation nephritis, sarcoidosis, Balkan nephritis, and some instances of obstructive uropathy.

Analgesic Nephropathy

Analgesic nephropathy typically occurs in patients with chronic and recurrent headaches or with chronic arthritis who habitually consume large amounts of the drugs. Phenacetin was implicated initially, but with elimination of
phenacetin from the mixtures, the incidence of analgesic nephropathy has not decreased. Chronic use of nonsteroidal anti-inflammatories is a frequent cause of this condition. The ensuing damage to the kidneys usually is detected late, after renal insufficiency has developed. Careful history taking or the detection of analgesic metabolites in the urine can lead to this diagnosis. The history of excessive use of analgesics may be concealed by the patient.

The kidney lesion is pathologically nonspecific, consisting of peritubular and perivascular inflammation with degenerative changes of the tubular cells (chronic interstitial nephritis). There are often no glomerular changes. Renal papillary necrosis extending into the medulla may involve many papillae.

Hematuria can be a common presenting complaint but it is usually microscopic. Renal colic occurs when necrotic renal papillae slough away. Polyuria may be prominent. Signs of acidosis (hyperpnea), dehydration, and pallor are common. Infection is a frequent complication. The urine usually is remarkable only for the presence of blood and small amounts of protein. Elevated blood urea nitrogen and creatinine and the electrolyte changes characteristic of metabolic acidosis and renal failure are typically present. Urinary concentrating impairments are usually present. Urograms show cavities and ring shadows typical of areas of destruction of papillae.

**Uric Acid Nephropathy**

Crystals of urate may produce an interstitial inflammatory reaction. Urate may precipitate out in acid urine in the calyces to form uric acid stones. Patients with myeloproliferative disease under treatment are subject to occlusion of the upper urinary tract by uric acid crystals. Alkalization of the urine and a liberal fluid intake help prevent crystal formation. Allopurinol is a useful drug to prevent hyperuricemia and hyperuricosuria. Recently, it has been suggested that many instances considered to be chronic “gouty nephropathy” are instead related to chronic lead renal injury and not due to primary uric acid depositions.

**Obstructive Uropathy**

Interstitial nephritis due to obstruction may not be associated with infection. Tubular conservation of salt and water is impaired. Partial to full renal recovery follows relief of the obstruction but is inversely related to the duration of obstruction.

**Myelomatosis**

Features of myelomatosis that contribute to renal disease include proteinuria (including filtrable Bence-Jones protein and κ and λ chains) with precipitation in the tubules leading to accumulation of abnormal proteins in the tubular lumen. A Fanconi-like syndrome may develop.

Plugging of tubules, tubular atrophy, and, occasionally, the accumulation of amyloid may also be present. Renal failure may occur acutely or develop slowly. Hemodialysis may rescue the patient during efforts to control the myeloma with chemical agents.

A primary renal disorder that is related is termed light chain nephropathy. Patients have nephrosis and usually progressive renal failure. In contrast to multiple myeloma, there is no malignant hematopoietic process. Either κ or λ light chains are measurable in the urine and light chain deposits are seen in the glomeruli. There is no effective treatment.

**HEREDITARY RENAL DISEASES**

The importance of inheritance and the familial incidence of disease warrant the inclusion of a classification of hereditary renal diseases. Although relatively uncommon in the population at large, hereditary renal disease must be recognized to permit early diagnosis for detection and genetic counseling.

**Chronic Hereditary Nephritis**

Evidence of the disease usually appears in childhood, with episodes of hematuria. Renal insufficiency commonly develops in males but only rarely in females. Survival beyond age 40 is rare.

In many families, deafness and abnormalities of the eyes accompany the renal disease (so-called Alport disease). Another form of the disease is accompanied by polyneuropathy. Infection of the urinary tract is a common complication.

There is splitting and thickening of the GBM or podocyte proliferation and thickening of Bowman’s capsule. Recently, kindreds have been described that have “thin-membrane disease.” This condition is characterized by microscopic hematuria and, often, later progression to chronic renal failure. This, like Alport disease, may represent inherited abnormalities or deficiencies in type IV collagen in the GBM. This affects both genders.

Laboratory findings with these conditions are commensurate with existing renal function. Treatment is symptomatic.

**Cystic Diseases of the Kidney**

Congenital structural anomalies of the kidney must be considered in any patient with hypertension, pyelonephritis, or renal insufficiency. Many of these patients are at increased risk of urinary tract infection.

**A. POLYCYSTIC KIDNEYS**

Polycystic kidney disease is familial and often involves not only the kidney but the liver and pancreas as well. It is clear that at least 2 genetic loci can lead to autosomal dominant polycystic kidney disease.
The formation of cysts on the cortex of the kidney is thought to result from failure of union of the collecting tubules and convoluted tubules of some nephrons. Intrarenal cysts may be of a proximal or a distal luminal type, differing on analysis by their cyst electrolyte content. This is important if one or more of these cysts become infected, and an antibiotic (with varying cyst-type penetrance) is chosen. New cysts do not form, but those present enlarge and, by exerting pressure, cause destruction of adjacent renal tissue. The incidence of cerebral vessel aneurysms and cardiac valve prolapse is higher than normal.

Cases of polycystic disease are discovered during the investigation of hypertension, by diagnostic study in patients presenting with pyelonephritis or hematuria, or by investigation of families of patients with known polycystic disease. At times, flank pain due to hemorrhage into a cyst occurs. Otherwise the symptoms and signs are those commonly seen in hypertension or renal insufficiency. On physical examination, the enlarged, irregular kidneys are often easily palpable.

The urine may contain leukocytes and erythrocytes. With bleeding into the cysts, there may also be bleeding into the urinary tract. The blood chemistry findings reflect the degree of renal insufficiency. Examination by sonography, CT scan, or x-ray shows the enlarged kidneys, and urography demonstrates the classic elongated calyces and renal pelves stretched over the surface of the cysts.

No specific therapy is available, and surgical interference is only indicated to decompress very large cysts in patients with severe pain.

Patients with polycystic kidney disease live in reasonable comfort with slowly advancing uremia. Both hemodialysis and renal transplantation extend the life of these patients. Nephrectomy is indicated only in patients with recurrent infections, severe recurrent bleeding, or markedly enlarged kidneys.

**B. CYSTIC DISEASE OF THE RENAL MEDULLA**

1. **Medullary cystic disease**—Medullary cystic disease is a familial disease that may become symptomatic during adolescence. Anemia is usually the initial manifestation, but azotemia, acidosis, and hyperphosphatemia soon become evident. Urine findings are not remarkable, although there is often an inability to concentrate and renal salt wasting often occurs. Many small cysts are scattered through the renal medulla. Renal transplantation is indicated by the usual criteria.

2. **Medullary sponge kidney**—Medullary sponge kidney is asymptomatic and is discovered by the characteristic appearance of tubular ectasia in the urogram. Enlargement of the papillae and calyces and small cavities within the pyramids is demonstrated by the contrast media in the excretory urogram. Many small calculi often occupy the cysts, and infection may be troublesome. Life expectancy is not affected and only therapy for ureteral stone or for infection is required.

**ANOMALIES OF THE PROXIMAL TUBULE**

**Defects of Amino Acid Reabsorption**

**A. CONGENITAL CYSTINURIA**

Increased excretion of cystine results in the formation of cystine calculi in the urinary tract. Ornithine, arginine, and lysine are also excreted in abnormally large quantities. There is also a defect in absorption of these amino acids in the jejunum. Nonopaque stones should be examined chemically to provide a specific diagnosis.

Treatment goals include a large fluid intake and keeping the urine pH above 7 by giving sodium bicarbonate and sodium citrate plus acetazolamide at bedtime to ensure an alkaline night urine. In refractory cases, a low-methionine (cystine precursor) diet may be necessary. Penicillamine has proved useful in some cases.

**B. AMINOACIDURIA**

Many amino acids may be poorly absorbed, resulting in unusual losses. Failure to thrive and the presence of other tubular deficits suggest the diagnosis. There is no treatment.

**C. HEPATOLENTICULAR DEGENERATION (WILSON’S DISEASE)**

In this congenital familial disease, aminoaciduria and renal tubular acidosis (RTA) are associated with cirrhosis of the liver and neurologic manifestations. Hepatomegaly, evidence of impaired liver function, spasticity, ataxia, emotional disturbances, and Kayser-Fleischer rings around the cornea constitute a unique syndrome. There is a decrease in synthesis of ceruloplasmin, with a deficit of plasma ceruloplasmin and an increase in free copper that may be etiologically specific.

Penicillamine is given to chelate and remove excess copper. Edathamil (EDTA) may also be used to remove copper.

**D. MULTIPLE DEFECTS OF TUBULAR FUNCTION (DE TONI-FANCONI-DEBRÉ SYNDROME)**

Aminoaciduria, phosphaturia, glycosuria, and a variable degree of RTA characterize this syndrome. Osteomalacia is a prominent clinical feature; other clinical and laboratory manifestations are associated with specific tubular defects described previously.

Treatment consists of replacing cation deficits (especially potassium), correcting acidosis with bicarbonate or citrate, replacing phosphate loss with isoinonic neutral phosphate (mono- and disodium salts) solution, and ensuring a
liberal calcium intake. Vitamin D is useful, but the dose
must be controlled by monitoring levels of serum calcium
and phosphate.

E. DEFECTS OF PHOSPHORUS &
CALCIUM REABSORPTION

Several sporadic, genetically transmitted, and acquired dis-
orders are grouped under this category and are character-
ized by persisting hypophosphatemia because of excessive
phosphaturia and an associated metabolic bone disorder,
rickets in childhood, and osteomalacia in adulthood.
Response to vitamin D therapy (1,25-dihydroxycholecalciferol, the active analog of vitamin D) is variable.

F. DEFECTS OF GLUCOSE ABSORPTION
(RENAL GLYCOUSURIA)

Renal glycosuria results from an abnormally poor ability to
reabsorb glucose and is present when blood glucose levels
are normal. Ketosis is not present. The glucose tolerance
response is normal. There is no treatment for renal glyco-
suria, just reassurance.

G. DEFECTS OF BICARBONATE REABSORPTION

Proximal RTA, type II, is due to reduced bicarbonate
reclamation in the proximal tubule, with resultant loss of
bicarbonate in the urine and decreased bicarbonate con-
centration in extracellular fluid. There are increased K+
losses into the urine and retrieval of Cl− instead of
HCO3. The acidosis is therefore associated with hypoka-
elia and hyperchloremia. Transport of glucose, amino
acids, phosphate, and urate may be deficient as well
(Fanconi syndrome).

ANOMALIES OF THE DISTAL TUBULE

Defects of Hydrogen Ion Secretion
& Bicarbonate Reabsorption (Classic
Renal Tubular Acidosis, Type I)

Failure to secrete hydrogen ion and to form ammonium ion
results in loss of “fixed base” sodium, potassium, and
calcium. There is also a high rate of excretion of phos-
phate. Vomiting, poor growth, and symptoms and signs of
chronic metabolic acidosis are accompanied by weakness
due to potassium deficit and bone discomfort due to
osteomalacia. Nephrocalcinosis, with calcification in the
medullary portions of the kidney, occurs in about one-half
of cases. The urine is alkaline and contains larger than nor-
mal quantities of sodium, potassium, calcium, and phos-
phate. An abnormality in urinary anion gap (U.Na+ +
U.K+ − U.Cl−) is noted (low), which is associated with the
reduced NH4+ production. This abnormality differentiates
this condition from type II RTA and from the metabolic
acidosis seen with diarrhea. The blood chemistry findings
are those of metabolic acidosis with low serum potassium.

Treatment consists of replacing deficits and increasing
the intake of sodium, potassium, calcium, and phospho-
rus. Sodium and potassium should be given as bicarbonate
or citrate. Additional vitamin D may be required.

Excess Potassium Secretion (Potassium
“Wastage” Syndrome)

Excessive renal secretion or loss of potassium may occur
in 4 situations: (1) moderate renal insufficiency with
diminished H+ secretion; (2) RTA (proximal and distal
RTA); (3) hyperaldosteronism and hyperadrenocortici-
cism; and (4) tubular secretion of potassium, the cause of
which is unknown. Hypokalemia indicates that the defi-
cit is severe. Muscle weakness and polyuria and dilute
urine are signs attributable to hypokalemia. Treatment
consists of correcting the primary disease and giving sup-
plemenary potassium.

Reduced Potassium Secretion

Reduced potassium secretion is noted in conditions in
which extrarenal aldosterone is reduced or when intrare-
nal production of renin (and secondary hypoaldoster-
onism) occurs. The latter condition is termed RTA, type
IV, and is associated with impaired H+ and K+ secretion
in the distal tubule. Drug-induced interstitial nephritis,
gout, and diabetes mellitus are clinical circumstances
that may produce type IV RTA and resulting hyperkale-
emia and mild metabolic acidosis. Treatment is to pro-
mote kaliuresis (with loop diuretics) to prescribe potas-
sium-binding gastrointestinal resins (Kayexalate), or to
provide the patient with a mineralocorticoid, fludrocort-
sisone acetate.

Defects of Water Absorption
(Renal Diabetes Insipidus)

Nephrogenic diabetes insipidus occurs more frequently in
males than females. Unresponsiveness to antidiuretic hor-
mone is the key to differentiation from pituitary diabetes
insipidus.

In addition to congenital refractoriness to antidiuretic
hormone, obstructive uropathy, lithium, methoxyflurane,
and demeclocycline also may render the tubule refractory
to vasopressin.

Symptoms are related to an inability to reabsorb water,
with resultant polyuria and polydipsia. The urine volume
approaches 12 L/d, and osmolality and specific gravity are
low.

Treatment consists primarily of an adequate water
intake. Chlorothiazide may ameliorate the polyuria; the
mechanism of action is unknown, but the drug may act by
increasing isosmotic reabsorption in the proximal segment
of the tubule.
UNSPECIFIED RENAL TUBULAR ABNORMALITIES

In idiopathic hypercalciuria, decreased reabsorption of calcium predisposes to the formation of renal calculi. Serum calcium and phosphorus are normal. Urine calcium excretion is high; urine phosphorus excretion is low. Microscopic hematuria may be present. See treatment of urinary stones containing calcium (Chapter 16).

REFERENCES

Oliguria literally means “reduced” urine volume—less than that necessary to remove endogenous solute loads that are the end products of metabolism. If the patient concentrates urine in a normal fashion, oliguria (for that person) is present at urine volumes under 400 mL/day, or approximately 6 mL/kg body weight. If the kidney concentration is impaired and the patient can achieve a specific gravity of only 1.010, oliguria is present at urine volumes under 1000–1500 mL/day.

Acute renal failure is a condition in which the glomerular filtration rate is abruptly reduced, causing a sudden retention of endogenous and exogenous metabolites (urea, potassium, phosphate, sulfate, creatinine, administered drugs) that are normally cleared by the kidneys. The urine volume is usually low (under 400 mL/day). If renal concentrating mechanisms are impaired, the daily urine volume may be normal or even high (high-output or nonoliguric renal failure). Rarely, there is no urine output at all (anuria) in acute renal failure.

The causes of acute renal failure are listed in Table 33–1. Prerenal renal failure is reversible if treated promptly, but a delay in therapy may allow it to progress to a fixed intrinsic renal failure (eg, acute tubular necrosis). The other causes of acute renal failure are classified on the basis of their involvement with vascular lesions, intrarenal disorders, or postrenal disorders.

**PRERENAL RENAL FAILURE**

The term prerenal denotes inadequate renal perfusion or lowered effective arterial circulation. The most common cause of this form of acute renal failure is dehydration due to renal or extrarenal fluid losses from diarrhea, vomiting, excessive use of diuretics, and so on. Less common causes are septic shock, “third spacing” with extravascular fluid pooling (eg, pancreatitis), and excessive use of antihypertensive drugs. Heart failure with reduced cardiac output also can reduce effective renal blood flow. Careful clinical assessment may identify the primary condition responsible for prerenal renal failure, but many times several conditions can coexist. In the hospital setting, these circulatory abnormalities often lead to more fixed, acute renal failure (acute tubular necrosis).

Acute reductions in glomerular filtration rate may also be noted in patients with cirrhosis (hepatorenal failure) or in patients taking cyclosporine, tacrolimus, nonsteroidal anti-inflammatory drugs, or angiotensin-converting enzyme inhibitors. It is felt that these conditions represent significant intrarenal hemodynamic functional derangements. In these clinical circumstances, the urinary findings may mimic prerenal renal failure, but the patient’s clinical assessment does not demonstrate the extrarenal findings seen in common prerenal conditions, as noted in the following section. Improvements in glomerular filtration rate are usually noted after drug discontinuance or, in cases of hepatorenal renal failure, with management of the liver disease or liver transplantation.

**Clinical Findings**

**A. SYMPTOMS AND SIGNS**

Except for rare cases with associated cardiac or “pump” failure, patients usually complain of thirst or of dizziness in the upright posture (orthostatic dizziness). There may be a history of overt fluid loss. Weight losses reflect the degree of dehydration. Physical examination frequently reveals decreased skin turgor, collapsed neck veins, dry mucous membranes, and, most important, orthostatic or postural changes in blood pressure and pulse.

**B. LABORATORY FINDINGS**

1. Urine—The urine volume is usually low. Accurate assessment may require bladder catheterization followed by hourly output measurements (which will also rule out lower urinary tract obstruction; see discussion following). High urine specific gravity (>1.025) and urine osmolality >600 mOsm/kg) also are noted in this form of acute apparent renal failure. Routine urinalysis usually shows no abnormalities.

2. Urine and blood chemistries—The blood urea nitrogen-creatinine ratio, normally 10:1, is usually increased with prerenal renal failure. Other findings are set forth in Table 33–2. Because mannitol, radiocontrast dyes, and diuretics affect the delivery and tubular handling of urea, sodium, and creatinine, urine and blood chemistry tests performed after these agents have been given to produce misleading results.

3. Central venous pressure—A low central venous pressure indicates hypovolemia. If severe cardiac failure is
CHAPTER 33

Table 33–1. Causes of Acute Renal Failure.

<table>
<thead>
<tr>
<th>I. Prerenal renal failure:</th>
<th>Acute Renal Failure</th>
<th>Prerenal Azotemia</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Dehydration</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Vascular collapse due to sepsis, antihypertensive drug therapy, &quot;third spacing&quot;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Reduced cardiac output</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

II. Functional–hemodynamic:

1. Angiotensin-converting enzyme inhibitor drugs
2. Nonsteroidal anti-inflammatory drugs
3. Cyclosporine; tacrolimus
4. Hepatorenal syndrome

III. Vascular:

1. Atheroembolism
2. Dissecting arterial aneurysms
3. Malignant hypertension

IV. Parenchymal (intrarenal):

1. Specific:
   a. Glomerulonephritis
   b. Interstitial nephritis
   c. Toxin, dye-induced
   d. Hemolytic uremic syndrome
2. Nonspecific:
   a. Acute tubular necrosis
   b. Acute cortical necrosis

V. Postrenal:

1. Calculus in patients with solitary kidney
2. Bilateral ureteral obstruction
3. Outlet obstruction
4. Leak, posttraumatic

the principal cause of prerenal renal failure (it is rarely the sole cause), reduced cardiac output and high central venous pressure are apparent.

4. Fluid challenge—An increase in urine output in response to a carefully administered fluid challenge is both diagnostic and therapeutic in cases of prerenal renal failure. Rapid intravenous administration of 300–500 mL of physiologic saline is the usual initial treatment. Urine output is measured over the subsequent 1–3 hours. A urine volume increase of more than 50 mL/h is considered a favorable response that warrants continued intravenous infusion. If the urine volume does not increase, the physician should carefully review the results of blood and urine chemistry tests, reassess the patient’s fluid status, and repeat the physical examination to determine whether an additional fluid challenge (with or without furosemide) might be worthwhile.

Treatment

In states of dehydration, fluid losses must be rapidly corrected to treat oliguria. Inadequate fluid management may cause further renal hemodynamic deterioration and eventual renal tubular ischemia (with fixed acute tubular necrosis; see discussion following). If oliguria and hypotension persist in a well-hydrated patient, vasopressor drugs are indicated in an effort to correct the hypotension associated with sepsis or cardiogenic shock. Pressor agents that restore systemic blood pressure while maintaining renal blood flow and renal function are most useful. Dopamine, 1–5 µg/kg/min, may increase renal blood flow without systemic pressor responses. Higher doses of 5–20 µg/kg may be necessary if systemic hypotension persists after volume correction. Discontinuance of antihypertensive medications or diuretics can, by itself, cure the apparent acute renal failure resulting from prerenal conditions.

Table 33–2. Acute Renal Failure versus Prerenal Azotemia.

<table>
<thead>
<tr>
<th></th>
<th>Acute Renal Failure</th>
<th>Prerenal Azotemia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urine osmolarity (mOsm/L)</td>
<td>&lt; 350</td>
<td>&gt; 500</td>
</tr>
<tr>
<td>Urine/plasma urea</td>
<td>&lt; 10</td>
<td>&gt; 20</td>
</tr>
<tr>
<td>Urine/plasma creatinine</td>
<td>&lt; 20</td>
<td>&gt; 40</td>
</tr>
<tr>
<td>Renal failure index* =</td>
<td></td>
<td></td>
</tr>
<tr>
<td>( \frac{U_\text{Na}}{U/P_\text{cr}} )</td>
<td>&gt; 1</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>( \frac{U/P_\text{Na}}{U/P_\text{cr}} \times 100 = )</td>
<td>&gt; 1</td>
<td>&lt; 1</td>
</tr>
</tbody>
</table>


VASCULAR RENAL FAILURE

Common causes of acute renal failure due to vascular disease include atheroembolic disease, dissecting arterial aneurysms, and malignant hypertension. Atheroembolic disease is rare before age 60 and in patients who have not undergone vascular procedures or angiographic studies. Dissecting arterial aneurysms and malignant hypertension are usually clinically evident.

Rapid assessment of the arterial blood supply to the kidney requires arteriography or other noncontrast blood flow studies (eg, magnetic resonance imaging or Doppler ultrasound). The cause of malignant hypertension may be identified on physical examination (eg, scleroderma). Primary management of the vascular process is necessary to affect the course of these forms of acute renal failure.

INTRARENAL DISEASE STATES; INTRARENAL ACUTE RENAL FAILURE

Diseases in this category can be divided into specific and nonspecific parenchymal processes.
1. Specific Intrarenal Disease States

The most common causes of intrarenal acute renal failure are acute or rapidly progressive glomerulonephritis, acute interstitial nephritis, toxic nephropathies, and hemolytic uremic syndrome.

Clinical Findings

A. Symptoms and Signs

Usually the history shows some salient data such as sore throat or upper respiratory infection, diarrheal illness, use of antibiotics, or intravenous use of drugs (often illicit types). Bilateral back pain, at times severe, is occasionally noted. Gross hematuria may be present. It is unusual for pyelonephritis to present as acute renal failure unless there is associated sepsis, obstruction, or involvement of a solitary kidney. Systemic diseases in which acute renal failure occurs include Henoch-Schönlein purpura, systemic lupus erythematosus, and scleroderma. Human immunodeficiency virus (HIV) infection may present with acute renal failure from HIV nephropathy.

B. Laboratory Findings

1. Urine—Urinalysis discloses variably active sediments: many red or white cells and multiple types of cellular and granular casts. Phase contrast microscopy usually reveals dysmorphic red cells in the urine. In allergic interstitial nephritis, eosinophils may be noted. The urine sodium concentration may range from 10 to 40 mEq/L.

2. Blood test—Components of serum complement are often diminished. In a few conditions, circulating immune complexes can be identified. Other tests may disclose systemic diseases such as lupus erythematosus. Thrombocytopenia and altered red cell morphologic structure are noted in peripheral blood smears in the hemolytic uremic syndrome. Rapidly progressive glomerulonephritis can be evaluated with tests for ANCA (antineutrophil cytoplasmic antibodies) and anti-GBM titers (anti-glomerular basement membrane antibodies).

3. Renal biopsy—Biopsy examination shows characteristic changes of glomerulonephritis, acute interstitial nephritis, or glomerular capillary thrombi (in hemolytic uremic syndrome). There may be extensive crescents involving Bowman’s space.

C. X-Ray Findings

Dye studies should be avoided because of the risk of dye-induced renal injury. For this reason, sonography is preferable to rule out obstruction.

Treatment

Therapy is directed toward eradication of infection, removal of antigen, elimination of toxic materials and drugs, suppression of autoimmune mechanisms, removal of autoimmune antibodies, or a reduction in effector-inflammatory responses. Immunotherapy may involve drugs or the temporary use of plasmapheresis. Initiation of supportive dialysis may be required (see discussion below).

2. Nonspecific Intrarenal States

Nonspecific intrarenal causes of acute renal failure include acute tubular necrosis and acute cortical necrosis. The latter is associated with intrarenal intravascular coagulation and has a poorer prognosis than the former. These forms of acute renal failure usually occur in hospital settings. Various morbid conditions leading to septic syndrome–like physiologic disturbances are often present.

Degenerative changes of the distal tubules (lower nephron nephrosis) are believed to be due to ischemia. With dialysis, most of these patients recover—usually completely—provided intrarenal intravascular coagulation and cortical necrosis does not occur.

Elderly patients, who are more prone to have this form of oliguric acute renal failure, develop following hypotensive episodes. It appears that exposure to some drugs such as nonsteroidal anti-inflammatory agents may increase the risk of acute tubular necrosis. Although the classic picture of lower nephron nephrosis may not develop, a similar nonspecific acute renal failure is noted in some cases of mercury (especially mercuric chloride) poisoning and following exposure to radiocontrast agents, especially in patients with diabetes mellitus or myeloma.

Clinical Findings

A. Symptoms and Signs

Usually the clinical picture is that of the associated clinical state. Dehydration and shock may be present concurrently, but the urine output and acute renal failure fail to improve following administration of intravenous fluids, in contrast to patients who have prerenal renal failure (see preceding discussion). On the other hand, there may be signs of excessive fluid retention in patients with acute renal failure following radiocontrast exposure. Symptoms of uremia per se (eg, altered mentation or gastrointestinal symptoms) are unusual in acute renal failure (in contrast to chronic renal failure).

B. Laboratory Findings

(See also Table 33–2.)

1. Urine—The specific gravity is usually low or fixed in the 1.005–1.015 range. Urine osmolality is also low (<450 mOsm/kg and U/P osmolar ratio <1.5:1). Urinalysis often discloses tubular cells and granular casts; the urine may be muddy brown. If the test for occult blood is positive, one must be concerned about the presence of myoglobin or
hemoglobin. Tests for differentiating myoglobin pigment are available.

2. Central venous pressure—This is usually normal to slightly elevated.

3. Fluid challenges—There is no increase in urine volume following intravenous administration of mannitol or physiologic saline. Occasionally, following the use of furosemide or “renal doses” of dopamine (1–5 µg/kg/min), a low urine output is converted to a high fixed urine output (low-output renal failure to high-output renal failure).

Treatment

If there is no response to the initial fluid or mannitol challenge, the volume of administered fluid must be sharply curtailed to noted losses. An assessment of serum creatinine and blood urea nitrogen and of the concentrations of electrolytes is necessary to predict the possible use of dialysis. With appropriate regulation of the volume of fluid administered, solutions of glucose and essential amino acids to provide 30–35 kcal/kg are used to correct or reduce the severity of the catabolic state accompanying acute tubular necrosis.

Serum potassium must be closely monitored to ensure early recognition of hyperkalemia. This condition can be treated with (1) intravenous sodium bicarbonate administration, (2) Kayexalate, 25–50 g (with sorbitol) orally or by enema, (3) intravenous glucose and insulin, and (4) intravenous calcium preparations to prevent cardiac irritability.

Peritoneal dialysis or hemodialysis should be used as necessary to avoid or correct uremia, hypokalemia, or fluid overload. Hemodialysis in patients with acute renal failure can be either intermittent or continuous (with arteriovenous or venovenous hemofiltration techniques). Vascular access is obtained with percutaneous catheters. The continuous dialysis techniques allow for easier management in many hemodynamically unstable patients in intensive care units.

Prognosis

Most cases are reversible within 7–14 days. Residual renal damage may be noted, particularly in elderly patients.

POSTRENAL ACUTE RENAL FAILURE

The conditions listed in Table 33–1 involve primarily the need for urologic diagnostic and therapeutic interventions. Following lower abdominal surgery, urethral or ureteral obstruction should be considered as a cause of acute renal failure. The causes of bilateral ureteral obstruction are (1) peritoneal or retroperitoneal neoplastic involvement, with masses or nodes; (2) retroperitoneal fibrosis; (3) calculous disease; and (4) postsurgical or traumatic interruption. With a solitary kidney, ureteral stones can produce total urinary tract obstruction and acute renal failure. Urethral or bladder neck obstruction is a frequent cause of renal failure, especially in elderly men. Posttraumatic urethral tears are discussed in Chapter 17.

Clinical Findings

A. SYMPTOMS AND SIGNS

Renal pain and renal tenderness often are present. If there has been an operative ureteral injury with associated urine extravasation, urine may leak through a wound. Edema from overhydration may be noted. Ileus is often present along with associated abdominal distention and vomiting.

B. LABORATORY FINDINGS

Urinalysis is usually not helpful. A large volume of urine obtained by catheterization may be both diagnostic and therapeutic for lower tract obstruction.

C. X-RAY FINDINGS

 Radionuclide renal scans may show a urine leak or, in cases of obstruction, retention of the isotope in the renal pelvis. Ultrasound examination often reveals a dilated upper collecting system with deformities characteristic of hydronephrosis.

D. INSTRUMENTAL EXAMINATION

Cystoscopy and retrograde ureteral catheterization demonstrate ureteral obstruction.

Treatment

For further discussion of ureteral injuries, see Chapter 17.

REFERENCES


Overview
In chronic renal failure, reduced clearance of certain solutes principally excreted by the kidney results in their retention in the body fluids. The solutes are end products of endogenous metabolism as well as exogenous substances (eg, drugs). The most commonly used indicators of renal failure are blood urea nitrogen and serum creatinine. The clearance of creatinine can be used as a reasonable measure of glomerular filtration rate (GFR).

Renal failure may be classified as acute or chronic depending on the rapidity of onset and the subsequent course of azotemia. An analysis of the acute or chronic development of renal failure is important in understanding physiologic adaptations, disease mechanisms, and ultimate therapy. In individual cases, it is often difficult to establish the duration of renal failure. Historical clues such as preceding hypertension or radiologic findings such as small, shrunken kidneys tend to indicate a more chronic process. Acute renal failure may progress to irreversible chronic renal failure. For a discussion of acute renal failure, see Chapter 32.

A new classification has been made to delineate chronic kidney disease (CKD) by varying degrees of reduced GFR (or creatinine clearance). This is presented in Table 34–1. This has been useful in studies of the progression of CKD, especially in varying drug regimens to reduce the rate of worsening of GFRs.

The incidence of end-stage renal disease (ESRD) is 330 cases per million population. These patients with ESRD require chronic dialysis or renal transplantation for life support. All age groups are affected. The severity and the rapidity of development of uremia are hard to predict. The use of dialysis and transplantation is expanding rapidly worldwide. A large increase in CKD in the past 20 years has been due to type 2 diabetes. Over 330,000 ESRD patients in the United States are currently treated with dialysis. Besides diabetes, increasingly older patients are being treated for other renal diseases. At the present time, 128,000 patients have functioning kidney transplants.

Historical Background
There are various causes of progressive renal dysfunction leading to end-stage or terminal renal failure. In the 1800s, Bright described several dying patients who presented with edema, hematuria, and proteinuria. Chemical analyses of sera drew attention to retained nitrogenous compounds and an association was made between this and the clinical findings of uremia. Although the pathologic state of uremia was well described, long-term survival was not achieved until chronic renal dialysis and renal transplantation became available after 1960–1970. Significant improvements in patient survival have been made in the past 50 years.

Etiology
A variety of disorders are associated with CKD. Either a primary renal process (eg, glomerulonephritis, pyelonephritis, congenital hypoplasia) or a secondary one (owing to a systemic process such as diabetes mellitus or lupus erythematosus) may be responsible. Once there is kidney injury, it is now felt that hyperfiltration to undamaged nephron units produces further stress and injury to remnant kidney tissue. The patient will show progression from one stage of CKD severity to the next. Superimposed physiologic alterations secondary to dehydration, infection, obstructive uropathy, or hypertension may put a borderline patient into uncompensated chronic uremia.

Clinical Findings
A. Symptoms and Signs
With milder CKD, there may be no clinical symptoms. Symptoms such as pruritus, generalized malaise, lassitude, forgetfulness, loss of libido, nausea, and easy fatigability are frequent and nonfocal complaints in moderate to severe CKD. Growth failure is a primary complaint in preadolescent patients. Symptoms of a multisystem disorder (eg, systemic lupus erythematosus) may be present coincidentally. Most patients with CKD have elevated blood pressure secondary to volume overload or from hyperreninemia. However, the blood pressure may be normal or low if patients have marked renal salt-losing tendencies (eg, medullary cystic disease). The pulse and respiratory rates are rapid as manifestations of anemia and metabolic acidosis. Clinical findings of uremic fetor, pericarditis, neurologic findings of asterixis, altered mentation, and peripheral neuropathy are present only with severe, stage V CKD. Palpable kidneys suggest polycystic disease. Ophthalmoscopic
Table 34–1. Chronic Kidney Disease (CKD) Stages.

<table>
<thead>
<tr>
<th>Stage</th>
<th>GFR (cc/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>&gt;90 with microalbuminuria</td>
</tr>
<tr>
<td>II</td>
<td>60–89 with microalbuminuria</td>
</tr>
<tr>
<td>III</td>
<td>30–59</td>
</tr>
<tr>
<td>IV</td>
<td>15–29</td>
</tr>
<tr>
<td>V</td>
<td>&lt;15 or dialysis</td>
</tr>
</tbody>
</table>


Examination may show hypertensive or diabetic retinopathy. Alterations involving the cornea have been associated with metabolic disease (eg, Fabry disease, cystinosis, and Alport hereditary nephritis).

B. HISTORY

In 20% of cases, there is a family history of CKD. A report of antecedent nephritis episodes or a history of previous proteinuria may be elicited. It is important to review drug usage and possible toxic exposures (eg, lead).

C. LABORATORY FINDINGS

1. Urine composition—The urine volume varies depending on the type of renal disease. Quantitatively normal amounts of water and salt losses in urine can be associated with polycystic and interstitial forms of disease. Usually, however, urine volumes are quite low when the GFR falls below 5% of normal. The urinary concentrating and acidification mechanisms are impaired. Daily salt losses become more fixed, and, if they are low, a state of positive sodium balance occurs with resulting edema. Proteinuria can be variable. Urinalysis examinations may reveal mononuclear white blood cells (leukocytes) and occasionally broad waxy casts, but usually the urinalysis is nonspecific and inactive.

2. Blood studies—Anemia is the rule with normal platelet counts. Platelet dysfunction or thrombasthenia is characterized by abnormal bleeding times. Several abnormalities in serum electrolytes and mineral metabolism become manifest when the GFR drops below 30 mL/min. Progressive reduction of body buffer stores and an inability to excrete titrable acids result in progressive acidosis characterized by reduced serum bicarbonate and compensatory respiratory hyperventilation. The metabolic acidosis of uremia is associated with a normal anion gap, hyperchloremia, and normokalemia. Hyperkalemia is not usually seen unless the GFR is below 5 mL/min. In patients with interstitial renal diseases, gouty nephropathy, or diabetic nephropathy, hyperchloremic metabolic acidosis with hyperkalemia (renal tubular acidosis, type IV) may develop. In these cases the acidosis and hyperkalemia are out of proportion to the degree of renal failure and are related to a decrease in renin and aldosterone secretion. In moderate to severe CKD, multiple factors lead to an increase in serum phosphate and a decrease in serum calcium. The hyperphosphatemia develops as a consequence of reduced phosphate clearance by the kidney. In addition, vitamin D activity is diminished because of reduced conversion of vitamin D_{2} to the active form of vitamin D_{3} in the kidney. These alterations lead to secondary hyperparathyroidism with skeletal changes of both osteomalacia and osteitis fibrosa cystica. Uric acid levels are frequently elevated but rarely lead to calculi or gout during chronic uremia.

D. X-RAY FINDINGS

Patients with reduced renal function should not be routinely subjected to contrast studies. Renal sonograms are helpful in determining renal size (usually small) and cortical thickness (usually thin) and in localizing tissue for percutaneous renal biopsy. Bone x-rays may show retarded growth, osteomalacia (renal rickets), or osteitis fibrosa. Soft-tissue or vascular calcification may be noted on plain films. Patients with polycystic kidney disease will have variably large kidneys with evident cysts (on sonograms or plain abdominal CT scans).

E. RENAL BIOPSY

Renal biopsies may not reveal much except nonspecific interstitial fibrosis and glomerulosclerosis. There may be pronounced vascular changes consisting of thickening of the media, fragmentation of elastic fibers, and intimal proliferation, which may be secondary to uremic hypertension or due to primary arteriolar nephrosclerosis. Percutaneous or open biopsies of end-stage shrunken kidneys are associated with a high morbidity rate, particularly bleeding.

Treatment

Recent studies indicate some benefit of drugs to reduce progression of CKD. These approaches include the use of angiotensin-converting enzyme inhibitors, angiotensin receptor blockers, lipid-lowering agents, and aldosterone antagonists. Patients need to be followed closely for potential hyperkalemia.

Overall, management should be conservative until it becomes impossible for patients to continue their customary lifestyles. Restriction of dietary protein (0.5 g/kg/day), potassium, and phosphorus is recommended. As well, maintenance of close sodium balance in the diet is necessary so that patients become neither sodium-expanded nor -depleted. This is best done by the accurate and frequent monitoring of the patient’s weight. Use of oral bicarbonate can be helpful when moderate acidemia occurs. Anemia can be treated with recombinant erythropoietin given subcutaneously. Prevention of possible uremic osteodystrophy and secondary hyperparathyroidism requires close atten-
tion to calcium and phosphorus balance. Phosphate-retaining antacids and calcium or vitamin D supplements may be needed to maintain the balance. Cinacalcet can directly reduce parathyroid hormone secretion. If severe secondary hyperparathyroidism occurs, subtotal parathyroidectomy may be needed.

A. CHRONIC PERITONEAL DIALYSIS

Chronic peritoneal dialysis is used electively or when circumstances (ie, no available vascular access) prohibit chronic hemodialysis. Ten percent of dialysis is done with this treatment. Improved soft catheters can be used for repetitive peritoneal lavages. In comparison to hemodialysis, small molecules (such as creatinine and urea) are cleared less effectively than larger molecules, but excellent treatment can be accomplished. Intermittent thrice-weekly treatment (IPPD), continuous cycler-assisted peritoneal dialysis (CCPD), or chronic ambulatory peritoneal dialysis (CAPD) is possible. With the latter, the patient performs 3–5 daily exchanges using 1–2 L of dialysate at each exchange. The dialysate contains a high glucose concentration and the peritoneal surface serves as the semipermeable membrane. Bacterial contamination and peritonitis are becoming less common with improvements in technology.

B. CHRONIC HEMODIALYSIS

Chronic hemodialysis using semipermeable dialysis membranes is now widely performed. Access to the vascular system is provided by an arteriovenous fistula, vascular grafts (with autologous saphenous vein or synthetic material), or by a percutaneous permcatheter (placed either surgically or with interventional radiology). The actual dialyzers are of various geometries. Body solutes and excessive body fluids can be easily cleared by using dialysate fluids of known chemical composition. Newer, high-efficiency membranes (high/flux) are serving to reduce dialysis treatment time.

Treatment is intermittent—usually 3–5 hours three times weekly. Computer modeling, using measurements of urea kinetics, has provided more precise hemodialysis prescriptions. Treatments may be given in a kidney center, a satellite unit, or the home. Home dialysis is optimal because it provides greater scheduling flexibility and is generally more comfortable and convenient for the patient, but only 20% of dialysis patients meet the requirements for this type of therapy.

More widespread use of dialytic techniques has permitted greater patient mobility. Treatment on vacations and business trips can be provided by prior arrangement.

Common problems with either type of chronic dialysis include infection, bone symptoms, technical accidents, persistent anemia, and psychological disorders. The excessive morbidity and mortality associated with atherosclerosis often occurs with long-term treatment. It is now recognized that occasionally uremic patients, despite dialysis, can develop wasting syndrome, cardiomyopathy, polyneuropathy, and secondary dialysis-amyloidosis so that kidney transplant must be urgently done. Routine bilateral nephrectomy should be avoided because it increases the transfusion requirements of dialysis patients. Nephrectomy in dialysis patients should be performed in cases of refractory hypertension, reflux with infection, and cystic disease with recurrent bleeding and pain. The dialysis patient can occasionally have acquired renal-cystic disease. Such patients need close monitoring for the development of in situ renal cell carcinoma.

Yearly costs range from an average of $50,000 for patients who receive dialysis at home to as much as $50,000–$75,000 for patients treated at dialysis centers, but much of this is absorbed under HR-1 (Medicare) legislation. If the patient has no other systemic problems (eg, diabetes), the mortality rates are 8–10%/year once maintenance dialysis therapy is instituted. Despite these medical, psychological, social, and financial difficulties, most patients lead productive lives while receiving dialysis treatment.

C. RENAL TRANSPLANTATION

After immunosuppression techniques and genetic matching were developed, renal homotransplantation became an acceptable alternative to maintenance hemodialysis. Improved transplantation results are now noted owing to the development of newer immunosuppressant drugs. Currently employed posttransplant drugs include prednisone, azathioprine, mycophenolate mofetil, cyclosporine, tacrolimus, sirolimus, and a variety of injectable bioagents. The great advantage of transplantation is reestablishment of nearly normal and constant body physiology and chemistry. Diet can be less restrictive. The disadvantages include bone marrow suppression, susceptibility to infection, oncogenesis risks, and the psychological uncertainty of the homograft’s future. Most of the disadvantages of transplantation are related to the medicines given to counteract the rejection. Later problems with transplantation include recurrent disease in the transplanted kidney and an increased incidence of cancer. Genitourinary infection appears to be of minor importance if structural urologic complications (eg, leaks) do not occur.

Nephrology centers, with close cooperation between medical and surgical staff, attempt to use these treatment alternatives of dialysis and transplantation in an integrated fashion.

For a more detailed review, see Chapter 35.

REFERENCES


The 50th anniversary of the first successful kidney transplant from a live donor to his identical twin was celebrated in 2004. During this interval, kidney transplantation has progressed from an experimental procedure to the preferred method of renal replacement therapy worldwide. This is a result of continually improving outcomes producing a better quality of life, and a prolongation of survival compared to dialysis (Wolfe et al, 1999). At the end of 2005, in the United States, there were about 325,000 patients receiving renal replacement therapy, with an incident rate of about 330 per million population (USRDS, 2006). In 2005, there were 16,477 kidney transplants performed in the United States, 9914 from deceased donors and 6562 from live donors (www.optn.org, 2006). However, over 65,000 patients were actively waiting for a kidney, and the gap between the number waiting and available organs widens every year (Port et al, 2006). Currently, over 65,000 patients were actively waiting for a kidney, and the gap between the number waiting and available organs widens every year (Port et al, 2006). Currently, 1- and 5-year kidney graft survival ranges between 89–95% and 66–80%, depending on donor source (Cohen et al, 2006) (Figure 35–1). The major reasons leading to improved outcomes are more potent yet selective immunosuppression, better surgical techniques, more sensitive cross-matching, and better prophylaxis and treatment of morbidity infections. There is also an emerging consensus that preemptive transplantation, immediately prior to the need to dialysis, is advantageous in reducing much morbidity and even mortality (Kasiske et al, 2002).

A. GENITOURINARY TRACT EVALUATION

It is important that the native urinary tract will function properly after transplant, and an accurate urologic history is essential. Potential recipients without a history of urologic symptoms or prior interventions do not need an extensive evaluation. Upper tract ultrasound and urine cultures usually suffice; some recommend voided cytology. An age-appropriate screening PSA in males. Patients with a history of urologic symptoms (especially hematuria, infections, stones, and incontinence), prior interventions, or a neurogenic bladder should have a full urologic evaluation including upper tract/pelvic imaging, a voiding cystogram, cystoscopy and retrograde studies, cytology, and, if indicated, a urodynamics study. If patients are dialyzed, upper tract computed tomography (CT) scans with intravenous (IV) contrast can be done, while contrast is avoided in late stages of renal failure. The use of gadolinium for MRI in late stages of renal failure should be avoided due to the risk of nephrogenic systemic fibrosis/nephrogenic fibrosing dermopathy (Grobner, 2007).

Upper Tract Abnormalities—Removal of the native kidneys, once advocated, is uncommon today and needed in 10% or fewer patients. Residual urine output and potassium excretion, even if small, as well as production of erythropoietin and vitamin D3 via the retained kidneys are considered beneficial. Medical indications for nephrectomies are rare, and include heavy proteinuria (>10 g/day), intractable hypertension (4–5 drugs), and persistent hematuria. Kidneys with chronic hydronephrosis, high-grade reflux, stones,
abscesses, filling defects, enhancing masses, complex or very large cysts, etc. that may lead to persistent infections or harbor potential cancers should be removed prior to transplant. In addition, very large polycystic kidneys may need removal for relief of symptoms or size considerations. Potential recipients with acquired renal cystic disease, found in 1/3–1/2 of all dialyzed patients, need intervention for contrast enhancing lesions. In most instances nephrectomy or nephroureterectomy can be done laparoscopically with diminished morbidity (Ghasemian et al, 2005; Ismail et al, 2005). Lesions not removed prior to transplant will need surveillance after the transplant.

**Lower Tract Abnormalities**—It is important to remember that dialyzed patients often have a diminished urine volume, resulting in a small-capacity bladder with low compliance. Such bladders will resume normal function, even 25 years later, once urine volume is restored (Serrano et al, 1996). However, small capacity bladders that are fibrotic and scarred from prior surgery, radiation, old TB, congenital anomalies (posterior urethral valves, meningomyelocele, etc.) will not recover. In these rare cases, often children, the preferred option is a bladder augmentation with bowel (ileum, stomach, colon, or dilated ureter) or a continent neobladder to produce a compliant reservoir with adequate volume (Nahas et al, 2002; Mendizabal et al, 2004; Rigamonti et al, 2005). Bladder augmentation is not without risk, as mucous production, residual urine, and infection often require subsequent intermittent catheterization. If the bladder is absent or destroyed, an ileal conduit can be created for transplantation (Hatch et al, 1993). It is advisable that such major reconstructions be done and healed prior to transplantation. Experience has taught that operations on a dry urinary tract, ie, bladder neck incisions, urethral stricture repair, and prostatectomy will lead to restructuring and further scarring. Therefore, they should only be done when urine volume is more than a liter per day; or if not, delayed for about 3 months after the transplant. This includes older males who may experience progressive prostatic growth absent urine output while on dialysis. These recipients can develop symptoms of prostatism or even urinary retention after the transplant, which will require treatment. During this interval for immunosuppression reduction and complete engraftment, recipients can be managed with a suprapubic tube, or preferably intermittent clean catheterization (Flechner et al, 1983).

**B. INFECTION**

**Bacterial**—Active infections are a contraindication to transplantation, which need to be appropriately treated and resolved. This may include surgical drainage of abscesses or removal of a chronic nidus such as bone, teeth, sinus, etc. The urinary tract should be sterile for transplantation. Recurrent urinary tract infections require a full urologic evaluation including upper tract imaging, a voiding cystogram, cystoscopy, and retrograde studies. Recipients with a prior history of tuberculous disease or exposure should receive a year of isoniazid prophylaxis. ESRD patients may be anergic to skin testing.

**Viral**—Herpes family DNA viruses such as cytomegalovirus (CMV), Epstein-Barr Virus (EBV), varicella zoster (VCZ), and herpes simplex (HSV) can be transmitted with the donor organ or reactivated from a latent state in the recipient. Therefore, recipients are usually given prophylaxis with a nucleoside inhibitor such as oral valganciclovir for 3 months, especially...
when they are seronegative and the donor is seropositive. Those patients with serologic evidence of prior hepatitis B or C exposure have diminished outcomes, especially if their liver has evidence of cirrhosis. However, those recipients with inactive liver disease who have antibodies to either virus may receive organs from donors that are also positive for either hepatitis B core or hepatitis C antibody (Akalin et al, 2005; Arolfdi et al, 2005). A relationship between hepatitis C infection and posttransplant diabetes is also emerging (Bloom and Lake, 2006). Renal failure patients with active and untreated human immunodeficiency virus (HIV) should not be further immunosuppressed by transplant. However, those stable HIV-positive individuals treated with current antiretroviral drug therapy can do well for up to 5 years after kidney transplant (Qiu et al, 2006).

C. MALIGNANT DISEASE

Active or recently recurrent malignant disease is an absolute contraindication to renal transplantation. The bulk of evidence suggests that immunosuppressive therapy facilitates the growth of residual cancers. The safe waiting period for transplantation after surgical removal of solid tumors varies and depends on the grade and stage of tumor on presentation and the associated risk of recurrence. Penn (1997) reported that of 1137 neoplasms treated prior to transplantation, the overall recurrence rate was 21%. Fifty-four percent recurred for those waiting only 2 years before transplantation, 33% in those waiting 2–5 years before transplantation, and 13% among those waiting more than 5 years. The highest recurrence rates occurred with breast carcinomas (23%), symptomatic renal carcinomas (27%), sarcomas (29%), bladder carcinomas (29%), nonmelanoma skin cancers (53%), and multiple myeloma (67%).

Therefore, with some exceptions, a minimum waiting period of 2 years for cancers with a favorable prognosis is desirable. A waiting period of 5 years is desirable for lymphomas, most carcinomas of the breast, colon, or for large (>5 cm) symptomatic renal carcinomas. No waiting period is necessary for incidentally discovered small renal carcinomas, in situ carcinomas, and possibly tiny focal neoplasms. More recently it has been suggested that rather than using fixed waiting times, it is more logical to use cancer recurrence nomograms to establish risk. This has been well established for localized prostate cancer, where risk for recurrence can be compared to mortality risk on dialysis to establish an individualized assessment (Secin et al, 2004).

D. SYSTEMIC AND METABOLIC DISEASE

Patients with certain metabolic diseases affecting the kidney such as Fabry’s disease, hemolytic uremic syndrome, vasculitis, systemic lupus erythematosus, amyloidosis, etc. as well as various forms of glomerulonephritis and focal sclerosis may experience recurrence, and patients should be counseled regarding this possibility (Couser, 2005). Those with severe metabolic stone disease that resulted in kidney loss will often experience recurrent stones and a poor outcome. A combined hepatic and kidney transplant is now commonly recommended for primary hyperoxaluria (Jamieson, 2005) and less so for cystinosis (Rogers et al, 2001).

E. CARDIOVASCULAR STATUS

Cardiovascular disease represents the leading cause of death after kidney transplantation and is ubiquitous among renal failure patients, especially diabetics and those over age 50. Potential recipients should be thoroughly screened, and have symptomatic lesions corrected prior to transplant since those with ESRD are at high risk for ischemic events (Pilmore, 2006). Since many dialysis patients are sedentary, already have abnormal EKG patterns, and diabetics may not experience angina with exertion, provocative stress tests are necessary. However, subjects should reach their target heart rate for these tests to have an accurate predictive value. If any uncertainty exists, the gold standard remains coronary angiography. In an analysis of dialysis patients undergoing coronary revascularization, Herzog et al (2002) found that although the in-hospital mortality was greatest for those undergoing coronary artery bypass grafting (CABG) (8.6%) compared to patients having stents (4.1%) or PTCA (6.4%); the 2-year patient all-cause survival was significantly superior for those after CABG (56.4%) than after stenting (48.4%) or PTCA (48.2%). Patients with a history of strokes or transient ischemic attacks should be screened with a carotid ultrasound and receive neurology clearance. Those with adult polycystic kidney disease need a brain MR angiogram to screen for aneurysms. Peripheral vascular disease is common in renal failure, especially diabetics, and ultrasound screening can be helpful. A pelvic CT scan without contrast can be helpful to determine the degree of calcification of the pelvic vessels and aid in kidney placement. Active claudication, femoral bruises, or diminished pulses demands a complete vascular surgical assessment.

F. GASTROINTESTINAL DISEASE

Patients with ESRD often have a history of gastrointestinal (GI) problems such as peptic ulcer disease, gastroesophageal reflux, cholecystitis, pancreatitis, inflammatory bowel disease, diverticulosis, chronic diarrhea or constipation, or hemorrhoids. If present, these should be evaluated and resolved prior to transplant. Upper or lower GI endoscopy and/or contrast imaging of the bowel may be required. Routine cholecystectomy for asymptomatic cholelithiasis is no longer advised (Jackson et al, 2005).
G. MODIFIABLE RISK FACTORS

Obesity—In North America obesity is affecting a greater number of patients with renal failure each year. Numerous reports have identified obesity (BMI >30 kg/m²) and morbid obesity (BMI >35 kg/m²) as an independent risk factor for increased cardiovascular mortality, decreased graft survival, delayed graft function (DGF), wound complications, posttransplant diabetes, proteinuria, and prolonged hospitalization (Modlin et al, 1997; Armstrong et al, 2005; Gore et al, 2006). Weight reduction to under the morbidly obese range is desirable, and may require bariatric surgery in extreme circumstances (Alexander et al, 2004).

Smoking—Tobacco smoking is particularly deleterious for transplant recipients, and patients need to stop prior to transplantation. Smoking both accelerates the progression of atherosclerotic cardiovascular disease and is nephrotoxic to the kidney resulting in proteinuria (Tozawa et al, 2002; Orth, 2004).

H. BLOOD TRANSFUSION

The use of intentional third party blood transfusions to modulate the immune system is no longer done. In fact, transfusions are generally avoided; both to prevent the possibility of disease transmission (hepatitis, HIV, etc.), and to prevent recipient sensitization to human leukocyte antigen (HLA) phenotypes that may diminish the chance of a negative cross-match with a potential donor. Anemia of renal failure is effectively treated with recombinant erythropoietin for most patients (Cody et al, 2005).

I. TRANSPLANT ALLOGRAFT NEPHRECTOMY

After a failed transplant, immunosuppression is weaned off and the patient returns to dialysis. If graft loss occurs after a year it is usually not necessary to remove the failed graft, as a new kidney can be placed on the contralateral side. In a few cases, when graft failure is early or is due resistant rejection, the kidney tissue may undergo necrosis and the graft needs to be removed. Indications for allograft nephrectomy include fevers, graft tenderness, gross hematuria, malaise, infection, and uncontrolled hypertension. The subcapsular allograft nephrectomy is the safest approach to prevent iliac vessel injury.

SELECTION OF DONORS

Living Donors

A. DIRECTED LIVING KIDNEY DONORS

Living kidney donation provides a better patient and allograft survival when compared with deceased-donor transplantation, especially when the live donor transplant is performed before the onset of dialysis (Figures 35–1 and 35–2) (Meier-Kriesche et al, 2002). Living donation rates vary worldwide, but in many Western countries, Asia, and the Middle East it has recently increased to be the predominant form of kidney transplantation. In the United States, the annual number of live kidney donors has surpassed the number of deceased donors since 2001, although the absolute number of transplants from deceased donors still outnumbers those from living donors (LDs) (Delmonico et al, 2005). Based on tissue typing disparities (HLA mismatches), an immunologic hierarchy can be established for the best “match” (Table 35–1). The advantages for identical twins and HLA identical siblings are quite significant; while all other live donor combinations are simi-
B. NONDIRECTED LIVING KIDNEY DONORS

The extreme shortage of kidneys to meet the demand of waiting recipients coupled with the success of LURD kidney transplantation has opened up creative ways to expand the pool of live donors. In particular, there are individuals who wish to be anonymous donors, i.e., “nondirected or altruistic donor.” However, in the United States, living-donor exchanges must adhere to Section 301 of the National Organ Transplant Act of 1984 (NOTA), which states, “It shall be unlawful for any person to knowingly acquire, receive, or otherwise transfer any human organ for valuable consideration for use in human transplantation.” Valuable consideration according to this act has traditionally been considered to be monetary transfer or a transfer of valuable property between the donor and the recipient. The donation of an organ is properly considered to be a legal gift. With these constraints any person who is competent, willing to donate, free of coercion, and found to be medically and psychosocially suitable may be a live kidney donor (Adams et al, 2002). Three protocols of nondirected living donation have been developed to accommodate such donors: (1) a live-donor paired exchange, (2) a live-donor/deceased-donor exchange, and (3) altruistic donation.

C. LIVE-DONOR PAIRED EXCHANGE

This approach involves exchanging donors who are ABO or cross-match incompatible with their intended recipients so that each donates a kidney to a compatible recipient (Delmonico, 2004). The exchange derives the benefit of live donation but avoids the risk of incompatibility; several computer algorithms have been modeled to execute the exchange (Roth et al, 2004). The best example is two families, one with an A donor to a B recipient and the second with B donor to and A recipient. Swapping donors solves the dilemma. Live-donor exchange procedures have been performed worldwide and are best performed with large sharing pools (Kranenburg et al, 2004).

D. LIVE-DONOR/DECEASED-DONOR EXCHANGE

Another system of exchange of donors was devised by centers in UNOS region 1, by permitting the live donor to be used by another compatible individual on the waiting list in “exchange” for the next blood type compatible deceased donor in the region, for the live donor’s recipient. With this method two patients will be transplanted instead of only one, although some fine tuning of donor organ quality and age is necessary.

E. ALTRUISTIC LIVING DONORS

Altruistic kidney donation (to a complete stranger) is developmental in several centers and must be approached with utmost sensitivity, especially today when organ exchanges are advertised on the internet. Participating centers usually offer the kidney to the highest wait-listed patient at their center after a match run. The motives of the nondirected donor should be established with care to avoid a prospective donor’s intention of remedying a psychological disorder via donation. Many who inquire about altruistic donation have only a limited understanding of these issues, and upon learning these basic realities about 60% withdraw from the process (Jacobs et al, 2004).

F. LIVING DONOR SAFETY

From its inception, the removal of a kidney from a healthy individual to benefit another has been problematic. The practice is based upon the belief that the removal of one kidney does not diminish survival or significantly harm long-term kidney function. This notion derives from fol-
low-up of patients up to 45 years after nephrectomy for trauma (Narkun-Burgess et al, 1993), and after kidney donation (Najarian et al, 1992; Fehrman-Ekholm et al, 1997). The effect of reduced renal mass by uninephrectomy on 3124 patients was compared to 1703 matched controls in a meta-analysis (Kasiske et al 1995). The reason for nephrectomy included organ donation in 60.5%, cancer 10.1%, infection 8.1%, stones/obstruction 6.8%, agenesis 3.4%, trauma 2.5%, and other 8.4%. Unilateral nephrectomy caused an average decrease of 17 mL/min in the GFR that tended to improve with each 10 years of follow-up (average increase 1.4 mL/min/decade). A small, progressive increase in proteinuria was also noted (average 76 mg/decade) but was negligible after nephrectomy for trauma or kidney donation, and nephrectomy did not affect the prevalence of hypertension. Thus, the published evidence indicates that there is little long-term medical risk to a healthy donor after unilateral nephrectomy. Nevertheless, Ellison et al (2002), identified 56 live kidney donors who were subsequently listed for a kidney transplant. The rate of ESRD in kidney donors was calculated to be 0.04%, comparable to the rate of ESRD in the general US population (0.03%). The renal diagnosis in these patients was hypertension, focal sclerosis, chronic glomerulonephritis, familial nephropathy, diabetes, and other. Recently some have advocated use of donors with isolated medical abnormalities such as hypertension, obesity, dyslipidemia, or stones, which may not result in the safety profiles previously reported.

Deceased Donors

The imbalance between the supply of brain-dead deceased donors and the growing demand for kidneys has created many innovative uses of organs that were excluded in the past. These generally include kidneys from donors over the age of 60, the presence of systemic disease such as atherosclerosis, hypertension or early diabetes, donors with cardiac arrest or significant hypotension, and some with prior exposure to virus and/or infections that have resolved (Ismail and Flechner, 2006). While kidneys that are severely traumatized or come from donors with active cancer, sepsis, or HIV-AIDs, are excluded, a number of donor organs with extended criteria that convey about a 10% worse overall graft survival have been incorporated into the donor pool. To maximize kidney usage, the following categories have been developed.

A. STANDARD CRITERIA DONORS

Most individuals that meet the criteria for brain death from age 5–60 years with normal kidney function and no history of systemic or infectious disease.

B. EXPANDED CRITERIA DONORS

Kidneys from brain-dead donors with a 1.7 times relative risk of graft failure. These criteria were developed from a consensus conference that analyzed registry survival data (Rosengard et al, 2002). These include any donor > age 60 or > age 50 with a history of hypertension, CVA death or creatinine >1.5mg/dL. (Table 35–2). Informed consent of the recipient is requested to receive an expanded criteria donor (ECD) kidney.

C. DONATION AFTER CARDIAC DEATH

When a potential donor does not meet brain-death criteria but has an irretrievable head injury, viable organs for transplant can be procured after a controlled cardiac arrest. Such kidneys experience a greater incidence of DGF, but long-term function is comparable to standard donor kidneys (Rudich et al, 2002).
D. Dual Transplants

At the extremes of life, one kidney may not be sufficient to deliver an adequate glomerular filtration rate (GFR) (nephron dose) to an adult recipient. In these instances using both kidneys from a single donor can overcome these limitations.

1. Pediatric en-bloc—Kidneys from donors under age 5 (often <6 cm in length) have a historically higher failure rate from technical problems and develop hyperfiltration injury (proteinuria) when transplanted into adults (Bresnahan et al, 2001). Both kidneys can be transplanted en-bloc, attached to the donor aorta and vena cava, in a more reliable fashion (Hobart et al, 1998). Such kidneys will grow to adult size in a year.

2. Adult dual transplants—When kidneys have extremely unfavorable risk factors for graft success due to insufficient nephron mass, both may provide for successful outcome (Bunnapradist et al, 2003). Such adult dual transplants can be placed in either iliac fossa, or preferably on the same side through one incision. The criteria establish for dual kidney allocation appear in Table 35–3. This approach utilizes kidneys that in the past were often discarded.

E. Extracorporeal Renal Preservation

1. Simple hypothermic storage and flush solutions—Once removed, kidneys are flushed and stored in a hyperosmolar, hyperkalemic, and hyponatremic solution at (4–10° C) to minimize ischemic injury (cellular swelling). This is usually sufficient for up to 24 hours of preservation although longer cold ischemic times (up to 40 hours) have been reported, but result in higher rates of DGF. A commercial storage solution from the University of Wisconsin is frequently used, which contains inert substrates like lactobionate, raffinose, hydroxyethyl starch, and adenosine as an energy substrate. Recently, a less viscous alternative Histidine-Tryptophan-Ketoglutarate (HTK) solution has been shown to yield similar results with cold ischemia times < and >24 hours. (Agarwal et al, 2006).

2. Pulsatile perfusion—Hypothermic pulsatile perfusion is an alternative method of preservation, which takes advantage of a continuous pulsatile flow through the graft. Some feel such hydrodistention is therapeutic in dilating the ischemic renal microcirculation, and permits the delivery of vasodilator drugs (ie, verapamil, beta-blockers). It also permits measurement of flow, pulse pressure, and resistance through the graft, which is an accurate method to determine viability of the kidney (Schold et al, 2005). Pulsatile perfusion is more costly and requires investment in a preservation unit (Waters Co, Rochester, MN) and a technologist, but has been gaining popularity due to the increasing number of expanded criteria donors that are considered for transplant (Matsuoka et al, 2006).

Table 35-2. The Expanded Criteria for Kidney Donors. The Decision Matrix Using Relative Risk for Graft Failure >1.7 for Donors Older Than 10 Years of Age. Used for Organ Allocation in the United States.

<table>
<thead>
<tr>
<th>Donor Age Category (Years)</th>
<th>Donor Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;10</td>
<td>CVA + HTN + Cr &gt;1.5</td>
</tr>
<tr>
<td>10–39</td>
<td>X</td>
</tr>
<tr>
<td>40–49</td>
<td>X</td>
</tr>
<tr>
<td>50–59</td>
<td>X</td>
</tr>
<tr>
<td>&gt;60</td>
<td>X</td>
</tr>
<tr>
<td>CVA + HTN</td>
<td>X</td>
</tr>
<tr>
<td>CVA + Cr &gt;1.5</td>
<td>X</td>
</tr>
<tr>
<td>HTN + Cr &gt;1.5</td>
<td>X</td>
</tr>
<tr>
<td>CVA</td>
<td>X</td>
</tr>
<tr>
<td>HTN</td>
<td>X</td>
</tr>
<tr>
<td>Cr &gt;1.5</td>
<td>X</td>
</tr>
<tr>
<td>None of above</td>
<td>X</td>
</tr>
</tbody>
</table>

CVA = cerebrovascular accident; HTN = hypertension; Cr >1.5 = creatinine over 1.5 mg/dL.
Source: OPTN.

Table 35-3. Criteria for Adult Dual Cadaveric Kidney Transplants.

Kidneys from adult donors must be offered singly unless the donor meets at least 2 of the following conditions and the OPO would not otherwise use the kidneys singly.

(A) Donor age >60 years.
(B) Estimated donor creatinine clearance <65 mL/min based upon serum creatinine upon admission.
(C) Rising serum creatinine (>2.5 mg/dL) at time of retrieval.
(D) History of medical disease in donor (defined as either longstanding hypertension or diabetes mellitus).
(E) Adverse donor kidney histology (defined as moderate to severe glomerulosclerosis (>15% and <50%).

Kidneys from donors older than 60 years, and kidneys from donors with rising serum creatinine (>2.5 mg/dL) at time of retrieval, and kidneys from donors with history of medical disease (either longstanding hypertension or diabetes mellitus), and kidneys from donors with adverse donor kidney histology (defined as moderate to severe glomerulosclerosis (>15% and <50%) must be offered singly unless used for Adult Dual Transplantation. In these instances, both kidneys can be transplanted en-bloc, attached to the donor aorta and vena cava, in a more reliable fashion (Hobart et al, 1998). Such kidneys will grow to adult size in a year.
THE MAJOR HISTOCOMPATIBILITY COMPLEX (MHC)

Tissue Typing—The MHC describes a region of genes located on chromosome 6 in man which encode proteins that are responsible for the rejection of tissue between different species or members of the same species (Flechner, Finke, and Fairchild, 2006). The cell surface MHC markers are called human leukocyte antigens, because they were first identified on white blood cells. There are two major types of HLA antigens termed class I and class II. Virtually all nucleated cells express HLA class I antigens, while class II antigens are primarily found on B cells, monocytes, macrophages, and antigen-presenting cells. Each individual inherits two serologically defined class I (called A and B) and one class II (called Dr) antigen from each parent; so six HLA antigens constitute an individual’s tissue type. One set of HLA A, B, and Dr antigens inherited from a parent is called a haplotype, so HLA-identical siblings have inherited both haplotypes. The HLA molecules are polymorphic (over 150 defined), so it is very unusual if two unrelated individuals have the same tissue type of six HLA antigens. HLA antigens not shared between two individual will generate an immune response. Therefore, the term HLA matching describes the number of shared antigens. One can generate a hierarchical rating of genetic HLA similarities, which roughly correlate to the risk for rejection and eventual kidney transplant outcomes ranging from identical twins to DD (Table 35–1). In clinical practice, the impact of HLA on graft survival is small the first years, but plays an important role after 5–10 years. No doubt other factors affect survival; especially donor organ quality (age, function, size, etc.) as well as recipient age and comorbidities. However, at the present time 6-Ag matched (or zero HLA mismatched) deceased donor kidneys are shared nationally due to the beneficial effect on immunological outcomes (Takemoto et al, 1993). In addition, HLA antigen matches also play a role in the algorithm for distribution of deceased donor kidneys with more points assigned for better matches.

Cross-matching—Preformed circulating anti-HLA antibodies against the specific phenotype of the donor will lead to acute (if not hyperacute) rejection. Such antibodies (usually IgG) are detected by cross-matching the sera of the recipient with lymphocytes of the donor and adding complement. Such complement-dependent cytotoxicity (CDC) will kill the donor cells and is indicative of deleterious clinical outcome. A similar yet more sensitive test has been developed using flow cytometry to identify the presence of anti-HLA antibodies bound to the surface of donor lymphocytes. A cross-match against both donor T and B lymphocytes is performed within 24 hours of surgery, and transplants are not done if these antibodies are present. In addition, the ABO system will trigger CDC against the mismatched blood group antigens (glycoproteins) present on many tissues. Therefore, transplants are usually done only between ABO-compatible individuals.

Serum Screening—At monthly intervals waiting patients have their serum screened for the presence of anti-HLA antibodies against a panel of HLA phenotypes (lymphocytes) that represent the general population. The result is reported as a percent of the total referred to as percent reactive antibody (PRA). Those with high titer (>50%) of anti-HLA antibody against the broad population are said to be sensitized and will find it very hard to find a cross-match-negative donor. Sensitized patients waiting for an organ depend on better HLA matches to find a cross-match-negative donor (McCune et al, 2002). Sensitization to HLA can occur from prior blood transfusions, viral infections, pregnancy, or previous transplants.

Posttransplant Antibodies—The development of de novo donor-specific or non-donor-specific anti-HLA antibodies after the transplant has a deleterious effect on outcomes. Both a greater frequency of acute and chronic rejection as well as lower graft survival have been reported among those patient with these antibodies detected by flow cytometry (El Fettouh et al, 2001; Hourmant et al, 2005). The presence of these antibodies may identify those recipients that need more rather than less immunosuppression.

DONOR NEPHRECTOMY FOR TRANSPLANTATION

Removal of a kidney for transplant depends upon minimizing both surgical injury and warm ischemia, which will hasten the recovery of function in the recipient. It is best to ensure a brisk diuresis in the donor before the kidney is removed, which can be enhanced by the use of volume expansion with saline and albumin, osmotic diuretics (mannitol), and loop diuretics (furosemide) in order to maximize immediate graft function in the recipient. Minimal dissection of the renal hilum is preferred.

A. LIVING DONORS

1. Evaluation—All donors should be evaluated both medically and surgically to ensure donor safety. An outline of the usual donor evaluation is shown in Table 35–4. First a thorough history and physical exam is needed to rule out hypertension, diabetes, obesity, infections, cancers, and specific renal/urologic disorders. Then laboratory testing of blood and urine, chest x-ray, electrocardiogram, and appropriate cardiac stress testing is done. Different methods to measure GFR and urine protein excretion are incorporated. Finally, radiographic assessment of the kidneys and vessels is ordered, which is usually accomplished by a CT angiogram (Kapoor et al, 2004). A catheter angiogram is reserved for complex anatomy. The donor is always left with the better kidney. If the two kidneys are
equal, the left is preferred for transplant due to its longer and often thicker renal vein. However, in cases when one kidney has multiple renal arteries, the kidney with the single artery is selected. In younger fertile female donors, concern about physiologic hydronephrosis of the right kidney is taken into consideration.

2. Surgical technique—Today, the most commonly used approach is intraperitoneal laparoscopic donor nephrectomy, primarily due to patient choice (Moinzadeh and Gill, 2006). This technique has all but supplanted open donor nephrectomy via an extraperitoneal flank incision due to reports of reduced pain and shorter recovery time. An alternative is the hand-assisted laparoscopic approach, where the extraction incision is used during the dissection (Fisher et al, 2006). Nevertheless, in cases with a short right vein or 3 or more arteries, we prefer an open nephrectomy using 12th rib-sparing flank incision (Turner-Warwick, 1965). When multiple renal arteries are encountered, they should be conjoined ex vivo while the kidney is on ice, in order to minimize the number of anastomoses in the recipient and reduce ischemia times (Flechner and Novick, 2002). Smaller upper pole arteries (<2 mm) often can be sacrificed, while lower pole vessels should be retained because of a risk to the ureteral blood supply.

B. DECEASED DONORS

Today, most donors are multiple-organ donors, and they require removal of the liver, heart and lungs, and pancreas, in addition to the kidneys. The retrieval needs to be coordinated and is often performed by several teams representing each organ for transplant. Usually the thoracic organs are removed first while the abdominal organs are cooled and perfused with UW or HTK perfusion solution. The kidneys are removed en bloc with the aorta and vena cava and a large amount of retroperitoneal tissue. They are separated on the bench by dividing the great vessels with the renal vessels attached.

### STANDARD RENAL TRANSPLANT SURGERY

There are several different methods for surgical revascularization of the kidney; the following is one reliable method (Goldfarb, Flechner, and Modlin, 2006). While either iliac fossa is acceptable for the transplant, the right side is often preferred due to the longer and more horizontal segments of external iliac artery and vein compared to the left side. A lower quadrant curvilinear (Gibson) incision is made, and the iliac vessels are exposed through a retroperitoneal approach, a self-retaining retractor is used. The renal-to-iliac-vein anastomosis is usually performed first, in an end-to-side fashion with 5-0 nonabsorbable monofilament suture, using a running quadrant technique. The renal artery can be anastomosed end-to-end to the internal iliac using 6-0 nonabsorbable monofilament suture. However, in older recipients and diabetics this vessel often has significant arterial plaque causing poor runoff. In addition, concern about compromising arterial flow to the penis via the pudendal artery with subsequent erectile dysfunction limits this approach in older males. Because of these factors, an end-to-side anastomosis of the renal artery to the external iliac artery is more frequently done with 6-0 nonabsorbable monofilament suture using a running quadrant technique. An extravesical ureteroneocystostomy (variation of Lich technique) is the preferred method to reimplant the ureter. When healthy-appearing ureter is short or the bladder is dysfunctional and small, a native to transplant uretero-ureterostomy can be done. An internal double J ureteral stent is always placed; and a closed suction drain is left in the deep pelvis.
IMMEDIATE POSTTRANSPLANT CARE

IMAGING OF THE TRANSPLANT KIDNEY

Immediately after the transplant, it is advisable to obtain a baseline duplex Doppler ultrasound to confirm patency of the renal vessels, blood flow to the parenchyma, and to identify large fluid collections, hematomas, or hydrenephrosis. This is especially important when the graft is oliguric. Similar information can be obtained using an isotopic (mercaptoacetyltriglyceride, 99mTc-MAG-3) renal scan, which is especially helpful to identify urinary extravasation. Kidneys with DGF demonstrate a typical pattern of isotopic uptake with little clearance or excretion. If fluid collections or intraperitoneal problems are suspected, finer definition can be obtained with a CAT scan. The use of 3-D CAT scans or MR angiography can delineate actual vascular lesions (stenoses, aneurysms, a-v fistula). Catheter angiography is reserved for interventions that require access to the renal vessels such as angioplasty. Imaging with IV-iodinated contrast should be limited when the creatinine is over 1.8 mg/dL, but cystograms and antegrade nephrostograms can be helpful to identify urinary fistulas or obstructions.

IMMEDIATE POSTTRANSPLANT CARE

A. HEMODYNAMIC MANAGEMENT

Initial postsurgical care the first hours and days focuses on the urine output and eventual recovery of GFR. It is important to avoid hypotension, dehydration, or use of alpha-adrenergic drugs, which will exacerbate surgical and preservation injury. It is helpful to monitor central venous pressures to maintain adequate preload (10–15 cm water). Urine outputs >1 cc/kg/hr are desirable, and hourly IV replacement at cc/cc of urine is usually sufficient. Some live donor kidneys may generate outputs up to a liter per hour, which will drop the blood pressure and should be managed with only 1/2–2/3 volume replaced. Alternatively, fluid overload and pulmonary edema may cause renal hypoperfusion and should be avoided. Treatment with fluid restriction, diuretics, and even dialysis may be needed. Even when hemodynamically stable, many DD recipients (and a few LD recipients) will experience delayed recovery of graft function, which is a consequence of extended cold preservation time in the recipient.

B. DELAYED RECOVERY OF GRAFT FUNCTION

DGF is more formally defined as the need for dialysis the first week after transplant and occurs in about a third of DD recipients. The term slow graft function (SGF) is said to occur if the recipient creatinine is not under 3 mg/dL by day 5, and occurs in another third of DD recipients (Humar et al., 2002). Patients with DGF may produce liters of urine a day (non-oliguric DGF), but have a rising creatinine and need dialysis. Others produce under 300 cc a day of urine and are described as oliguric DGF, which is usually an indication of a more prolonged recovery time. These clinical events are associated with specific histological findings referred to as acute tubular necrosis (ATN), the hallmark of which is tubular epithelial swelling, necrosis, and regeneration with mitotic figures. If kidneys are in oliguric DGF for over a week and imaging studies demonstrate good blood flow, a biopsy should be done to rule out rejection and confirm ATN. Transplant DGF resolves in most cases, but may take up to several weeks; while about 1–2% of grafts never function (primary nonfunction). DGF does have a negative impact on both short- and long-term graft survival compared to kidneys that function immediately (Shoskes et al., 1997). During DGF it is helpful to delay the introduction of calcineurin inhibitor (CNI) drugs for 7–10 days until some recovery of function is evident. This usually requires the use of an induction antibody as an umbrella of protection until the graft heals.

C. SUDDEN DROP IN URINE OUTPUT

During the first few days, a sudden loss of urine output after an initial diuresis demands prompt attention to ensure patency of the Foley catheter, and if easily obtainable, a repeat ultrasound to confirm vascular flow and exclude hydrenephrosis. If there is any question of abnormal blood flow or a delay in obtaining an imaging study, the kidney should be promptly reexplored since vascular compromise of a few hours will result in allograft necrosis. Loss of urine output from the bladder catheter with increased drain output may suggest a urine fistula. The drainage fluid can be sent for creatinine, and if 5–10 times the serum level suggests urine. If the above problems are excluded with imaging studies, renal biopsy is needed to rule out acute rejection or thrombotic microangiopathy, and to ensure graft viability.

TRANSPLANT REJECTION

The disparate HLA phenotypes on donor tissue trigger an immune response that leads to renal dysfunction and histological changes in the transplanted kidney called rejection. These responses are both humoral and cellular, and depend upon the presentation of processed donor HLA antigens via either donor (direct) or host (indirect) antigen-presenting cells to the recipient’s immunocompetent T cells (Flechner, Finke, and Fairchild, 2006). The clinical signs and symptoms of acute renal allograft rejection include fever, chills, lethargy, hypertension, pain and swelling of the graft, diminished urine output, edema, an elevated serum creatinine and BUN, and proteinuria. Immunosuppression is designed to prevent these events. Rejection can also be divided in three distinct clinical entities based on the timing and mechanism responsible for triggering these events.
Hyperacute rejection—occurs immediately after revascularization of a kidney when preformed cytotoxic anti-HLA antibody is present. It will lead to graft thrombosis, and the kidney must be removed. While there is no treatment, it can be prevented almost completely by using the sensitive cross-matching techniques available today.

Acute rejection—episodes can occur at anytime after the transplant, but most occur in the first 3 months. Such episodes can be mild or severe and cause the symptoms previously described to a variable degree. With the currently available immunosuppression about 20% or less of transplant recipients experience acute rejection and most episodes are reversible with treatment. Less than 5% of recipients lose their graft due to unresponsive acute rejection. These episodes are predominantly cellular and cause graft infiltration of cytotoxic cells, but humoral mechanisms contribute to the process.

Chronic rejection—defines a process of gradual, progressive, decline in renal function over time. It is associated with hypertension and proteinuria, and is accompanied by histological features of tubular atrophy, interstitial fibrosis, and an occlusive arteriolopathy (Figure 35–4). It can be detected as early as 6 months after transplant, and is thought to have a strong humoral response against the graft. Some, but not all recipients have had prior acute rejections or have donor-specific antibody detected. There is a role for alloimmunity (antigen dependent factors), since it does not occur in identical twins, is rare in HLA-identical sibling transplants, and is most common among DD recipients (Kreiger et al, 2003). However, many of these histologic changes are found with older donor age, ischemic injury, viral infections, and other systemic comorbidities, referred to as antigen-independent factors. Therefore the process remains less well characterized, is no doubt multifactorial, and is often given the name chronic allograft nephropathy (CAN). Treatment is often not effective, and consists of tight control of blood pressure, the use of ACE/ARB drugs for proteinuria, and sparing or elimination of CNI drugs.

IMMUNOSUPPRESSION

The goal in transplantation is to develop methods that permit a recipient to keep a transplanted organ in a state of “tolerance” or donor-specific unresponsiveness. Until that day arrives, clinical practice is dependent on our ability to interrupt the host immune response using agents that are not precise. It is a constant struggle to deliver enough immunosuppression to prevent rejection, but not too much to render the patient susceptible to infections and cancers. In addition, immunosuppressive drugs have unique mechanisms of action and their own specific toxicities. (Halloran, 2004). Immunosuppressive agents can be used in one of three ways: (1) high dose or induction therapy to prevent a primary immune response immediately after transplantation, (2) low dose or maintenance therapy initiated once the graft function has stabilized, or (3) additional high dose therapy to treat acute rejection.

A. CHEMICAL IMMUNOSUPPRESSION WITH SMALL MOLECULES

1. Corticosteroids—Since the initial observations more than 40 years ago that corticosteroids could prevent and treat renal allograft rejection (Hume et al, 1963), they have become the cornerstone of immunosuppressive therapy. Corticosteroids have numerous effects on the immune sys-
2. Antiproliferative drugs—

a. Azathioprine—Introduced first in the 1960s, 6-mercaptopurine and its imidazole derivative azathioprine represent antimetabolites that blocks purine biosynthesis and cell division. The developers of azathioprine, Gertrude Elion and George Hitchings, received the 1988 Nobel Prize. Azathioprine is most effective if given immediately after antigen presentation to prevent rejection and is ineffective in treating established rejection. Adverse effects of azathioprine include bone marrow suppression (primarily leukopenia), alopecia, hepatotoxicity, and increased risk of infection and neoplasia. When compared directly with another antiproliferative agent, mycophenolate mofetil (MMF), azathioprine is not as potent in rejection prophylaxis. Therefore, its use has been diminishing rapidly over the past few years, but serves as a secondary agent replacing MMF for intractable toxicity.

b. Mycophenolate mofetil—MMF is a morpholinoethyl ester of the fungal antibiotic mycophenolic acid, which is a noncompetitive inhibitor of the enzyme inosine monophosphate dehydrogenase. MMF inhibits purine biosynthesis preventing the proliferation of activated T and B cells, thereby blocking both cellular and humoral immune responses. It is thought to be more specific for those lymphocytes that rely primarily on de novo purine synthetic pathways, and has replaced azathioprine as an antimetabolite. MMF is usually well tolerated at dosages up to 2 g (divided dosing), with GI disorders (nausea, vomiting, cramps, and diarrhea) and bone marrow suppression (leukopenia, anemia) being its major toxicities. Recently therapeutic drug monitoring of blood levels have been reported to address interpatient variability, efficacy, and some reduction in GI toxicity.

c. Cyclophosphamide—Cyclophosphamide has historically been used in place of azathioprine, although it is much less commonly used today. It is an alkylating agent that is biotransformed by the hepatic microsomal oxidase system to active alkylating metabolites. It inhibits DNA replication and, like azathioprine, affects rapidly dividing cells and is most effective immediately after antigen presentation. Cyclophosphamide has a narrower therapeutic-toxic ratio than azathioprine, and adverse effects include myelosuppression with leukopenia, fertility disorders, and hemorrhagic cystitis.

d. Leflunomide—Leflunomide is an oral agent that inhibits the enzyme dihydro-orotate dehydrogenase, essential for de novo pyrimidine synthesis. The drug exhibits both antiproliferative and antiinflammatory activity, and was initially approved for the treatment of rheumatoid and psoriatic arthritis. Its use in organ transplantation as an adjunctive agent is limited. The most common side effects include diarrhea, nausea, dyspepsia, rash, abnormal liver function test results, or marrow suppression. Interestingly, the major metabolite has antiviral activity against CMV and polyoma virus, which can infect transplant recipients (Josephson et al, 2006).

3. Antilymphocytic drugs—

a. Calcineurin inhibitor drugs—Cyclosporine, a lipophilic small molecule, has been the cornerstone of transplant immunosuppression since the early 1980s and is the prototype CNI drug. It binds to a specific intracellular immunophilin (cyclophilin) causing conformational changes and subsequent engaging of the enzyme calcineurin phosphatase; thereby preventing the downstream gene transcription of IL-2 and other cytokines required for T-cell activation and proliferation. The adverse effects of cyclosporine, which are related to the concentration of the drug, include nephrotoxicity, hypertension, hyperlipidemia, gingival hyperplasia, hirsutism, and the hemolytic uremic syndrome. CNI drugs are metabolized by the hepatic cytochrome P-450 (3A4) system, and other drugs that inhibit or stimulate this enzyme system (ie, diltiazem and ketoconazole, or phenytoin and isoniazid) can significantly affect blood levels, thus favoring therapeutic drug monitoring. Recent developments include monitoring of the peak cyclosporine levels 2 hours after administration to better reflect exposure to the drug. A microemulsion that exhibits more
reproducible absorption and metabolism has replaced the initial oral formulation.

Tacrolimus is another CNI drug that engages a different immunophilin, FK-binding protein 12 (FKBP-12), to create a complex that inhibits calcineurin with greater molar potency than does cyclosporine. Some centers report better rejection prophylaxis with tacrolimus, but recent analyses suggest that with the current dosing strategies the efficacy of cyclosporine and tacrolimus are similar. Tacrolimus can also result in nephrotoxicity and the hemolytic uremic syndrome. It is more likely to induce new onset diabetes after transplant and neurological irritability (seizures, tremors). Compared to cyclosporine it seems less likely to cause hyperlipidemia, hypertension, and cosmetic problems. The use of tacrolimus has increased steadily and is now the dominant CNI, but many transplantation programs selectively use both agents, depending on individual patient risks. Hypertension, hyperlipidemia, and cosmetic changes argue for tacrolimus, whereas a high risk of diabetes (eg, older age or obesity), seizure risk argues for cyclosporine.

However, the most distressing feature of continuous CNI use is acute and chronic nephrotoxicity. Acute CNI nephrotoxicity is mediated by pronounced vascular and to a lesser degree tubular alterations, manifested by oligoanuria and azotemia, with associated hyperkalemia, hyperuricemia, hypertension, hypomagnesia, and renal tubular acidosis. A dose-dependent reduction in renal blood flow and glomerular filtration is well documented. Chronic CNI nephrotoxicity is more insidious, associated with progressive deterioration of graft histology (scarring) in over 50% by 5 years and virtually all treated patients by 10 years (Nankivell et al, 2003). CNI-treated recipients have a profile of upregulated genes associated with profibrotic/fibrotic activity and tissue remodeling (Flechner et al, 2004). Dosage reduction will often mitigate against some these effects, and numerous regimens have been tested to try to minimize or eliminate CNI drugs; although it must be done carefully to avoid increased risk of rejection (Russ et al, 2005; Abramowicz et al, 2005). In a carefully controlled comparison of monitored exposure to cyclosporine versus tacrolimus (Rowshani et al, 2006) reported a similar degree of scarring at 1 year after transplant. Calcium channel blockers are often used to ameliorate CNI nephrotoxicity due to their ability to reduce the dosage requirements, treat the associated hypertension, and reverse the calcium-dependent afferent arteriolar vasoconstriction.

b. Target-of-rapamycin inhibitors—Sirolimus and everolimus form a class of immunosuppressive agents that have similar molecular structure to the CNIs, and bind to the same immunophilin protein (FKBP-12) as tacrolimus. However, their mode of action appears to be distinct, as the sirolimus complex does not inhibit calcineurin. Instead, the sirolimus-FKBP complex appears to engage a distinct p70 kinase called mTOR (molecular target of rapamycin). The inhibition of mTOR blocks IL-2 signal transduction pathways that prevent cell-cycle progression from G to S phase in activated T cells. The principal nonimmune toxic effects of sirolimus and everolimus include hyperlipidemia, marrow suppression, and impaired wound healing and lymphocytes. Other reported side effects include delayed recovery from ATN, reduced testosterone concentrations, aggravation of proteinuria, mouth ulcers, and pneumonitis. However, sirolimus and everolimus may reduce CMV disease. Sirolimus and everolimus were developed for use with cyclosporine, but the combination increased nephrotoxicity, the hemolytic–uremic syndrome, and hypertension. Sirolimus has been combined with tacrolimus, but this combination also produced renal dysfunction and hypertension; which indicates that sirolimus potentiates CNI nephrotoxicity. Practitioners can reduce the toxicity of the combination of a TOR and CNI inhibitors by withdrawing one of the drugs (Russ et al, 2005). TOR inhibitors may have antineoplastic and arterial-protective effects. Since these agents slow the growth of established experimental tumors, they have potential applications in oncology (Guba et al, 2002). The possibility that sirolimus and everolimus can protect arteries is suggested by two observations: TOR inhibitors that are incorporated into coronary stents inhibit restenosis (Morice et al, 2002), and TOR inhibitors plus CNI inhibitors reduce the incidence of graft coronary artery disease associated with heart transplantation (Eisen et al, 2003).

B. Antilymphocyte Antibodies

1. Polyclonal antibodies—Polyclonal antibodies are produced by injecting (immunizing) animals such as horses, goats, sheep, or rabbits with cells from human lymphoid tissue. Immune sera from several animals are pooled and the gamma globulin fractions extracted and purified. A rabbit-derived antithymocyte antibody (Thymoglobulin, Genzyme) is the most frequently used preparation. Once injected, the antibodies bind to lymphocytes resulting in a rapid lymphopenia or depletion due to complement-mediated cell lysis; as well as masking of surface antigens or induction of suppressor populations that block cell function. Polyclonal antibodies have been used primarily in cadaveric renal transplantation, initially as induction therapy, and to treat vascular or antibody-mediated rejection. Because of their strong immunosuppressive effects, polyclonal antibodies are limited to short courses of 3–10 days, but their depletion may last 6–12 months. Adverse effects include fever, chills, and arthralgias related to the injection of foreign proteins and the release of cytokines. These effects can be minimized by pretreatment with corticosteroids and antihistamines. More serious adverse effects include increased susceptibility to infections (especially viral), and neoplasia.
2. Monoclonal antibodies that deplete lymphocytes—The introduction of murine hybridoma technology opened the door to the development of highly specific antibodies directed against functional cell surface targets. These antibodies, like polyclonal antibodies, exert their effects through a variety of immune mechanisms. In addition to complement-mediated lysis, blockade and inactivation of cell surface molecules, and opsonization with phagocytosis, these antibodies can induce cytotoxicity and modulation of cell surface molecules on target tissues.

a. Muromonab-CD3—Muromonab-CD3, a mouse monoclonal antibody against CD3, was the first commercially available monoclonal antibody used in transplantation for induction and to treat rejection. Muromonab-CD3 binds to the T-cell receptor-associated CD3 complex, which first triggers a massive cytokine-release syndrome before both depleting and functionally modulating T cells. Humans can make neutralizing (human antimouse) antibodies against muromonab-CD3 that terminate its effect and limit its reuse. Adverse effects from a typical 5-mg dose include a first-dose response that simulates a severe flu-like syndrome, consisting of fever, chills, nausea, vomiting, diarrhea, myalgias, headache, and in severe cases, aseptic meningitis and pulmonary edema. These effects can be minimized (but not eliminated) by pretreatment with corticosteroids and antihistamines. Prolonged courses of muromonab-CD3 increase the risk of posttransplantation lymphoproliferative disease (PTLD). The use of muromonab-CD3 has declined due to the introduction of humanized and/or chimeric antibodies that are better tolerated.

b. Alemtuzumab—Alemtuzumab is a humanized monoclonal antibody (IgG1) that specifically interacts with the 21- to 28-kd lymphocyte cell surface glycoprotein CD52, which is predominantly expressed on peripheral blood lymphocytes, monocytes, and macrophages. Once engaged with CD52, it produces a profound depletion of lymphocyte populations (T, B, and NK) that can persist for over a year. Although multiple doses are approved for treating B-cell chronic lymphocytic leukemia, one or two 30-mg doses have been cautiously introduced as an induction agent in organ transplantation. Side effects of alemtuzumab include first-dose reactions, bone marrow suppression, and autoimmunity. Worries concerning immunodeficiency complications (infections and cancer) with alemtuzumab persist until long-term data emerge. Early predictions that the agent would induce proper or “almost” tolerance were not confirmed, as some reports suggested a higher than expected incidence of rejection episodes, including antibody-mediated rejection.

c. Rituximab—Rituximab is chimeric anti-CD20 monoclonal antibody that eliminates most B cells, and was initially approved for treating refractory non-Hodgkin’s B-cell lymphomas. Interestingly it was introduced in transplantation to treat a similar tumor, PTLD. Rituximab is currently being evaluated to treat donor-specific alloantibody responses such as antibody-mediated rejection or in transplanting sensitized recipients. It is used in combination with maintenance immunosuppressive drugs, plasmapheresis, and intravenous immune globulin. While plasma cells are usually CD20-negative, some precursors are CD20-positive and their elimination may reduce some antibody responses. Such therapy may provide the first of future tools to control humoral rejection.

3. Monoclonal antibodies that are nondepleting—

a. Daclizumab and basiliximab—Another selective site for monoconal antibody targeting of the immune response is the IL-2 receptor (CD25), present on the surface of activated T cells and responsible for further signal transduction and T-cell proliferation. Both a chimeric (basiliximab) and a humanized (daclizumab) anti-CD25 have been genetically engineered to produce a hybrid IgG that retains the specific anti-CD25 binding characteristics with a less xenogenic (murine) backbone. These agents cause minimal cytokine release upon first exposure, and exhibit a prolonged elimination half-life resulting in weeks to months of CD25 suppression. Because expression of CD25 (interleukin-2 receptor a chain) requires T-cell activation, anti-CD25 antibody causes little depletion of T cells. Anti-CD25 antibodies are useful as safe induction agents in low- to moderate-risk recipients, but have little effect in treating an established rejection episode. Their use appear to offer a favorable risk-benefit compared to depleting agents, providing for improved graft survival with a lower risk of posttransplant cancers (Opelz et al, 2006).

b. Belatacept—Basic immunology generated the concept that blocking costimulation (signal 2) could prevent the activation of antigen-primed T cells, thus providing a new avenue for control of allograft rejection. A first generation of monoclonal antibodies designed to block costimulation proved the concept in animals, but lacked sufficient efficacy in initial clinical trials. Belatacept is a second-generation cytotoxic T-lymphocyte-associated antigen 4 (CTLA-4) immune globulin, engineered as a fusion protein combining CTLA-4 with the Fc portion of an IgG molecule. This biological agent engages CD80 and CD86 on the surface of antigen presenting cells, thereby blocking costimulation through T cell CD28. The one-year results of a phase 2 trial in renal transplant recipients given MMF, steroids, and anti-CD25 antibody demonstrated that belatacept was as effective as cyclosporine in preventing acute rejection (Vincenti et al, 2005). If proven durable the use of a nondepleting biological agent to control rejection is a novel form of therapy that may be desirable for many patients. Belatacept is given at intervals of 2–4 week as an intravenous preparation, which may be limiting. A subcutaneous preparation of belatacept is under development.
C. BASELINE IMMUNOSUPPRESSION

Current regimens vary according to center preference, and are often subject to center experience and willingness to participate in clinical trials. Two areas of current investigative interest include CNI-sparing or avoidance (to minimize CNI nephrotoxicity) and steroid-sparing or avoidance trials (to minimize steroid side effects). A very typical regimen applicable to HLA mismatched deceased or live donor recipients would include an induction agent, either a non-depleting (basiliximab/daclizumab), or a depleting (thymoglobulin/alemtuzumab) antibody. Maintenance therapy would include an antilymphocytic agent (tacrolimus, cyclosporine, or sirolimus), an antiproliferative agent (MMF or azathioprine), and steroids. Delayed introduction of CNI drugs for 7–10 days is often selected for recipients with DGF to permit early healing of the ischemic injury, assuming an induction antibody has been administered.

D. TREATMENT OF REJECTION

Acute rejection leads to graft injury and eventual CAN if untreated. Therefore, it requires prompt and accurate diagnosis, which is best provided from a percutaneous transplant renal biopsy often done under ultrasound guidance. One of the remarkable achievements of the last 10 years has been the universal acceptance of the Banff Schema to diagnose and characterize renal allograft rejection (Racusen et al, 1999). The scoring system is semiquantitative, based on light microscopy, and describes features for acute rejection and chronic/sclerosing nephropathy as well as features attributed to both cellular and antibody-mediated mechanisms. For patients with Banff I or II acute rejections, high-dose IV steroid pulses of 5–7 mg/kg/day for 3 days will reverse about 85%. Some clinicians also prefer to add a 10–14 day recycle of oral prednisone at 2 mg/kg tapered to baseline. If rejections are unresponsive to steroids or histology confirms a component of Banff II or III vascular changes, a depleting antibody such as thymoglobulin is given at 7–8 mg/kg over a week. If repeat flow cross-matching identifies new donor-specific antibody, more extensive treatments such as plasmapheresis, blocking IV immune globulin (2 g/kg), or even anti-CD20 monoclonal antibody (Rituximab) can be used. It is not generally prudent to treat more than 2–3 acute rejections in any one recipient.

RESULTS OF KIDNEY TRANSPLANTATION

There have been dramatic improvements in short-term kidney transplant outcomes since the inception of clinical practice 4 decades ago. For recipients of LD kidneys 1-year patient and graft survival has increased to about 97.6% and 95.1%; and for DD recipients 94.5% and 89% (Figure 35–1). The major reasons for this improvement are a reduction of acute rejection episodes (better immunosuppression and cross-matching techniques) with fewer complications from its treatment; and better prophylaxis and treatment of the common posttransplant infections. However, long-term graft loss beyond 5–10 years has not changed much, with stagnant survival half-lives of 7–8 years for DD and 10–11 years for LD kidneys. Factors that are statistically associated with graft failure are listed in Table 35–5. Ultimately, these factors lead to a multifaceted process of graft scarring (Figure 35–4) resulting in decline of function termed chronic allograft nephropathy (CAN), which is the major reason for late graft loss. The etiologies of CAN include processes that are immune related as well as those associated with nonspecific renal injury (Colvin, 2003). The second leading cause of late graft loss is death with a functioning graft, primarily due to the consequences of atherosclerotic cardiovascular disease; less so infections and cancers. Some risk factors for CAN and cardiovascular disease overlap (hypertension, hyperlipidemia, smoking, diabetes, etc). Graft loss secondary to patient noncompliance with medications has been estimated at 5–10%.

Complications of Kidney Transplantation

A. SURGICAL

The majority of significant surgical problems posttransplant are either vascular or urologic. They include renal artery thrombosis, disruption, stenosis, or mycotic aneurysm; renal vein thrombosis or disruption; urinary fistula or ureteral stenosis; lymphocele or hematoma; scrotal hydrocele or abscess; wound abscess, dehiscence, or hernia (Flechner and Novick, 2002). Prevention is the best way to avoid these problems using meticulous surgical and antiseptic techniques, including the routine use of prophylactic broad-spectrum antibiotics.

1. Vascular problems—In the early posttransplant period, vascular problems may prevent a new kidney from ever functioning, and questions raised from imaging studies often require surgical reexploration. Anastomotic bleeding requires immediate repair; twisting or compression of the vessels may require reanastomosis, while complete thrombosis necessitates nephrectomy. Early large hematomas should be surgically drained and hemostasis obtained. Significant transplant renal artery stenosis can occur from
3. Wound problems—Wound complications are poorly controlled hypertension, renal dysfunction (especially after ACE inhibitors or beta-blockers), or a new pelvic aneurysm or unusual lesions. Percutaneous transluminal angioplasty is the treatment of choice and restores kidney perfusion in 60–90% of cases. The risk of restenosis can be minimized with an internal stent (Bruno et al, 2004). Pseudoaneurysms of the renal or iliac artery and a-v fistula after biopsy are often amenable to embolization or endovascular stenting. Large >5 cm or mycotic aneurysms, perfusion in 60–90% of cases. The risk of restenosis can be minimized with an internal stent (Bruno et al, 2004). Suprafascial dehiscence or infection can resolve upon which pelvic structures are being compressed. Simple aspiration tends to recur; definitive treatments include prolonged tube drainage, sclerotherapy (Povidine iodine, fibrin glue, tetracycline, etc.), or marsupialization and drainage into the peritoneal cavity via laparoscopy or open surgery (Karaalltincaba, 2005; Khauli et al, 1993). Clinical presentation may include abdominal swelling, ipsilateral leg edema, renal dysfunction, or lower urinary voiding symptoms depending upon which pelvic structures are being compressed. Simple aspiration tends to recur; definitive treatments include prolonged tube drainage, sclerotherapy (Povidine iodine, fibrin glue, tetracycline, etc.), or marsupialization and drainage into the peritoneal cavity via laparoscopy or open surgery (Karaalltincaba, 2005; Khauli et al, 1993).

2. Urologic problems—Urologic complications are reported in 2–10% of kidney transplants (Streeter et al, 2002), and usually do not result in graft loss if promptly treated (van Rijen et al, 2001). Recent meta-analysis has confirmed that the routine placement of an indwelling ureteral stent will aid healing and reduce early ureteral fistula or obstruction (Wilson et al, 2005). It is advisable to leave a Foley catheter for 10–14 days for thin-walled, poorly vascularized, or small defunctionalized bladders. Ureteral fistulas and stenoses are usually a consequence of ischemia to the distal ureter from surgical dissection, overzealous electrocautery, or immune injury. Recently cases of CMV and BK virus infection had been attributed to ureteral stenosis (Mylonakis et al 2001, Fusaro et al, 2003). For large fistulas rapid surgical repair and drainage is advised, either by reimplantation to the bladder, or native uretero-ureterostomy or uretero-pyelostomy. Small fistulas are occasionally amenable to long-term stenting with or without a proximal diverting nephrostomy, or bladder catheter. Ureteral stenoses are often amenable to balloon dilation and stenting, but if recurrent require open repair. Urinary retention is more common in recent years as older males with prostatism are transplanted. It is advisable to wait a few months if prostatectomy is needed to ensure healing of the graft. Hydroceles, usually ipsilateral to the transplant and a consequence of spermatic cord transection, may cause discomfort or may enlarge. They are best repaired by hydrocelectomy, although successful aspiration and sclerotherapy has been reported.

3. Wound problems—Wound complications are reported in 5–20% of transplants, and are best prevented since they can cause significant morbidity and take many months to resolve. Since immunosuppression delays wound healing, especially sirolimus and MMF, the use of nonabsorbable sutures in the fascia and more conservative surgical technique in the obese are warranted (Humar et al, 2001; Flechner et al, 2003). A closed suction pelvic drain is also helpful immediately posttransplant. Early fascial defects or late incisional hernias require operative repair, synthetic mesh or AlloDerm may be required (Buinewicz and Rosen, 2004). Suprafascial dehiscence or infection can resolve slowly by secondary intention, which may be hastened by the use of vacuum-assisted closure (Argenta et al, 2006). Lymphocele formation in the retroperitoneum can develop from disruption of small lymphatic channels in the pelvis or around the kidney. The reported incidence of symptomatic lymphoceles ranges from 6% to 18%, and is influenced by obesity, immunosuppression (mTor inhibitors, steroids), and treatment of rejection (Goel et al, 2004). Most are asymptomatic, and resolve spontaneously over several months (Khauli et al, 1993). Clinical presentation may include abdominal swelling, ipsilateral leg edema, renal dysfunction, or lower urinary voiding symptoms depending upon which pelvic structures are being compressed. Simple aspiration tends to recur; definitive treatments include prolonged tube drainage, sclerotherapy (Povidine iodine, fibrin glue, tetracycline, etc.), or marsupialization and drainage into the peritoneal cavity via laparoscopy or open surgery (Karaalltincaba, 2005; Khauli et al, 1992).

B. MEDICAL COMPLICATIONS

1. Bacterial infections—Renal failure and immunosuppression make recipients more susceptible to infections after the transplant that includes bacterial, viral, fungal, and opportunistic pathogens. It is not surprising that such infections occur more often during the first 6 months when doses of immunosuppression are greatest. It is therefore common practice to prophylax recipients against those infective agents that occur with the greatest frequency. Bacterial urinary tract infections are the most common, and are controlled by the use of daily prophylaxis with oral trimethoprim/sulfa for the first year. This antibiotic is particularly useful since it also provides excellent prophylaxis of Pneumocystis carinii pneumonia, an opportunistic infection that is usually restricted to transplant patients, or others immunocompromised by HIV-AIDS, cancer chemotherapy, etc. Breakthrough infections and transplant pyelonephritis need further workup to identity, obstruction, reflex, foreign body, or stones.

2. Viral infections—One of the most significant advances in transplant practice in the last decades has been the control of viral infections, in particular the Herpes viruses (CMV, EBV, VCZ, and HSV), which caused major morbidity and even mortality in past years. These DNA viruses are characterized by transmission from donor to host resulting in primary infections, as well reactivation of latent virus in the host (Rubin, 2001). Therefore, recipients that have had no prior exposure (serologically negative at transplant) are at the greatest risk for infections. CMV is the most frequently encountered pathogen (10–50% of recipients), and Donor and Recipient serology (anti-CMV IgG) define risk of infection (D+R– > D+R+ > D–R+ > D–R–) and treatment strategies (Flechner et al, 1998). The virus can cause an asymptomatic infection (viral DNA copies in the blood); CMV syndrome with fever and leukopenia; and tissue-invasive disease with the liver, lung, GI tract-colon, and
retina often infected. The introduction of the potent nucleoside inhibitors acyclovir, ganciclovir, and valganciclovir has largely controlled these infections. Those who receive organs from CMV-positive donors or have had prior exposure are routinely given 3 months of prophylaxis with oral acyclovir or valganciclovir. Some prefer the use of preemptive therapy, awaiting detection by screening for virus (Khoury et al, 2006). The use of IV ganciclovir is often coadministered with anti-T-cell antibodies for patients at risk. The BK virus, one of the Polyoma virus family, has been encountered as an infectious agent with increasing frequency in kidney recipients. It is often transferred with the donor kidney, shed in the urine, and can cause inflammation and stricture in the ureter. When advanced it can cause polyoma virus associated nephropathy (PVAN), which results in cellular infiltrates and graft damage (Hirsch et al, 2005). The treatment is immunosuppressive drug reduction, and possibly the use of cidofovir or leflunomide, which have some antiviral activity.

3. Fungal infections—Candida urinary infections or esophagitis occur with some frequency, especially in diabetics. The use of oral fluconazole or Mycelex troche provides prophylaxis the first few months. Systemic fungal infections are uncommon, but sporadic cases of aspergillosis, cryptococcosis, histoplasmosis, mucormycosis, etc. are reported. Invasive fungal infections usually require treatment with Amphotericin B, or its liposomal formulation.

4. Posttransplant diabetes—New onset diabetes after renal transplantation is a growing problem (10–20% of adults) that mimics the features of diabetes type 2. It is a result of both impaired insulin production as well as peripheral insulin resistance, and includes patients that have hyperglycemia responsive to oral agents as well as those that require exogenous insulin. It can be diagnosed up to several years after transplant and is attributed to the use of CNI drugs (tacrolimus > cyclosporine) as well as glucocorticoids. Family history, old age, weight gain, hyperlipidemia, sedentary lifestyle, and viral infections are contributing factors (Duclos et al, 2006).

5. Posttransplant cancer—Immunosuppression impairs immune surveillance, and not surprisingly is associated with an increased incidence of de novo cancers. Kasiske et al (2004) examined malignancy rates among first-time recipients of deceased or LD kidney transplantations in 1995–2001 (n = 35,765) using Medicare billing claims. They found that compared to the general population, a 20-fold increase for non-Hodgkin’s lymphomas (including PTLD), nonmelanoma skin cancers, and Kaposis’s sarcoma; 15-fold for kidney cancers, fivefold for melanoma, leukemia, hepatobiliary tumors, cervical and vulvovaginal tumors; threefold for testicular and bladder cancers; and twofold for most common tumors, eg, colon, lung, prostate, stomach, esophagus, pancreas, ovary, and breast. Posttransplant lymphoproliferative disorders (PTLD) comprise a spectrum of diseases characterized by lymphoid proliferation ranging from benign lymphoid hyperplasia to high-grade invasive lymphoma. Most PTLD are B-cell lymphomas arising as a result of immunosuppression and many of these are associated with EBV infections. PTLD is reported to occur in up to 3% of adults and up to 10% of children after kidney or liver transplantation (Opelz et al, 2003). Recently, registry data has emerged that identify the use of a depleting anti-T-cell antibody for induction therapy as a significant risk factor for PTLD (Opelz et al, 2006). Since the rates for most malignancies remain higher after kidney transplantation compared with the general population, cancer should continue to be a major focus of prevention.

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Disorders of the Ureter & Ureteropelvic Junction

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The ureter is a complex functional conduit carrying urine from the kidneys to the bladder. Any pathologic process that interferes with this activity can cause renal abnormalities, the most common sequels being hydronephrosis (see Chapter 11) and infection. Disorders of the ureter can be classified as congenital or acquired.

■ CONGENITAL ANOMALIES OF THE URETER

Congenital ureteral malformations are common and range from complete absence to duplication of the ureter. They may cause severe obstruction requiring urgent attention, or they may be asymptomatic and of no clinical significance. The nomenclature can be confusing and has been standardized to prevent ambiguity (Glassberg et al, 1984).

URETERAL ATRESIA

The ureter may be absent entirely, or it may end blindly after extending only part of the way to the flank. Either anomaly is caused during embryologic development, either by failure of the ureteral bud to form from the mesonephric duct or by an arrest in its development before it comes in contact with the metanephric blastema. The genetic determinants of ureteral bud development and the causes of bud abnormalities are being elucidated and so far the PAX-2 and RET genes have been shown to play an important role (Brophy et al, 2001; Tang et al, 2002). In any event, the end result of an atretic ureteral bud is an absent or multicystic, dysplastic kidney. The multicystic kidney is usually unilateral and asymptomatic and of no clinical significance. In rare cases it can be associated with hypertension (Javadpour et al, 1970), infection (Yoshida and Sakamoto, 1986), or tumor. Contralateral vesicoureteral reflux is common, and many clinicians recommend a voiding cystourethrogram as part of the initial workup (Selzman and Elder, 1995). There is a natural tendency of these kidneys to involute (Rottenberg, Gordon, and DeBruyn, 1997); hence, most clinicians feel observation is the best treatment. A few recommend nephrectomy owing to the small risk of neoplasia and the relatively small morbidity (Homsy et al, 1997). However, the preponderance of evidence now suggests that no treatment and indeed no follow-up is needed from a urological standpoint (Onal and Kogan, 2006).

DUPLICATION OF THE URETER

Complete or incomplete duplication of the ureter is one of the most common congenital malformations of the urinary tract. Nation (1944) found some form of duplication of the ureter in 0.9% of a series of autopsies. The condition occurs more frequently in females than in males and is often bilateral. The mode of inheritance is autosomal dominant, although the gene is of incomplete penetrance (Atwell et al, 1974).

Incomplete (Y) type of duplication is caused by branching of the ureteral bud before it reaches the metanephric blastema. In most cases, this anomaly is associated with no clinical abnormality. However, disorders of peristalsis may occur near the point of union (Figure 36–1) (O’Reilly et al, 1984).

In complete duplication of the ureter, the presence of 2 ureteral buds leads to the formation of 2 totally separate ureters and 2 separate renal pelves. Because the ureter to the upper segment arises from a cephalad position on the mesonephric duct, it remains attached to the mesonephric duct longer and consequently migrates farther, ending medial and inferior to the ureter draining the lower segment (Weigert-Meyer law). Thus, the ureter draining the upper segment may migrate too far caudally and become ectopic and obstructed, whereas the ureter draining the lower segment may end laterally and have a short intravesical tunnel that leads to vesicoureteral reflux (Figure 36–2) (Tanagho, 1976).

Although many patients with duplication of the ureter are asymptomatic, a common presentation is persistent or recurrent infections. In females, the ureter to the upper pole may be ectopic, with an opening distal to the external sphincter or even outside the urinary tract. Such
patients have classic symptoms: incontinence characterized by constant dribbling, and at the same time, a normal pattern of voiding. In males, because the mesonephric duct becomes the vas and seminal vesicles, the ectopic ureter is always proximal to the external sphincter, and associated incontinence does not occur. In recent years, prenatal ultrasonography has led to the diagnosis in many asymptomatic neonates.

Excretory urography and voiding cystourethrography have been the classic studies for detecting duplication of the ureter. The excretory urogram shows the duplication in most cases. Occasionally, one segment of the kidney functions so poorly that it is not visualized. In such cases, the diagnosis can be inferred from the displacement of the visualized calyces or ureter or from the discrepancy between the amount of renal parenchyma and the relatively small number of visualized calyces. The voiding cystourethrogram discloses vesicoureteral reflux and may demonstrate the presence of a ureterocele. At the present time, the excretory urogram has been supplanted by sonography, which usually can visualize a hydronephrotic upper pole and a dilated distal ureter and can readily evaluate parenchymal thickness and the presence of bladder anomalies. Renal scanning (especially with $^{99m}$Tc-dimercaptosuccinic acid) is helpful for estimating the degree of renal function in each renal segment (Carter, Malone, and Lewington, 1998) (Figure 36–3).

The treatment of reflux alone should not be influenced by the presence of ureteral duplication (Lee et al, 1991). Lower grades of reflux are generally treated medically and higher grades of reflux surgically. Because of anatomic variations, many surgical options are available (Decter, 1997). If upper-pole obstruction or ectopy is present, surgery is almost always required. Numerous operative approaches have been recommended (Belman, Filmer, and King, 1974). If renal function in one segment is very poor, heminephrectomy is the most appropriate procedure (Barrett, Malek, and Kelalis, 1975). In an effort to preserve renal parenchyma, treatment by pyeloureterostomy, ureteroureterostomy, or ureteral reimplantation are all appropriate (Amar, 1970; Amar, 1978; Bieri et al, 1998).

URETEROCELE

A ureterocele is a saculation of the terminal portion of the ureter (Figure 36–4). It may be either intravesical or ectopic; in the latter case, some portion is located at the bladder neck or in the urethra. Intravesical ureteroceles are associated most often with single ureters, whereas ectopic ureteroceles nearly always involve the upper pole of duplicated ureters. Ectopic ureteroceles are four times more common than those that are intravesical (Snyder and Johnston, 1978).
Figure 36–3. Duplication of the ureter and the ureterocele. **Upper left:** Excretory urogram shows duplication of the right kidney (arrowheads on upper pole) and visualization of only the lower pole (arrows on lower pole) of the left kidney (white arrow). There is a filling defect on the left side of the bladder. **Upper right:** Cystogram confirms the filling defect. There is no reflux. **Lower left:** Renal scan with $^{99m}$Tc-dimercaptosuccinic acid shows some functioning parenchyma in upper pole to left kidney. **Lower right:** After excision of ureterocele and reimplantation of both ureters on left, repeat excretory urogram shows improved excretion of contrast medium from upper pole of left kidney.
Ureterocele occurs seven times more often in girls than in boys, and about 10% of cases are bilateral. Mild forms of ureterocele are found occasionally in adults examined for unrelated reasons.

Ureterocele has been attributed to delayed or incomplete canalization of the ureteral bud leading to an early prenatal obstruction and expansion of the ureteral bud prior to its absorption into the urogenital sinus (Tanagho, 1976). The cystic dilation forms between the superficial and deep muscle layers of the trigone. Large ureteroceles may displace the other orifices, interfere with the muscular backing of the bladder, or even obstruct the bladder outlet. There is nearly always significant hydronephrosis, and a dysplastic segment of the upper pole of the kidney may be found in association with a ureterocele.

Clinical findings vary considerably. Patients commonly present with infection, but bladder outlet obstruction or incontinence may be the initial complaint. Occasionally, a ureterocele may prolapse through the female urethra (Ahmed, 1984). Calculi can develop secondary to urinary stasis and are often seen in the distal ureter. Currently, many cases are diagnosed by antenatal maternal ultrasound (Gloor, Ogburn, and Matsumoto, 1996). Although excretory urography (Figures 36–3 and 36–5) is usually diagnostic, sonography has replaced the excretory urogram in most centers. Voiding cystourethrography should always be part of the workup (Bauer and Retik, 1978). It may demonstrate reflux into the lower pole or contralateral ureter and occasionally shows eversion of the ureterocele during urination, in which case the ureterocele has the appearance of a diverticulum. Renal scanning is helpful for estimating renal function (Geringer et al, 1983).

Treatment must be individualized. Transurethral incision was used previously only in very ill children with pyohydronephrosis; however, it has been recognized as the definitive procedure in many instances, particularly in patients with intravesical ureteroceles (Blyth et al, 1993; Pfister et al, 1998) and especially in neonates (Coplen, 2001; Upadhyay et al, 2002). When an open operation is needed, the procedure must be chosen on the basis of the anatomic location of the ureteral meatus, the position of the ureterocele, and the degree of hydroureronephrosis and impairment of renal function. In general, choices range from heminephrectomy and ureterectomy (Husmann et al, 1995) to excision of the ureterocele, vesical reconstruction, and ureteral reimplantation. Often, a second procedure is necessary (Caldamone, Snyder, and Duckett, 1984).

### Ectopic Ureteral Orifice

Although an ectopic ureteral orifice most commonly occurs in association with duplication of the ureter (see preceding sections), single ectopic ureters do occur (Gotoh et al, 1983). They are caused by a delay or failure of separation of the ureteral bud from the mesonephric duct during embryologic development. Again, the genetic determin-
nants of these ureteral bud abnormalities are currently being determined, but at least the PAX-2 and RET genes are involved (Brophy et al, 2001; Tang et al, 2002). In anatomic terms, the primary anomaly may be an abnormally located ureteral bud; that explains the high incidence of dysplastic kidneys associated with single ectopic ureters. The clinical picture varies according to the sex of the patient and the position of the ureteral opening. Boys are seen because of urinary tract infection or epididymitis. In these cases, the ureter may drain directly into the vas deferens or seminal vesicle (Umeyama et al, 1985). In girls, the ureteral orifice may be in the urethra, vagina, or perineum. Although infection may be present, incontinence is the rule. Continual dribbling despite normal voiding is pathognomonic. Urgency and urge incontinence may confound the diagnosis (Johnson and Perlmutter, 1980).

Sonography and voiding cystourethrography help delineate the problem. However, because an ectopic kidney may be both tiny and in an abnormal location, it may be difficult to find by ultrasound; and magnetic resonance imaging, cystoscopy, or laparoscopy may be necessary to confirm the diagnosis (Borer et al, 1998). During cystoscopy, a hemi-trigone may be seen and the ectopic orifice may be visualized directly or demonstrated by retrograde catheterization (Figures 36–6 and 36–7). Renal scanning is also helpful in estimating relative renal function. As in ureteroceles and duplication of the ureter, the clinical picture and the degree of renal function dictate the therapeutic approach.

ABNORMALITIES OF URETERAL POSITION

Retrocaval ureter (also called circumcaval ureter and postcaval ureter) is a rare condition in which an embryologically normal ureter becomes entrapped behind the vena cava because of abnormal persistence of the right subcardinal (as opposed to the supracardinal) vein. This forces the right ureter to encircle the vena cava from behind. The ureter descends normally to approximately the level of L3, where it curves back upward in the shape of a reverse J to pass behind and around the vena cava. Obstruction generally results.

Traditionally, the diagnosis of retrocaval ureter was made by excretory urography. However, since sonography is now usually the first test performed, the radiologist must be suspicious of the anomaly based on a dilated proximal (but not distal) ureter. Currently, magnetic resonance imaging
imaging is the best single study to delineate the anatomy clearly and noninvasively. Surgical repair for retrocaval ureter, when indicated, consists of dividing the ureter (preferably across the dilated portion), bringing the distal ureter from behind the vena cava, and reanastomosing it to the proximal end. The procedure has been performed laparoscopically to reduce morbidity (Polascik and Chen, 1998).

**OBSTRUCTION OF THE URETEROPELVIC JUNCTION**

In children, primary obstruction of the ureter usually occurs at the ureteropelvic junction or the ureterovesical junction (Figure 36–8). Obstruction of the ureteropelvic junction is probably the most common congenital abnor-
Figure 36–7. Ectopic ureter. Left: Cystoscopy in a 6-year-old girl with a lifelong history of urinary incontinence revealed 2 ureteral orifices on the right and one on the left; these were catheterized and urograms obtained. Right: Same patient. An ectopic ureteral orifice near the urethral meatus was catheterized. Retrograde urogram demonstrates second hydronephrotic renal pelvis on left. Resection of upper pole and ureter cured incontinence.

Figure 36–8. Congenital ureteral obstruction. Left: Right ureteropelvic junction obstruction with hydronephrosis. Right: Left ureterovesical junction obstruction (obstructed megaureter) with hydroureteronephrosis.
mality of the ureter. It is seen more often in boys than in
girls (5:2 ratio) and, in unilateral cases, more often on the
left than on the right side (5:2 ratio). Bilateral obstruction
occurs in 10–15% of cases and is especially common in
infants (Johnston et al, 1977). The abnormality may occur
in several members of the same family, but it shows no
clear genetic pattern.

The exact cause of obstruction of the ureteropelvic
junction often is not clear. Ureteral polyps and valves have
been reported but are very rare (Punjabi, 1983; Sant, Bar-
ballas, and Klauber, 1985). There is almost always an
angulation and kink at the junction of the dilated renal
pelvis and ureter. This by itself can cause obstruction, but
it is unclear whether this is primary or merely secondary to
another obstructive lesion. True stenosis is found rarely; how-
ever, a thin-walled, hypoplastic proximal ureter is
observed frequently. Characteristic histologic and ultra-
structural changes are observed in this area and could
account for abnormal peristalsis through the ureteropelvic
junction and consequent interference with pelvic emptying
(Hanna et al, 1976). Two other findings sometimes seen at
operation are a high origin of the ureter from the renal pel-
vis and an abnormal relationship of the proximal ureter to
a lower-pole renal artery. It is debatable whether these
findings are the result or the cause of pelvic dilatation, but
Stephens (1982) has suggested that abnormal rotation of
the renal pelvis allows the ureter to become entrapped in
the blood vessels of the lower pole of the kidney, ultimately
leading to obstruction. Using careful studies at the time of
operation, it is possible to define whether the principal
lesion is intrinsic or extrinsic (Koff et al, 1986; Johnston,
1969).

Clinical findings vary depending on the patient’s age at
diagnosis. Recent improvements in prenatal ultrasonogra-
phy now allow most cases to be diagnosed in utero (Man-
dell et al, 1991). Later, pain and vomiting are the most
common symptoms; however, hematuria and urinary
infection also may be seen. A few patients have complica-
tions such as calculi (Figure 36–9), trauma to the enlarged
kidney, or (rarely) hypertension.

The diagnosis is made most often by sonography. In
equivocal cases, diuretic renography or (rarely) antegrade
urography with pressure-flow studies is helpful (Thrall,
consider a voiding cystourethrogram a routine part of the
preoperative workup, since radiographic findings in ves-
coureteral reflux may be similar to those in ureteropelvic
junction obstruction. This fact is especially relevant when
the ureter is well seen or dilated (or both) below the ure-
teropelvic junction (Maizels, Smith, and Firlit, 1984).

Symptomatic obstruction of the ureteropelvic junction
should be treated surgically. Because most cases are now
detected by hydronephrosis on prenatal ultrasonography
and the infants are asymptomatic, it becomes important to
assess the significance of the hydronephrosis. On the one
hand, early surgery may prevent future urinary tract infec-
tions, stones, or other complications; on the other hand,
many of the patients could live their whole lives without
experiencing a consequence of the hydronephrosis. This
remains an area of considerable controversy. Early surgery
is recommended for patients who have kidneys with
diminished function, massive hydronephrosis, infection, or
stones. Nonoperative surveillance with good follow-up is
thought to be safe (Onen et al, 2002), although about
25% of patients will ultimately require an operative repair
for pain, urinary infection, or reduced renal function on
repeat nuclear scan (Palmer et al, 1998). This subject
remains particularly controversial (Peters, 2002).

Because of anatomic variations, no single procedure is
sufficient for all situations (Smart 1979). Regardless of the
technique used, all successful repairs have in common the
creation of a dependent and funnel-shaped ureteropelvic
junction of adequate caliber. Although preservation of the
intact ureteropelvic junction is feasible in some circum-
stances (Perlberg and Pfau, 1984), when the obstruction
appears to be caused by a dyskinetic segment of proximal
ureter, the most popular operation is a dismembered pye-
loureteroplasty (Anderson, 1963). Dismembered pyel-
oureteroplasty is also favored when the proximal ureter is
hooked over a lower-pole blood vessel. When there is a
dilated extrarenal renal pelvis, dismembered pylouretero-
plasty can be combined with a Foley Y-V plasty to create a
more funnel-shaped ureteropelvic junction (Foley, 1937).
Pelvic flap procedures (Culp and DeWeerd, 1951; Scar-
dino and Prince, 1953) are suited ideally to cases in which
the ureteropelvic junction has remained in a dependent
dissection despite significant pelvic dilatation. They also
have the advantage of interfering less with the ureteral
blood supply; this is particularly relevant when distal ure-
teral surgery (eg, ureteral reimplantation) is contemplated
in the future. In most centers, the dismembered pyel-
oureteroplasty is the mainstay of repairs.

Both the Y-V plasty and the flap techniques are useful
in managing ureteropelvic junction obstructions in horse-
shoe or pelvic kidneys, in which the anatomy may prevent
creation of a dependent ureteropelvic junction if a dis-
membered technique is attempted. The use of stenting
catheters and proximal diversion at the time of pyeloplasty
has been the subject of debate, and the issue has not been
resolved. Excellent results have been reported both with
and without stents and diversions (Smith et al, 2002).

The prognosis is generally good. In several large series,
the reported reoperation rate has been only 2–4%, but the
postoperative radiographic appearance of the area may be
disappointing. There can be marked improvement when a
large extrarenal renal pelvis has prevented massive calyceal
distortion; however, in most cases considerable deformity
persists despite adequate drainage of the kidney. Further-
more, it is usually many years before the radiographic
appearance improves (Amling et al, 1996).
Figure 36–9. Ureteropelvic junction obstruction with calculi. Upper left: Plain film of abdomen shows radiopacities in region of right kidney. Bottom left: Early film from excretory urogram demonstrates dilation of calyces on right and layering of calculi in large right renal pelvis. Bottom Right: Delayed film from excretory urogram shows a typical right ureteropelvic junction obstruction.
The recent explosion in the field of endourology as a subspecialty of urology has encouraged use of percutaneous techniques for the repair of ureteropelvic junction obstruction in selected patients (Ramsay et al, 1984; Badlani, Eshghi, and Smith, 1986; Van Cangh et al, 1989). The technique is similar to that reported by Davis (1943) but is done entirely endoscopically. The technique may be applied antegrade, via a nephrostomy tract, or retrograde, using either a ureteroscope (for direct vision) or an Acusize (Applied Urology, Laguna Beach, CA) balloon catheter with fluoroscopic visualization. In adults the procedure is clearly an option, with an anticipated success rate of 80–85% and a marked reduction in morbidity (Aslan and Preminger, 1998). In children, retrograde endopyelotomy also has an 85% success rate, but this is still considerably less than 98% and the benefit in terms of reduced morbidity is less significant (Bogaert et al, 1996). An intriguing new option that is becoming commonplace in some centers is laparoscopic pyeloplasty (Moore et al, 1997; Yeung et al, 2001).

**OBSTRUCTED MEGAURETER**

Obstruction at the ureterovesical junction is four times more common in boys than in girls. It may be bilateral and is usually asymmetric. The left ureter is slightly more often involved than the right.

The embryogenesis of the lesion is uncertain. It is clear that in most cases there is no stricture at the ureterovesical junction. At operation, a retrograde catheter or probe can usually be passed through the area of obstruction. Close observation either at operation or by fluoroscopy reveals a failure of the distal ureter to transmit the normal peristaltic wave, resulting in a functional obstruction. Moreover, on fluoroscopy, retrograde peristalsis is seen. This transmits abnormal pressures up to the kidney, resulting in calyceal dilation out of proportion with the renal pelvic dilation. Histologic findings include an excess of circular muscle fibers and collagen in the distal ureter that may account for the problem (Tanagho, Smith, and Guthrie, 1970). Ultrastructural studies show that this obstruction is similar in appearance to obstruction of the ureteropelvic junction.

Currently, most cases are discovered on prenatal sonography; however, a few come to light because of hematuria or infection. Sonography usually shows the pathognomonic configuration of a dilated distal ureter, a less dilated proximal ureter, a relatively normal-appearing renal pelvis, and calyces blunted out of proportion to the renal pelvis (Figure 36–10).

It was assumed previously that surgery was indicated in most cases. Ureteral reimplantation with excision of the distal ureter is curative. Because of the excessive dilation of the ureter, ureteral tapering or folding may be necessary (Hendren, 1969; Hanna, 1982; Ehrlich, 1985). Because the ureteral muscle is generally healthy, these cases have an excellent prognosis (Peters et al, 1989). However, in recent years it has become obvious that at least 50% of cases will undergo spontaneous resolution. A period of observation is nearly always appropriate when the diagnosis is made in an asymptomatic patient (Baskin et al, 1994). Because of the high risk of infection, 1–2 years of prophylactic antibiotics are recommended in neonates.

**UPPER URINARY TRACT DILATATION WITHOUT OBSTRUCTION**

It should not be assumed that every dilated upper urinary tract is obstructed. A voiding cystourethrogram is an essential part of the evaluation, not only to rule out reflux but also to ensure that no abnormality of the lower urinary tract is responsible for the upper urinary tract dilatation. Other cases in which diagnosis may be difficult include residual dilatation in a previously obstructed system, dilatation associated with bacterial infection (presumably related to a direct effect of endotoxin on the ureteral musculature), neonatal hydronephrosis (Homsy, Williot, and Danaïs, 1986), and prolonged polyuria in patients with diabetes insipidus.
In such cases, the usual investigations may not provide sufficient information. A radionuclide diuretic renogram is especially helpful in distinguishing nonobstructive from obstructive dilation and in determining whether renal functional injury has occurred (Figure 36–11) (Thrall, Koff, and Keyes, 1981). However, the procedure must be performed carefully, as technical problems may confuse the results (Nguyen et al, 1997; Gungor et al, 2002). Use of percutaneous renal puncture is occasionally beneficial; in the dilated system, it carries minimal risk, making antegrade urography and pressure-flow studies feasible in selected cases. Measurement of the renal pelvic pressure during infusion of saline into the renal pelvis at high rates (10 mL/min) (the Whitaker test) may help differentiate nonobstructive from obstructive dilation (Wolk and Whitaker, 1982). Unfortunately, there is no true “gold standard,” and these studies do not always agree; clinical judgment is the final arbiter (Lupton et al, 1985).

ACQUIRED DISEASES OF THE URETER

Nearly all acquired diseases of the ureter are obstructive in nature. Although they are seen frequently, their actual incidence is unknown. Their clinical manifestations, effects on the kidney, complications, and treatment are similar to those described previously. The lesions can be broadly categorized as either intrinsic or extrinsic.

Intrinsic Ureteral Obstruction

The most common causes of intrinsic ureteral obstruction are as follows:

1. Ureteral stones (see Chapter 16)
2. Transitional cell tumors of the ureter (see Chapter 20)
3. Chronic inflammatory changes of the ureteral wall (usually due to tuberculosis or schistosomiasis) leading to contracture or insufficient peristalsis (see Chapter 14 and Figures 14–2 and 14–4)

Extrinsic Ureteral Obstruction

The most frequent causes of extrinsic ureteral obstruction are

1. Severe constipation, sometimes with bladder obstruction, seen primarily in children but in adult women as well.
2. Secondary obstruction due to kinks or fibrosis around redundant ureters. The primary process is either distal obstruction or massive reflux.

Figure 36–11. Upper urinary tract dilatation. Left: Three months after resection of posterior urethral valves, hydronephrosis in the right kidney has completely resolved. The left collecting system remains dilated. (Dashed lines outline kidneys.) Right: Radionuclide diuretic renography was performed to determine if there was secondary ureteropelvic or ureterovesical obstruction. Renogram demonstrates clear-cut “washout” of radionuclide following injection of furosemide (arrow). There is no significant obstruction.
3. Benign gynecologic disorders such as endometriosis or right ovarian vein syndrome (Gourdie and Rogers, 1986).
4. Local neoplastic infiltration associated with carcinoma of the cervix, bladder, or prostate (Richie, Withers, and Ehrlich, 1979).
5. Pelvic lymphadenopathy associated with metastatic tumors.
6. Iatrogenic ureteral injuries, primarily after extensive pelvic surgery (Figure 36–12) and also after extensive radiotherapy.
7. Retroperitoneal fibrosis.

**RETROPERITONEAL FIBROSIS**
(RETROPERITONEAL FASCIITIS,
CHRONIC RETROPERITONEAL FIBROPLASIA, ORMOND DISEASE)

One or both ureters may be compressed by a chronic inflammatory process that involves the retroperitoneal tissues over the lower lumbar vertebrae. There are numerous causes of retroperitoneal fibrosis. Malignant diseases (most commonly Hodgkin’s disease, carcinoma of the breast, and carcinoma of the colon) should always be suspected and ruled out. Some medications have been implicated, most notably methysergide (Sansert), an ergot derivative used to treat migraine headaches. Rarely, membranous glomerulonephritis (Shirota et al, 2002), inflammatory bowel disease (Siminovitch and Fazio, 1980) or an aortic aneurysm (Brock and Soloway, 1980; Peters and Cowie, 1978) is responsible. The remainder of cases are idiopathic, a condition sometimes referred to as Ormond disease.

The symptoms are nonspecific and include low back pain, malaise, anorexia, weight loss, and, in severe cases, uremia. Infection is uncommon. The diagnosis is usually made by excretory urography (Figure 36–13). There is medial deviation of the ureters with proximal dilation. A long segment of ureter is usually involved, and in some cases there is a pipestem appearance caused by aperistalsis related to the fibrosis. A retrograde ureterogram is necessary when renal function is poor and, in any case, helps to delineate the length of the affected segment of ureter. Ultrasonography is useful, not only for diagnosis but also for monitoring the response to therapy. Computed tomography scanning or magnetic resonance imaging is essential for evaluating the retroperitoneum itself, as well as for imaging the ureters (Hricak, Higgins, and Williams, 1983). Recently, positron emission tomography scanning has been advocated (Cheung et al, 2002).
Spontaneous regression has been reported (Kume and Kitamura, 2001); however, treatment is usually surgical. A course of corticosteroids may be tried first and in an occasional series, remarkable success has been reported (Kardar et al, 2002). When the response to corticosteroids is poor or the obstruction is severe, the ureter must be dissected surgically from the fibrous plaque. After it is freed, it should either be placed intraperitoneally or wrapped in omentum in an attempt to prevent recurrence (Lepor and Walsh, 1979). Rarely, autotransplantation is necessary (Deane, Gingell, and Pentlow, 1983). Numerous biopsies of the fibrous tissue should be obtained at the time of operation to determine whether there is a malignant tumor.

**URETERAL OBSTRUCTION SECONDARY TO MALIGNANT DISEASE**

Ureteral obstruction associated with widespread malignant disease was at one time a terminal event. Because therapy for malignant diseases has improved, however, urinary diversion is indicated more frequently in such cases. Diversion usually is necessary for relatively short periods of time; either the tumor is progressive, or if therapy is effective, the obstruction resolves. Thus, the goal of treatment is to leave the urinary tract intact and effect as little morbidity as possible. This can be accomplished with indwelling stents passed either retrogradely during cystoscopy (Hepperlen, Mardis, and Kammandel, 1979) or antegrade using percutaneous techniques (Elyaderani et al, 1982).

**REFERENCES**

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General


Retroperitoneal Fibrosis


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Ureteral Obstruction Secondary to Malignant Disease


CONGENITAL ANOMALIES OF THE BLADDER

EXSTROPHY

Exstrophy of the bladder is a complete ventral defect of the urogenital sinus and the overlying skeletal system (see Chapter 2). Other congenital anomalies are frequently associated with it. The lower central abdomen is occupied by the inner surface of the posterior wall of the bladder, whose mucosal edges are fused with the skin. Urine spurts onto the abdominal wall from the ureteral orifices.

The rami of the pubic bones are widely separated. The pelvic ring thus lacks rigidity, the femurs are rotated externally, and the child “waddles like a duck.” Since the rectus muscles insert on the rami, they are widely separated from each other inferiorly. A hernia, made up of the exstrophic bladder and surrounding skin, is therefore present. Epispadias almost always accompanies it.

Many untreated exstrophic bladders reveal fibrosis, derangement of the muscularis mucosae, and chronic infection. These changes tend to defeat efforts to form a bladder of proper capacity. About 60 instances of adenocarcinoma developing in such bladders have been reported.

Renal infection is common, and hydronephrosis caused by ureterovesical obstruction may be found on urography. These films also reveal separation of the pubic bones. During the last few years, there have been encouraging reports of complete reconstruction of this defect. Earlier, urinary diversion and resection of the bladder, with later repair of the epispadiac penis, was usually accomplished. With improved techniques and early surgery before the bladder deteriorates, however, good results are being obtained with complete reconstruction. Lattimer et al (1978), pioneers in this field, followed up their 17 patients with reconstructed bladders for as long as 20 years. They reported that the quality of life of these patients was good.

Ansel (1979) performed reconstruction in 28 patients in the neonatal period in an attempt to protect the bladder from later serious changes. Half of these patients did well, and most were continent. DeMaria et al (1980) found the renal function and urine cultures of their patients to be normal. Eight of their patients had complete continence, while 12 had enuresis. Toguri et al (1978) reported that all of their 23 patients were continent.

Lima et al (1981) reconstructed the bladder with human dura mater to increase vesical capacity; they were successful in 8 cases. They perform osteotomy as part of the first stage and recommend that the surgery be performed when patients are 3–18 months old. Enterocystoplasty is currently the method of choice to augment bladder capacity and aid reservoir function. Mollard (1980) recommends the following steps for satisfactory repair of bladder exstrophy: (1) bladder closure with sacral osteotomy in order to close the pelvic ring at the pubic symphysis, plus lengthening of the penis; (2) antiureteral reflux procedure and bladder neck reconstruction; and (3) repair of the epispadiac penis. He completed 16 such 3-step procedures, with satisfactory results in 11. In 1983 and 1989, respectively, Jeffs and Gearhart reported results of staged reconstruction: 86% of patients who underwent primary repair were continent, and renal function was preserved in approximately 90% (Gearhart et al, 1993). Urethral and genital reconstructions have been equally successful. These are the best-reported results. In small-capacity bladders, augmentation cystoplasty might be needed (Oesterling and Jeffs, 1987; Gearhart and Jeffs, 1988; Gearhart, 1999). Recently, long-term follow-up in large series of patients with bladder exstrophy has demonstrated satisfactory outcomes in 70–80% with regard to continence and renal function.

When the bladder is small, fibrotic, and inelastic, functional closure becomes inadvisable, and urinary diversion with cystectomy is the treatment of choice. Some physicians perform ureterointestinostomy, while others prefer to use the colon for the diversion. A continent reservoir is a current consideration and is preferable. Spence, Hoffman, and Pate (1975) employ ureterosigmoidostomy. Turner, Ransley, and Williams (1980) noted that, although untreated newborns have normal upper urinary tracts, urinary diversion often causes hydronephrosis or pyelonephritis in these patients.
The common complication of total reconstruction is urinary incontinence, but Light and Scott (1983) reported on the implantation of an artificial sphincter in 11 patients who were still incontinent after total reconstruction. They claimed 10 perfect results. Ikeme (1981) reported on 2 patients who became pregnant after repair of bladder extrophy; one woman had 3 successful pregnancies, and the other had one. Complete primary closure appears to be the best choice for improved continence with reduced morbidity (Grady, Carr, and Mitchell, 1999).

**PERSISTENT URACHUS**

Embryologically, the allantois connects the urogenital sinus with the umbilicus. Normally, the allantois is obliterated and is represented by a fibrous cord (urachus) extending from the dome of the bladder to the navel (see Chapter 2). Urachal formation is directly related to bladder descent. Lack of descent is more commonly associated with patent urachus than with bladder outlet obstruction.

Incomplete obliteration sometimes occurs. If obliteration is complete except at the superior end, a draining umbilical sinus may be noted. If it becomes infected, the drainage will be purulent. If the inferior end remains open, it will communicate with the bladder, but this does not usually produce symptoms. Rarely, the entire tract remains patent, in which case urine drains constantly from the umbilicus. This is apt to become obvious within a few days of birth. If only the ends of the urachus seal off, a cyst of that body may form and may become quite large, presenting a low midline mass (Figure 37–1). If the cyst becomes infected, signs of general and local sepsis will develop.

Adenocarcinoma may occur in a urachal cyst, particularly at its vesical extremity, and tends to invade the tissues beneath the anterior abdominal wall. It may be seen cystoscopically. Stones may develop in a cyst of the urachus. These can be identified on a plain x-ray film.

Treatment consists of excision of the urachus, which lies on the peritoneal surface. If adenocarcinoma is present, radical resection is required. Unless other serious congenital anomalies are present, the prognosis is good. The complication of adenocarcinoma offers a poor prognosis.

**CONTRACTURE OF THE BLADDER NECK**

There is considerable debate about the incidence of congenital narrowing of the bladder neck. Some feel that its presence is a common cause of vesicoureteral reflux, vesical diverticula, a bladder of large capacity, and the syndrome of irritable bladder associated with enuresis. A few observers consider this contracture a rare phenomenon and believe that the diagnosis is purely presumptive. The diagnosis is based on endoscopic observation, which is an unreliable method. Voiding cystourethrography has been used to depict such narrowing, but interpretation of the films varies from urologist to urologist and radiologist to radiologist.

Nunn (1965) studied the intravesical and urethral pressures during voiding in patients with the signs mentioned previously and found no evidence of bladder neck obstruction. The 2 recorded pressures were essentially equal. It appears that the bladder neck would have to be extremely stenotic to truly obstruct urine flow. It is becoming increasingly clear that in young girls, the obstructive lesion is spasm of the periurethral striated muscle, which develops secondary to distal urethral stenosis (see Chapter 39).

Empirical treatment is often employed; this consists of suprapubic bladder neck revision or transurethral resection. Making the bladder neck incompetent in young boys may cause later retrograde ejaculation and, therefore, infertility. Revision of the bladder neck in females may cause urinary incontinence and is never advised. The diagnosis must therefore be made with caution.

Genuine functional bladder neck obstruction can be detected only in the presence of already high voiding pressures combined with lower resistance in the external sphincteric segment associated with a low flow rate. This condition is highly suggestive of functional bladder neck obstruction, although not 100% diagnostic.

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**Figure 37–1.** Types of persistent urachus. **Left:** Communicating urachus continuous with the bladder. This is a “pseudodiverticulum” and usually causes no symptoms. **Center:** Urachal cyst; usually causes no symptoms or signs unless it becomes larger or infected. **Right:** Patent urachus. There is constant drainage of urine from the umbilicus.
CHAPTER 37

ACQUIRED DISEASES OF THE BLADDER

INTERSTITIAL CYSTITIS (HUNNER’S ULCER, SUBMUCOUS FIBROSIS)

Interstitial cystitis is primarily a disease of middle-aged women. It is characterized by fibrosis of the vesical wall, with consequent loss of bladder capacity. Frequency, urgency, and pelvic pain with bladder distention are the principal symptoms.

PATHOGENESIS & PATHOLOGY

Infection does not appear to be the cause of fibrosis of the bladder wall, because the urine is usually normal. It has been postulated that the fibrosis is due to obstruction of the vesical lymphatics secondary to pelvic surgery or infection, but many of these patients fail to give such a history. Fibrosis may be secondary to thrombophlebitis complicating acute infections of the bladder or pelvic organs; may be the result of prolonged intrinsic arteriolar spasm secondary to vasculitis or psychogenic impulses; or could be of neuropathic origin. Endocrinologic factors are also suggested. Investigators are currently studying the role of mast cells and bladder surface glycosaminoglycans (GAGs) in the pathogenesis of interstitial cystitis. Currently, it is believed that interstitial cystitis is a neuroimmunoendocrine disorder. It might be primarily a neurogenic inflammation that leads to the release of neuropeptides that activate the differential secretion of potent mast cell mediators. It is thought that mast cells, through their vasoactive and nociceptive secretions, have a major role in the etiology of interstitial cystitis.

The primary change is fibrosis in the deeper layers of the bladder. The capacity of the organ is decreased, sometimes markedly. The mucosa is thinned, especially where mobility is greatest as the bladder fills and empties (ie, over the dome), and small ulcers or cracks in the mucous membrane may be seen in this area. In the most severe cases, the normal mechanism of the ureterovesical junctions is destroyed, leading to vesicoureteral reflux. Hydroureteronephrosis and pyelonephritis may ensue.

Microscopically, the mucosa may be thinned or even denuded. The capillaries of the tunica propria are often engorged, and signs of inflammation are apparent. The muscle is replaced by varying amounts of fibrous tissue, which is often quite avascular. The lymphatics may be engorged. Increased mast cells and lymphocytic infiltration are seen.

There has been a tendency recently to overdiagnose interstitial cystitis, particularly in patients with excessive frequency, urgency, and suprapubic or pelvic pain, even though they lack the pathologic manifestations and usually have normal or large bladder capacity. These patients have voiding dysfunction. Although we may not know the exact cause of their symptoms, these patients should not be labeled as having interstitial cystitis and should not be treated as such.

CLINICAL FINDINGS

Interstitial cystitis should be considered when a middle-aged woman with clear urine complains of severe frequency and nocturia and suprapubic pain on vesical distention. In addition, chronic pelvic pain in the absence of pelvic pathology can be suggestive.

A. Symptoms

There is a long history of slowly progressive frequency and nocturia, both of which may be severe. The history does not suggest infection (burning on urination, cloudy urine). Suprapubic pain is usually marked when the bladder is full. Pain may also be experienced in the urethra or perineum; it is relieved on voiding. Gross hematuria is occasionally noted, usually when urination has had to be postponed (ie, following vesical overdistention). The patient is tense and anxious. Whether the anxiety is secondary to the prolonged and severe symptoms or is the primary cause of the vesical changes is not clear. A history of allergy may be obtained.

B. Signs

Physical examination is usually normal. Some tenderness in the suprapubic area may be noted. There may be some tenderness in the region of the bladder when it is palpated through the vagina.

C. Laboratory Findings

If the patient has had no previous treatment (eg, instrumentation), the urine is almost always free of infection. Microscopic hematuria may be noted. Results of renal function tests are normal except in the occasional patient in whom vesical fibrosis has led to vesicoureteral reflux or obstruction.

D. X-Ray Findings

Excretory urograms are usually normal unless reflux has occurred, in which case hydroureteronephrosis is found. The accompanying cystogram reveals a bladder of small capacity; reflux into a dilated upper tract may be noted on cystography.

E. Instrumental Examination

Cystoscopy is usually diagnostic. As the bladder fills, increasing suprapubic pain is experienced. The vesical capacity may be as low as 60 mL. In a patient not previously treated (by fulguration or hydraulic overdistention),...
the bladder lining may look fairly normal. However, if a second distention is done (Messing and Stamey, 1978), punctate hemorrhagic areas may appear over the most distensible portion of the wall. With further distention, an arcuate split in the mucosa will occur and may bleed profusely. Mucosal changes are usually diffuse. Congestion, edematous reaction, and petechial hemorrhages (glomerulitis) are common findings.

**DIFFERENTIAL DIAGNOSIS**

Tuberculosis of the bladder may cause true ulceration but is most apt to involve the region of the ureteral orifice that drains the tuberculous kidney. Typical tubercles may be identified, pyuria is present, and tubercle bacilli usually can be found. Furthermore, urograms often show the typical lesion of renal tuberculosis.

Vesical ulcers due to schistosomiasis cause symptoms similar to those of interstitial cystitis. The diagnosis is suggested if the patient lives in an area in which schistosomiasis is endemic. Most patients are males. The typical ova found in the urine and the pathognomonic appearance of the bladder confirms the diagnosis. Nonspecific vesical infection seldom causes ulceration. Pus and bacteria are found in the urine. Antimicrobial treatment is effective.

**Complications**

Gradual ureteral stenosis or reflux and its sequelae (eg, hydronephrosis) may develop.

**Treatment**

**A. SPECIFIC MEASURES**

There appears to be no definitive treatment for interstitial cystitis. The therapy usually employed frequently affords partial relief, but it may be completely ineffective. Hydrational distention, with or without anesthesia, sometimes gradually improves the bladder capacity. Vesical lavage with increasing strengths of silver nitrate (1:5000–1:100) may have the same effect. Superficial (transcystoscopic) electrocau­gulation of the split mucosa will occur and may bleed profusely.

Occasionally, symptomatic relief follows the instillation of 50 mL of 50% dimethyl sulfoxide (DMSO) into the bladder every 2 weeks. It is left in for 15 minutes. Messing and Stamey (1978) claim their best results were obtained with vesical irrigations of 0.4% oxychlorosene sodium (Clorpactin WCS-90). At 10 cm of water pressure, the bladder is repeatedly filled to capacity until 1 L has been used. This must be done under anesthesia. Cystography should be done before instituting this therapy. The presence of vesicoureteral reflux has caused ureteral fibrosis.

Parsons, Schmidt, and Pollen (1983) observed the results obtained in patients who failed to respond to hydraulic distention or the instillation of DMSO. They found that the bladder mucosa needs a layer of sulfonated GAGs on its surface to protect the transitional cells from the effect of urine, and this substance was absent from the mucosa of these patients. They administered sodium pentosanpolysulfate (Elmiron) orally, in doses of either 50 mg 4 times a day or 150 mg twice daily, for 4–8 weeks. Of 24 patients, 20 noted at least 80% relief of urgency, frequency, and nocturia, and 2 noted 50–80% relief. These 22 patients continued to improve. Two patients experienced no apparent relief.

Cortisone acetate, 100 mg, or prednisone (Meticorten), 10–20 mg/day, in divided doses orally for 21 days, followed by decreasing amounts for an additional 21 days, has also been found effective. Transcystoscopic injection of the lesions with prednisone has its proponents. Antihistamines (eg, tripelennamine [Pyribenzamine], 50 mg 4 times a day) may also afford some relief. Heparin sodium (long-acting), 20,000 units intravenously daily, also blocks the action of histamine, and its use in the treatment of interstitial cystitis is encouraging. Newer treatments being tested (eg, resiniferotoxin, botulinum toxin, gene therapy, and neuromodulation) may potentially be effective.

If the bladder becomes fibrotic and the capacity small, ceco- or ileocystoplasty can be done to augment vesical capacity. Most patients are cured or greatly improved; those who are not may require urinary diversions. Denervation by presacral and sacral neurectomy and perivesical procedures (cystolysis, cystoplasty, transvaginal neurotomy) is to be condemned, as it is rarely of lasting benefit. In severe contracture, augmentation cystoplasty is indicated.

**B. GENERAL MEASURES**

General or vesical sedatives may be prescribed but seldom afford relief. If urinary infection is found (usually following instrumentation), it should be treated with appropriate antibiotics. If senile urethritis is discovered, diethylstilbestrol vaginal suppositories may prove helpful.

**Prognosis**

Most patients respond to one of the conservative measures mentioned previously. Those who do not may require surgery.

**INTERNAL VESICAL HERNIATION**

One side of the bladder may become involved in an inguinal hernia (in men) or a femoral hernia (in women) (Figure 37–2). Such a mass may recede on urination. It is most often found as a previously unsuspected complication during surgical correction of a hernia (Bell and Witherington, 1980). Open or laparoscopic surgery with mesh repair is equally successful (McCormack et al, 2005).
Partial or complete urinary incontinence may develop after prostatectomy, particularly radical or transurethral prostatectomy. Intrinsic damage to the smooth muscle urethral sphincter is implied. Although it is common to incriminate damage to or resection of the external voluntary sphincter, this is very rare. Such a patient can stop the voiding stream by contraction of the latter sphincter, but prolonged control is impossible because of fatigue of striated muscle. Only the smooth muscle with its constant tone can afford continence.

Scott, Bradley, and Timm (1974) and Light and Scott (1983) have described an ingenious method for affording urinary control by means of an artificial sphincter. It consists of a reservoir of fluid in a Silastic bag placed deep to the abdominal wall near the bladder and a collar of Silastic material that can encircle either the bladder neck or the bulbar urethra. The former is used in females, the latter in males. One Silastic bulb is implanted in one scrotal (or labial) sac. This bulb has a special pressurized valve that inflates or deflates the cuff around the urethra; compressed fluid passes from the cuff to the reservoir, permitting free voiding. The cuff refills spontaneously after a delay of 2 minutes. This device has been successful in affording control in most instances. Results are perfect in 75% or more of cases. Most failures follow technical difficulties with the prosthesis, for example, leakage, which requires reoperation.

A rectangular flap of the heavy layer of the middle circular layer of the detrusor muscle, anteriorly, is formed into a tube, thus affording sphincteric action. This is anastomosed to the prostatic urethra. With this procedure, 44 of 50 patients who had post-prostatectomy incontinence were cured. Williams and Snyder (1976) have used this procedure successfully in children. Recently, bulking agents (Contigen) implanted around the bladder neck have been shown to be of some help. A perineal sling compressing the bulbar urethra is also a consideration.

Enuresis originally meant incontinence of urine, but usage has caused the term to be restricted to bedwetting after age 3 years. Most children have achieved normal bladder control by that time, girls earlier than boys. At age 6 years, 10% have enuresis. Even at age 14 years, 5% still wet the
Disorders of the Bladder, Prostate, & Seminal Vesicles

It is difficult to be sure, but it seems that more than 50% of cases are caused by delayed maturation of the nervous system or an intrinsic myoneurogenic bladder dysfunction; 30% are of psychological origin; and 20% are secondary to more obvious organic disease. Most children with functional enuresis spontaneously gain nocturnal control by age 10 years. The current thinking is that children with enuresis have high nocturnal urine production; some have normal bladder capacity, others reduced capacity.

Psychodynamics

Training in bladder control should begin after age 1 1/2 years; attempts made before this time are usually fruitless and may be harmful. If the parents fail in this teaching, the child may not develop cerebral inhibitory control over the infantile uninhibited bladder until much later in childhood. If the parents are emotionally unstable, their anxieties may be transmitted to the child, who may express tension through enuresis.

The birth of a sibling may cause loss of the child’s paramount position in the family. The child may then regress to an infancy pattern in an attempt to recapture the parents’ affection. An acute illness may be accompanied or followed by recurrence of incomplete nocturnal control. Physiologic or psychological stress (fear and anxiety) may reestablish an uninhibited bladder. Possibly 40% of enuretic children have electroencephalograms that are borderline or compatible with epilepsy or delayed maturation of the central nervous system.

Clinical Findings

A. Symptoms

A child may wet the bed occasionally or regularly. Careful questioning of the parents or observation by the physician reveals that the patient voids a free stream of normal caliber. This tends to rule out obstruction of the lower tract as a cause of the enuresis. Children with daytime incontinence are apt to have more than psychogenic enuresis. Many void frequently and are found to have a diminished vesical capacity, although capacity is normal under anesthesia. This is probably a reflection of delayed maturation. There is no burning, although frequency and urgency are common. The urine is clear. Observation of the parents often reveals that they are anxious and tense, traits that can only be aggravated by the child’s bedwetting.

B. Signs

General physical and urologic examinations are normal.

C. Laboratory Findings

In the emotional and delayed maturation groups, all tests, including urinalysis, are normal. An electroencephalogram may be abnormal, however.

D. X-Ray Findings

Excretory urograms show no abnormality. The accompanying cystogram reveals no trabeculation; a film of the bladder taken immediately after voiding shows no residual urine.

E. Instrumental Examination

A catheter of suitable size passes readily to the bladder, thereby ruling out stricture. If the catheter is passed after urination, no residual urine is found. Urethrocytostoscopy is normal. Cystometric studies are usually abnormal, and a curve typical of the "uninhibited" (hyperirritable) neuropathic bladder is often obtained. Unless infection or some more obvious organic disease is discovered, instrumentation, x-ray, and urodynamic studies are not necessary. Genetic factors are currently considered the primary cause of enuresis.

Differential Diagnosis

A. Obstruction

Lower tract obstruction (eg, posterior urethral valves, meatal stenosis) causes a urinary stream of decreased caliber. Painful, frequent urination during the day and night, pyuria, and fever (eg, pyelonephritis) are often present, and the bladder may be distended. Urinalysis usually reveals evidence of infection. Anemia and impairment of renal function may be demonstrated.

Excretory urograms may show dilatation of the bladder and the upper urinary tract. Incomplete vesical emptying may be seen on the postvoiding film. Cystography may demonstrate distal urethral stenosis or reflux. Urethrocytostoscopy reveals the organic cause. Severe obstruction from severe spasm of the entire pelvic floor musculature on a psychosomatic basis can cause damage to the bladder and kidneys; infection is the rule.

B. Infection

Chronic urinary tract infection not due to obstruction usually produces frequency both day and night and pain on urination, although such infections may occur without symptoms of vesical irritability. Recurrent fever with exacerbations is common.

General examination may be normal. Anemia may be noted. Urinalysis shows pus cells or bacteria, or both. Renal function may be deficient. Excretory urograms may be essentially normal, although changes compatible with healed pyelonephritis are often seen. Cystoscopy shows the changes caused by infection. Urine specimens obtained by ureteral catheter may reveal renal infection. Cystography may show vesicoureteral reflux.

C. Neurogenic Disease

Children who have sacral cord or root abnormality (eg, myelodysplasia) may have incomplete urinary control both day and night. Since they ordinarily have significant
amounts of residual urine, infection is usually found on urinalysis. The passage of a catheter or the postvoiding film taken in conjunction with excretory urograms demonstrates the presence of residual urine. A plain film of the abdomen may reveal spina bifida. The cystometrogram is usually typical of a flaccid neurogenic bladder. Cystoscopy demonstrates an atonic bladder with moderate trabeculation and evidence of infection.

D. DISTAL URETHRAL STENOSIS

Distal urethral stenosis, a congenital anomaly, is the cause of enuresis in many young girls, even in the absence of cystitis. Urethral calibration establishes this diagnosis.

Complications

The complications of functional enuresis are psychological, not organic. These children are particularly disturbed when they begin to attend school; even more pressure is brought to bear by their parents. These children find it impossible to stay overnight at the homes of their playmates. Unhealthy introversion may be their lot. Enuresis may be prolonged because of undue emphasis on dryness or as a result of punitive or shaming measures.

Late Sequels

Occasionally an adult is seen who, under stress, develops nocturnal frequency without comparable diurnal frequency. Thorough urologic investigation proves to be negative. Many of these people give histories of enuresis of long duration in childhood. It is suggested that their cerebrovesical pathways again break down with excessive emotional tension; nocturnal frequency may be the adult expression of enuresis.

Treatment

Treatment should be considered if enuresis persists after age 3 years.

A. GENERAL MEASURES

Fluids should be limited after supper. The bladder should be completely emptied at bedtime, and the child should be completely awakened a little before the usual time of bedwetting and allowed to void.

Drug therapy has its proponents.

1. Imipramine has been reported to cure 50–70% of patients and is probably the drug of choice. The starting dose is 25 mg before dinner, which is increased as needed to 50 mg. Usually, 25 mg is sufficient.

2. Parasympatholytic drugs such as atropine or belladonna, by decreasing the tone of the detrusor, may at times be of value. Methantheline bromide, 25–75 mg at bedtime, is more potent.

3. Sympathomimetic drugs, for example, dextroamphetamine sulfate, 5–10 mg at bedtime, may cause enough wakefulness so that the child perceives the urge to void.

4. Desmopressin is an antiuretic that increases renal reabsorption of water, reducing urine output in patients with a decreased nocturnal peak in antidiuretic hormone. Given as a nasal spray by night, it has been successful in 70% of patients with increased nocturnal urine output.

5. Phenytoin has been found to control symptoms in some children whose electroencephalograms are abnormal.

The use of mechanical devices such as metal-covered pads that when wet cause an alarm to ring may be of benefit in cases of delayed maturation by setting up a conditioned reflex. Urologic treatments (eg, urethral dilation, urethral instillations of silver nitrate), though often recommended, should be condemned in the absence of demonstrable local disease. They are physically and psychologically traumatic and can only cause further apprehension and fear in an already disturbed child.

B. PSYCHOTHERAPY

Analytic evaluation and treatment may be indicated for some enuretic children and their parents. Responsibility for correction of the patient’s feelings of insecurity rests with the parents, who must be cautioned not to punish the child or in any way increase existing feelings of guilt and insecurity. The handling of the parents may prove difficult, in which case psychiatric referral may be necessary.

Prognosis

Retraining the enuretic child and, above all, reeducating the parents are difficult and time-consuming tasks. Psychiatric referral for the parents and, at times, for the child may be necessary. Most patients conquer their enuresis by age 10 years. A few, however, do not, and they may later develop vesical irritability of the psychogenic type in response to acute or chronic tension or anxiety.

FOREIGN BODIES INTRODUCED INTO THE BLADDER & URETHRA

Numerous objects have been found in the urethra and bladder of both men and women. Some of them find their way into the urethra in the course of inquisitive self-exploration. Others are introduced (in the male) as contraceptive devices in the hope that plugging the urethra will block emission of the ejaculate.

The presence of a foreign body causes cystitis. Hematuria is not uncommon. Embarrassment may cause the victim to delay medical consultation. A plain x-ray of the
bladder area discloses metal objects. Nonopaque objects sometimes become coated with calcium. Cystoscopy visualizes them all.

Cystoscopic or suprapubic removal of the foreign body is indicated. If not removed, the foreign body will lead to infection of the bladder. If the infecting organisms are urea-splitting, the alkaline urine (which causes increased insolubility of calcium salts) contributes to rapid formation of stone on the foreign object (Figure 16–13).

VESICAL MANIFESTATIONS OF ALLERGY

So many mucous membranes are affected by allergens that the possibility of allergic manifestations involving the bladder must be considered. Hypersensitivity is occasionally suggested in cases of recurrent symptoms of acute “cystitis” in the absence of urinary infection or other demonstrable abnormality. During the attack, general erythema of the vesical mucosa may be seen and some edema of the ureteral orifices noted.

A careful history may reveal that these attacks follow the ingestion of a food not ordinarily eaten (eg, fresh lobster). Sensitivity to spermicidal creams is occasionally observed. If vesical allergy is suspected, it may be aborted by the subcutaneous injection of 0.5–1 mL of 1:1000 epi-nephrine. Control may also be afforded by the use of one of the antihistamines. Skin testing has not generally proved helpful in determining the source of allergy.

DIVERTICULA

Most vesical diverticula are acquired and are secondary to either obstruction distal to the vesical neck or the upper motor neuron type of neurogenic bladder. Increased intravesical pressure causes vesical mucosa to insonate itself between hypertrophied muscle bundles, so that a mucosal extravesical sac develops. Often this sac lies just superior to the ureter and causes vesicoureteral reflux (Hutch saccule; Figure 12–6). The diverticulum is devoid of muscle and therefore has no expulsive power; residual urine is the rule, and infection is perpetuated. If the diverticulum has a narrow opening that interferes with its emptying, transurethral resection of its neck will improve drainage. Carcinoma occasionally develops on its wall. Mic’ic´ and Ilic´ (1983) discovered 13 diverticula harboring malignant tumors: 9 transitional cell tumors, 2 squamous cell tumors, and 2 adenocarcinomas. Gerridzen and Futter (1982) saw 48 cases of vesical diverticula. Transitional cell tumors were found in 5 of these patients, but almost all the rest had abnormal histopathology: chronic inflammation and metaplasia. These authors stress the need for visualizing the interior of diverticula during endoscopy. At the time of open prostatectomy, resection of a diverticulum should be considered.

VESICAL FISTULAS

Vesical fistulas are common. The bladder may communicate with the skin, intestinal tract, or female reproductive organs. The primary disease is usually not urologic. The causes are as follows: (1) primary intestinal disease—diverticulitis, 50–60%; cancer of the colon, 20–25%; and Crohn disease, 10% (Badlani et al, 1980); (2) primary gynecologic disease—pressure necrosis during difficult labor; advanced cancer of the cervix; (3) treatment for gynecologic disease following hysterectomy, low cesarean section, or radiotherapy for tumor; and (4) trauma.

Malignant tumors of the small or large bowel, uterus, or cervix may invade and perforate the bladder. Inflammations of adjacent organs may also erode through the vesical wall. Severe injuries involving the bladder may lead to perivesical abscess formation, and these abscesses may rupture through the skin of the perineum or abdomen. The bladder may be inadvertently injured during gynecologic or intestinal surgery; cystotomy for stone or prostatectomy may lead to a persistent cutaneous fistula.

Clinical Findings

A. VESICOINTESTINAL FISTULA

Symptoms arising from a vesicointestinal fistula include vesical irritability, the passage of feces and gas through the urethra, and usually a change in bowel habits (eg, constipation, abdominal distention, diarrhea) caused by the primary intestinal disease. Signs of bowel obstruction may be elicited; abdominal tenderness may be found if the cause is inflammatory. The urine is always infected.

A barium enema, upper gastrointestinal series, or sigmoidoscopic examination may demonstrate the communication. Following a barium enema, centrifuged urine should be placed on an x-ray cassette and an exposure made. The presence of radiopaque barium establishes the diagnosis of vesicocolonic fistula. Cystograms may reveal gas in the bladder or reflux of the opaque material into the bowel (Figure 37–3). Cystoscopic examination, the most useful diagnostic procedure, shows a severe localized inflammatory reaction from which bowel contents may exude. Catheterization of the fistulous tract may be feasible; the instillation of radiopaque fluid often establishes the diagnosis.

B. VESICOVAGINAL FISTULA

This relatively common fistula is secondary to obstetric, surgical, or radiation injury or to invasive cancer of the cervix. The constant leakage of urine is most distressing to the patient. Pelvic examination usually reveals the fistulous opening, which also can be visualized with the cystoscope. It may be possible to pass a ureteral catheter through the fistula into the vagina. Vaginography often successfully shows ureterovaginal, vesicovaginal, and rectovaginal fistu-
Figure 37–3. Vesical fistulas. **Above left:** Primary carcinoma of the sigmoid, with perforation through bladder wall. **Above right:** Injury to base of bladder following delivery by forceps. **Below:** Cystogram showing radiopaque fluid entering sigmoid containing multiple diverticula; right ureteral reflux, gallbladder calculi.
A 30-mL Foley catheter is inserted into the vagina, and the balloon is distended. A radiopaque solution is then instilled, and appropriate x-rays are taken. Biopsy of the edges of the fistula may show carcinoma.

**Differential Diagnosis**

It is necessary to differentiate ureterovaginal from vesicovaginal fistula. Phenazopyridine (Pyridium) is given by mouth to color the urine orange. One hour later, 3 cotton pledgets are inserted into the vagina, and methylene blue solution is instilled into the bladder. The patient should then walk around, after which the pledgets are examined. If the proximal cotton ball is wet or stained orange, the fistula is ureterovaginal. If the deep cotton pledget contains blue fluid, the diagnosis is vesicovaginal fistula. If only the distal pledget is blue, the patient probably has urinary incontinence (Raghavaiah, 1974).

**Treatment**

**A. VESICOINTESTINAL FISTULA**

If the lesion is in the rectosigmoid, treatment consists of proximal colostomy. When the inflammatory reaction has subsided, the involved bowel may be resected, with closure of the opening in the bladder. The colostomy can be closed later. Some authors recommend that the entire procedure be performed in one stage, thus avoiding the need for preliminary colostomy. Small bowel or appendiceal vesical fistulas require bowel or appendiceal resection and closure of the vesical defect.

**B. VESICOVAGINAL FISTULA**

Tiny fistulous openings may become sealed following the introduction of an electrode into the fistula. As the electrode is withdrawn, the fistula is coagulated with the electrosurgical unit to destroy the epithelium of the tract. An indwelling catheter should be left in place for 2 weeks or more. Occasionally, good results are noted in cases of small vesicovaginal fistulas treated by inserting a metal screw through the vaginal end of the fistula. It is moved up and down to act as a curet. The vaginal mucosa is then closed and an indwelling catheter placed for 3 weeks.

Larger fistulas secondary to obstetric or surgical injuries respond readily to surgical repair, which may be done either through the vagina or transvesically. Persky, Herman, and Guerrier (1979) advise repairing such fistulas immediately rather than waiting for 3–6 months as counseled by most surgeons. Fistulas that develop following radiation therapy for cancer of the cervix are much more difficult to close because of the avascularity of the tissues. Surgical closure of fistulas that arise from direct invasion of the bladder by cervical carcinoma is impossible; diversion of the urinary stream above the level of the bladder (eg, ureterosigmoidostomy) is therefore necessary.

**C. VESICOADNEXAL FISTULA**

These fistulas are cured by removal of the involved female reproductive organs, with closure of the opening in the bladder.

**Prognosis**

The surgical repair of fistulas caused by benign disease or operative trauma is highly successful. Postirradiation necrosis offers a more guarded prognosis. Fistulas secondary to invading cancers present difficult problems.

**PERIVESICAL LIPOMATOSIS**

The cause of perivesical lipomatosis is not known. The disorder seems to affect principally black men in the 20- to 40-year age group. There are no pathognomonic symptoms. There may be some dysuria or mild urinary obstructive symptoms. Examination may demonstrate a distended or enlarged pear-shaped bladder. Excretory urograms and cystography may show dilatation of both upper tracts and an upward displacement and lateral compression of the bladder. In the perivesical area, x-ray reveals areas of radiolucency compatible with fatty tissue. A barium x-ray may show extrinsic pressure on the rectosigmoid. Angiography shows no evidence of neoplastic vessels.

Computed tomography scan in association with the preceding findings establishes the diagnosis by clearly demonstrating the fatty nature of the perivesical tissue. Church and Kazam (1979) found sonography equally helpful.

On surgical exploration, lipomatous tissue is found surrounding the bladder and rectosigmoid. Though it is tempting to proceed with its resection, there are no cleavage planes. Such dissections usually fail to relieve the ureteral obstruction. Ballesteros (1977) believes that surgical excision is feasible and reported excellent results in one such case. Crane and Smith (1977) found, after a 5-year follow-up, that hydronephrosis progressed in most. Many patients finally required urinary diversion.

**RADIATION CYSTITIS**

Many women receiving radiation treatment for carcinoma of the cervix develop some degree of vesical irritability. These symptoms may develop months after cessation of treatment. The urine may or may not be sterile. Vesical capacity is usually appreciably reduced. Cystoscopy reveals a pale mucous membrane with multiple areas of telangiectatic blood vessels. Vesical ulceration may be noted, and vesicovaginal fistulas may develop. If symptoms are severe and prolonged, diversion of urine from the bladder may be necessary.
NONINFECTIOUS HEMORRHAGIC CYSTITIS

Some patients, following radiotherapy for carcinoma of the cervix or bladder, are prone to intermittent, often serious vesical hemorrhage. The same is true of those given cyclophosphamide.

In the case of the latter, the drug must be stopped. To control bleeding, cystoscopic fulguration can be tried, though it usually fails. The instillation of 3.9% formalin (prepared by diluting the standard 39% solution 10 times) is more efficacious. The catheter is clamped for 30 minutes and the bladder lavaged with 10% alcohol. A second or third instillation may be necessary on subsequent days. Holstein et al (1973) recommend the transurethral placement of a large balloon in the bladder. The balloon is filled to a pressure level equal to the systolic blood pressure and left in place for 6 hours.

Giulani et al (1979) have reported success by selective transcatheter embolization of the internal iliac arteries. Ostroff and Chenault (1982) believe that the best and least harmful method of treatment is continuous irrigation with 1% alum solution (the ammonium or potassium salt) through a 3-way Foley catheter.

Despite these measures, the mortality rate is significant. Droller, Saral, and Santos (1982) have evolved a plan for reducing the incidence of cyclophosphamide-induced hemorrhagic cystitis: they produce diuresis and have the patient void frequently (or use open catheter drainage). This reduces the concentration of cyclophosphamide metabolites and the duration of their contact with bladder mucosa. Before the institution of this regimen, 8 of 97 such patients died; afterward, 1 of 198 patients died.

EMPYEMA OF THE BLADDER

If supravesical diversion of the urine is performed without cystectomy, severe infection of the bladder may develop because of lack of washout. In males, cystostomy or cutaneous vesicostomy may be necessary. In females, the formation of a vesicovaginal fistula permits drainage (Spence and Allen, 1971). Occasionally, cystectomy may be necessary.

CONGENITAL ANOMALIES OF THE PROSTATE & SEMINAL VESICLES

Congenital anomalies of the prostate are rare. Cysts of the prostate and the seminal vesicles have been reported. Enlargements of the prostatic utricle are often found in association with penoscrotal or perineal hypospadias. The cysts are usually small, lying in the midline posterior to the prostate and emptying through the verumontanum. These cysts represent embryologic remnants of the distal end of the Müllerian ducts (see Chapter 2). Rarely, they become large enough to be easily palpable rectally or even abdominally. Through local pressure, they may cause symptoms of obstruction of the bladder neck.

BLOODY EJACULATION

Hemospermia is not an uncommon complaint of middle-aged men. It is the wife who usually recognizes the symptom. It is thought by some to be caused by hyperplasia of the mucosa of the seminal vesicles. For this reason, the use of diethylstilbestrol, 5 mg/day for 1 week, has been suggested. In my hands, it has worked well. Thorough urologic investigation of men without other symptoms rarely reveals a pathologic lesion. The cause is therefore not clear.

REFERENCES

Exstrophy


**Persistent Urachus**


**Contracture of the Bladder Neck**


**Interstitial Cystitis**


**Internal Vesical Herniation**


**Urinary Incontinence**


**Enuresis**


DISORDERS OF THE BLADDER, PROSTATE, & SEMINAL VESICLES / 587


**Foreign Bodies Introduced into the Bladder & Urethra**


**Vesical Manifestations of Allergy**


**Diverticula**


**Vesical Fistulas**


**Perivesical Lipomatosis**


**Radiation Cystitis**


**Noninfectious Hemorrhagic Cystitis**

Giulani L et al: Gelatin foam and isobutyl-2-cyanoacrylate in the treatment of life-threatening bladder haemorrhage by selective tran-


**Empyema of the Bladder**


**Congenital Anomalies of the Prostate & Seminal Vesicles**


**Bloody Ejaculation**

Male Sexual Dysfunction

Anthony J. Bella, MD, & Tom F. Lue, MD

In the United States, it is estimated that more than half of men aged 40–70 are unable to attain or maintain a penile erection sufficient for satisfactory sexual performance. Advances in pharmacologic therapy for erectile dysfunction (ED), coupled with a better understanding of male sexual dysfunction made possible by innovative laboratory and clinical research in the mechanism, neurophysiology, and pharmacology of penile erection, have resulted in greater numbers of patients seeking primary and specialty care for sexual concerns. Oral phosphodiesterase type-5 inhibitors have emerged as the preferred first-line treatment of ED worldwide due to their efficacy, ease-of-use, and patient safety. Erectile function can now be evaluated by the response to these agents at home or intracavernous injection (ICI) of vasoactive agents in the office, and improved diagnostic tests can differentiate among types of impotence. Patient satisfaction with penile prostheses is high, as the latest generation of devices is more sophisticated and durable than ever. Current treatments continue to evolve and together with novel molecular, stem cell, and gene therapies will represent the next generation of more physiologic and disease-specific solutions to various types of ED.

PHYSIOLOGY OF PENILE ERECTION

Innervation of the Penis

The autonomic spinal erection center is located in the intermediolateral nuclei of the spinal cord at levels S2–S4 and T12–L2. Nerve fibers from the thoracolumbar (sympathetic) and sacral (parasympathetic) spinal segments join to form inferior hypogastric and pelvic plexuses, which send branches to the pelvic organs. The fibers innervating the penis (cavernous nerves) travel along the posterolateral aspect of the seminal vesicles and prostate and then accompany the membranous urethra through the genitourinary diaphragm (Figure 38–1). Some of these fibers enter the corpora cavernosa and corpus spongiosum with the cavernous and bulbourethral arteries. Others travel distally with the dorsal nerve and enter the corpus cavernosum and corpus spongiosum in various locations to supply the mid- and distal portions of the penis. The terminal branches of the cavernous nerves innervate the helicine arteries and trabecular smooth muscle, and are responsible for vascular events during tumescence and detumescence.

The center for somatic motor nerves is located at the ventral horn of the S2–S4 segment (Onuf’s nucleus). The motor fibers join the pudendal nerve to innervate the bulbocavernous and ischiocavernous muscles. The somatic sensory nerves originate at receptors to innervate the bulbocavernous and ischiocavernous muscles. The somatic sensory nerves originate at receptors to innervate the bulbocavernous and ischiocavernous muscles.

Anatomy & Hemodynamics of Penile Erection

The tunica of the corpora cavernosa is a bilayered structure with multiple sublayers. The inner circular bundles sup-
port and contain the cavernous tissue. From this inner layer, intracavernosal pillars that act as struts radiate to augment the septum; both structures provide essential support to the erectile tissue. The outer-layer bundles are oriented longitudinally and extend from the glans penis to the proximal crus. These fibers insert into the inferior pubic ramus but are absent between the 5- and 7-o’clock positions. In contrast, the corpus spongiosum lacks an outer layer or intracorporeal struts, ensuring a lower pressure structure during erection. The tunica is composed of elastic fibers forming a network on which the collagen fibers rest. Emissary veins run between the inner and outer layers for a short distance, often piercing the outer bundles obliquely. Branches of the dorsal artery take a more direct perpendicular route and are surrounded by a periarterial fibrous sheath (Hsu et al, 2004).

The paired internal pudendal artery is the major carrier of the blood supply to the penis, dividing into 3 branches: the bulbourethral artery, dorsal artery, and the cavernous artery (deep artery). The cavernous artery supplies the corpora cavernosa; the dorsal artery, the skin, subcutaneous tissue, and the glans penis; and the bulbourethral artery, the corpus spongiosum. In some cases, accessory pudendal arteries from external iliac or obturator arteries may supply a major portion of the penis, with collaterals among the 3 branches often observed. The venous drainage of the glans is mainly through the deep dorsal vein. The corpus spongiosum is drained via the circumflex, urethral, and bulbar veins, but the drainage of the corpora cavernosa is more complex: the mid- and distal shaft are drained by the deep dorsal vein to the preprostatic plexus while the proximal portion is drained by the cavernous and crural veins to the preprostatic plexus and internal pudendal vein. The drainage of all 3 corpora originates in the subtunical venules, which unite to form emissary veins. The glans penis possesses numerous large and small veins that communicate freely with the dorsal veins. The penile skin and subcutaneous tissue are drained by superficial dorsal veins, which then empty into the saphenous veins.

Activation of the autonomic nerves produces a full erection secondary to filling and trapping of blood in the cavernous bodies. After full erection is achieved, contraction of the ischiocavernosus muscle (from activation of the somatic nerves) compresses the proximal corpora and raises the intracorporal pressure well above the systolic blood pressure, resulting in rigid erection (Table 38–1). This rigid phase occurs naturally during masturbation or sexual intercourse but can also occur from slight bending of the penis, without muscular action. The erection process can be divided into phases as shown in Table 38–1 and Figure 38–2. The

\[ Figure 38–1. \text{Location of cavernous nerves in relation to urethra.} \]
MALE SEXUAL DYSFUNCTION

Hemodynamics of the penile glans is somewhat different. Arterial flow increases in a manner similar to that in the shaft. Because it lacks the tunica albuginea, however, the glans functions as an arteriovenous fistula during the full erection phase. Nevertheless, during rigid erection, most of the venous channels are temporarily compressed, and further engorgement of the glans can be observed (Lue, 2000).

Mechanism of Penile Erection

The penile erectile tissue, specifically cavernous, arteriolar, and arterial wall smooth musculature, is key to the erectile process. In the flaccid state, these smooth muscles are tonically contracted due to intrinsic smooth-muscle tone and possibly tonic adrenergic discharge, allowing only a small amount of arterial flow for nutritional purposes. The blood partial pressure of oxygen (PO$_2$) is about 35 mmHg. When smooth muscles relax due to the release of neurotransmitters, resistance to incoming flow drops to a minimum. Arterial and arteriolar vasodilatation occurs, and sinusoids expand to receive a large increase of flow. Trapping of blood causes the penis to lengthen and widen rapidly until the capacity of the tunica albuginea is reached. Expansion of the sinusoidal walls against one another and the tunica albuginea results in compression of the subtunical venous plexus. As well, uneven stretching of the layers of the tunica albuginea compresses the emissary veins and effectively reduces the venous flow to a minimum (Lue, 2000; Figure 38–3A and B). Intracavernous pressure (ICP) and PO$_2$ increase to about 100 and 90 mm Hg, respectively, raising the penis from a dependent position to the erect state; further pressure increases due to contraction of the ischiocavernous muscles (to several hundred millimeters of mercury) result in the rigid-erection phase (Dean and Lue, 2005).

Hormones and Sexual Function

Androgens are essential for male sexual maturity. Testosterone (T) regulates gonadotropin secretion and muscle development; dihydrotestosterone mediates male sexual maturation, including hair growth, acne, male pattern baldness, and spermatogenesis. In adults, androgen deficiency results in decreased libido (sexual interest) and impaired seminal emission. Aging is associated with a progressive decline of testosterone, dehydroepiandrosterone, thyroxine, melatonin and growth hormone, and increased levels of sex hormone-binding globulin, pituitary gonad-
tropins, and prolactin (Morales, 2005). In a longitudinal study of middle-aged men for 7–10 years, total T levels declined at 0.8%/year of age, whereas both free and albumin-bound T declined at about 2%/year. Sex hormone-binding globulin increased at 1.6%/year (Feldman et al, 2002). Testosterone levels do not correspond to severity of ED, however lower levels are observed in men with reduced libido. Although frequency, magnitude, and latency of nocturnal penile erections are reduced with decreased T, erectile response to visual sexual stimulation is preserved in men with hypogonadism, suggesting that androgen is not essential for erection. Due to the inhibitory action of prolactin on central dopaminergic activity and resultant decreases in gonadotropin-releasing hormone secretion, hyperprolactinemia of any cause results in both reproductive and sexual dysfunction (Corona et al, 2004).

**Neurotransmitters and Pharmacology of Erection**

Neural control of penile erection involves adrenergic, cholinergic, and nonadrenergic-noncholinergic (NANC) neurotransmitter systems. Adrenergic nerves mediate intracavernous smooth-muscle contraction, maintaining the penis in a nonerect state through the release of norepinephrine, the principal neurotransmitter controlling penile flaccidity and detumescence. Currently, it is suggested that sympathetic contraction is mediated by activation of postsynaptic alpha-1a- and alpha-1d-adrenergic receptors and modulated by presynaptic alpha-2-adrenergic receptors (Giuliano et al, 2004). Cholinergic nerves may contribute to smooth-muscle relaxation and penile erection through inhibition of adrenergic nerves via inhibitory interneurons and the release of nitric oxide (NO) from the endothelium by acetylcholine (Saenz de Tejada et al, 2004). Endothelin, a potent vasoconstrictor produced by the endothelial cells, has also been suggested to be a mediator for detumescence.

The principal neurotransmitter for penile erection is NO from parasympathetic, NANC nerve terminals. Once blood rushes into the sinusoids, shear stress can also release NO from endothelium to augment smooth muscle relaxation and erection. In addition, oxygen tension and substances secreted by endothelium lining the sinusoidal spaces, prostaglandins, endothelins, and angiotensin may also be involved in penile erection and detumescence (Musicki and Burnett, 2006). The agents capable of inducing erection and causing detumescence are summarized in Table 38–2. Although mechanisms of action vary, erection-inducing substances cause smooth muscle to relax and detumescing agents cause them to contract.

**Molecular Mechanism of Smooth-Muscle Contraction & Relaxation**

Smooth-muscle contraction is regulated by Ca\(^{2+}\). As cytosolic free Ca\(^{2+}\) increases from resting levels of 120–
270 to 500–700 nM, calmodulin-4 Ca2+ complex binds to myosin light-chain kinase. The activated kinase then phosphorylates the light chain, which ceases to inhibit the myosin-actin interaction and initiates a contraction cycle (Andersson, 2003). Once cytosolic Ca2+ returns to basal levels, calcium-sensitizing pathways take over. Activation of excitatory receptors coupled to G proteins causes contraction by increasing calcium sensitivity without changes in cytosolic Ca2+ levels. This pathway involves RhoA, a small monomeric G protein that activates Rho-kinase. Activated Rho-kinase phosphorylates, and thereby inhibits, the regulatory subunit of smooth muscle myosin phosphatase, preventing dephosphorylation of myofilaments and maintaining contractile tone (Jin and Burnett, 2006). The emerging consensus is that phasic contraction of penile smooth muscle is regulated by increased cytosolic Ca2+ and tonic contraction is governed by calcium-sensitizing pathways.

**Signal Transduction in Penile Erection**

During sexual stimulation, NO released from nerve endings and endothelium diffuses into the trabecular and arterial smooth muscle cells to activate guanylyl cyclase, catalyzing the formation of second messenger cyclic guanosine monophosphate (cGMP). cGMP in turn activates protein kinase G, phosphorylating potassium and calcium channels; the end result is hyperpolarization, reduced intracytosolic calcium, and dissociation of the myosin head from actin as smooth muscle relaxes. Cyclic adenosine monophosphate (cAMP) is another second messenger involved in smooth muscle relaxation and is activated by cAMP-signaling molecules including adenosine, calcitonin gene-related peptides, and prostaglandins (Lin et al, 2005).

**Intercellular Communication**

Gap junctions are aqueous intercellular channels that have been demonstrated to interconnect the cytoplasm of adjacent cells in many tissues. In the penis, smooth-muscle cells are sparsely innervated by the terminal branches of the cavernous nerves. Therefore, gap junctions play a vital role in the intercellular communication within the corpus cavernosum, enabling the penis to function as a unit (Schiff and Melman, 2006).

**MALE SEXUAL DYSFUNCTION**

Male sexual dysfunction, denoting the inability to achieve a satisfactory sexual relationship, may involve inadequacy of erection or problems with emission, ejaculation, or orgasm. Premature (rapid) ejaculation refers to persistent or recurrent occurrence of ejaculation with minimal sexual stimulation before, on, or shortly after penetration and before the person wishes it. Retarded ejaculation is undue delay in reaching a climax during sexual activity. Retrograde ejaculation denotes backflow of semen into the bladder during ejaculation owing to an incompetent bladder neck mechanism. Anorgasmia is the inability to achieve an orgasm during conscious sexual activity, although nocturnal emission may occur.

**Epidemiology**

In the Massachusetts Male Aging Study, a community-based survey of men between 40 and 70 years of age, 52% of respondents reported some degree of ED: 17% mild, 25% moderate, and 10% complete. Although the preva-
lence of mild ED remained constant (17%) between the age of 40 and 70, there was a doubling in the number of men reporting moderate ED (from 17% to 34%) and a tripling in the number of men reporting complete ED (from 5% to 15%) (Feldman et al, 1994). More than 70% of men over 65 years report that they are sexually active; however, 40% are dissatisfied with their sexual function (Braun et al, 2000). Among the major predictors of ED are hypertension, hyperlipidemia, diabetes mellitus, and heart disease. Risk of ED appears to increase with smoking, and may occur in a dose-dependent manner (Polsky et al, 2005). There is a higher prevalence of ED in men who have undergone radiation or surgery for prostate cancer or other pelvic malignancies. The psychological correlates of ED include decreased self-esteem, depression, anxiety, anger, and relationship dissatisfaction (Althof et al, 2006). Other male sexual dysfunctions have also been found to be highly prevalent: premature ejaculation and decreased libido (lack of sexual interest) are common patients’ concerns.

Classification and Pathogenesis

The classification system of ED most commonly used encompasses organic, psychogenic, and mixed etiologies of ED and is endorsed by the International Society of Impotence Research (Table 38–3). In the 1950s, 90% of cases of ED were believed to be psychogenic. Most authors now believe that mixed organic and psychogenic ED is the most common.

Psychological Disorders

Many psychologic conditions (performance anxiety, strained relationship, lack of sexual arousal, depression, and schizophrenia) can either cause or aggravate ED. Sexual behavior and penile erection are controlled by the hypothalamus, cerebral cortex, and limbic systems. Given the number and complexity of known and as yet unidentified factors involved, it is not surprising that the pathogenesis of psychogenic ED is still speculative. Possible mechanisms proposed include an imbalance of central neurotransmitters, overinhibition of spinal erection center by the brain, inadequate NO release, and sympathetic overactivity (Bodie et al, 2003).

Neurogenic Disorders

It has been estimated that up to 20% of all ED is neurogenic in origin, resulting from peripheral (cavernous and pudendal nerve) or central pathologies (Saenz de Tejada et al, 2005). In men with spinal cord injury, the degree of erectile function depends on the nature, location, and the extent of the lesion. Brain lesions associated with ED include dementias, Parkinson’s disease, stroke, tumors, trauma, and Shy-Drager syndrome (Papatsoris et al, 2006). Peripheral neuropathy due to diabetes mellitus, chronic alcohol abuse, or vitamin deficiency may affect the nerve endings and result in a deficiency of neurotransmitters. Direct injury to the cavernous or pudendal nerves from trauma, radical pelvic surgeries for malignancy, or pelvic irradiation can also cause ED. Iatrogenic impotence resulting from common procedures has been reported at the following rates: radical prostatectomy 30–100%, abdominal-peroneal resection 15–100%, and external sphincterotomy at 3- and 9-o’clock positions, 2–49% (Dean and Lue, 2005). It is important to note that even with nerve-sparing approaches to surgery, erectile recovery can take up to 24 months; newer data suggest functional improvement continues up to 48 months postoperatively.

Hormonal Disorders

Historically, hypogonadism as a cause of ED was thought to be rare, but recent data support a significant increase of hypogonadism with age. Hypogonadism due to hypothalamic or pituitary tumors, estrogen or antiandrogen therapy, or orchietomy can suppress sexual interest and nocturnal erections. As mentioned earlier, erections are usually

Table 38–3. New Classification of ED Recommended by the International Society of Impotence Research.

<table>
<thead>
<tr>
<th>I. Psychogenic</th>
</tr>
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<tbody>
<tr>
<td>1. Generalized type</td>
</tr>
<tr>
<td>A. Generalized unresponsiveness</td>
</tr>
<tr>
<td>a. Primary lack of sexual arousability</td>
</tr>
<tr>
<td>b. Aging-related decline in sexual arousability</td>
</tr>
<tr>
<td>2. Situational type</td>
</tr>
<tr>
<td>A. Partner related</td>
</tr>
<tr>
<td>a. Lack of arousability in specific relationship</td>
</tr>
<tr>
<td>b. Lack of arousability due to sexual object preference</td>
</tr>
<tr>
<td>c. High central inhibition due to partner conflict or threat</td>
</tr>
<tr>
<td>B. Performance related</td>
</tr>
<tr>
<td>a. Associated with other sexual dysfunction/s (eg, rapid ejaculation)</td>
</tr>
<tr>
<td>b. Situational performance anxiety (eg, fear of failure)</td>
</tr>
<tr>
<td>C. Psychological distress or adjustment related</td>
</tr>
<tr>
<td>a. Associated with negative mood state (eg, depression) or major life stress (eg, death of partner)</td>
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</tbody>
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<table>
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<th>II. Organic</th>
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</thead>
<tbody>
<tr>
<td>1. Neurogenic</td>
</tr>
<tr>
<td>2. Hormonal</td>
</tr>
<tr>
<td>3. Arterial</td>
</tr>
<tr>
<td>4. Cavernosal (venogenic)</td>
</tr>
<tr>
<td>5. Drug induced</td>
</tr>
</tbody>
</table>

| III. Mixed organic/psychogenic (most common type) |
preserved to some extent. Hyperprolactinemia, Cushing's syndrome, and Addison's disease can cause decreased libido and ED. Hyperthyroidism is commonly associated with decreased libido, likely due to elevated estrogen levels, while hypothyroidism can contribute to ED through diminished testosterone secretion and elevated prolactin levels (Veronelli et al, 2006).

**Arterial Disorders**

Although arteriogenic ED may be congenital or due to trauma, most often it is part of a generalized systemic arterial disease. The distribution and severity of the disease, however, differ from person to person. Traumatic arterial occlusive or atherosclerotic disease of the hypogastric (iliac)-cavernous-helicine arterial tree can decrease flow to the sinusoidal spaces and perfusion pressure, thus decreasing the rigidity or prolonging time to maximal erection. Some patients with severe arterial disease may retain potency as long as arterial flow exceeds venous flow; conversely, some patients with minimal arterial disease may be partially or completely impotent because of relatively large venous outflow, cavernous smooth-muscle dysfunction, or inadequate neurotransmitter release (Dean and Lue, 2005).

The incidence and age at onset are parallel for coronary disease and ED. Common risk factors associated with arterial insufficiency include hypertension, hyperlipidemia, diabetes mellitus, and cigarette smoking. Long-distance cycling is also a likely risk factor for vasculogenic and neurogenic ED (Huang et al, 2005).

Arterial disease is classified as extra- or intrapenile arterial insufficiency. Extrapenile disease may be amenable to surgical repair in selected patients, and comprises diseases of the internal pudendal artery, internal and common iliac arteries, and aorta, the pelvic steal syndrome, and pelvic trauma. Intrapenile arterial disease secondary to aging, arteriosclerosis, or diabetes mellitus does not respond well to currently available surgical techniques (Milbank and Montague, 2004).

**Cavernosal Disorders**

Cavernous veno-occlusive dysfunction (CVOD) may result from a variety of pathophysiologic processes. Degenerative changes (Peyronie's disease, aging, and diabetes) and traumatic injury to the tunica albuginea (penile fracture) can impair the compression of the subtunical and emissary veins. Fibroelastic alteration of the trabeculae, cavernous smooth muscle, and endothelium may result in venous leakage (Deveci et al, 2006). Men with diabetes mellitus and atherosclerosis are at increased risk of smooth-muscle atrophy, fibrous replacement, and endothelial disruption.

CVOD impotence can be divided into 5 types according to cause: In type 1, large veins exit the corpus cavernosum (etiology is likely congenital); in type 2, venous channels are enlarged as a result of distortion of the tunica albuginea (secondary to Peyronie’s disease or weakening associated with aging); in type 3, the cavernous smooth muscle is unable to relax because of fibrosis, degeneration, or dysfunctional gap junctions; in type 4, there is inadequate neurotransmitter release (neurologic or psychologic impotence, or endothelial dysfunction); and in type 5, there is abnormal communication between the corpus cavernosum and the spongiosum or glans (congenital, traumatic, or secondary to shunt procedures for priapism) (Dean and Lue, 2005).

**Medication-Induced Erectile Dysfunction**

Many drugs have been reported to cause ED, although the mechanism of action is often unknown and there are few controlled studies on the sexual side effects of a particular agent. As ED is common among older men, it will coexist with other conditions that are themselves risk factors for ED, such as cardiovascular disease, diabetes, or depression. Sexual symptoms related to medications can also involve a combination of complaints concerning desire, arousal, and orgasm rather than being limited to impaired function.

In general, drugs that interfere with central neuroendocrine or local neurovascular control of penile smooth muscle have the potential to cause ED. Central neurotransmitter pathways, including serotonergic, noradrenergic, and dopaminergic pathways involved in sexual function, may be disturbed by antipsychotics, antidepressants, and centrally acting antihypertensive drugs (Balon, 2005; Papazisis and Korantzopoulos, 2006). Selective serotonin reuptake inhibitors are the most common class of drugs currently used to treat depression; it is estimated that up to 50% of patients using these agents experience a change in sexual function (Kellner et al, 2002). Beta-adrenergic blocking drugs may cause ED by potentiating alpha-1 adrenergic activity in the penis. Conversely, alpha-1 blockers and angiotensin-II-receptor blockers both tend to improve sexual function during treatment and may therefore be useful when commencing antihypertensive therapy in men with preexisting ED (Khan et al, 2002). Thiazide and nonthiazide diuretics have been reported to cause ED; spironolactone can also cause a decrease in libido and gynecomastia. Alpha-adrenergic blocking drugs, such as doxazosin, terazosin, and tamsulosin, may cause retrograde ejaculation owing to relaxation of the bladder neck (Guillian, 2006). Other drugs thought to cause ED include opiates, antiretroviral agents, and histamine H2 receptor antagonists (cimetidine) (Colson et al, 2002).

Antiandrogens modify sexual behavior by varying degrees, ranging from complete loss to normal function, chiefly by modulating sexual desire via central nervous system androgen receptors. Finasteride, a 5-alpha-reductase inhibitor commonly used to treat benign prostatic hypertrophy, is the antiandrogen with the least effect on circulating testosterone and sexual function. Sexual symptoms are
reported in approximately 5% of men treated with a 5-mg dose (Miner et al, 2006). Estrogens and drugs with antiandrogenic action such as ketoconazole, LHRH agonists nonsteroidal (bicalutamide) and steroidal (cyproterone acetate) acetate can diminish sexual function. The near-complete androgen deprivation achieved by medical castration with LHRH agonists results in a profound loss of sexual desire, which is usually accompanied by ED.

Cigarette smoking may induce vasoconstriction and penile venous leakage because of its contractile effect on the cavernous smooth muscle and is seen to approximately double the rate of ED in coronary artery disease, hypertension, and atherosclerosis (Korenman, 2004). Alcohol in small amounts improves erection and increases libido because of its vasodilatory effect and the suppression of anxiety; however, large amounts can cause central sedation, decreased libido, and transient ED. Chronic alcoholism may cause hypogonadism and polyneuropathy, which may affect penile nerve function (Ravaglia, 2004).

### Aging and Systemic Disease

Sexual function progressively declines in “healthy” aging men. Longitudinal studies demonstrate a nonlinear decline for most aspects of sexual function as age increases, with a more pronounced decline in older groups (Araujo et al, 2004). The latent period between sexual stimulation and erection increases, erections are less turgid, ejaculation is less forceful, ejaculatory volume decreases, and the refractory period between erections lengthens. There is also a decrease in penile sensitivity to tactile stimulation, a decrease in serum testosterone concentration, and an increase in cavernous muscle tone. While relational, psychologic, and organic issues are important contributors to ED across age groups, organic issues tend to play a more profound role as men age.

ED in men with DM occurs approximately threefold that of the general population, approaching 55% at 60 years of age, and can be the presenting symptom for DM and/or predict later neurologic sequelae (Fonseca and Java, 2005). Diabetes may affect small vessels, cavernous nerve terminals, smooth muscle and endothelial cells; neurovascular sequelae of long-term diabetes results in decreased responsiveness to oral PDE-5 inhibitor therapy.

Men with severe pulmonary disease may have ED because of fear of aggravating dyspnea during sexual intercourse (Koseoglu et al, 2005). Patients with angina, myocardial infarction (MI), or heart failure may have ED from anxiety, depression, or concomitant penile arterial insufficiency, which is quite common in these patients. Chronic renal failure has frequently been associated with diminished erectile function, impaired libido, and infertility (Shamsa et al, 2005). In men with chronic renal failure and ED, many were found to have cavernous artery occlusive disease and veno-occlusive dysfunction. The mechanism is likely multifactorial: low serum testosterone concentration, diabetes mellitus, vascular insufficiency, multiple medications, auto-nomic and somatic neuropathy, and psychological stress. Other systemic disorders such as cirrhosis, chronic debilitation, and cachexia can cause ED due to loss of libido or neurovascular dysfunction.

### DIAGNOSIS & TREATMENT

The management of ED is built on a patient-centered and evidence-based principle. A detailed medical, sexual, and psychosocial history, and a thorough physical examination are the most important steps in the differential diagnosis of sexual dysfunction. Interviewing the partner, if available, is helpful in eliciting a reliable history, planning treatment, and obtaining a successful outcome.

#### Medical, Sexual, & Psychosocial History

The goals of the medical history are to evaluate the potential role of underlying medical conditions, differentiate between potential organic and psychogenic causes, and to assess the potential role of medication(s), both causative and therapeutic. The patient’s past surgical history may similarly yield insights. A sexual history confirms the diagnosis, and should ascertain the severity, onset, and duration of the problem, as well as the presence of concomitant medical or psychosocial factors. It is necessary to determine whether the presenting complaint (eg, ED, premature ejaculation) is the primary sexual problem or if some other aspects of the sexual response cycle (desire, ejaculation, orgasm) are involved (Rosen, 2004). Psychosocial assessment of past and present partner relationships is essential given the interpersonal context of sexual problems. Sexual dysfunction may affect the patient’s self-esteem and coping ability, as well as social relationships and occupational performance.

#### Physical & Laboratory Examination

A focused physical examination is performed on each patient, assessing the genitourinary, endocrine, neurologic, and vascular systems, and includes a complete genital examination including digital rectal, and measurement of blood pressure and heart rate. Examination may yield a diagnosis of Peyronie’s disease, prostatic enlargement, malignancy, or evidence of hypogonadism (decreased testosterone, altered secondary sex characteristics).

Recommended tests include fasting glucose, lipid profile, and morning testosterone (calculated free testosterone is more reliable to establish hypogonadism). Diabetics should have hemoglobin A1C measured and additional hormonal testing (prolactin, follicle-stimulating hormone [FSH], and luteinizing hormone [LH]) is required when low testosterone levels are noted or for clinical suspicion of abnormality. Optional tests, including prostate-specific antigen (PSA), thyroid-stimulating hormone (TSH), complete blood count, and creatinine, must be tailored to the patient’s complaints and risk factors (Wespes et al, 2006).
**Self-Reported Questionnaires & Laboratory Investigations**

The most commonly used validated questionnaires are the 15-item International Index of Erectile Function (IIEF) or an abridged 5-item version (IIEF-5) more suited for office use. Sexual domain functions measured by the IIEF include erectile function, orgasmic function, sexual desire, intercourse satisfaction, and overall satisfaction. These tools are useful to determine baseline erectile function and to assess the impact of a specific treatment modality. ED severity is classified into 5 categories based on the IIEF-5: severe (5–7), moderate (8–11), mild to moderate (12–16), mild (17–21), no ED (22–25) (Rosen, 2004).

The physician should review the findings, inquire about the goals and preferences of the man (and his partner), and discuss further diagnostic and therapeutic options such that his (or their) participation in the decision-making process is well informed (Burnett, 2006). For a minority of patients, referral for further testing and/or assessment may be appropriate. Indications for referral include patient request, treatment failure, complex gonadal or other endocrine disorders, neurologic deficits suggestive of brain or spinal cord disease, deep-seated psychologic or psychiatric problems, and active cardiovascular disease. Patients deemed to be at intermediate or higher cardiac risk (cardiac status uncertain, moderate to severe symptoms, or unable to perform exercise of modest intensity) should be assessed by a cardiologist/internist and sexual activity deferred until cardiac condition is stabilized or resumption of sexual activity is deemed safe by the consultant. Low-risk patients, those able to perform exercise of modest intensity (6 or more metabolic equivalents [METS]) without symptoms, do not generally require cardiologic assessment (Jackson et al, 2006).

**Follow-up Strategy**

Regardless of the treatment regimen chosen, follow-up is essential to ensure optimal outcomes. Monitoring adverse events, assessing satisfaction or failure with a given treatment, identifying a partner’s sexual dysfunction, and assessing overall health and psychosocial function are key considerations. In patients who do not respond to first-line therapy (oral PDE-5 inhibitor therapy), consideration should also be given to whether an alteration in dose or treatment might be of value, as most second- and third-line options have demonstrated reasonable response and satisfaction rates in controlled studies (Lue et al, 2004).

**ADVANCED TESTING FOR ERECTILE DYSFUNCTION**

For patients with more complex problems, including penile deformity, history of pelvic or perineal trauma, ED of unknown etiology, cases requiring vascular or neurosurgical intervention, complicated endocrinopathy, complicated psychiatric disorder, complex relationship problems, medicolegal concerns or at the patient’s request, a variety of vascular and neurologic diagnostic tests are available to identify the cause of ED or plan treatment (Table 38–4).

### A. TESTS FOR PENILE VASCULAR FUNCTION

The goal of vascular evaluation is to identify and evaluate arterial and veno-occlusive dysfunction. The most commonly utilized tests include combined injection and stimulation (CIS), duplex ultrasound, dynamic infusion cavernosometry and cavernosography (DICC), and selective penile angiography.

1. **CIS (Combined intracavernous injection and stimulation) test**—This most commonly performed diagnostic procedure for ED, office pharmacotesting, consists of an ICI, visual or manual sexual stimulation, and a rating of the subsequent erection. Before injection, the patient should be informed about the purpose, alternatives, risks, and benefits of the test. Neurogenic and hormonal influences are bypassed as the vascular status of the penis is assessed directly and objectively. The most commonly used vasodilator is 10 mcg of alprostadil or 0.3 mL of a mixture of papaverine and phentolamine. A rigid erection lasting >10 minutes is indicative of normal venous function. However, the same conclusion cannot be made for arterial function as some men with mild arterial insufficiency can also have the same response.

**Table 38–4. Tests Suggested for Various Treatment Options.***

<table>
<thead>
<tr>
<th>Treatment Options</th>
<th>Tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Oral medication, transurethral therapy, or vacuum constriction device</td>
<td>No further testing</td>
</tr>
<tr>
<td>2. Intracavernous injection therapy</td>
<td>CIS test</td>
</tr>
<tr>
<td>3. Penile prosthesis</td>
<td>CIS test or NPT test or duplex scanning</td>
</tr>
<tr>
<td>4. Venous surgery</td>
<td>CIS test</td>
</tr>
<tr>
<td></td>
<td>Duplex scanning or cavernous arterial occlusion pressure test</td>
</tr>
<tr>
<td></td>
<td>Cavernosometry and cavernosography</td>
</tr>
<tr>
<td>5. Arterial surgery (or combined arterial and venous surgery)</td>
<td>CIS test</td>
</tr>
<tr>
<td></td>
<td>Duplex scanning or cavernous arterial occlusion pressure test</td>
</tr>
<tr>
<td></td>
<td>Cavernosometry and cavernosography</td>
</tr>
<tr>
<td></td>
<td>Pharmacologic arteriography</td>
</tr>
</tbody>
</table>

*CIS, combined injection and stimulation; NPT, nocturnal penile tumescence.

*Regardless of desired treatment, all patients must undergo history, physical examination, and basic laboratory testing.
2. **Duplex ultrasonography (gray scale or color)**—Color duplex ultrasound, the most reliable and least invasive evidence-based assessment of ED, is utilized when further vascular diagnostic testing is indicated. It consists of an intracavernous pharmaco-test and measurement of blood flow by duplex Doppler ultrasound, (Golijanin et al, 2006; Figure 38–4). High-resolution (7–10 MHz) real-time ultrasonography and color pulsed Doppler enables the ultrasonographer to visualize the dorsal and cavernous arteries selectively, perform dynamic blood flow analysis, and is the best tool available for the diagnosis of high-flow priapism and localization of a ruptured artery. Normal arterial response is a peak flow velocity measured at the base of penis of >30 cm/sec, a sharp upstroke of the waveform and absence of diastolic flow after sexual stimulation. Ultrasound can also be used to detect penile abnormalities such as Peyronie’s plaque, calcifications, thickened vessel walls, and intracavernous fibrosis. When the Doppler waveform exhibits high systolic flow (>30 cm/sec peak systolic velocity [PSV]) and persistent end-diastolic flow velocity (EDV) >5 cm/sec accompanied by quick detumescence after self-stimulation, the patient is considered to have venogenic impotence. The parameters useful in diagnosing venous leakage include a diastolic venous flow velocity of >5 cm/sec and/or a resistive index (RI) of <0.75. RI = peak systolic velocity (PSV)–EDV / PSV.

The Pulsed Doppler Study is performed using the Midus portable unit, which is specifically designed for use in the urologist’s office. Penile blood flow is evaluated in a fashion similar to duplex ultrasound by recording the waveform of the cavernous arteries without providing a real-time ultrasound image (UroMetrics, Anoka, MN).

3. **Cavernosometry and cavernosography**—Pharmacologic cavernosometry involves ICI of a potent vasodilator combination (papaverine + phentolamine + alprostadil) followed by saline infusion and simultaneous monitoring of ICP to assess the penile outflow system. In men with normal venous function, the maintenance flow should be <10 mL/min at an ICP of 100 mm Hg and the rate ICP drops after infusion ceases should be <50 mm Hg in 30 seconds. Veno-occlusive dysfunction is indicated by either the inability to increase ICP to the level of the mean systolic blood pressure with saline infusion or a rapid drop of ICP after cessation of infusion.

Cavernosography is used to visualize the site of venous leakage. Following the induction of an artificial erection by ICI of vasodilators (activation of the veno-occlusive mechanism), diluted radiocontrast solution is infused into the corpora cavernosa (Mulhall et al, 2004). Minimal or no contrast is seen outside the corpora cavernosa with normal veno-occlusive function. In patients with venous leakage of congenital or traumatic origin, the leakage is seen most often in the crura or at the site of injury respectively (Figure 38–5). Leakage sites to the glans, corpus spongiosum, superficial dorsal veins, and cavernous and crural veins can occur.
be detected. In the majority of patients, more than one site is visualized. The typical finding in men with intrinsic disease of the corpus cavernosum or the tunica albuginea is a diffuse leakage through all penile venous channels.

Dicc is reserved for young men who might be candidates for penile vascular operations, specifically those with a history of pelvic trauma or life-long ED (primary ED) (Rahman et al, 2005). DICC is invasive, requiring 2 needles to remain in the penis for saline infusion and pressure recording. Cavernosography is performed after cavernometry and should reveal opacification of the corpora cavernosa but absent or minimal visualization of venous structures or corpus spongiosum in men with normal veno-occlusive function (Montorsi, 2005).

4. Arteriography—Arteriography is reserved for the evaluation of the complex patient for whom revascularization surgery is contemplated; indications include the young patient with ED secondary to a traumatic arterial disruption or a patient with a history of perineal compression injury. The study is performed by ICI of a vasodilating agent (papaverine, papaverine + phentolamine, or alprostadil) followed by selective cannulation of the internal pudendal artery and injection of a diluted contrast solution of low osmolarity. The anatomy and radiographic appearance of the cavernous arteries is then evaluated (Figure 38–6), as are the iliac vessels, dorsal penile arteries, and the size and length of the inferior epigastric arteries. Variation of intrapenile arterial anatomy often confounds interpretation as “normal” paired common penile arteries are documented in only 50% of normal potent volunteers.

5. Cavernous arterial occlusion pressure—Cavernous arterial occlusion pressure, a variation of penile blood pressure measurement, involves infusing saline solution into the corpora after ICI of vasodilators to raise the ICP above the systolic blood pressure. A pencil Doppler transducer is then applied to the side of the penile base, saline infusion is stopped, and the intracavernous pressure is allowed to fall. The pressure at which the cavernous arterial flow becomes detectable is defined as the cavernous artery systolic occlusion pressure (CASOP). A gradient between the cavernous and brachial artery pressures of <35 mm Hg and equal pressure between the right and left cavernous arteries are defined as normal. Results correlate well with arteriography and PSV obtained by high-resolution duplex Doppler ultrasound. However, this more invasive procedure is prone to psychological inhibition and is not feasible if the ICP cannot be raised above the systolic blood pressure (eg, in patients with severe venous leakage).

B. NEUROLOGIC TESTS

Physiologically, 3 types of erections can occur: nocturnal, psychogenic, and reflexogenic. In a broader sense, neurologic testing should assess peripheral, spinal, and supraspinal centers, as well as somatic and autonomic pathways associated with all types of erection and sexual arousal.
However, the effect of a neurologic insult on penile erection is a complicated phenomenon and with few exceptions, neurologic testing rarely changes management. Testing is recommended for specific research protocols or medicolegal investigations, including cases of trauma or surgical complications, or can be used in selected cases to (1) uncover reversible neurologic disease such as dorsal nerve neuropathy secondary to long-distance bicycling, (2) assess the extent of neurologic deficit from a known neurologic disease such as diabetes mellitus or pelvic injury, and (3) determine whether a referral to a neurologist is necessary (eg, workup for possible spinal cord tumor). Based on best-available evidence, neurologic tests lack adequate sensitivity and reliability for routine clinical diagnosis.

Somatic nerves are evaluated by testing nerve conduction velocities and evoked potentials, and these tests have well-known reproducibility and validity. Autonomic function tests are less reliable because they simultaneously measure a chain of events or reactions involving receptors, small fibers, and target organs. The complex interactions among central and peripheral sympathetic and parasympathetic nerve systems make autonomic testing difficult. Currently available tests are not well standardized, and lack validity, reproducibility, and comparability.

1. Biothesiometry—This test is designed to measure the sensory perception threshold to various amplitudes of vibratory stimulation produced by a hand-held electromagnetic device (biothesiometer) placed on the pulp of the index fingers, both sides of the penile shaft, and the glans penis.

2. Bulbocavernosus reflex latency—This test is performed by placing 2 stimulating ring electrodes around the penis, one near the corona and the other 3 cm proximal. Concentric needle electrodes are placed in the right and left bulbocavernous muscles to record the response to square-wave impulses delivered via a direct current stimulator. The latency period for each stimulus response is measured from the onset of the stimulus to the beginning of the response. An abnormal bulbocavernosus reflex (BCR) latency time, defined as a value greater than three standard deviations above the mean (30–40 msec), carries a high probability of neuropathology.

3. Penile thermal sensory testing—Thermal threshold measurements quantify conductance of small sensory nerve fibers, which can indirectly reflect autonomic disturbances in the context of diffuse neuropathies, such as diabetic polyneuropathy. Penile thermal sensory testing correlates strongly with the clinical evaluation of erectile function, and is a promising tool for the diagnosis of neurogenic impotence (Bleustein et al, 2003).

C. Nocturnal Penile Tumescence Test

Nocturnal erections, 80% of which occur during REM sleep, occur in healthy males of all ages and are relatively free of psychologically mediated effects. The average man has 3–5 episodes of nocturnal penile tumescence (NPT) each night, with episodes ranging from 30 to 60 minutes. Total NPT time declines with increasing age. Classically, NPT has been measured by several methods including the stamp test, snap or strain gauges, and sleep lab NPTR. Contemporary NPT testing is performed with simpler outpatient devices such as Rigiscan NPTR (Rigiscan, Timm Medical Technologies). These newer devices electronically record the number, duration, rigidity, and circumference of penile erections; although sleep lab NPTR concurrently records nasal air flow, oxygen saturation, and electroencephalographic, -myographic, and -oculographic data to document REM sleep, hypoxia, and/or abnormal limb movement, it has been replaced by outpatient Rigiscan due to prohibitive expense.

NPT was originally designed to differentiate psychogenic from organic ED, as a full erection indicates a
functionally intact neurovascular axis. In the United States, it is now mostly used in medicolegal or complicated cases, as well as for research purposes (testing of erectogenic agents or functional outcomes following pelvic surgery) (Bannowsky et al, 2006).

D. PSYCHOLOGICAL EVALUATION

Psychogenic ED is defined as the persistent inability to achieve or maintain an erection satisfactory for sexual performance, which is due predominantly or exclusively to psychological or interpersonal factors. Clinical subtypes of psychogenic ED include generalized versus situational and lifelong (primary) versus acquired (secondary, including substance abuse or major psychiatric illness). The psychological and interpersonal dimensions of sexual function and dysfunction are complex; therefore, a skillful diagnostic interview is the mainstay of a good psychological evaluation. A suggestive history of psychogenic ED includes sudden onset, selective dysfunction (eg, rigid erection with one partner and poor erection with others, or normal erection during masturbation or fantasy but not during intercourse), and a normal pattern of nocturnal erections coupled with an abnormal pattern during waking hours. This is often associated with anxiety, guilt, fear, emotional stress, religious or parental inhibition. If the medical and sexual histories suggest a combination of organic and psychological risk factors, these patients should be diagnosed as having mixed organic/psychogenic ED; successful treatment must address both components.

Although psychological consultation is not indicated for most patients, it is very useful in evaluating and treating men with deep-seated psychological problems. Three groups of psychometric instruments are commonly used: (1) personality questionnaires, (2) the depression inventory, and (3) questionnaires for sexual dysfunction and relationship factors. The Minnesota Multiphasic Personality Inventory (MMPI)-2 is a valuable tool for evaluating the patient’s personality and its relevance to sexual dysfunction. The self-reported Beck Depression Inventory is a validated test for which a score exceeding 18 is considered indicative of significant clinical depression. Overall quality of relationships can be assessed using the Short Marital Adjustment Inventory (married couples) and the Dyadic Adjustment Inventory (unmarried individuals) (Mallis et al, 2005).

NONSURGICAL TREATMENT OF ERECTILE DYSFUNCTION

Advances in pharmacologic therapy for ED have resulted in greater numbers of patients seeking primary and specialty care for sexual concerns. Oral phosphodiesterase type-5 inhibitors have emerged as the preferred first-line treatment of ED worldwide due to their efficacy, ease-of-use, and patient safety. Prior to their release, efficacious noninvasive therapies for ED were unavailable and patient options were limited to nonspecific modalities including lifestyle modification, medication changes, hormonal or psychotherapy, and vacuum devices, intraurethral agents, intracavernosal injection therapy, or penile prosthesis insertion. Although it remains one of the most effective treatments for all types of ED, nonsurgical approaches have replaced prosthetic surgery as the preferred choice of management over the last decade. In most cases, nonspecific therapies appear to be more effective for ED, however, the patient should also be aware of specific therapies so that an informed treatment decision can be made.

Lifestyle Changes

The beneficial effect of lifestyle change (total body weight loss of 10% or more by reducing caloric intake and increasing physical activity) was demonstrated in a randomized, single-blind trial of 110 obese men (body mass index ≥30) age 35–55 years, without diabetes, hypertension, or hyperlipidemia, but with ED (a score of 21 or less on the IIEF). The mean IIEF score improved in the intervention group (from 13.9±4.0 to 17±5; P <0.001), but remained stable in the control group who were given general information about healthy food choices and exercise (Esposito et al, 2004).

It is well known that ED is intimately related to the atherosclerotic coronary and peripheral vascular diseases, as well as the metabolic syndrome, characterized by central obesity, abnormal lipids, insulin dysregulation, and borderline hypertension. Regular exercise, a healthy diet, smoking cessation, and limiting use of alcohol can reduce the risk of ED or improve underlying dysfunction. A Mediterranean-style diet (fruits, vegetables, nuts, whole grain, olive oil, and decreased saturated lipids) has recently been shown to improve endothelial function scores and inflammatory markers (C-reactive protein) when compared to men on a control diet (Esposito et al, 2006). Obesity and smoking have been prospectively identified as risk factors for ED, while physical activity is inversely associated with the development of ED; reducing the risk of ED may serve as more concrete motivation for men to engage in health-promoting behaviors (Bacon et al, 2006).

Perineal compression on penile arteries from long-distance cycling may also represent a modifiable risk factor for ED. Changing the bicycle seat or riding practices will often improve erectile function if penile vascular compromise is identified; specific strategies include replacing a protruding nose with a noseless seat, changing posture to a more upright/reclining position, using a gel saddle, and tilting the seat downwards (Huang, Munarriz, Goldstein, 2005).

Changing Medications

When a patient complains of sexual dysfunction after taking a particular medication, it is important to determine
Psychosexual Therapy

Therapies such as a phosphodiesterase-5 inhibitors or ICIs may provide more rapid relief for patients with ED compared to a prolonged course of psychosexual therapy. However, in patients with evidence of psychological problems, referral to a psychologist or sex therapist is highly recommended because the elimination of a specific underlying cause may result in a cure. Recent approaches to sex therapy have included cognitive-behavioral interventions focused on challenging or correcting maladaptive cognitions, behavioral techniques (desensitization and assertiveness exercises), exploration of past developmental experiences on present behavior, and couples’ therapy. Moreover, in some patients with mixed psychogenic and organic ED, psychosexual therapy may help relieve the anxiety and remove unrealistic expectations associated with medical or surgical therapy.

Hormonal Therapy

Referral to an endocrinologist is recommended for patients with thyroid, adrenal, pituitary or hypothalamic dysfunction. Aging men with ED may show a variety of symptoms when hypogonadism coexists: low libido, depression, decreased intellectual abilities, lean body mass, bone mineral density, or skin turgor, changes in body hair distribution, changes in sleep patterns, and increased visceral fat. In the patient with documented hypogonadism and ED, it is reasonable to initiate androgen therapy; for hypogonadal patients unresponsive to PDE-5 inhibitor therapy alone, the addition of testosterone may enhance the treatment effect and improve erectile function. 

Parenteral depo-preparations of testosterone (T), such as testosterone cypionate and enanthate, are the least expensive form of androgen supplementation and are effective in restoring serum levels to normal. A normal circadian rhythm is not replicated by these agents. Rather, they are administered through deep intramuscular injection (200–250 mg q 2 weeks) and result in supraphysiologic levels of T for 72 hours with steady exponential decline to subphysiologic levels in 10–12 days; the initial supraphysiologic “rush” may be disconcerting to some patients, but others enjoy an improved sense of well-being, aggression, and libido.

Transdermal delivery more closely simulates normal circadian levels of testosterone if patients apply a patch in the morning, as the higher initial absorption mimics normal diurnal variations. Several U.S. Food and Drug Administration (FDA)-approved preparations are available in the United States. Testoderm TTS, which has largely replaced the Testoderm scrotal patch, is convenient to use as it is applied daily to the arm, back, or upper buttocks as a 5-mg patch. Another product, Androderm, comes as a patch that delivers 2.5 or 5 mg T per day. The most common adverse reactions to both are itching, chronic skin irritation, and allergic contact dermatitis. Patients should alternate application sites and avoid sun-exposed areas. Local application of cortisone cream may relieve irritated skin. AndroGel is a 1% gel pack (containing 50 mg, 75 mg, or 100 mg of T), which is also applied daily in the morning to clean, dry skin over the shoulders, upper arms, or abdomen. Hands should be washed thoroughly after application, as skin contact can transmit testosterone. Testim topical gel also contains 1% T, providing continuous transdermal delivery for 24 hours after a single application to intact, clean, dry skin of the shoulders and upper arms. One 5-g tube of Testim contains 50 mg of testosterone.

When taken orally, testosterone preparations are largely rendered metabolically inactive during “first pass” circulation through the liver. The large dosages (exceeding 200 mg/day) required to achieve therapeutic levels can be hepatotoxic, leading to hepatitis, cholestatic jaundice, hepatomas, hemorrhagic liver cysts, and hepatocarcinoma. Although unavailable in the United States, the only orally active and safe form of T is Andriol (testosterone-undecenoate in oleic acid [TU]), which owing to its lipophilic side chain, is partly taken up by the lymph and partially escapes hepatic inactivation. The maximal plasma concentration is generally observed within 2–3 hours, but after 6–8 hours levels have returned to pretreatment values. A dosage of 40 mg three times a day generally provides adequate androgen replacement, yielding T levels within the (low) normal range, whereas DHT levels are moderately increased (2–4 nMol/L). Absorption varies with food consumption and the dose required should be based upon plasma levels and clinical effects. FDA-approved delivery systems recently introduced to clinical use include Testo- pel subcutaneous pellets (dosage of 2–6 pellets [150–450 mg T]) and Striant, a tablet-like mucoadhesive buccal system used to deliver 30 mg of T twice daily.

In patients with hyperprolactinemia with or without hypogonadism, androgen therapy does not improve sexual function while taking antidepressants may benefit from watchful waiting, substitution (bupropion, nefazodone, buspirone, mirtazapine), drug holidays, selective serotonin-reuptake inhibitor (SSRI) dosage reduction, and/or PDE-5 inhibitors.
function. Treatment should first be aimed at eliminating the offending drugs, such as estrogens, sedatives, neuroleptics, or morphine. Bromocriptine, a dopamine agonist that lowers prolactin levels and restores T to normal, is used to reduce the size of prolactin-secreting adenomas. Surgical ablation may occasionally be required if the bromocriptine response is unsatisfactory or if visual field changes are secondary to optic nerve compression.

Potential Adverse Effects of Testosterone Replacement

Testosterone replacement is clearly the treatment of choice for young hypogonadal men without contraindications. However, the potential risks of androgen therapy may outweigh the benefits for some patients. Supraphysiologic levels of T suppress LH and FSH production and can lead to infertility, breast tenderness, and gynecomastia. Erythrocytosis is the most common laboratory alteration noted with long-term therapy: increases in red cell mass, thromboxane A2, and platelet aggregation may increase cardiovascular risk. Androgens may also induce or worsen sleep apnea. Long-term therapy requires a commitment from the patient and the specialist for continued follow-up, as outlined below.

Regarding prostate safety, a number of studies in the literature suggest that androgen replacement does not induce prostate cancer in men with normal prostates, and placebo-controlled studies show little difference in prostate volume, PSA, and obstructive symptoms. No increased risk of prostate cancer has been noted in (1) clinical trials of T supplementation, (2) longitudinal population-based studies, or (3) in a high-risk population of hypogonadal men receiving T treatment (Morgentaler, 2006). Although the fear of exacerbating an occult cancer of the prostate remains a concern, many older hypogonadal patients whose libido and erectile function can be restored by T therapy likely should not be denied this treatment option. When a patient desires T replacement, we routinely perform a digital rectal examination and obtain a serum PSA level. When in doubt, ultrasound-guided biopsy is performed before T therapy is given. The presence of prostate or breast cancer is an absolute contraindication to androgen supplementation. Patients are followed every 6 months with a rectal examination and serum PSA indefinitely while on therapy. Laboratory surveillance should also include: hemoglobin/hematocrit levels, liver function tests, cholesterol and lipid profile. The efficacy of supplementation is reasonably determined by clinical response rather than blood levels of testosterone.

Oral Pharmacologic Therapy

A. PHOSPHODIESTERASE (PDE) INHIBITORS

Sildenafil (Viagra), vardenafil (Levitra), and tadalafil (Cialis), the 3 selective phosphodiesterase-5 inhibitors (PDE-5Is) currently approved for clinical use, have become the preferred first-line therapy for most men with ED due to their efficacy, safety, and ease-of-use. All are highly effective in enhancing erectile function across a wide range of outcome measures, causes of ED, patient subgroups, and regional populations. Because of differences in trial designs, comparisons among these agents across published studies are not feasible. However, the 3 PDE-5Is appear to have equivalent efficacy in the treatment of ED, are generally well tolerated, and have similar contraindications and warnings (Carson and Lue, 2005).

1. Mechanism of action—Sexual stimulation results in the release of NO from nerve endings and vascular endothelial cells in the penis, which then diffuses into vascular and cavernous smooth muscle cells of the corpus cavernosum. Stimulation of guanylyl cyclase elevates levels of cGMP, lowering cytoplasmic calcium and resulting in smooth muscle relaxation and subsequent penile erection. PDE-5Is potentiate rather than trigger the physiologic erectile response to NO and amplify the NO-cyclic cGMP pathway through competitive inhibition of second-messenger cGMP enzymatic degradation (by phosphodiesterase-5), regardless of the underlying etiology of ED. Without sexual stimulation and resultant NO release however, these inhibitors are ineffective; therefore, a positive treatment effect would not be expected in patients with bilateral non-nerve-sparing pelvic surgery.

2. Clinical efficacy—The clinical efficacy and safety of sildenafil, vardenafil, and tadalafil has been evaluated in many placebo-controlled, double-blind trials and open label studies (Carson and Lue, 2005; Brock et al, 2002; Porst et al, 2003; Goldstein et al, 1998). Treatment effect was principally assessed by items 3 and 4 of the IIEF-15 questionnaire (ability to attain and maintain erection). For sildenafil, improvements in erectile function were reported by 56–84% of subjects taking 25–100 mg of sildenafil versus 25% in the placebo group. Little ED benefit and considerably more side effects were noted above 100 mg. An overall beneficial treatment effect was seen in 70–80% of patients; for specific etiologies, sildenafil was effective in 70% of hypertensive patients, 57% of diabetics, 43% of radical prostatectomy patients, and 80% of spinal cord injury patients. The efficacy of vardenafil was similarly evaluated: 73% of patients randomized to 10 mg and 81% randomized to a 20-mg dose reported improved erections. Mean IIEF scores increased from 12.8 at baseline to 21 at week 12 of treatment (compared to 13.6 to 15.0 for placebo). Similarly, integrated phase III studies of tadalafil involving 1112 patients demonstrated IIEF erectile domain scores of 24 in men receiving a 20-mg dose versus 15 for placebo. More than 70% of intercourse attempts were successfully completed from >30 minutes to 36 hours after dosing. For difficult to treat groups, including diabetics, severe ED, and postradical prostatectomy, the 3 PDE-5Is represent effective...
therapies for most men. Ongoing, longer term, randomized-control trials will better define the role (including early or prophylactic treatment after nerve-sparing radical prostatectomy) and efficacy of these agents, including efficacy and safety data for cohorts previously not studied.

3. Time of onset—The onset of activity, in reports with similar methods, is 14 minutes with sildenafil, 10 minutes with vardenafil, and 16 minutes with tadalafil. However, success rates after 20 minutes are much less than after 1 hour; therefore, if patients do not experience a rapid beneficial effect, they should be advised to delay sexual intercourse for one (sildenafil or vardenafil) or 2 hours (tadalafil)—when serum concentrations have peaked. High-fat meal intake has been shown to delay absorption of vardenafil and sildenafil; this effect is not seen with tadalafil (Carson and Lue, 2005).

4. Period of efficacy—Tadalafil therapy has a broader window of clinical responsiveness than either sildenafil or vardenafil because of its longer half-life (17.5 versus 4–5 hours for sildenafil or vardenafil). Tadalafil enhances erectile function in men with ED for up to 36 hours and may mean less planning and pressure to have sexual intercourse according to a schedule, which can result in greater patient and/or partner convenience.

5. Adverse events—The biochemical selectivity of an inhibitor is a key factor in determining its side effect profile. For PDE-5Is, selectivity is usually expressed in terms of potency (IC50) to inhibit PDE-5 as opposed to inhibiting nontarget PDEs (or other proteins). Eleven distinct families (PDE-1–PDE-11) are known to have or are implicated in a broad range of cellular functions. PDE-5 is present in high concentrations in the smooth muscle of the penile corpora cavernosa. Sildenafil and vardenafil cross-react slightly with PDE-6, that is, the IC50s for PDE-5 are only 4- to 10-fold lower than those for PDE-6. This may explain why some patients using sildenafil complain of visual disturbances. Tadalafil minimally cross-reacts with PDE-11, but the consequences of this effect are unknown (Weeks et al, 2005). Most side effects associated with PDE-5Is result from inhibition of PDE-5 in other tissue or organs. In randomized controlled trials, flushing (10%) and visual side effects were more common in patients receiving sildenafil or vardenafil, and back pain/myalgia (1–4%) was more common for tadalafil users. These events were mostly mild, abated with time (within 2–4 weeks), and prompted treatment discontinuation in only a small number of patients (Brock et al, 2002; Porst et al, 2003; Goldstein et al, 1998). Except for visual disturbances, the other reported side effects of PDE-5Is (headaches 15%, flushing, rhinitis 5–10% for vardenafil and sildenafil, slight lowering of blood pressure, dyspepsia, etc.) are likely caused by PDE-5 inhibition in vascular or gastrointestinal smooth muscle.

ED is common among men who have atherosclerotic coronary artery disease. With regard to overall cardiac safety, controlled and postmarketing studies of the 3 FDA-approved PDE-5Is demonstrated no increase in MI or death rates in double-blind, placebo-controlled, or open-label studies when compared with expected study population rates. Patients who develop angina during sexual activity with a PDE-5I should discontinue sexual activity, relax for 5–10 minutes and if pain persists, seek emergency care and inform emergency medical personnel that a PDE-5I was taken. Patients who have an acute MI after a PDE-5I may be given standard therapies (except for organic nitrates). Nitroglycerin should not be given with 24 hours of sildenafil or vardenafil use, or 48 hours for tadalafil. Patients who develop hypotension after organic nitrates and PDE-5Is should be placed in the Trendelenburg position and given IV fluids, with alpha-agonists (such as phenylephrine) added as needed. In patients with refractory hypotension, intra-aortic balloon counterpulsation should be administered, as suggested by the American College of Cardiology/American Heart Association (ACC/AHA) guidelines. At present, there is no pharmacologic antidote to the PDE-5 inhibitor/nitrate interaction (Kostis et al, 2005).

PDE-5Is have a minimal effect on QTc interval (Morganroth et al, 2004). However, only vardenafil is not recommended for patients who take type-1A antiarrhythmics (such as quinidine or procainamide), type-3 antiarrhythmics (such as sotalol or amiodarone), or with congenital prolonged QT syndrome.

Finally, there have been recent reports of the development of nonarteritic anterior ischemic optic neuropathy (NAION) in men using PDE-5Is. Epidemiologically, NAION is the second most common acquired optic neuropathy in men aged 50 years and older. Risk factors for NAION, cardiovascular disease, and ED are shared and include age, dyslipidemia, diabetes, hypertension, and cigarette smoking. To date, <50 cases of NAION associated with PDE-5I use have been reported to the FDA: 38 sildenafil, 4 tadalafil, and 1 vardenafil. Given the large number of men safely using these agents and a limited number of events, it is not possible to determine whether NAION is directly linked to the use of PDE-5Is, underlying cardiovascular risk factors, ocular anatomical defects, a combination of these variables, or as yet unidentified factors (Laties and Sharlip, 2006). Reasonable and informed consent regarding the possible but low risk of NAION with the use of PDE-5Is is recommended. PDE-5Is should not be prescribed for patients with prior episode(s) of NAION. Loss or decreased vision, whether painful or painless, demands urgent patient assessment and immediate cessation of PDE-5 inhibitor use.

6. Warnings and drug interaction—PDE-5Is are contraindicated for patients using nitrates, as a precipitous and potentially life-threatening hypotensive episode may occur with concurrent use. Package inserts also warn against use in patients with severe cardiovascular diseases, left ventric-
ular outflow obstruction, and patients not studied in clinical trials including those with hereditary degenerative retinal disorders and tendency to develop priapism (sickle cell anemia, leukemia, etc) (United States prescribing information of Viagra, Cialis and Levitra, July 2006). PDE-5Is are either not recommended or are to be used with caution in men with unstable angina, cardiac failure, recent MI, uncontrolled or life-threatening arrhythmia, or poorly controlled blood pressure (resting BP <90/50 mm Hg or >170/100–110 mm Hg). Certain drugs, such as ketoconazole anditraconazole, and protease inhibitors, such as ritonavir, can impair the metabolic breakdown of PDE-5Is by blocking the CYP3A4 pathway and may require a dose reduction (cytochrome P450 isoenzyme). On the other hand, agents such as rifampin may induce CYP3A4, enhancing the breakdown of inhibitors and requiring higher PDE-5I doses. Older age (>65 years), hepatic impairment, and severe renal insufficiency are associated with increased plasma levels of PDE-5Is and therefore the dosage should be reduced in these patients. Caution is also advised when an alpha-blocker and a PDE-5 inhibitor are given together, as interaction can lead to excessive vasodilation and hypotension.

7. Starting doses—The recommended starting doses are 50 mg for sildenafil and 10 mg for vardenafil and tadalafil. The dose may be increased to 100 mg (sildenafil) or 20 mg (vardenafil and tadalafil) or decreased to 25 mg or 5 mg, respectively, based on individual efficacy and tolerability. Patients should also be counseled to trial a PDE-5I several times before declaring it “ineffective”; for instance, the cumulative probability of success with sildenafil increases with the first nine to ten attempts, after which it stabilizes (McCullough et al, 2002).

B. CENTRALLY ACTING ORAL AGENTS

Apomorphine is a proerectile D1/D2 dopamine receptor antagonist. Uprima, a sublingual form designed for buccal absorption, has been approved for ED in Europe (but not in the United States) and reported to induce erection in 67% of psychogenically impotent patients. It is not an opiate and is chemically unrelated to morphine; this agent acts in the brain within the paraventricular nucleus, which functions as the sexual drive center in mammals. Sexual arousal is necessary to enhance its effects and efficacy is lost if the tablet is swallowed. Apomorphine has a rapid onset of action, with a window of sexual opportunity of approximately 2 hours from ingestion. Maximal plasma concentrations are reached in 50 minutes. In a double-blind placebo-controlled study of 2- and 4-mg dosages, erections firm enough for intercourse were reported by 45% and 55% of patients, respectively, with placebo responses of 35% and 36%. Self-assessment of success was 47% and 59.9% at these doses. Adverse events included nausea 16.9%, dizziness 8.3%, sweating 5%, somnolence 5.8%, yawning 7.9%, and emesis 3.7%. At the highest recommended dosage, syncope occurred in 0.6% of patients and was accompanied by a clear prodrome suggestive of a vasovagal event: nausea, vomiting, sweating, dizziness and light-headedness (Heaton, 2001). There were no documented food/drug interactions in clinical trials (with the exception of ethanol) and specifically, no documented pharmacologic interactions for subjects using nitrate drugs.

Yohimbine is a centrally acting alpha-2-adrenergic antagonist which is not recommended for use in the treatment of ED by the 2005 American Urological Association (AUA) guidelines. Side effects include gastrointestinal intolerance, palpitations, headache, agitation, anxiety, and increased blood pressure (precautions are advised in men with cardiovascular disease). Trazodone is also not recommended; efficacy in pooled analyses was statistically equivalent to placebo. Side effects include drowsiness, nausea, emesis, blood pressure changes (both hypotension and hypertension are reported), urinary retention, and priapism (especially at therapeutic antidepressant levels) (Montague et al, 2005).

Transurethral Therapy

Alprostadil, a synthetic formulation of PGE-1, is the only FDA-approved pharmacologic agent approved for the management of ED via intracavernous and transurethral routes. After absorption from the urethra to the corpus spongiosum and then corpus cavernosum, alprostadil stimulates adenyly cyclase to increase intracellular levels of cAMP and lower levels of intracellular calcium, thereby relaxing arterial and trabecular smooth muscle. MUSE (medicated urethral system for erection) consists of a very small semisolid pellet (3 × 1 mm) administered into the distal urethra (3 cm) by a proprietary applicator (MUSE, VIVUS Inc, Menlo Park, CA, USA). Clinical studies showed that 66% of men responded to in-office trials; however, postmarketing studies have produced less successful results of about 50% (Mulhall et al, 2001). Penile rigidity can be enhanced by placing an elastic ring at the base of the penis (ACTIS, VIVUS Inc.) to mechanically assist veno-occlusion. Penile and/or scrotal pain or discomfort is a ubiquitous side effect of alprostadil-based therapies and is clearly dosage related and was reported in 33% of men in MUSE trials. Hypotension and syncope have been noted in 1–5.8% of patients, thus mandating initial trial dose administration in the office setting. Some female partners also report vaginal discomfort (about 10%) after ejaculation by a man using MUSE.

Intracavernous Injection

ICI of vasoactive drugs is considered the most effective nonsurgical therapy for ED. It remains the first-line therapy for select patients and a valuable treatment option for PDE-5I nonresponders or those that cannot tolerate side effects of oral agents as ICI treatment offers several poten-
tial advantages to the patient, including a rapid onset of action, reduced incidence of systemic complications and drug interactions compared to systemic treatments, and dependable efficacy for vascular and nonvascular (hormonal, neurogenic, or psychogenic) forms of ED. Men who have failed first-line oral pharmacotherapy constitute the largest group of ICI-treated patients, with a significant erectile response rate of >85% demonstrated among PDE-5I nonresponders indicating that progression to second-line injection therapy is appropriate. A list of drugs that have been used clinically is presented in Table 38–5, and the most commonly used agents and combinations are discussed below.

**A. PAPAVERINE**

Papaverine, an alkaloid isolated from the opium poppy, induces relaxation of cavernous smooth muscle and penile vessels via nonspecific inhibition of phosphodiesterase, elevates cAMP, and impairs calcium influx through blockage of voltage-dependent calcium channels. It is metabolized by the liver, with a plasma half-life of 1–2 hours. Monotherapy doses range from 15 to 60 mg. Advantages include low cost and stability at room temperature, while adverse effects include a higher incidence of priapism (up to 6%), corporal fibrosis (6–30%; thought to be associated with poor technique, minimal injection site compression time, >1 cc injection volume, pH 3–4), and occasional elevation of liver enzymes (Bella and Brock, 2004).

**B. PHENTOLAMINE METHYLATE (REGITINE)**

Monotherapy with phentolamine, an alpha-adrenergic antagonist with equal affinity for alpha-1 and alpha-2 receptors, has been disappointing as increases in corporal blood flow are not accompanied by a significant rise in intracorporal pressure. Systemic hypotension, reflex tachycardia, nasal congestion, and gastrointestinal upset are the most common systemic side effects. Plasma half-life is 30 minutes.

**C. ALPROSTADIL (PROSTAGLANDIN E1)**

Alprostadil causes smooth muscle relaxation, vasodilation, and inhibition of platelet aggregation through elevation of intracellular cAMP. It is metabolized by the enzyme prostaglandin 15-hydroxydehydrogenase, which has been shown to be active in human corpus cavernosum. After ICI, 96% of alprostadil is locally metabolized within 60 minutes and no change in peripheral blood levels is observed. Alprostadil have been approved by the FDA for intracavernous therapy as Caverject (Pharmacia & Upjohn, Peapack, NJ) and Edex (Schwarz Pharma, Milwaukee, WI). Cumulative data yield a success rate of 70–75% across ED etiologies, utilizing a median dose of 12–15 mg. Common adverse effects include pain at the injection site or during erection (11–15%), small hematoma or bruising, penile fibrosis (1–3%), and burning sensation at time of injection. Rates of priapism are low (1–3%) and systemic side effects are rare (Bella and Brock, 2004).

**D. DRUG COMBINATIONS**

The most commonly used drug combinations for ICI are bimix (papaverine/phentolamine) and trimix (papaverine/phentolamine/alprostadil) in various concentrations. Multiple series have demonstrated patient satisfaction rates of >75% and low rates of priapism or fibrosis. Side-effects are reduced, as smaller amounts of each agent are required, and the targeting of multiple pathways increases therapeutic efficacy. The main advantage of the bimix combination versus trimix is stability without refrigeration.

1. **Bimix**—The most commonly used formula contains 30 mg/mL of papaverine and 1 mg/mL of phentolamine. Efficacy and safety data were reviewed for a total 13,030 bimix injections in 160 men (Armstrong, Convery, Dinsmore, 1993). Erections sufficient for sexual intercourse were achieved in 72% of men, with etiology-specific response rates as follows: vasculogenic (48%), psychogenic (93%), neurogenic (92%), diabetic (68%), idiopathic (63%), traumatic (60%), alcohol-related (80%), and drug-related (75%). About half of men were continued with bimix ICI at 14 months. Priapism was rare, as 22 episodes occurred in 16 patients, and one patient developed corporal fibrosis.

2. **Trimix**—The 3-agent formulation is more potent than its bimix predecessor, demonstrating a level of patient satisfaction approaching 90%. In a randomized crossover study of 228 patients, trimix was compared to bimix or alprostadil alone (McMahon, 1991). Statistically, trimix was more effective in patients with severe arteriogenic or mild veno-occlusive dysfunction. The incidence of prolonged erection was lower when compared with bimix but not significantly different from alprostadil alone. Trimix

| Table 38–5. Intracavernous Vasodilator Injection Therapy.* |
|-----------------|-----------------|
| **Drug**        | **Test Dose**   | **Therapeutic Dose** |
| Papaverine      | 15–30 mg        | 15–60 mg            |
| Alprostadil     | 5–10 mcg        | 5–60 mcg            |
| Papaverine (30 mg) + Phentolamine (1 mg) | 0.1–0.3 mL | 0.2–1 mL |
| Papaverine (30 mg) + Phentolamine (1 mg) + Alprostadil (10 mcg) | 0.1–0.3 mL | 0.2–1 mL |

*Lower doses for management of neurogenic and psychogenic impotence.
combination has been shown to be as effective as alprostadil alone or more so (up to 89% response rate), but has a much lower incidence of painful erection. It is generally reserved for men in whom PGE1 or papaverine/phenolamine therapy has failed or who have significant penile pain with PGE1.

**E. PATIENT ACCEPTANCE AND DROP-OUT RATE**

In several studies, the percentage of patients accepting injection therapy when offered in the office ranges from 49% to 84%. However, patient discontinuation of therapy remains high, with most series describing long-term drop-out rates of 20–60%. Major determining factors include lack of patient motivation, cost, loss/disinterest of partner, and dissatisfaction with drug-induced erection.

**F. SERIOUS ADVERSE EFFECTS**

Priapism and fibrosis are the 2 more serious side effects associated with ICI therapy. Priapism occurred in 1.3% of 8090 patients in 48 studies with alprostadil, which is about five times lower than with papaverine or bimix (1.5 versus 10 versus 7%) (Linet and Ogrinc, 1996). Fibrosis can occur as a nodule, diffuse scarring, plaque or curvature. The incidence is about 10 times lower with alprostadil than with papaverine or bimix (1 versus 12 versus 9% of patients), although rates up to 12% have been reported. Incidence of both priapism and fibrosis with trimix therapy is similar to, or slightly less than, alprostadil.

**G. DOSAGE AND ADMINISTRATION**

Patients must have the first injection performed by medical personnel and receive appropriate training and education before home injection. An initial dose of 2.5 mcg for alprostadil is recommended. If the response is inadequate, increases in 2.5-mcg increments can be given until a full erection is achieved or a maximum of 60 mcg is reached. For drug combinations, treatment is initiated with a small dose (eg, 0.1 mL) and titrated according to erectile response. The goal is to achieve a full erection of <1-hour duration to avoid priapism. Compression of the needle puncture site for at least 5 minutes is recommended to prevent bleeding and fibrosis.

**H. TREATMENT OF PROLONGED ERECTION OR PRIAPISM**

The best treatment is prevention; prolonged erections, a potentially devastating adverse effect of ICI, are often secondary to rapid dose escalation by the patient, missed initial injection with second attempt, or use among neurogenic and/or young patients. Gradual and progressive dose increases by the patient will prevent most occurrences of priapism. It is imperative that the clinician prescribing intracavernous therapies emphasizes to the patient that priapism represents a urologic emergency, and any erection lasting >4 hours necessitates urgent medical evaluation.

Most episodes can be prevented by careful patient instruction at the outset of an ICI program. If an erection lasts >4 hours, the patient should either contact their physician or present to the emergency room for treatment. The best regimen for averting priapism is ICI of diluted phenylephrine 250–500 mcg every 3–5 minutes until detumescence. In patients with cardiovascular disease, monitoring of blood pressure and pulse is recommended (Montague et al, 2003).

**I. CONTRAINDICATIONS**

ICI is contraindicated in patients with sickle cell anemia, schizophrenia or a severe psychiatric disorder, and severe venous incompetence. For patients using an anticoagulant or aspirin, compressing the injection site for 7–10 minutes after injection is recommended. In patients with poor manual dexterity, the sexual partner can be instructed to perform the injection.

**Vacuum Constriction Device**

The vacuum constriction device consists of a plastic cylinder connected directly or by tubing to a vacuum-generating source (manual or battery-operated pump). Only devices containing a vacuum limiter should be used, as injury to the penis avoided by preventing extremely high negative pressures (Montague et al, 2005). After the penis is engorged, a constricting ring is applied to the base to maintain the erection. The ring may be uncomfortable or painful; to avoid injury, it should not be left in place for >30 minutes. The erection produced differs from a physiologic or ICI-induced erection as the portion of the penis proximal to the ring is not rigid, which may result in a pivoting effect. The penile skin may be cold and dusky, and ejaculation may be trapped by the constricting ring. Complications include penile pain and numbness, difficult ejaculation, ecchymosis, and petechiae. Patients taking aspirin or Coumadin should exercise caution when using these devices.

In some patients, the device can produce an erection that is of sufficient rigidity for coitus or engorge the glans for men with glanular insufficiency. In patients with severe vascular insufficiency, the device may not produce adequate erection. Although it is a safe and less costly means of treating ED when used properly, low patient acceptability limits the application or use of this therapy.

**PENILE VASCULAR SURGERY**

Isolated stenosis or occlusion of extrapenile arteries may be amenable to surgical repair. Arterial reconstructive surgery is a treatment option for healthy men, usually aged 55 or younger, with acquired ED secondary to focal arterial occlusion and the absence of generalized vascular disease secondary to hyperlipidemia, diabetes mellitus, chronic hypertension, and so on, or cavernous myopathy due to
cavernous ischemia (Montague et al, 2005). The most commonly used technique for penile revascularization is a bypass from the inferior epigastric artery to the dorsal artery or deep dorsal vein of the penis.

Penile venous surgery is also indicated only in young men with congenital or traumatic venous leakage. In congenital venous leakage, the venous insufficiency is typically through abnormal crural veins or superficial dorsal vein and is amenable to surgical cure. Traumatic venous leak is usually due to localized damage to the tunica albuginea or formation of a “fistula” between the corpus cavernosum and corpus spongiosum. Repair of the tunica or closure of the fistula can result in significant improvement of erectile function. In older men with chronic systemic diseases, venous leakage is usually caused by atrophy of the cavernous smooth muscle and intracavernous fibrosis; ligation of penile vein will only produce transient improvement and is not recommended.

**PENILE PROSTHESIS**

Patients considered for prosthesis implantation should be made aware of types of prosthesis available, efficacy, and potential complications including infection, mechanical failures, cylinder or tubing leaks, perforation, persistent pain, penile shortening, and autoinflation (Mulcahy et al, 2004). Penile prostheses are divided into 3 general types: malleable (semirigid), mechanical, and inflatable devices. The malleable devices are made of silicone rubber and several models contain a central intertwined metallic core. The mechanical device is also made of silicone rubber but contains polytetrafluoroethylene-coated interlocking polysulfone rings in a rod column, which provides rigidity when the rings are lined up in a straight line and flaccidity when the penis is bent. Inflatable (hydraulic) devices are further divided into 2-piece and 3-piece devices. Two-piece inflatable prostheses consist of a pair of cylinders attached to a scrotal pump reservoir. The most commonly used device, the three-piece inflatable penile prostheses, consist of paired penile cylinders, a scrotal pump, and a suprapubic fluid reservoir (Table 38–6).

In general, the malleable devices last longer than the inflatable ones. Modern 3-piece prostheses are extremely durable and reliable. However, patients should be informed that a 5–15% failure rate is expected within the first 5 years for inflatable implants, and the majority of devices will fail in 10–15 years and need replacement. Patient satisfaction with the 3-piece device is high, exceeding 85–90% in appropriately selected patients (Milbank and Montague, 2004). Recent innovations in penile prosthetics include antibiotic and hydrophilic-coated devices (infection rate <1%), lock-out valves to prevent autoinflation, and a more patient-friendly tactile pump (Delk et al, 2005; Droggin et al, 2005).

**Table 38-6. Types of Penile Protheses.**

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Semirigid</strong></td>
<td>American Medical System (AMS) Dura II and 600/650M (malleable)</td>
</tr>
<tr>
<td></td>
<td>Mentor Acu-Form</td>
</tr>
<tr>
<td><strong>Two-piece inflatable</strong></td>
<td>AMS Ambicor</td>
</tr>
<tr>
<td><strong>Three-piece inflatable</strong></td>
<td>Mentor Titan and Alpha-1, Alpha-1 narrow base</td>
</tr>
<tr>
<td></td>
<td>AMS 700 MS series with Inhibizone: CX, CXM, Ultrex</td>
</tr>
</tbody>
</table>

**MALE SEXUAL DYSFUNCTION INVOLVING EMISSION, EJACULATION, & ORGASM**

**Physiology of Emission, Ejaculation, & Orgasm**

Different mechanisms are involved in erection, emission, ejaculation, and orgasm, and these events can be dissociated from one another (eg, a frequent complaint of impotent patients is ejaculating through a “limp penis”). Except for nocturnal emissions, or “wet dreams,” emission and ejaculation require stimulation of the external genitalia. Impulses traveling from the pudendal nerves reach the upper lumbar spinal sympathetic nuclei. Efferent signals traveling in the hypogastric nerve activate secretions and transport sperm from the distal epididymis, vasa deferentia, seminal vesicles, and prostate to the prostatic urethra. Coordinated closing of the internal urethral sphincter and relaxation of the external sphincter direct the semen into the bulbous urethra (emission). Subsequent rhythmic contractions of the bulbocavernous muscles force the semen through a pressurized conduit—the much narrowed urethral lumen compressed by the engorged corpora cavernosa—to produce the 2- to 5-mL ejaculate. The external ejaculation process involves the somatomotor effent of the pudendal nerve to contract the bulbocavernous muscle. Since this action is involuntary, however, integrated autonomic and somatic action is required.

The mechanism of orgasm is the least understood of the sexual processes. It probably involves cerebral interpretation and response to sexual stimulation. Along with emission and ejaculation, several nongenital responses also occur. These include involuntary rhythmic contractions of the anal sphincter, hyperventilation, tachycardia, and elevation of blood pressure.

**Disorders Affecting Ejaculation, Emission, & Orgasm**

Premature or rapid ejaculation (PE), a persistent or recurrent occurrence of ejaculation with minimal sexual stimulation before, on, or shortly after penetration and before the person wishes it, is reported by up to 20–30% of men
(Althof 2006). In addition to psychotherapy or behavioral therapy, current guidelines suggest an SSRI, such as paroxetine (10–40 mg daily or 20 mg 3–4 hours before intercourse) may be used as pharmacotherapy for PE, although this indication is not FDA approved (Montague et al., 2004). Penile sensitivity may be reduced with use of a condom or topical anesthetizing creme (lidocaine–prilocaine). Although time-to-ejaculation is prolonged, a significant percentage of men experience decreased pleasure (penile numbness) or loss of erection.

Successful emission and ejaculation without orgasm occur in some patients with spinal cord injury. A history of disease or surgery is helpful in differentiating emission failure from retrograde ejaculation. If microscopic examination confirms the presence of sperm in bladder urine after a dry ejaculation, retrograde ejaculation can be diagnosed. If no sperm is found, emission failure is the cause. Bilateral sympathectomy at the L2 level may result in ejaculatory dysfunction for about 40% of patients while high bilateral retroperitoneal lymphadenectomy causes an even higher percentage of emission failures. Retrograde ejaculation is usually the result of dysfunction of the internal sphincter or the bladder neck, as seen after prostatectomy, with alpha-blocker therapy, and in autonomic neuropathy due to diabetes.

Elimination of alpha-adrenergic blockers may cure some patients with emission failure or retrograde ejaculation. Alpha-sympathomimetics, such as ephedrine or a combination of chlorpheniramine maleate and phenylpropanolamine hydrochloride (Ornala), have been used successfully in patients with retrograde ejaculation (McMahon et al., 2004). Electroejaculation via a rectal probe has been applied in patients suffering from spinal cord injury with some success. Psychosexual counseling is appropriate for patients who have normal wet dreams but cannot achieve orgasm and ejaculation.

REFERENCES


INTRODUCTION

Female urology encompasses urinary incontinence as well as pelvic reconstructive medicine for prolapse. It is common for urinary incontinence and pelvic organ prolapse to coexist in the same woman or to develop subsequently. This chapter will concentrate on pelvic organ prolapse as urinary incontinence has been discussed in detail in Chapter 29.

Pelvic organ prolapse is the protrusion of the pelvic organs (uterus, bladder, and bowel) into or past the vaginal introitus. Estimates of prevalence vary widely depending on definition used, whether the patient is symptomatic, the epidemiologic methods used, and the population studied. The U.S. National Center for Health Statistics estimates over 250,000 operations performed for genital prolapse apart from hysterectomy. With aging of the population, these quality of life issues and their treatment assume additional importance.

ANATOMY

Bony Pelvis

The maintenance of continence and prevention of pelvic organ prolapse rely on the support mechanisms of the pelvic floor. The bony pelvis is the rigid foundation to which all of the pelvic structures are ultimately anchored. These bones are the ilium, ischium, pubic rami, sacrum, and coccyx. It is important to understand and discuss the bony pelvis from the perspective of a standing woman. In the upright position, the bony arches of the pelvic inlet are oriented in an almost vertical plane (Figure 39–1). This directs the pressure of the intra-abdominal and pelvic contents toward the bones of the pelvis instead of the muscles and fascial attachments of the pelvic floor. This dispersion of forces minimizes the pressure on the pelvic musculature and transmits them to the bones that are better suited to the long-term cumulative stress of daily life.

Musculofascial Support

The muscles of the pelvic floor, particularly the levator ani muscles, have a critical role in supporting the pelvic organs and play an integral role in urinary, defecatory, and sexual function. The levator muscle complex consists of the pubococcygeus, the puborectalis, and the iliococcygeus (Figure 39–2). The pubococcygeus originates on the posterior inferior pubic rami and inserts on the midline organs and the anococcygeal raphe. The puborectalis also originates on the pubic bone, but its fibers pass posteriorly and form a sling around the vagina, rectum, and perineal body, resulting in the anorectal angle and promoting closure of the urogenital hiatus. The iliococcygeus originates from the arcus tendineus levator ani (ATLA) and inserts in the midline onto the anococcygeal raphe.

The ATLA is a linear fascial covering of the obturator internus muscle, and extends from the ischial spine to the posterior area of the superior ramus (Figure 39–2). The levator ani muscle group forms a broad hammock upon which the bladder, proximal vagina, and intrapelvic rectum lie, providing the musculofascial support for a large portion of the anterior pelvis. The space between the levator ani musculature through which the urethra, vagina, and rectum pass is called the urogenital hiatus. The fusion of levator ani where they meet in the midline creates the so-called levator plate.

The pelvic diaphragm has investing connective tissue which is often referred to as “fascia”; it is however, less organized and less distinct than traditional fascia (eg, rectus abdominis fascia). This visceropelvic fascia consists of collagen, smooth muscle, and elastin. Microscopic studies suggest that this fascia may be histologically indistinct from the deep vaginal wall, and not a separate “fascia.”

The pelvic fascia consists of 2 leaves—the endopelvic fascia (abdominal side) and the perivesical fascia (vaginal side). The urethra, bladder, vagina, and uterus are all contained within these 2 layers of fascia. The 2 leaves fuse laterally to insert along the arcus tendineus.

There are 3 important components of the pelvic fasciae. Anteriorly, the pubourethral ligaments attach to the lower portion of the pubis and insert on the proximal third of the urethra. Laterally, the arcus tendineus fascia pelvis extends from the pubourethral ligament (inferior portion of pubic symphysis) to the ischial spine. The arcus tendineus fascia pelvis is a thickening of the fascia overl-
Figure 39–1. Orientation of the bony pelvis in an upright woman. (Reproduced, with permission, from Drake RL et al (eds): Gray’s Anatomy for Students. Philadelphia, Churchill Livingston, 2005.)

Figure 39–2. Muscular support of the pelvis. The pubococygeus, the puborectalis, and the iliococcygeus comprise the levator ani muscles. (Reproduced, with permission, from Walsh PC, Retik AB, Vaughan ED et al (eds): Campbell’s Urology. 8th edition. Philadelphia, WB Saunders, 2002; p 49.)
ing the iliococcygeus muscle. It corresponds to the lateral attachment of the anterior bladder wall to the pelvic side wall (Figure 39–3). Fascia extending medially from this arc carries a variety of names (pubourethral, pubocervical, urethropelvic, vesicopelvic) and provides important support to the urethra and anterior vaginal wall. Posterior to the ischial spine, the fascia fans out to either side of the rectum and attaches to the pelvic side wall as the strong cardinal and uterosacral ligaments (Figure 39–4). The cardinal and uterosacral ligaments hold the uterus and upper vagina in their proper place over the levator plate.

**Innervation**

Many anatomic and surgical texts suggest that the levator ani muscles are dually innervated from the pudendal nerve on the perineal surface and direct branches of the sacral nerves on the pelvic surface. However, recent anatomic, neurophysiologic, and experimental evidence indicates that the levator ani muscles are innervated solely by a nerve traveling on the intrapelvic surface of the muscles without contribution of the pudendal nerve. This nerve originates from S3, S4, and/or S5 and innervates both the coccygeus and the levator ani muscle complex. After exiting the sacral foramina, it travels 2–3 cm medial to the ischial spine and ATLA across the coccygeus, iliococcygeus, pubococcygeus, and puborectalis. Given its location, the levator ani nerve is susceptible to injury during parturition and pelvic surgery.

The pudendal nerve innervates the striated urethral and anal sphincters as well as the deep and superficial perineal muscles and provides sensory innervation to the external genitalia. This nerve follows a complex course that originates from S2–S4 and travels behind the sacropinous ligament just medial to the ischial spine, exiting the pelvis through the greater sciatic foramen. It then enters the ischiorectal fossa through the lesser sciatic foramen and travels through the pudendal canal (Alcock’s canal) on the medial aspect of the obturator internus muscles before separating into several terminal branches that terminate within the muscles and skin of the perineum.

**PATHOPHYSIOLOGY**

Pelvic prolapse is prevented by several mechanisms. The most important support is from the continuous contraction of the levator ani pelvic muscles. The activity of skeletal muscle is a combination of basic tone, reflex contraction or relaxation, and voluntary contraction or relaxation. In patients with multiple deliveries, there is widening and descent of the levator plate. Therefore, the musculature becomes less important, and the “fascial” structures become the more important elements of support as the organs cross the pelvic floor.

Because of the complexity of pelvic organ support, vaginal prolapse is likely multifactorial: myopathic or neuropathic disorders, aging, atrophy, chronic increase in abdominal pressures, multiple deliveries, hysterectomy, and hormonal changes. However, intrinsic collagen abnormalities and other individual predisposing factors, such as genetics, differences in pelvic architecture, inherent quality of the pelvic musculature, and tissue response to injury, might explain why patients with known risk factors do not develop prolapse and many patients without risk factors do.
CLASSIFICATION

A number of classification systems have been published to facilitate standardization of clinical findings and enable communication about patients. Until recently, none of the systems have been validated. In 1996, the International Continence Society accepted standardization of the terminology for prolapse, known as the POPQ system for pelvic organ prolapse quantification. Although most clinicians still use the Baden-Walker system (grade 1–4), the POPQ system is the accepted standard for clinical studies and published data on prolapse (Figure 39–5).

The POPQ system is a site-specific quantitative description of support that locates 6 defined points around the vagina (2 anterior, 2 posterior, and 2 apical) with respect to their relationship to the hymenal ring. Negative numbers (in centimeters) are assigned to structures that have not prolapsed and positive numbers to those that protrude, with the plane of the hymen defined as zero. Points Aa and Ap are 3 cm above the hymenal ring. Points Ba and Bp are defined as the lowest points of the prolapse. The apex anteriorly is C (cervix), and posteriorly is D (pouch of Douglas). When the uterus is absent, point C is the vaginal cuff and D is omitted. TVL is total vaginal length at rest. GH is the genital hiatus measured from the middle of the urethral meatus to the posterior hymenal ring. PB is the perineal body measured from the posterior aspect of GH to the midanal opening (Figures 39–6 and 39–7).

The terminology avoids assigning a specific label, such as cystocele or rectocele, to the prolapsing part of the vagina, acknowledging that the actual organ(s) frequently cannot be determined on physical examination. Although it is more difficult to learn than traditional systems, reproducibility studies document interobserver and intraobserver reliability.

DIAGNOSIS

Symptoms

Pelvic organ prolapse is often asymptomatic until it is severe. Many women present with symptoms in addition to the vaginal bulge, as a result of the associated organ dysfunction.

The most common complaint due to anterior compartment prolapse (cystocele) is vaginal bulging, with or without suprapubic pressure and pain. Other symptoms include urgency, frequency, urge incontinence, and recurrent urinary tract infections. Obstructive voiding symptoms are due to urethral kinking when the bladder descends beyond the pubic ramus but the urethra remains fixed. This is commonly seen in the setting with previous surgery (eg, bladder neck suspension, urethropexy, sling). Patients may describe using unusual positions to void, such as pelvic tilting, squatting, or standing. Acute urinary retention and hydronephrosis is rare in patients with pelvic prolapse.
Many women with severe prolapse report that their stress incontinence improved as the prolapse worsened. Reduction of the prolapse during examination can produce stress incontinence in over 50% of clinically continent patients. This unmasking of occult stress incontinence warrants attention when considering surgical therapy.

Constipation and difficult defecation with distal stool trapping or excessive straining are symptoms commonly attributed to posterior compartment prolapse (rectocele). The patient may report having to manually splint the perineum or vagina to assist evacuation.

**Physical Examination**

A thorough physical examination should be done with a comfortably full bladder, at rest and with straining, in the supine (lithotomy) position. Depending on the patient’s ability to strain, examination in the upright position may be performed to accentuate the degree of prolapse to match that reported by the patient. The goal of examination is to determine the degree of prolapse, the specific anatomic defects, and the presence of concomitant organ prolapse in other compartments or incontinence. In the supine position, the origin of prolapse should be determined. Using a half-speculum blade, the posterior vaginal wall is retracted; the patient is asked to strain while the anterior defect is evaluated. After characterizing the anatomic defects, an attempt can be made to reduce the cystocele in order to elicit occult stress urinary incontinence and hypermobility. Similarly, the anterior vaginal wall is retracted to determine the presence of any posterior defect. A digital rectal exam assesses rectal tone, presence of impacted stool, attenuation of the prerectal fascia, and perineal laxity. Using both blades of speculum, the vaginal vault or cervix can be examined for uterine descent, vault prolapse, and enterocele.

**EVALUATION**

The basic evaluation for prolapse includes a history, physical examination, measurement of postvoid residual volume, and urinalysis. Additional testing may be needed in the setting of a large introital bulge where it may be difficult to differentiate between a severe cystocele, an enterocele, or high rectocele by physical examination only. Imaging studies can be performed to identify which organs are prolapsing. An ideal study should provide precise information about which structures are prolapsed, the presence of urinary retention and obstruction, urethral hypermobility, and urinary incontinence.

**Cystourethrography**

The patient is upright with a full bladder. Films are taken in a lateral position during both rest and strain. This exam provides information about bladder position, bladder neck funneling, urethral mobility, stress incontinence, and postvoid residual. The normal position of the base of the bladder should be above the pubococcygeal line (Figure 39–8). The presence of a rectocele can also be inferred.
when bowel gas is identified below the pubic symphysis. This exam is static and does not provide information about other pelvic organs or soft tissues of the pelvic floor.

**Ultrasonography**

Ultrasound is an attractive imaging modality because it is easy to perform, minimally invasive, and avoids radiation exposure. Tubo-ovarian and renal disease can also be assessed during the exam. There is evidence that ultrasound is useful in evaluating bladder neck hypermobility; however, transvaginal imaging for pelvic prolapse does not provide adequate visualization of the soft tissues. Transabdominal ultrasound can be used to quantify prolapse, although it appears to be better for the anterior compartment and uterine descent than for the posterior compartment.

**Dynamic Magnetic Resonance Imaging**

Recently, magnetic resonance imaging (MRI) has been used to evaluate pelvic prolapse. It is performed quickly,

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*Figure 39–6.* The pelvic organ prolapse quantification system. Aa—anterior vaginal wall 3 cm from hymen. Ba—lowest point of anterior vaginal wall prolapse. C—distance from hymen to cervix. D—distance from hymen to posterior fornix (pouch of Douglas). Ap—posterior vaginal wall 3 cm from hymen. Bp—lowest point of posterior vaginal wall prolapse. TVL—total vaginal length. GH—genital hiatus measured from midurethral meatus to posterior hymen. PB—perineal body measured from posterior hymen to mid-anus. (Reproduced, with permission, from Bump RC, Mattiasson A, Bo K et al: The standardization of terminology of female pelvic organ prolapse and pelvic floor dysfunction. Am J Obstet Gynecol 1996;175:10.)

*Figure 39–7.* A: Example of complete vaginal prolapse (eversion) using POPQ classification. This occurs after a hysterectomy, therefore there is no point D. Points Aa and Ap are maximally distal. Points Ba, C, and Bp are maximally everted. B: Normal support with no vaginal wall descent. (Reproduced, with permission, from Bump RC, Mattiasson A, Bo K et al: The standardization of terminology of female pelvic organ prolapse and pelvic floor dysfunction. Am J Obstet Gynecol 1996;175:10.)

*Figure 39–8.* Example of a cystocele. Lateral image of contrast-filled bladder extending well past the pubococcygeal line (line drawn from inferior edge of pubis to coccyx).
without contrast or ionizing radiation, and permits visualization of the soft tissues as well as the upper urinary tract. Gousse and colleagues have shown that in comparison with findings at surgery, MRI has a 100% sensitivity, 83% specificity, and a positive predictive value of 97% when assessing cystoceles. MRI does not require intravesical catheterization, does not require contrast, is fast, and non-invasive. The drawback is that it may underestimate the extent of cystocele and enterocele because the exam must be performed in the supine position, which diminishes the downward forces that can be generated with abdominal straining. Standing MRI’s will be the ultimate modality in obtaining an even more precise study for prolapse. Cost is the main reason that prohibits the wide use of MRIs currently (Figure 39–9A & B).

**Video Urodynamic Study**
A urodynamic study can be done with or without video assistance. Both methods provide information about bladder compliance, capacity, sensation of filling, detrusor instability, and contractility. An advantage in performing videourodynamics is the capability to watch the position and funneling of the bladder neck during filling and straining. This combines cystourethrography with the urodynamic study. Choosing to use fluoroscopy should be based on cost, availability, and familiarity with this method.

The importance of documenting the presence of urinary incontinence in patients with large cystoceles is controversial. The incidence of occult stress urinary incontinence is felt to be as high as 22–80% among patients with high-stage vaginal prolapse. Due to the known masked urinary incontinence, and the high incidence of postoperative “de novo” stress incontinence, many pelvic reconstruction surgeons routinely perform a concomitant anti-incontinence surgery in all anterior vaginal reconstruction, independent of the continence status.

Determining the presence of bladder instability is important in preoperative counseling because it can affect the postoperative result. In the majority of cases (60–80%), urgency may resolve after surgery. However, some patients may have no change in urgency or a worsening of their urgency with sling placement and/or bladder neck elevation. Urodynamic study may also suggest urinary obstruction with elevated voiding pressure, low urinary flow, or radiographic evidence of urethral kinking.

**Cystourethroscopy**
This exam is performed to rule out concurrent pathology in the bladder and urethra, such as bladder carcinoma, urethral diverticulum, stones or foreign bodies (ie, suture material) from previous surgery. Cystoscopic illumination

![Figure 39–9. A: Sagittal MRI image of cystocele and enterocele. Small intestine protrudes posterior to the prolapsed bladder (white). B: Sagittal MRI image of an enterocele only. Bladder does not prolapse past the pubococcygeal line.](image-url)
can also be used to differentiate an enterocele from a cystocele. A pelvic exam is performed with the cystoscope in the bladder. The bladder transilluminates through the anterior vaginal wall so that the extent of the bladder prolapse is demarcated. It can also be used as a bedside urodynamic exam, assessing filling sensation, postvoid residual and visual cystometrics for bladder contractions. With a full bladder, a supine valsava stress test can be performed, looking for urethral leakage.

**Upper Urinary Tract Evaluation**

Patients with a high stage cystocele should have upper tract imaging because there is a 4–7% incidence of moderate hydroureteronephrosis amongst patients with severe vaginal prolapse. This risk is greater in patients with procidentia, compared to vault prolapse. Ultrasound, excretory urography, computed tomography, or MRI can be used. An advantage of MRI is the ability to simultaneously evaluate the upper urinary tract, tubo-ovarian disease, and pelvic organ prolapse simultaneously.

**Laboratory Evaluations**

Patients must have sterile urine prior to proceeding with an operative procedure. In preparation for surgery, a complete blood count, basic metabolic panel, coagulation profile, and urine culture should be obtained.

**TREATMENT**

**Nonsurgical Therapy**

When surgery is contraindicated or must be deferred, a patient can be made comfortable with a vaginal pessary. They are of primarily 2 types, ring and support. However, a word of caution regarding the use of pessaries. Without close monitoring of the patient and frequent examinations, ulcerations of the vaginal mucosa and fistula formation to the bladder can occur. Proper sizing, care, cleansing, and estrogen replacement can minimize complications. Vaginal estrogen cream has been shown to have beneficial effects on vaginal epithelium by improving vascularity and total skin collagen content, especially in preparation for surgical correction.

Other nonsurgical treatments include pelvic muscle exercises as well as measures to improve the associated factors such as chronic cough, obesity, and constipation. These may help alleviate symptoms and prevent worsening of the prolapse, but the actual prolapse will not spontaneously disappear.

**Surgical Repairs**

The goal of repair is to restore pelvic anatomical support. This is rarely an independent surgery. Often, surgery entails addressing incontinence as well as prolapse of all the compartments. The end result must restore anatomy and function by restoring normal vaginal axis and depth while preserving urinary, bowel, and sexual function. Because the various forms of organ prolapse are interrelated owing to shared support mechanisms, some recommend that all defects be repaired at the same time because occult weaknesses in other sites may be acquired.

**Anterior Compartment**

Anterior colporrhaphy was initially introduced in 1914. It is a transvaginal approach that reduces the herniation of the bladder by plicating the redundant tissue, imbricating the detrusor, and approximating the tissue in the midline (Figure 39–10). The long-term results have been disappointing. There have been numerous modifications to the anterior colporrhaphy, including the use of synthetic or allograft materials, variations in suture placement and anchoring techniques. One method places a piece of mesh into the fold of the imbricated bladder wall. Another

places the prosthetic layer over the plication sutures and anchors it in place laterally in the arcus tendineus fasciae pelvis or obturator fascia bilaterally. These repairs supported by a piece of biomaterial appear to be much more durable than just a simple anterior colporrhaphy.

The vaginal paravaginal repair is another technique with various modifications. It involves reapproximation of the weakened lateral vaginal attachments to the arcus tendineus. Two to three interrupted permanent sutures are placed on each side between the arcus tendineus laterally and the pubocervical fascia medially from the back of the pubis to the ischial spine.

Cystocele can also be repaired abdominally by suturing the endopelvic fascia and lateral vaginal sulcus to the arcus tendineus with interrupted permanent sutures. As a result of multiple modifications to these techniques and the addition of anti-incontinence procedures that are performed concomitantly, the durability of each procedure is uncertain. However, most pelvic reconstruction surgeons have evolved to repairing anterior compartment defects with a graft material to support the bladder with or without anterior colporrhaphy.

More recent innovative approaches for anterior vaginal wall repair anchor an allograft, xenograft, or polypropylene mesh without tension via strips placed through the obturator foramen with a special device (Perigee, American Medical Systems; Anterior Prolift, Gynecare). These techniques await safety and efficacy studies but are increasing in use.

**Apical Compartment**

An enterocele has a hernia sac behind vaginal epithelium that lacks musculofascial support of the vaginal vault. Repair involves transvaginal dissection of the peritoneal sac from the vaginal wall laterally, the bladder anteriorly, and the rectum posteriorly. Obliteration and ligation of the sac is performed by the use of pursestring sutures. Because an enterocele presents due to weakened vault support, the vaginal vault must also be resuspended.

Vaginal vault suspension can be performed by transvaginal reattachment of the uterosacral ligaments to the vaginal apex. Excellent anatomic outcomes have been described but ureteral injury is a limiting factor, with rates as high as 11%. Therefore, cystoscopy after intravenous indigo carmine is essential.

The vaginal vault can also be elevated to the sacrospinous ligament that extends from the ischial spine to the sacrum. Transvaginal placement of sutures is important as the pudendal nerve, artery, and vein are located immediately deep to the sacrospinous ligament, and damage to these structures can cause significant morbidity.

Fixation of the vaginal apex to the iliococcygeus fascia and/or muscle is another method to resuspend the vaginal vault. One or two sutures are placed into the iliococcygeus just anterior to the ischial spine. If the patient is not sexually active, this can be performed without a vaginal incision by placing a monofilament permanent suture at full thickness through the vaginal wall into the muscle either unilaterally or bilaterally. A potential benefit is the absence of critical structures in the area. Very recently, the use of polypropylene mesh to support vaginal vault suspensions has been reported with very good success.

Abdominal sacrocolpopexy is considered by most surgeons to be the gold-standard procedure for vaginal vault prolapse. The procedure entails suspension of the vaginal apex to the sacral promontory with or without using a graft bridge. Autologous, allogenic, and synthetic material have all been described (Figure 39–11). Although this procedure requires an abdominal incision and there is the risk of bleeding from the sacral promontory and postoperative ileus, the resultant anatomy carries the greatest longevity and least risk of sexual dysfunction and dyspareunia. The laparoscopic approach appears to be just as successful in experienced hands.

**Posterior Compartment**

The aims of posterior colporrhaphy for the repair of rectocele are to plicate the prerectal and pararectal fascia in the midline, narrow the posterior aspect of the levator hiatus, and repair the perineal body. Rectocele repair is indicated only if the patient is symptomatic because increased dyspareunia as well as worsening defecatory symptoms occurs...
in some patients. There is some evidence and opinion that asymptomatic rectoceles should be repaired at the time of repair of other defects because new defects may be acquired if a weakness exists. Until there is compelling evidence of benefit, the decision to repair asymptomatic rectoceles is left to the surgeon’s clinical judgment.

Proper treatment of pelvic organ prolapse entails a thoughtful preoperative evaluation, strong knowledge of pelvic floor anatomy, and restoration of the vagina to its normal position and axis while maintaining normal physiologic and sexual function.

**FEMALE SEXUAL DYSFUNCTION**

**Introduction**

Current definitions of sexual dysfunction in women reflect a change in our understanding of normal sexual response. Rather than the traditional view of a sexual response progressing through discrete phases in sequence (desire, arousal, orgasm, and resolution), it is now recognized that these phases overlap and that the sequence can vary. Also recognized is the importance to sexual satisfaction of the subjective experience and of an environment and stimuli that are conducive to sexual feelings.

Disorders of female sexual function are summarized in Table 39–1. The prevalence of the sexual desire/interest disorder, diagnosed when a woman fails to feel desire at any stage during the sexual experience, is uncertain. Studies have focused on a lack of desire at the initiation of and between sexual experiences, as well as on a lack of sexual thoughts. However, sexual thoughts are infrequent in many women without apparent sexual dissatisfaction, and the frequency of sexual fantasies or sexual thoughts has little correlation with sexual satisfaction in women. Arousal disorders are categorized according to whether there is a lack of subjective arousal alone or a lack of both subjective arousal and awareness of genital congestion. No objective measurements are used to establish diagnoses. Arousal disorders also have an uncertain prevalence; most studies focus only on vaginal lubrication.

**Table 39–1. Definitions of Sexual Dysfunction.**

<table>
<thead>
<tr>
<th></th>
<th>APA Definition*</th>
<th>AUA Definition†</th>
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<tbody>
<tr>
<td>Hypoactive sexual desire disorder</td>
<td>Persistent or recurrent deficiency or absence of sexual fantasies and desire for sexual activity. Judgment of deficiency is made by the clinician, taking into account factors that affect sexual functioning</td>
<td>Absent or diminished feelings of sexual interest or desire, absent sexual thoughts or fantasies, and a lack of responsive desire. Motivations for attempting to become sexually aroused are scarce or absent. Lack of interest goes beyond a normal lessening with increasing age and relationship duration</td>
</tr>
<tr>
<td>Lack of subjective arousal</td>
<td>No DSM-IV definition addresses the lack of subjective arousal</td>
<td>Absent or markedly reduced feelings of sexual arousal (sexual excitement and sexual pleasure) from any type of stimulation, and absent or impaired genital sexual arousal (vulvar swelling and lubrication)</td>
</tr>
<tr>
<td>Subjective arousal disorder</td>
<td>No DSM-IV definition addresses the lack of subjective arousal</td>
<td>Absent or markedly reduced feelings of sexual arousal (sexual excitement and sexual pleasure) from any type of stimulation. Vaginal lubrication and other signs of physical response still occur</td>
</tr>
<tr>
<td>Female sexual arousal disorder</td>
<td>Persistent or recurrent inability to attain, or to maintain until completion of sexual activity, adequate lubrication, and swelling response of sexual excitement</td>
<td>Absent or impaired genital sexual arousal (minimal vulvar swelling or vaginal lubrication from any type of sexual stimulation, and reduced sexual sensations when genitalia are caressed)</td>
</tr>
<tr>
<td>Female orgasmic disorder</td>
<td>Persistent or recurrent delay or absence of orgasm after a normal sexual excitement phase</td>
<td>Lack of orgasm, markedly diminished intensity of orgasmic sensations, or marked delay of orgasm from any kind of stimulation, despite self-reported high sexual arousal or excitement</td>
</tr>
</tbody>
</table>

*Data from the American Psychiatric Association (APA).
†Data from the international committee sponsored by the American Urological Association (AUA).
PHYSIOLOGY

The basis of desire and perceived arousal in women is poorly understood, but it appears to involve interactions among multiple neurotransmitters, sex hormones, and environmental factors.

Genital vasocongestive responses occur in women within seconds after erotic stimulation. Both parasympathetic and sympathetic nerves release nitric oxide and vasointestinal polypeptide, which mediate vasodilatation, and acetylcholine, which promotes endothelial release of nitric oxide.

The effect of estrogen levels on sexual function is complex. Although low estrogen levels and vaginal atrophy are associated with reduced measures of vaginal congestion when the woman is not receiving sexual stimulation, the percent increase in congestion in response to erotic stimuli is similar in the presence of low and high estrogen levels. Similarly, changes in the volume of the vaginal wall and clitoris and the relative volume of regional blood in response to sexual stimulation are similar before and after menopause. Estrogen deficiency does not necessarily preclude adequate lubrication, provided that stimulation is sufficient.

Subjective arousal is poorly correlated with genital response. For example, increases in genital vascongestion in response to erotic videos are similar among women who report problems with arousal and women who report no problems with arousal.

Indirect evidence suggests that testosterone and dopamine play a role in modulating sexual response, since testosterone supplementation or treatment with a dopaminergic agonist can augment response. However, large population studies have failed to find the expected positive correlations between sexual function and serum testosterone levels. One possible explanation is that serum levels do not reflect the intracellular production of testosterone from adrenal and ovarian precursors.

Several other factors have been associated with reduced subjective arousal. These include distractions, expectations of a negative experience (eg, as a result of dyspareunia, the partner’s sexual dysfunction, or negative experiences in the past), sexual anxiety, fatigue, and depression. Medications including selective serotonin-reuptake inhibitors and oral contraceptives have also been implicated. Oral contraceptives increase levels of sex hormone–binding globulin, which in turn reduces free testosterone levels; it is hypothesized that some women are particularly sensitive to these effects, which may be prolonged.

On the basis of survey data, several factors have been closely linked to women’s sexual satisfaction and desire. These include stable past and current mental health, positive emotional well-being and self-image, rewarding past sexual experiences, positive feelings for the partner, and positive expectations for the relationship. Certain diseases such as multiple sclerosis, renal failure, and premature menopause induced by chemotherapy are associated with a high incidence of sexual dysfunction. In women, unlike men, vascular disease related to age does not appear to correlate with reduced sexual satisfaction.

EVALUATION

A detailed history is the main tool in the assessment and diagnosis of sexual dysfunction and is usually obtained from both partners. Important aspects of the history include the quality of the couple’s relationship, the woman’s mental and emotional health, the quality of past sexual experiences, specific concerns related to sexual activity (such as insufficient nongenital and nonpenetrative genital stimulation), and the woman’s thoughts and emotions during sexual activity.

A physical examination, including a pelvic examination, is part of routine care, but it infrequently identifies a cause of sexual dysfunction. Its usefulness may be greater when there is associated dyspareunia. Table 39–2 details features that are potentially relevant to sexual dysfunction that need to be assessed during an examination.

The possibility that laboratory testing will identify causes of sexual dysfunction is low. Estrogen deficiency, for example, is best detected by taking a history and performing an examination. In addition, serum levels of testosterone do not correlate with sexual function. Measurement of prolactin or thyrotropin is warranted if other symptoms or signs suggest the presence of abnormal levels.

Table 39–2. Physical Examination Findings Potentially Relevant to Sexual Dysfunction.

<table>
<thead>
<tr>
<th>External genitalia</th>
<th>Introitus</th>
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<tbody>
<tr>
<td>Sparsity of pubic hair, suggestive of low adrenal androgen levels</td>
<td>Vulvar atrophy</td>
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<tr>
<td>Vulvar skin disorders (eg, lichen sclerosis)</td>
<td>Lichen sclerosus</td>
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<tr>
<td>Cracks or fissures in interlabial folds suggestive of candidiasis</td>
<td>Splitting of posterior fourchette</td>
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<tr>
<td>Labial abnormalities that may cause embarrassment</td>
<td>Abnormalities of the hymen</td>
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<table>
<thead>
<tr>
<th>Internal</th>
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<tbody>
<tr>
<td>Labial adhesions</td>
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<tr>
<td>Swelling of vestibular glands</td>
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<td>Vestibulitis</td>
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<tr>
<td>Pelvic organ prolapse</td>
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<tr>
<td>Abnormal vaginal discharge</td>
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<tr>
<td>Hypertonicity of pelvic muscles</td>
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<tr>
<td>Presence of tender “trigger points” on palpation of levator ani muscles</td>
</tr>
<tr>
<td>Fixed retroversion of uterus, tenderness of vaginal fornix on bimanual exam causing deep dyspareunia</td>
</tr>
</tbody>
</table>
TREATMENT

The management of sexual dysfunction in women is guided by the history. Data from randomized trials that support the use of any particular intervention are limited.

Psychological

Cognitive behavioral therapy focuses on identifying and modifying factors that contribute to sexual dysfunction, such as maladaptive thoughts, unreasonable expectations, behaviors that reduce the partner’s interest or trust, insufficient erotic stimuli, and insufficient nongenital physical stimulation. Strategies are suggested to improve the couple’s emotional closeness and communication and to enhance erotic stimulation. Sex therapy for couples is focused on similar issues but also includes sensate focus techniques, consisting initially of nonsexual physical touch, with gradual progression toward sexual touch; partners are encouraged to alternately touch each other and to provide feedback about what touches are pleasurable. These techniques help change the undue focus on a performance goal.

Pharmacologic

Other than estrogen therapy for dyspareunia related to genitourinary atrophy, no medications are currently approved by the Food and Drug Administration for the treatment of sexual dysfunction in women. Several off-label uses of drugs have been considered, although data about effectiveness are sparse.

Nonhormonal

The involvement of nitric oxide in neurogenic vasodilatation suggests that phosphodiesterase inhibitors may ameliorate genital arousal disorder. In a small, laboratory-based, randomized trial, a single 50-mg dose of sildenafil (Viagra, Pfizer) increased subjective arousal, genital sensations, and ease of orgasm in some women with genital arousal disorder. The benefit was observed only among women who had a marked reduction in the normal vasocongestive response to subjectively arousing visual erotic stimulation. In women in whom arousal and desire disorders were diagnosed (rather than genital arousal disorder), sildenafil improved no measure of sexual desire, sensation, lubrication, or satisfaction.

Hormonal

Supraphysiologic androgen therapy has been prescribed for sexual dysfunction since the 1930s, but more recently, testosterone at lower doses have been studied in randomized trials. The results of 4 recent randomized trials show increase in sexually satisfying events, sexual desire, and response. Important limitations of these studies include their brevity (which is of particular importance, given the expected long-term use of the drug) and that their results are generalizable only to women in whom menopause was surgically induced and who also receive estrogen therapy. In some women who have undergone natural menopause, the ovaries continue to be an important source of androgens, and thus, the effects of androgen supplementation may differ from those whose ovaries have been surgically removed. Furthermore, risks associated with the long-term use of conjugated estrogens arouse concern about the use of any postmenopausal estrogen therapy over time. Prescribing testosterone alone to women who lack estrogen would raise their already high ratios of androgen to estrogen. There are no safety or efficacy data for testosterone supplementation for estrogen-deficient women. A chief concern with long-term androgen use is a potential increase in insulin resistance, which could predispose a woman to the metabolic syndrome or exacerbate the syndrome if it is already present.

Some researchers suggest supplementation with steroid because older women have a physiologic decrease of dehydroepiandrosterone produced. However, rigorous data that support such supplementation are lacking.

The role of systemic estrogen in increasing desire and subjective arousal remains unclear. In the Women’s Health Initiative trial, no significant differences were found between the estrogen and placebo groups in reported satisfaction after sexual activity. However, sexual dysfunction was not a primary focus of the trial, and the assessment tool was inadequate.

The prevalence of sexual disorders that are associated with the use of antidepressants in women is estimated at 22–58%, with higher rates reported for selective serotonin-reuptake inhibitors and lower rates reported for bupropion. A recent metaanalysis of strategies to ameliorate dysfunction associated with antidepressants did not recommend any particular drug, although the potential advantages of adding bupropion were noted.

RECOMMENDATIONS AND CONCLUSIONS

Recommendation guidelines for the evaluation and management of sexual dysfunction in women advocate attention to mental and overall health and to both interpersonal and personal psychological issues. Local estrogen therapy is recommended for dyspareunia that is associated with vulvar atrophy that results in reduced sexual motivation. Testosterone therapy should be viewed as investigational and should be prescribed only by clinicians who are knowledgeable about sexual dysfunction in women.

A better understanding is needed of the endogenous and environmental factors that mediate sexual desire and arousal. Randomized clinical trials are also needed to assess the effects of psychological and pharmacologic therapies.
alone and in combination. The risks and benefits of long-term testosterone therapy require further study.

REFERENCES

Introduction


Anatomy


Pathophysiology


Classification


Diagnosis


Evaluation


Treatment


Sexual Dysfunction


Physiology


Evaluation


Management


CONGENITAL ANOMALIES OF THE PENIS

APENIA
Congenital absence of the penis (apenia) is extremely rare. In this condition, the urethra generally opens on the perineum or inside the rectum.

Patients with apenia should be considered for assignment to the female gender. Castration and vaginoplasty should be considered in combination with estrogen treatment as the child develops.

MEGALOPENIS
The penis enlarges rapidly in childhood (megalopenis) in boys with abnormalities that increases the production of testosterone, for example, interstitial cell tumors of the testicle, hyperplasia, or tumors of the adrenal cortex. Management is by correction of the underlying endocrine problem.

MICROPENIS
Micropenis is a more common anomaly and has been attributed to a testosterone deficiency that results in poor growth of organs that are targets of this hormone. A penis smaller than 2 standard deviations from the norm is considered a micropenis (see Table 40–1). The testicles are small and frequently undescended. Other organs, including the scrotum, may be involved. Early evidence suggests that the ability of the hypothalamus to secrete luteinizing hormone-releasing hormone (LHRH) is decreased. The pituitary-gonadal axis appears to be intact, since the organs respond to testosterone, although this response may be sluggish at times. Studies have shown that topical application of 5% testosterone cream causes increased penile growth, but its effect is due to absorption of the hormone, which systemically stimulates genital growth. Patients with micropenis must be carefully evaluated for other endocrine and central nervous system anomalies. Retarded bone growth, anosmia, learning disabilities, and deficiencies of adrenocorticotropic hormone and thyrotropin have been associated with micropenis. In addition, the possibility of intersex problems must be carefully investigated before therapy is begun.

The approach to management of micropenis has undergone gradual change in recent years, but androgen replacement is the basic requirement. The objective is to provide sufficient testosterone to stimulate penile growth without altering growth and closure of the epiphyses. A regimen of 25 mg orally every 3 weeks for no more than 4 doses has been recommended. Penile growth is assessed by measuring the length of the stretched penis (pubis to glans) before and after treatment. Therapy should be started by age 1 year and aimed at maintaining genital growth commensurate with general body growth. Repeat courses of therapy may be required if the size of the penis falls behind as the child grows. For undescended testicles, orchiopexy should be done before the child is 2 years old. In the future, treatment with LHRH may correct micropenis as well as cause descent of the testicles, but at present, LHRH is not approved for such use.

ADULT PENILE SIZE
In recent years, penile augmentation and enhancement procedures have been done with increasing frequency, although no validation of success has been documented. Suspensory ligament release with pubic fat pad advancement, fat injections, and dermal fat grafts have been used in attempts to enhance penile size. Many consider that these procedures have not been proved safe or efficacious in normal men. Wessells, Lue, and McAninch (1996) evaluated penile size in the flaccid and erect state in otherwise normal adult men and found very good correlation between stretched and erect length ($R^2 = 0.793$; Table 40–2). This information can provide a guideline for physicians whose patients are concerned with their penile dimensions.
CONGENITAL ANOMALIES OF THE URETHRA

DUPICATION OF THE URETHRA

Duplication of the urethra is rare. The structures may be complete or incomplete. Resection of all but one complete urethra is recommended.

URETHRAL STRICTURE

Congenital urethral stricture is uncommon in infant boys. The fossa navicularis and membranous urethra are the 2 most common sites. Severe strictures may cause bladder damage and hydronephrosis (see Chapter 11), with symptoms of obstruction (urinary frequency and urgency) or urinary infection. A careful history and physical examination are indicated in patients with these complaints. Excretory urography and excretory voiding urography often define the lesion and the extent of obstruction. Retrograde urethrography (Figure 40–1) may also be helpful. Cystoscopy and urethroscopy should be performed in all patients in whom urethral stricture is suspected.

Strictures can be treated at the time of endoscopy. Diaphragmatic strictures may respond to dilation or visual urethrotomy. Other strictures should be treated under direct vision by internal urethrotomy with the currently available pediatric urethrotome. It may be necessary to repeat these procedures in order to stabilize the stricture. Single-stage open surgical repair by anastomotic urethroplasty, buccal mucosa graft, or penile flap is desirable if the obstruction recurs.

POSTERIOR URETHRAL VALVES

Posterior urethral valves, the most common obstructive urethral lesions in infants and newborns, occur only in males and are found at the distal prostatic urethra. The valves are mucosal folds that look like thin membranes; they may cause varying degrees of obstruction when the child attempts to void (Figure 40–2).

Clinical Findings

A. SYMPTOMS AND SIGNS

Children with posterior urethral valves may present with mild, moderate, or severe symptoms of obstruction. They often have a poor, intermittent, dribbling urinary stream. Urinary infection and sepsis occur frequently. Severe obstruction may cause hydronephrosis (see Chapter 11), which is apparent as a palpable abdominal mass. A palpable midline mass in the lower abdomen is typical of a distended bladder. Occasionally, palpable flank masses indicate hydronephrotic kidneys. In many patients, failure to thrive may be the only significant symptom, and examination may reveal nothing more than evidence of chronic illness.

B. LABORATORY FINDINGS

Azotemia and poor concentrating ability of the kidney are common findings. The urine is often infected, and anemia may be found if infection is chronic. Serum creatinine and blood urea nitrogen levels and creatinine clearance are the best indicators of the extent of renal failure.

C. X-RAY FINDINGS

Voiding cystourethrography is the best radiographic study available to establish the diagnosis of posterior urethral valves. The presence of large amounts of residual urine is
Figure 40–1. **Upper left:** Retrograde urethrogram showing congenital diaphragmatic stricture. **Upper right:** Posterior urethral valves revealed on voiding cystourethrography. Arrow points to area of severe stenosis at distal end of prostatic urethra. **Lower left:** Posterior urethral valves. Patient would not void with cystography. Retrograde urethrogram showing valves (arrow). **Lower right:** Cystogram, same patient. Free vesicoureteral reflux and vesical trabeculation with diverticula.
apparent on initial catheterization done in conjunction with radiographic studies, and an uncontaminated urine specimen should be obtained via the catheter and sent for culture. The cystogram may show vesicoureteral reflux and the severe trabeculations of long-standing obstruction, and the voiding cystourethrogram often demonstrates elongation and dilatation of the posterior urethra, with a prominent bladder neck (Figure 40–1). Excretory urograms may reveal hydrourereter and hydrenephrosis when obstruction is severe and long standing.

D. ULTRASONOGRAPHY

Ultrasonography can be used to detect hydronephrosis, hydroureter, and bladder distention in children with severe azotemia. It can also detect fetal hydronephrosis, which is typical of urethral valves, as early as 28 weeks of gestation; when the obstruction is from valves, an enlarged bladder with bilateral hydroureretonephrosis is usually present.

E. INSTRUMENTAL EXAMINATION

Urethroscopy and cystoscopy, performed with the patient under general anesthesia, show vesical trabeculation and cellules and, occasionally, vesical diverticula. The bladder neck and trigone may be hypertrophied. The diagnosis is confirmed by visual identification of the valves at the distal prostatic urethra. Supravesical compression shows that the valves cause obstruction.

Treatment

Treatment consists of destruction of the valves, but the approach depends on the degree of obstruction and the general health of the child. In children with mild to moderate obstruction and minimal azotemia, transurethral fulguration of the valves is usually successful. Occasionally, catheterization, cystoscopy, or urethral dilation by perineal urethrostomy destroys the valves.

The more severe degrees of obstruction create varying grades of hydronephrosis requiring individualized management. Treatment of children with urosepsis and azotemia associated with hydronephrosis includes use of antibiotics, catheter drainage of the bladder, and correction of the fluid and electrolyte imbalance. Vesicostomy may be of benefit in patients with reflux and renal dysplasia.

In the most severe cases of hydronephrosis, vesicostomy or removal of the valves may not be sufficient, because of
ureteral atony, obstruction of the ureterovesical junction from trigonal hypertrophy, or both. In such cases, percutaneous loop ureterostomies may be done to preserve renal function and allow resolution of the hydronephrosis. After renal function is stabilized, valve ablation and reconstruction of the urinary tract can be done.

The period of proximal diversion should be as short as possible, since vesical contracture can be permanent after prolonged suprapubic diversion.

It has been noted that approximately 50% of children with urethral valves have vesicoureteral reflux and that the prognosis is worse if the reflux is bilateral. After removal of the obstruction, reflux ceases spontaneously in about one-third of patients. In the remaining two-thirds of patients, the reflux should be corrected surgically.

Long-term use of antimicrobial drugs is often required to prevent recurrent urosepsis and urinary tract infection even though the obstruction has been relieved.

**Prognosis**

Early detection is the best way to preserve kidney and bladder function. This can be accomplished by ultrasonography in utero, by careful physical examination and observation of voiding in the newborn, and by thorough evaluation of children who have urinary tract infections. Children in whom azotemia and infection persist after relief of obstruction have a poor prognosis.

**ANTERIOR URETHRAL VALVES**

Signs of anterior urethral valves, a rare congenital anomaly, are urethral dilatation or diverticula proximal to the valve, bladder outlet obstruction, postvoiding incontinence, and infection. Enuresis may be present. Urethroscopy and voiding cystourethrography will demonstrate the lesion, and endoscopic electrofulguration will effectively correct the obstruction.

**URETHRORECTAL & VESICORECTAL FISTULAS**

Urethrectal and vesicocolic fistulas are rare and are almost always associated with imperforate anus. Failure of the urorectal septum to develop completely and separate the rectum from the urogenital tract permits communication between the 2 systems (see Chapter 2). The child with such a fistula passes fecal material and gas through the urethra. If the anus has developed normally (ie, if it opens externally), urine may pass through the rectum.

Cystoscopy and panendoscopy usually show the fistulous opening. Radiographic contrast material given by mouth will reach the blind rectal pouch, contrast material between the end of the rectum and the perineum can be seen on appropriate radiograms.

Imperforate anus must be opened immediately and the fistula closed, or if the rectum lies quite high, temporary sigmoid colostomy should be performed. Definitive surgery, with repair of the urethral fistula, can be done later.

**HYPOSPADIAS**

In hypospadias, the urethral meatus opens on the ventral side of the penis proximal to the tip of the glans penis (Figure 40–3).

Sexual differentiation and urethral development begin in utero at approximately 8 weeks and are complete by 15 weeks. The urethra is formed by the fusion of the urethral folds along the ventral surface of the penis, which extends to the corona on the distal shaft. The glandular urethra is formed by canalization of an ectodermal cord that has grown through the glans to communicate with the fused urethral folds (see Chapter 2). Hypospadias results when fusion of the urethral folds is incomplete.

Hypospadias occurs in 1 in every 300 male children. Estrogens and progestins given during pregnancy are known to increase the incidence. Although a familial pattern of hypospadias has been recognized, no specific genetic traits have been established.

**Classification**

There are several forms of hypospadias, classified according to location: (1) glandular, that is, opening on the proximal glans penis; (2) coronal, that is, opening at the coronal sulcus; (3) penile shaft; (4) penoscrotal; and (5) perineal. About 70% of all cases of hypospadias are distal penile or coronal.

Hypospadias in the male is evidence of feminization. Patients with penoscrotal and perineal openings should be considered to have potential intersex problems requiring appropriate evaluation. Hypospadiac newborns should not be circumcised, because the preputial skin may be useful for future reconstruction.

**Clinical Findings**

**A. Symptoms and Signs**

Although newborns and young children seldom have symptoms related to hypospadias, older children and adults may complain of difficulty directing the urinary stream and stream spraying. Chordee (curvature of the penis) causes ventral bending and bowing of the penis shaft, which may prevent sexual intercourse. Perineal or penoscrotal hypospadias necessitates voiding in the sitting position, and these proximal forms of hypospadias in adults can be the cause of infertility. An additional complaint of almost all patients is the abnormal (hooded) appearance of the penis, caused by deficient or absent ventral foreskin. The hypospadiac meatus may be stenotic and
Figure 40–3. Hypospadias and epispadias. **Upper left:** Hypospadias, penoscrotal type. Redundant dorsal foreskin that is deficient ventrally; ventral chordee. **Upper right:** Hypospadias, midscrotal type. Chordee more marked. Penis often small. **Lower left:** Epispadias. Redundant ventral foreskin that is absent dorsally; severe dorsal chordee. **Lower right:** Traction on foreskin reveals dorsal defect.
should be carefully examined and calibrated. (A meato- 

tomy should be done when stenosis exists.) There is an 

increased incidence of undescended testicles in children 

with hypospadias; scrotal examination is necessary to 

establish the position of the testicles.

B. LABORATORY, X-RAY, & ENDOSCOPIC FINDINGS

Since children with penoscrotal and perineal hypospadias 

often have a bifid scrotum and ambiguous genitalia, a bucc- 

cal smear and karyotyping are indicated to help establish 

the genetic sex. Urethroscopy and cystoscopy are of value 

to determine whether internal male sexual organs are nor- 
mally developed. Excretory urography is also indicated in 

these patients to detect additional congenital anomalies of 

the kidneys and ureters.

Some authors recommend routine use of excretory 

urography for all patients with hypospadias; however, this 

seems to be of little value in the more distal types of the 

disorder, because there appears to be no increased inci- 
dence of upper urinary tract anomalies.

Differential Diagnosis

Any degree of hypospadias is an expression of feminiza-

tion. Perineal and scrotal urethral openings should be care-

fully evaluated to ascertain that the patient is not a female 

with androgenized adrenogenital syndrome. Urethroscopy 

and cystoscopy will aid in evaluating the development of 

internal reproductive organs.

Treatment

For psychological reasons, hypospadias should be repaired 

before the patient reaches school age; in most cases, this 

can be done before age 2.

More than 150 methods of corrective surgery for 

hypospadias have been described. Currently, 1-stage 

repairs with foreskin island flaps and incised urethral 

plate are performed by more and more urologists. It now 
appears that buccal mucosa grafts are more advantageous 

than others and should be considered the primary graft-

ing technique when indicated. Fistulas occur in 15–30% 

of patients, but the fistula repair is considered a small, 

second-stage reconstruction.

All types of repair involve straightening the penis by 

removal of the chordee. The chordee removal can be con-

firmed by producing an artificial erection in the operating 

room following urethral reconstruction and advancement. 

Most successful techniques for repair of hypospadias use 

local skin and foreskin in developing the neourethra. In 

recent years, advancement of the urethra to the glans penis 
has become technically feasible and cosmetically acceptable.

Prognosis

After corrective surgery, most patients are able to void in 

the standing position as well as to deposit semen into the 
vagina. The overall cosmetic appearance and the preven-
tion of fistula formation remain the greatest challenges in 

these repairs.

CHORDEE WITHOUT HYPOSPADIAS

Congenital ventral chordee without hypospadias is seen 
ocasionally and is caused by a short urethra, fibrous tis-
sues surrounding the corpus spongiosum, or both. The 
urethral opening is in the normal position on the glans 
penis; only with erection does the penis bow, thus prevent-
ing satisfactory vaginal penetration. During examination, if 
the patient cannot achieve an erection naturally, erection 
can be induced by injecting saline solution into the corpus 
cavernosum after placing a tourniquet at the base of the 
penis. This technique should also be used during corrective 
surgery to be certain that the penis will be straight after 
the operation.

If the penis is adequate in length, the dorsal surface can 
be shortened (1) by excising elliptic portions of the tunica 
albuginea on the dorsum of the penis on either side of the 
midline or (2) by making transverse cuts in a similar posi-
tion and then closing them longitudinally, thus shortening 
the dorsum. Fibrous tissue found in association with the 
urethra and corpus spongiosum should be totally excised.

EPISPADIAS

The incidence of complete epispadias is approximately 1 in 
120,000 males and 1 in 450,000 females. The urethra is 
displaced dorsally, and classification is based on its position 
in males. In glandular epispadias, the urethra opens on the 
dorsal aspect of the glans, which is broad and flattened. In 
the penile type, the urethral meatus, which is often broad 
and gaping, is located between the pubic symphysis and 
the coronal sulcus. A distal groove usually extends from the 
meatus through the splayed glans. The penopubic type has 
the urethral opening at the penopubic junction, and the 
entire penis has a distal dorsal groove extending through 
the glans.

Patients with glandular epispadias seldom have urinary 
incontinence. However, with penopubic and penile epis-

padias, incontinence is present in 95% and 75% of cases, 
respectively.

Females with epispadias have a bifid clitoris and separa-
tion of the labia. Most are continent.

Urinary incontinence is a common problem because of 
maldevelopment of the urinary sphincters. Dorsal curva-
ture of the penis (dorsal chordee) is also present (Figure

40–3). The pubic bones are separated as in extrophy of 
the bladder. Epispadias is a mild form of bladder extro-

phy, and in severe cases, extrophy and epispadias coexist.

Surgery is required to correct the incontinence, remove 
the chordee to straighten the penis, and extend the urethra 
out onto the glans penis. Repair of the urinary sphincter 
has not been very successful. Chordee excision and ure-
ththroplasty with advancement of the meatus have been successful in achieving acceptable cosmetic and functional results. Bladder augmentation combined with the artificial sphincter may be required in patients in whom incontinence cannot be corrected.

### ACQUIRED DISEASES & DISORDERS OF THE PENIS & MALE URETHRA

#### PRIAPISM

Priapism is an uncommon condition of prolonged erection. It is usually painful for the patient, and no sexual excitement or desire is present. The disorder is idiopathic in 60% of cases, while the remaining 40% of cases are associated with diseases (eg, leukemia, sickle cell disease, pelvic tumors, pelvic infections), penile trauma, spinal cord trauma, or use of medications (trazodone). Currently, intracavernous injection therapy for impotence may be the most common cause. Although the idiopathic type often is initially associated with prolonged sexual stimulation, cases of priapism due to the other causes are unrelated to psychic sexual excitement.

Priapism may be classified into high- and low-flow types. High-flow priapism (nonischemic) usually occurs secondary to perineal trauma, which injures the central penile arteries and results in loss of penile blood-flow regulation. Aneurysms of one or both central arteries have been observed. Aspiration of penile blood for blood-gas determinations demonstrates high oxygen and normal carbon dioxide levels. Arteriography is useful to demonstrate aneurysms that will respond to embolization; erectile function is usually preserved.

The patient with low-flow priapism (ischemic) usually presents with a history of several hours of painful erection. The glans penis and corpus spongiosum are soft and unininvolved in the process. The corpora cavernosa are tense with congested blood and tender to palpation. The current theories regarding the mechanism of priapism remain in debate, but most authorities believe the major abnormality to be physiologic obstruction of the venous drainage. This obstruction causes buildup of highly viscous, poorly oxygenated blood (low O₂, high CO₂) within the corpora cavernosa. If the process continues for several days, interstitial edema and fibrosis of the corpora cavernosa will develop, causing impotence.

Ischemic priapism must be considered a urologic emergency. Epidural or spinal anesthesia can be used. The sludged blood can then be evacuated from the corpora cavernosa through a large needle placed through the glans. The addition of adrenergic agents administered via intracavernous irrigation has proved helpful. Monitoring intracavernous pressure ensures that recurrence is not imminent. Multiple wedges of tissue can be removed with a biopsy needle to create a shunting fistula between the glans penis and corpora cavernosa. This technique, which has been very successful, provides an internal fistula to keep the corpora cavernosa decompressed. To maintain continuous fistula drainage, pressure should be exerted intermittently (every 15 minutes) on the body of the penis. The patient can do this manually after he has recovered from anesthesia.

If the shunt described fails, another shunting technique may be used by anastomosing the superficial dorsal vein to the corpora cavernosa. Other effective shunting methods are corpora cavernosa to corpus spongiosum shunt by perineal anastomosis; saphenous vein to corpora cavernosa shunt; and pump decompression.

Patients with sickle cell disease have benefited from massive blood transfusions, exchange transfusions, or both. Hyperbaric oxygen also has been suggested for these patients. Patients with leukemia should receive prompt chemotherapy. Appropriate management of any underlying cause should be instituted without delay. Such treatment should not prevent aggressive management of the priapism if the erection persists for several hours.

Impotence is the worst sequel of priapism. It is more common after prolonged priapism (several days). Early recognition (within hours) and prompt treatment of priapism offer the best opportunity to avoid this major problem.

#### PEYRONIE’S DISEASE

Peyronie’s disease (plastic induration of the penis) was first described in 1742 and is a well-recognized clinical problem affecting middle-aged and older men. Patients present with complaints of painful erection, curvature of the penis, and poor erection distal to the involved area. The penile deformity may be so severe that it prevents satisfactory vaginal penetration. The patient has no pain when the penis is in the nonerect state.

Examination of the penile shaft reveals a palpable, dense, fibrous plaque of varying size involving the tunica albuginea. The plaque is usually near the dorsal midline of the shaft. Multiple plaques are sometimes seen. In severe cases, calcification and ossification are noted and confirmed by radiography. Although the cause of Peyronie’s disease remains obscure, the dense fibrous plaque is microscopically consistent with findings in severe vasculitis. The condition has been noted in association with Dupuytren’s contracture of the tendons of the hand, in which the fibrosis resembles that of Peyronie’s disease when examined microscopically.

Spontaneous remission occurs in about 50% of cases. Initially, observation and emotional support are advised. If remission does not occur, p-aminobenzoic acid powder or...
PHIMOSIS

Phimosis is a condition in which the contracted foreskin cannot be retracted over the glans. Chronic infection from poor local hygiene is its most common cause. Most cases occur in uncircumcised males, although excessive skin left after circumcision can become stenotic and cause phimosis. Calculi and squamous cell carcinoma may develop under the foreskin. Phimosis can occur at any age. In diabetic older men, chronic balanoposthitis may lead to phimosis and may be the initial presenting complaint. Children under 2 years of age seldom have true phimosis; their relatively narrow preputial opening gradually widens and allows for normal retraction of foreskin over the glans. Circumcision for phimosis should be avoided in children requiring general anesthesia; except in cases with recurrent infections, the procedure should be postponed until the child reaches an age when local anesthesia can be used.

Edema, erythema, and tenderness of the prepuce and inflammation has subsided.

A decrease in urinary stream is the most common complaint. Spraying or double stream is often noted, as is postvoiding dribbling. Chronic urethral discharge, occasionally a major complaint, is likely to be associated with chronic prostatitis. Acute cystitis or symptoms of infection are seen at times. Acute urinary retention seldom occurs unless infection or prostatic obstruction develops. Urinary frequency and mild dysuria may also be initial complaints. Induration in the area of the stricture may be palpable. Tender enlarged masses along the urethra usually represent

CIRCUMCISION

Although circumcision is routinely performed in some countries for religious or cultural reasons, it is usually not necessary if adequate penile cleanliness and good hygiene can be maintained. There is a higher incidence of penile carcinoma in uncircumcised males, but chronic infection and poor hygiene are usually underlying factors in such instances. Circumcision is indicated in patients with infection, phimosis, or paraphimosis (see preceding sections).

URETHRAL STRICTURE

Acquired urethral stricture is common in men but rare in women. (Congenital urethral stricture is discussed earlier in the chapter.) Most acquired strictures are due to infection or trauma. Although gonococcal urethritis is seldom a cause of stricture today, infection remains a major cause—particularly infection from long-term use of indwelling urethral catheters. Large catheters and instruments are more likely than small ones to cause ischemia and internal trauma. External trauma, for example, pelvic fractures (see Chapter 17) can partially or completely sever the membranous urethra and cause severe and complex strictures. Straddle injuries can produce bulbar strictures.

Urethral strictures are fibrotic narrowings composed of dense collagen and fibroblasts. Fibrosis usually extends into the surrounding corpus spongiosum, causing spongiolysis. These narrowings restrict urine flow and cause dilation of the proximal urethra and prostatic ducts. Prostatitis is a common complication of urethral stricture. The bladder muscle may become hypertrophic, and increased residual urine may be noted. Severe, prolonged obstruction can result in decompensation of the ureterovesical junction, reflux, hydronephrosis, and renal failure. Chronic urinary stasis makes infection likely. Urethral fistulas and periurethral abscesses commonly develop in association with chronic, severe strictures.

Clinical Findings

A. SYMPTOMS AND SIGNS

A decrease in urinary stream is the most common complaint. Spraying or double stream is often noted, as is postvoiding dribbling. Chronic urethral discharge, occasionally a major complaint, is likely to be associated with chronic prostatitis. Acute cystitis or symptoms of infection are seen at times. Acute urinary retention seldom occurs unless infection or prostatic obstruction develops. Urinary frequency and mild dysuria may also be initial complaints. Induration in the area of the stricture may be palpable. Tender enlarged masses along the urethra usually represent
periurethral abscesses. Urethrocystic fistulas may be present. The bladder may be palpable if there is chronic retention of urine.

**B. LABORATORY FINDINGS**

If urethral stricture is suspected, urinary flow rates should be determined. The patient is instructed to accumulate urine until the bladder is full and then begin voiding; a 5-second collection of urine should be obtained during midstream maximal flow and its volume recorded. After the patient repeats this procedure 8–10 times over several days in a relaxed atmosphere, the mean peak flow can be calculated. With strictures creating significant problems, the flow rate will be less than 10 mL/s (normal 20 mL/s).

Urine culture may be indicated. The midstream specimen is usually bacteria free, with some pyuria [8–10 white blood cells (leukocytes) per high-power field] in a carefully obtained first aliquot of urine. If the prostate is infected, bacteria will be present in a specimen obtained after prostatic massage. In the presence of cystitis, the urine will be grossly infected.

**C. X-RAY FINDINGS**

A urethrogram or voiding cystourethrogram (or both) will demonstrate the location and extent of the stricture. Sonography has also been a useful method of evaluating the urethral stricture. Urethral fistulas and diverticula are sometimes noted. Vesical stones, trabeculations, or diverticula may also be seen.

**D. INSTRUMENTAL EXAMINATION**

Urethroscopy allows visualization of the stricture. Small-caliber strictures prevent passage of the instrument through the area. Direct visualization and sonourethrography aid in determining the extent, location, and degree of scarring. Additional areas of scar formation adjacent to the stricture may be detected by urethroscopy.

The stricture can be calibrated by passage of bougies à boucle.

**Differential Diagnosis**

Benign or malignant prostatic obstruction can cause symptoms similar to those of stricture. After prostatic surgery, bladder neck contracture can develop and induce stricture-like symptoms. Rectal examination and panendoscopy adequately define such abnormalities of the prostate. Urethral carcinoma is often associated with stricture; urethroscopy demonstrates a definite irregular lesion, and biopsy establishes the diagnosis of carcinoma.

**Complications**

Complications include chronic prostatitis, cystitis, chronic urinary infection, diverticula, urethrocystic fistulas, periurethral abscesses, and urethral carcinoma. Vesical calculi may develop from chronic urinary stasis and infection.

**Treatment**

**A. SPECIFIC MEASURES**

1. **Dilation**—Dilation of urethral strictures is not usually curative, but it fractures the scar tissue of the stricture and temporarily enlarges the lumen. As healing occurs, the scar tissue reforms.

   Dilation may initially be required because of severe symptoms of chronic retention of urine. The urethra should be liberally lubricated with a water-soluble medium before instrumentation. A filiform is passed down the urethra and gently manipulated through the narrow area into the bladder. A follower can then be attached (see Chapter 10) and the area gradually dilated (with successively larger sizes) to approximately 22F. A 16F silicone catheter can then be inserted. If difficulty arises in passing the filiform through the stricture, urethroscopy should be used to guide the filiform under direct vision.

   An alternative method of urethral dilation employs Van Buren sounds. These instruments are best used by an experienced urologist familiar with the size and extent of the stricture involved. First, a 22F sound should be passed down to the stricture site and gentle pressure applied. If this fails, a 20F sound should be used. Smaller sounds should be used with care, because they can easily perforate the urethral wall and produce false passages. Bleeding and pain are major problems caused by dilation.

2. **Urethrotomy under endoscopic direct vision**—Lysis of urethral strictures can be accomplished using a sharp knife attached to an endoscope. The endoscope provides direct vision of the stricture during cutting. A filiform should be passed through the stricture and used as a guide during lysis. The stricture is usually incised circumferentially with multiple incisions. A 22F instrument should pass with ease. A catheter is left in place for a short time to prevent bleeding and pain. Results of this procedure have been satisfactory in short-term follow-up in 70–80% of patients, but long-term success rates are much lower. The procedure has several advantages: (1) Minimal anesthesia is required—in some cases, only topical anesthesia combined with sedation; (2) it is easily repeated if the stricture recurs; and (3) it is very safe, with few complications.

3. **Surgical reconstruction**—If urethroscopy under direct vision fails, open surgical repair should be performed. Short strictures (<2 cm) of the anterior urethra should be completely excised and primary anastomosis done. If possible, the segment to be excised should extend 1 cm beyond each end of the stricture to allow for removal of any existing spongiosfibrosis and improve postoperative healing.

   Strictures >2 cm in length can be managed by patch graft urethroplasty. The urethra is incised in the midline...
for the full length of the stricture plus an additional 0.5 cm proximal and distal to its ends. A full-thickness skin graft is obtained—preferably from the penile skin or buccal mucosa—and all subcutaneous tissue is carefully removed. The graft is then tailored to cover the defect and meticulously sutured into place (Figure 40–4).

In very long, densely fibrotic strictures, the distal penile fasciocutaneous flap technique has been successful in >80% cases. This single-stage procedure can be combined with buccal mucosa grafting in panurethral strictures. In adults, grafts from buccal mucosa or penile skin should be applied with an onlay technique in the bulbar region of the urethra to maximize graft vascularization from the corpus spongiosum.

Strictures involving the membranous urethra ordinarily result from external trauma (see Chapter 17) and present problems in reconstruction. Most can be corrected by a perineal approach with excision of the urethral rupture defect and direct anastomosis of the bulbar urethra to the prostatic urethra (Figure 40–5). At times, partial pubectomy from the perineal approach can be done to improve urethral approximation without tension on the anastomosis. Rarely, total pubectomy combined with the perineal approach is required to accomplish the direct end-to-end anastomosis.

These single-stage procedures have a high success rate and create a urethra free of hair—a major problem seen with 2-stage procedures. Although seldom required, 2-stage procedures are important reconstructive techniques to be considered in complex urethral strictures.

B. TREATMENT OF COMPLICATIONS

Urinary tract infection in patients with strictures requires specific antimicrobial therapy, followed by long-term prophylactic therapy until the stricture has been corrected. Periurethral abscesses require drainage and use of antimicrobial drugs. Urethral fistulas usually require surgical repair.

Prognosis

A stricture should not be considered “cured” until it has been observed for at least 1 year after therapy, since it may not be fully healed.

Figure 40–4. Left: Urethrogram demonstrating multiple anterior urethral strictures. Right: Voiding cystourethrogram following a patch skin graft of 14 cm in the same patient. There are no residual strictures.

Figure 40–5. Voiding urethrogram following repair of traumatic posterior urethral stricture. Arrow indicates that area of repair is stricture free.
recur at any time during that period. Urinary flow rate measurements and urethrograms are helpful to determine the extent of residual obstruction.

**URETHRAL CONDYLOMATA ACUMINATA (URETHRAL WARTS)**

Condylomata acuminata are uncommon in the urethra and are almost always preceded by lesions on the skin. They are wart-like papillomas caused by a papilloma virus and are usually transmitted by direct sexual contact but may be transmitted nonsexually.

Patients commonly complain of bloody spotting from the urethra and occasionally have dysuria and urethral discharge. Examination of the urethral meatus often reveals a small, protruding papilloma. If a lesion is not found in this location, the meatus should be separated with the examining fingers so that the distal urethra can be inspected. About 90% of such lesions are situated in the distal urethra. Complete urethroscopy must be done to be certain other lesions do not exist.

Lesions of the meatus can be treated by local excision. A local anesthetic is applied to the area at the base of the lesion, and the pedunculated lesions are sharply incised with small scissors. The area is then fulgurated by electrocautery. Meatotomy may be indicated for excision of lesions in the fossa navicularis and glandular urethra.

Deeper lesions may be fulgurated transurethrally with a resectoscope or Bugby electrode. Recently, lesions have been successfully destroyed using a carbon dioxide or holmium laser. Laser therapy does minimal damage to the urethral mucosa, and stricture formation seems less likely with its use.

Multiple lesions have also been treated with fluorouracil, 5% solution or cream. The drug is instilled in the urethra for 20 minutes twice a week for 5 weeks. Care must be taken to protect the penile skin and scrotum from coming in contact with the medication, since it may produce severe irritation.

Lesions may become infected and ulcerated. This suggests carcinoma, and histopathologic confirmation of the diagnosis should be obtained. Rarely, giant condylomata (Buschke-Löwenstein tumors) involving the glans penis and often the urethra may be seen. Such lesions suggest carcinoma and a biopsy must be done. Surgical excision is the treatment of choice.

To prevent recurrence of condylomata acuminata, the sexual partner must also be examined and treated if necessary.

**STENOSIS OF THE URETHRAL MEATUS**

Newborns are often suspected of having meatal stenosis of some degree. This condition is thought to be secondary to ammonia dermatitis following circumcision and resulting in prolonged irritative meatitis.

Calibration is important, since the visual appearance of the meatus does not correlate well with its actual size. The urethra should easily accept the tip of an 8F pediatric feeding tube. The significance of meatal stenosis is debated, but a meatal caliber <5F in children <10 years of age is an indication for meatotomy.

**PENILE PHLEbothROMBOSIS & LYMPHATIC OCCLUSION**

Superficial veins and lymphatic vessels of the dorsal penile shaft just proximal to the corona may become irritated and inflamed. A careful history usually indicates that minor trauma to the area (eg, from prolonged sexual intercourse) has occurred. Examination reveals a tender, indurated, cord-like structure on the distal penile shaft. Slight erythema may be present.

For clinical purposes, there is no need to distinguish lymphatic and venous causes, since both penile phlebothrombosis and lymphatic occlusion will resolve spontaneously. The patient must be reassured.

**REFERENCES**

**Congenital Anomalies**

**Penis & Urethra**


**Hypospadias**


**Epispadias**


**Acquired Diseases & Disorders**


**Priapism**


**Peyronie’s Disease**


**Paraphimosis**


**Circumcision**


**Urethral Stricture**


**Urethral Condylomata Acuminata**


**Penile Thrombophlebitis & Lymphatic Occlusion**


CONGENITAL ANOMALIES OF THE FEMALE URETHRA

DISTAL URETHRAL STENOSIS IN INFANCY & CHILDHOOD (SPASM OF THE EXTERNAL URINARY SPHINCTER) AND DYSFUNCTIONAL VOIDING

There has been considerable confusion about the site of lower tract obstruction in young girls who have enuresis, a slow and interrupted urinary stream, recurrent cystitis, and pyelonephritis, and who, on thorough examination, often exhibit vesicoureteral reflux. Treatment has been directed largely to the bladder neck on rather empiric grounds. Most of these children, however, have congenital distal urethral stenosis with secondary spasm of the striated external sphincter rather than bladder neck obstruction due to functional or organic causes.

At birth, calibration of the urethra with bougies à boule reveals no evidence of a distal ring of urethral stenosis (Fisher et al, 1969). Within a few months, however, such a ring develops as a normal anatomic structure. After puberty, the ring disappears. The inference is that the absence of estrogens leads to the development of this lesion. Lyon and Tanagho (1965) found that the ring calibrates at 14F at age 2 and at 16F between the ages of 4 and 10. Even though from the hydrodynamic standpoint such a stenotic area should not be obstructive, almost all observers agree that dilatation of the ring does relieve symptoms in these children and that it results in cure or amelioration of persistent infection or vesical dysfunction in 80% of cases. Lyon and Tanagho thought it possible that the basic cause of these urinary difficulties might be reflex spasm of periurethral striated sphincter and noted that voiding cystourethrograms supported that view (Figure 41–1).

Tanagho et al (1971) measured pressures in the bladder and in the proximal and mid urethra simultaneously in symptomatic girls and found high resting pressures, some as high as 200 cm of water (normal, 100 cm of water) in the midurethral segment. Attempts at voiding caused intravesical pressures as high as 225 cm of water (normal, 30–40 cm of water) to develop. Under curare, the urethral closing pressures dropped to normal (40–50 cm of water), proving that these obstructing pressures were caused by spasm of the striated sphincter muscle. If the distal urethral ring was treated and symptoms abated, repeat pressure studies showed normal midurethral and intravesical voiding pressures. If, on the other hand, symptoms persisted, pressures were found to remain at extremely high levels. It seems clear, therefore, that the major cause of urinary problems in young girls is spasm of the external sphincter and not vesical neck stenosis (Smith, 1969).

In addition to recurrent urinary tract infections, these patients have hesitancy in initiating micturition and a slow, hesitant, or interrupted urinary stream. Enuresis and involuntary loss of urine during the day are common complaints. Abdominal straining may be required in order to void. Small amounts of residual urine are found, which impair the vesical defense mechanism. A voiding cystourethrogram may reveal an open bladder neck and ballooning of the proximal urethra secondary to spasm of the external sphincter (Figure 41–1).

The voiding cystourethrogram may reveal evidence of the distal ring, but typical findings are not always seen, particularly if the flow rate is slow. Definitive diagnosis is made by bougienage.

Historically, the simplest and least harmful treatment is overdilatation with sounds up to 32–36F or with the Kollmann dilator (Lyon and Tanagho, 1965; Lyon and Marshall, 1971; Hendry, Stanton, and Williams, 1973). With either method, the ring “cracks” anteriorly, with some bleeding. Recurrence of the ring is rare. Internal urethrotomy has its proponents (Immergut and Gilbert, 1973; Hradec et al, 1973), but Kaplan, Sammons, and King (1973) found that results with urethrotomy were poor, since incising the urethra along its entire length does not cut the external sphincter, whose abnormal tone is the cause of the obstruction, whereas “cracking” the ring by overdilatation accomplishes this purpose.

It has been reported in the past that, in 80% of affected children, destruction of the ring of distal urethral stenosis helps to overcome enuresis and achieve a normal free voiding pattern and to cure recurrent cystitis or persistent bacteriuria (Lyon and Marshall, 1971). Spontaneous resolution of reflux is possible only in the case of “borderline” values that tend to give way in the presence of increased voiding pressure and infection.
Since the ring normally disappears at puberty, it is possible to await spontaneous cure; however, the ring should be broken if symptoms have been severe enough to bring the child to the attention of a urologist.

More recently, it has been recognized that many of these symptoms are due to functional, rather than neurologic or anatomic, causes of obstruction. Children must achieve adult patterns of urinary control and disturbances, especially around the time of toilet-training. This may cause a variety of symptoms that result from voiding against a voluntarily closed urethral sphincter (dysfunctional voiding). These range from severe functional obstruction with urinary retention, altered bladder anatomy, and vesicoureteral reflux (known as Hinman’s syndrome) to less severe incomplete control of urination. They are often accompanied by bowel symptoms, such as constipation or encopresis. Treatment requires bladder retraining, including psychologic help and biofeedback, as well as restoration of normal bowel habits, including dietary changes and laxatives.

LABIAL FUSION (SYNECHIA VULVAE)

Some children with recurring urinary infection are found to have fused labia minora, which are apt to obstruct the flow of urine, causing it to pool in the vagina. Local application of estrogen cream twice daily for 2–4 weeks usually causes spontaneous separation, with minimal side effects (Leung et al, 2005; Aribag, 1975). Forceful separation or dissection has its advocates (Christensen and Oster, 1971).

It is uncommonly seen as an acquired disease after puberty, caused by genital trauma (sexual abuse, vaginal delivery, surgery, etc) (Kumar et al, 2006). In cultures where “female circumcision” is performed, this may be a relatively common complication (Adekunle et al, 1999).
ACQUIRED DISEASES OF THE FEMALE URETHRA

ACUTE URETHRITIS

Acute urethritis frequently occurs with gonorrheal (Neisseria gonorrhoeae) or trichomoniasis (Trichomonas vaginalis) infection in women, and may less commonly occur with infection by Chlamydia trachomatis (approximately 25% of cases are symptomatic). Urinary symptoms are often present at the onset of the disease. Cultures and smears establish the diagnosis. Prompt cure can be achieved with antimicrobial drugs, usually to cover both gonorrhea and chlamydia, such as a combination of intramuscular ceftriaxone and oral azithromycin or doxycycline. Treatment is important, as 40% of women with untreated Chlamydia infections will have pelvic inflammatory disease, which may lead to ectopic pregnancy, pelvic pain, and infertility (Simms and Stephenson, 2000).

The detergents in bubble bath and some spermicidal jellies may cause vaginitis and urethritis. Symptoms of vesical irritability may occur (Bass, 1968; Marshall, 1965).

CHRONIC URETHRITIS

Chronic urethritis is one of the most common urologic problems of females. The distal urethra normally harbors pathogens, and the risk of infection may be increased by wearing contaminated diapers, by insertion of an indwelling catheter, by spread from cervical or vaginal infections, or by intercourse with an infected partner. Urethral inflammation may also occur from the trauma of intercourse or childbirth, particularly if urethral stenosis, either congenital or following childbirth, is present.

Clinical Findings

The urethral mucosa is reddened, quite sensitive, and often stenotic. Granular areas are often seen, and polypoid masses may be noted just distal to the bladder neck.

A. Symptoms

The symptoms resemble those of cystitis, although the urine may be clear. Complaints include burning on urination, frequency, and nocturia. Discomfort in the urethra may be felt, particularly when walking.

B. Signs

Examination may disclose redness of the meatus, hyper- sensitivity of the meatus and of the urethra on vaginal palpation, and evidence of cervicitis or vaginitis. There is no urethral discharge.

C. Laboratory Findings

When the initial and midstream urine are collected in separate containers, the first glass contains pus and the second does not (Marshall, Lyon, and Schieble, 1970). Ureaplasma urealyticum (formerly called T strains of mycoplasmas) is often identifiable in the first glass. These findings are similar to those of nongonococcal (chlamydial) urethritis in males. Clinically, the presence of white blood cells (leukocytes) in the absence of bacteria on a routine stain or culture suggests nongonococcal urethritis. In other cases, various bacteria (eg, Streptococcus faecalis, Escherichia coli) may be cultured from both the urethral washings and a specimen taken from the introitus.

D. Instrumental Examination

A catheter, bougie à boule, or sound may meet resistance because of urethral stenosis. Panendoscopy reveals redness and a granular appearance of the mucosa (Krieger, 1988). Inflammatory polyps may be seen in the proximal portion of the urethra. Cystoscopy may show increased injection of the trigone (trigonitis), which often accompanies urethritis.

Differential Diagnosis

Differentiation of urethritis from cystitis depends on bacteriologic study of the urine; panendoscopy demonstrates the urethral lesion. Both urethritis and cystitis may be present. Chronic noninflammatory urethritis may be a manifestation of psychic stressors. Patients with anxiety or other temporary or chronic psychologic disorders may present with symptoms that are very suggestive of urethritis. Alternatively, women with long-standing symptoms may have these symptoms as an adult version of childhood voiding dysfunction (see earlier discussion).

Treatment & Prognosis

Gradual urethral dilatations (up to 36F in adults) are indicated for urethral stenosis; this allows for some inevitable contracture. Immergut and Gilbert (1973) prefer internal urethrotomy (Farrar, 1980). U. urealyticum and Chlamydial urethritis usually respond to doxycycline or azithromycin. For ascending bacterial infections, Bruce et al (1973) recommend the regular, local application of an antiseptic (eg, hexachlorophene, chlorhexidine cream) to the introitus in order to prevent bacteria from the area of the perineum, vagina, and vulva from reinfecting the urethra. However, other studies suggest that these and other types of douching practices may increase the rate of infectious complications (Simpson et al, 2004).

SENILE URETHRITIS

After physiologic (or surgical) menopause, hypoestrogenism occurs and retrogressive (senile) changes take place
in the vaginal epithelium, so that it becomes rather dry and pale (Smith, 1972); atrophic vaginitis may affect 20–30% of postmenopausal women. Similar changes develop in the lower urinary tract, which arises from the same embryologic tissues as the female reproductive organs. Some eversion of the mucosa about the urethral orifice, from atrophy of the vaginal wall, is usually seen. This is commonly misdiagnosed as caruncle.

Clinical Findings

A. Symptoms

Many postmenopausal women have symptoms of vesical irritability (burning, frequency, urgency) and stress incontinence. Dysuria may occur due to urine contact with the inflamed atrophic tissues themselves or because of the increased incidence of urinary tract infections in these women. They may complain of vaginal and vulval itching, discharge, dyspareunia, and may have bloody vaginal spotting, especially after intercourse.

B. Signs

The vaginal epithelium is dry and pale. The mucosa at the urethral orifice is often reddened and hypersensitive; eversion of its posterior lip from atrophy of the urethrovaginal wall is common. Atrophic vaginitis also increases the risk for urinary tract infections, and approximately 10–15% of women over 60 years of age have frequent urinary tract infections.

C. Laboratory Findings

The urine is usually free of microorganisms. The diagnosis can be made by the following procedure: A dry smear of vaginal epithelial cells is stained with Lugol’s solution. The slide is then washed with water and immediately examined microscopically while wet. In hypoestrogenism, the cells take up the iodine poorly and are therefore yellow. When the mucosa is normal, these cells stain a deep brown because of their glycogen content. The diagnosis may also be confirmed by a Papanicolaou smear. Postmenopausal status is associated with a higher vaginal pH, a decrease in vaginal lactobacillus colonization, and increased colonization with *E. coli*.

D. Instrumental Examination

Panendoscopy usually demonstrates a reddened and granular urethral mucosa. Some urethral stenosis may be noted.

Differential Diagnosis

Senile urethritis is often mistaken for urethral caruncle. Eversion of the posterior lip of the urinary meatus is evident in both conditions; however, a hypersensitive vascular tumor is not present in senile urethritis. Before operations to relieve stress incontinence are performed, estrogen (or androgen) therapy should be tried.

**Treatment**

Senile urethritis responds well to diethylstilbestrol vaginal suppositories, 0.1 mg nightly for 3 weeks. Estrogen creams applied locally are also effective. Estrogen urethral suppositories have been recommended, but they offer no advantages and are difficult to insert. Three or more courses are occasionally indicated, depending on the symptoms and the appearance of the vaginal smear stained as outlined previously.

An estradiol vaginal ring which delivers low level of estradiol over a 3-month period is also available. Local application has the advantage of minimal change of systemic blood level and thus avoids the side effects associated with systemic hormone therapy.

**Prognosis**

Topical estriol vaginal cream is an effective treatment in postmenopausal women with recurrent infections (Quinlivan, 1965). In one study, patients treated with the estriol cream averaged 0.5 infections per year, compared with about 6.0 infections per year in women who were not treated (Raz and Stamm, 1993).

**URETHRAL CARUNCLE**

Urethral caruncle is a benign, red, raspberry-like, friable vascular tumor involving the posterior lip of the external urinary meatus. It is rare before the menopause. Microscopically, it consists of connective tissue containing many inflammatory cells and blood vessels and is covered by an epithelial layer (Lee, 1995).

Clinical Findings

Symptoms include pain on urination, pain with intercourse, and bloody spotting from even mild trauma. A sessile or pedunculated red, friable, tender mass is seen at the posterior lip of the meatus.

Differential Diagnosis

Carcinoma of the urethra may involve the urethral meatus. Palpation reveals definite induration. Biopsy establishes the true diagnosis. Senile urethritis is often associated with a polypoid reaction of the urinary meatus and in fact is the most common cause of masses in this region. The diagnosis can be made by verifying the patient’s hypoestrogenic status and by demonstrating a favorable response to estrogen replacement therapy. Biopsy should be done if doubt exists (Young, 1996; Neilson, 1989).

Thrombosis of the urethral vein presents as a bluish, swollen, tender lesion involving the posterior lip of the urin-
nary meatus. It has the appearance of the thrombosed hemorrhoid. It subsides without treatment.

**Treatment**

Local excision is indicated only if symptoms are troublesome.

**Prognosis**

True caruncle is usually cured by excision, but in a few instances it does recur.

**Prolapse of the Urethra**

Prolapse of the female urethra is not common. It usually occurs only in children or in paraplegics suffering from a lower motor neuron lesion. The protruding urethral mucosa presents as an angry red mass that may become gangrenous if it is not reduced promptly (Kleinjan, 1996). When a young girl has a protruding mass, urethral prolapse must be differentiated from prolapse of a ureterocele. (Valerie et al, 1999; Fernandes et al, 1993).

Again reduction, cystoscopy should be done to rule out ureterocele. Recurrences are rare following reduction; the accompanying inflammation probably “fixes” the tissue in place as healing progresses. If the prolapsed urethra cannot be reduced or if it recurs, an indwelling catheter should be inserted, traction placed on it, and a heavy piece of suture material tightly tied over the tissue and catheter just proximal to the mass. The tissue later sloughs off. Using this same technique, the tissue can be resected, preferably with an electrosurgical cautery (Devine, 1980).

**URETHROVAGINAL FISTULA**

Urethrovaginal fistulas may follow local injury secondary to fracture of the pelvis or obstetric or surgical injury (see Chapter 39). A common cause in the industrial world is accidental trauma to the urethra or its blood supply in the course of surgical repair of a cystocele or excision of urethral diverticula. In the developing world, obstructed labor is the most common cause (Elkins, 1994). Other causes may include radiation therapy for pelvic malignancy, trauma/pelvic fracture, and vaginal neo-plasms. Diagnosis can usually be made on physical examination and urethroscopy. Vaginal urethroplasty is indicated, and these surgeries may be very challenging due to poor quality local tissue; often, 2 or more procedures may be necessary (Webster et al, 1984).

**URETHRAL DIVERTICULUM**

The incidence of diverticulation of the urethral wall is between 0.6–5% (Andersen, 1967; Davis and Robinson, 1970). Diverticula are at times multiple. Most cases are probably secondary to obstetric urethral trauma or severe urethral infection. A few cases of carcinoma in such diverticula have been reported (Marshall, 1977; Nakamura et al, 1995; Kato, 1998). Urethral diverticula are usually associated with recurrent attacks of cystitis, irritative voiding symptoms, and urethral pain. A mnemonic is the 3 Ds: dribbling, dyspareunia, and dysuria. Purulent urethral discharge is sometimes noted as the infected diverticulum empties. On occasion, the diverticulum may be large enough to be discovered by the patient.

The diagnosis is usually made on feeling a rounded cystic mass in the anterior wall of the vagina that leaks pus from the urethral orifice when pressure is applied. Endoscopy may reveal the urethral opening, although the openings are often very difficult to locate. The postvoiding film of an excretory urographic series may demonstrate the lesion. It may be possible to introduce a small catheter through which radiopaque fluid can be instilled. Appropriate x-ray films are then exposed (Figure 41–2). The plain film may show a stone in the diverticulum (Presman, 1964). If these methods fail, the following procedures can be used:

1. Empty the diverticulum manually. Via a catheter, instill 5 mL of indigo carmine and 60 mL of contrast medium into the bladder. Remove the catheter and have the patient begin to void. Occlude the meatus with a finger. This maneuver usually causes the diverticulum to fill with the test solution. Take appropriate x-rays, and perform panendoscopy to look for leakage of blue dye from the mouth of the diverticulum.

2. Insert a Davis-TeLinde catheter. This looks like a Foley catheter but is surrounded by a second movable balloon. Pass the catheter to the bladder and inflate the proximal balloon. While exerting tension on the catheter, slide the second balloon against the urinary meatus and inflate it. Then inject contrast medium into the catheter. The radiopaque fluid will escape from the catheter through a hole between the balloons and will fill the urethra and diverticulum, after which x-rays can be exposed. Occasionally, urethral diverticulum is elusive and difficult to visualize. Transvaginal ultrasonography (Vargas-Serrano et al, 1997; Siegel et al, 1998; Baert, 1992; Mouritsen, 1996) or pelvic magnetic resonance imaging (Debaere et al, 1995; Kim, 1993) can be helpful in diagnosis; MRI is reported to be much more sensitive in detecting diverticulae than double-balloon urethrography (Neitlich et al, 1998).

Treatment consists of removal of the sac through an incision in the anterior vaginal wall, care being taken not to injure the urethral sphincteric musculature. Incision is carried down to the diverticular mucosa, and the plane of cleavage is followed all around to the neck of the diverticulum. The diverticular sac is completely excised and the defect in the urethra repaired. Elik (1957) recommends that the diverticulum be opened, stuffed with absorbable cellulose (Oxycel), and then closed; the resulting inflammatory reaction destroys the cyst. A suprapubic cys-
tostomy should be left in place for 15 days following surgical excision of the diverticulum.

The outcome is usually good unless the diverticulum is so situated that its excision injures the external urinary sphincter mechanism. In a few cases, urethrovaginal fistula may develop. If the fistula does not close with adequate suprapubic drainage, surgical repair will be necessary 2–3 months later.

URETHRAL STRICTURE

True organic stricture of the adult female urethra is not common. (Functional urethral obstruction is more common.) It may be congenital or acquired. The trauma of intercourse and especially of childbirth may lead to periurethral fibrosis with contracture, or the stricture may be caused by the surgeon during vaginal repair. It may develop secondary to acute or chronic urethritis.

Persistent hesitancy in initiating urination and a slow urinary stream are the principal symptoms of stricture. Burning, frequency, nocturia, and urethral pain may occur from secondary urethritis or cystitis. If secondary infection of the bladder is present, pus and bacteria will be found in the urine. A fairly large catheter (22F) may pass to the bladder only with difficulty. Panendoscopy may demonstrate the point of narrowness and disclose evidence of urethritis. Cystoscopy often reveals trabeculation (hypertrophy) of the bladder wall.

Chronic cystitis may cause similar symptoms, but urinalysis reveals evidence of infection. Cancer of the urethra causes progressive narrowing of the urethra, but induration and infiltration of the urethra are found on vaginal examination. Panendoscopy with biopsy establishes the diagnosis. Vesical tumor involving the bladder neck causes tenancy and impairment of the urinary stream. Cystoscopy is definitive. Chronic urethritis commonly accompanies urethral stenosis; either may be primary. Recurrent or chronic cystitis is often secondary to stenosis.

Treatment consists of gradual urethral dilatation (up to 36F) at weekly intervals. Slight overstretching is necessary, since some contracture will occur after therapy is discontinued. Measures to combat urethritis and cystitis also must be employed. Internal urethrotomy has its proponents (Essenhigh, 1968). With proper overdilatation of the urethra and specific therapy of the urethritis that is usually present, the prognosis is good.

REFERENCES

Distal Urethral Stenosis

**Labial Fusion**


**Acute Urethritis**


**Chronic Urethritis**


**Senile Urethritis**


**Urethral Caruncle**


**Prolapse of the Urethra**


**Urethrovaginal Fistula**


**Urethral Diverticulum**


**Urethral Stricture**

Skin Diseases of the External Genitalia

Timothy G. Berger, MD

INFLAMMATORY DERMATOSES

The patient with skin disease of the external genitalia should be questioned about and examined for other possible areas of involvement. Scabies must always be considered in cases of genital pruritus. Patients, fearing genital infection, frequently overtreat genital lesions. Repeated scratching and rubbing tend to prolong and complicate genital conditions.

CONTACT DERMATITIS

Contact dermatitis includes both irritant dermatitis and allergic contact dermatitis. True allergic contact dermatitis is pruritic, erythematous, edematous, and weepy. Possible causes are feminine hygiene products, condoms, and plants (poison oak and ivy). Twice-daily cool water compresses followed immediately by the application of a mild, nonfluorinated topical steroid (1% hydrocortisone ointment or 1% Pramosone [hydrocortisone 1% plus pramoxine] ointment) is usually effective.

CIRCUMSCRIBED NEURODERMATITIS (LICHEN SIMPLEX CHRONICUS)

The labia majora and scrotum are predisposed to this chronic skin condition, caused by an itch-scratch cycle. Stress exacerbates the condition. Anogenital pruritus may be caused by lumbosacral radiculopathy. The skin becomes thickened. The scratching may be subconscious and occur during sleep. The treatment is the same as that for contact dermatitis (described previously) plus stopping the scratching. The addition of pramoxine hydrochloride 1%, topical doxepin cream 5% (Zonalon) or eutectic mixture local anesthetic (EMLA—lidocaine 2.5% and prilocaine 2.5%) to topical steroid treatment may be of benefit. Capsaicin 0.006% topically can be effective.

INTERTRIGO

Intertrigo describes the moist plaques that occur in areas of persistent maceration. It is often complicated by bacterial and yeast overgrowth. It occurs in the groin, abdominal skin folds, inframammary areas, and axillae, usually in obese individuals. It is more common in hot, humid weather. Simply allowing the occluded areas to be exposed to air may result in healing. Further treatment consists of twice-daily cool soaks with Burrow’s solution 1:20, followed by application of nystatin ointment plus 1% hydrocortisone ointment (see Candidiasis given below).

DRUG ERUPTIONS

Most drug eruptions are widespread, but they may first appear in the groin. Fixed drug eruption, due usually to laxatives (phenolphthalein), sulfonamides, or nonsteroidal anti-inflammatory drugs (NSAIDs), commonly presents on the genitalia. These medications are often taken only intermittently, so the association with the offending medication is often missed. Two percent of all genital ulcers are fixed drug eruptions. Lesions often begin within a day of drug exposure and present as bright red to violaceous macules that quickly blister and erode. The erosion is superficial and broad (usually >1 cm). Fixed drug eruption occurs in the same site with each exposure to the drug. Treatment is to stop the offending medication.

PSORIASIS

Psoriasis may involve flexural surfaces (inverse psoriasis), such as the groin and the cleft of the buttocks. In moist surfaces the lesions of psoriasis are bright red and usually free of scale. Itching may be intense or nonexistent. A solitary plaque may present on the glans penis, leading to confusion with high-grade dysplasia (erythroplasia of Queyrat). The diagnosis usually can be made by inspection and by noting other areas of involvement such as the scalp, elbows, knees, and nails. Hydrocortisone cream, 1%, plus an imidazole cream ( clotrimazole, 1%; miconazole, 2%; or ketoconazole, 2%) is usually efficacious. The condition is chronic and relapsing and often flares after intercourse due to the exposure of the psoriatic lesion to Candida. Washing the glans after intercourse is critical in controlling penile psoriasis. The skin lesions of reactive arthritis in the genitalia are identical to that of psoriasis and favor the glans penis (cirkinite...
SEBORRHEIC DERMATITIS

Seborrhea favors hairy areas and usually involves the base of the penis, inner thighs, and pubic area. The treatment is identical to that of psoriasis in the groin.

LICHEN PLANUS

Lichen planus may affect the glans penis or the labia majora and minora and vaginal mucosa. The genitalia may be the only site of involvement. The lesions are polygonal, violet-hued, flat-topped papules about 0.5–1 cm in diameter, with milky striations over their shiny surfaces. Vaginal lesions are often erosive. Lesions may be asymptomatic, pruritic, or painful if eroded. Vulvar erosive disease often coexists with oral erosive disease (vulvovaginal-gingival syndrome). The condition may mimic lichen sclerosus (LS) completely, making clinical distinction impossible. A biopsy may be required. Superpotent topical corticosteroids and topical tacrolimus 0.1% ointment may be helpful in relieving symptoms and healing erosions. The disease may disappear after months to years.

LICHEN SCLEROSUS

LS almost inevitably involves the anogenital regions, where severe pruritus or painful erosions may develop. LS of the glans penis (balanitis xerotica obliterans) may lead to phimosis and urethral stenosis. Squamous cell carcinoma of the genitalia may complicate LS, but prophylactic surgery of these genital lesions is not called for. Superpotent topical steroids are the treatment of choice for all forms of genital LS, in children and adults. Initially treatment is twice daily, with gradual tapering to once daily, then several times weekly. An initial trial should be 6 weeks of treatment. Atrophy reverses despite the use of potent steroids on thin genital skin. Once the patient is in remission, milder steroids or bland emollients may be used for maintenance. Topical testosterone is not beneficial. Childhood LS may not improve during puberty or adulthood.

COMMON SUPERFICIAL INFECTIONS

ARTHROPODS

Pediculosis Pubis (Pubic Lice, Crabs)

Pediculosis pubis may be sexually or nonsexually transmitted. Itching may be intense, leading to scratching and pyoderma. The nits are found on the hair shafts in the pubic area and the eyelids. Treatment consists of application of permethrin 1% crème rinse applied for 10 minutes then washed off. All hairy areas contiguous with the genital area, which may be a significant portion of the male body (chest, abdomen, legs, and axilla), should be treated. Lindane 1% shampoo for 4 minutes or 5% permethrin cream for 8 hours are acceptable alternatives. The sexual partner(s) should also be examined and treated. All clothing, bedding, and towels should be washed and heat dried or dry cleaned. If lice are found 1 week later, the treatment should be repeated.

Scabies

Infestation with the human mite Sarcoptes scabiei usually causes a severely pruritic, widespread eruption. In males, very itchy papules or nodules with a central crust are common on the penile shaft or glans and on the scrotum and are virtually pathognomonic of scabies. These nodules may persist for weeks to months after treatment. In adults, scabies is a sexually transmitted disease. Treatment consists of overnight (8–12 hours) application of 5% permethrin cream to the whole body from the neck down. The treatment may be repeated in 1 week. All members of the household and all sexual partners of the index case should be treated simultaneously. All clothing, bedding, and towels should be washed and heat dried or dry cleaned. Potent topical steroids may be used to treat persistent genital nodules.

FUNGAL INFECTIONS (TINEA CRURIS)

Tinea cruris is characterized by marginated, slightly elevated, peripherally scaling patches on the inner thighs and in the groin. Tinea cruris does not affect the scrotum and is less intense deep in the inguinal folds. This is in contrast to candidiasis, which characteristically affects the scrotum and is accentuated deep in the inguinal folds (the moistest area). Pruritus may be intense. Direct microscopic examination of skin scrapings in potassium hydroxide solution reveals hyphae.

Miconazole, clotrimazole, ketoconazole, econazole, ciclopirox olamine, terbinafine, or butenafine creams are all effective. All are applied twice daily except the last 2, which are applied once daily. One percent hydrocortisone cream may be used concomitantly. In severe or refractory cases, oral antifungal treatment may be required. Griseofulvin ultramicronized (Grispeg), 250 mg twice daily for 4–6 weeks; itraconazole, 200 mg twice daily with food and an acid beverage for 7 days; or terbinafine, 250 mg once daily for 2–4 weeks, is usually adequate, even in the most severe cases.

CANDIDIASIS

Erythematous, weeping lesions with peripheral satellite vesiculopustules characterize infection with Candida albi-
cans. Lesions occur most commonly on the inner thighs and buttocks, with a predilection for the depths of the creases. Scrotal involvement is common in candidiasis. Pregnancy, diabetes mellitus, obesity, broad-spectrum antibiotics, and immunosuppression are predisposing factors. The skin involvement may be secondary to vaginal infection. Lesions occur under the prepuce (candidal balanitis). High-power microscopic examination of skin scrapings in potassium hydroxide solution shows clusters of tiny spores and fine mycelial filaments. Nystatin ointment or cream is effective in most instances. Miconazole, clotrimazole, ketoconazole, and econazole applied twice daily are alternatives to nystatin but may cause burning on application.

Nystatin can be applied to eroded areas. In severe cases in women, fluconazole tablets (100–200 mg) once daily for 1 week, followed by 150 mg once weekly, is often dramatically effective.

**BACTERIAL INFECTIONS (PYODERMA)**

*A Staphylococcus aureus* is the most common cause of primary bacterial infections in the genital area. Culture may confirm the diagnosis. *S. aureus* produces 2 types of primary lesions: a follicular pustule (folliculitis) and a superficial blister (impetigo).

Staphylococcal folliculitis begins as a superficial infection of the follicle but may extend deeply (furunculosis). It is usually acute but may be chronic or recurrent. Chronic folliculitis is usually due to nasal carriage of *Staphylococcus*. Deep, draining abscesses are rarely due to bacteria alone and suggest the presence of a chronic suppurative disorder, for example, inflammatory bowel disease, lymphogranuloma venereum, or hidradenitis suppurativa. Recurrent folliculitis in the groin is common in acquired immunodeficiency syndrome (AIDS).

Topical treatment alone is often inadequate for bacterial folliculitis. A penicillinase-resistant penicillin (dicloxacillin) or first-generation cephalosporin is the treatment of choice. Penicillin-allergic individuals may be treated with doxycycline. Community-acquired methicillin-resistant *Staphylococcus aureus* (MRSA) is also treated with doxycycline. Treatment is continued until all lesions are healed. Adding rifampin to the described treatment is recommended for frequent recurrences.

Staphylococcal impetigo starts as a very superficial blister that quickly breaks, leaving a crusted, weeping erosion. Treatment is the same as for staphylococcal folliculitis but usually of shorter duration.

**VIRAL INFECTIONS**

**Genital Warts**

External genital warts, caused by human papilloma viruses, are the most common sexually transmitted disease. One percent of all sexually active adults have genital warts, and up to 10% can be documented to have active human papilloma virus infection by sensitive techniques such as polymerase chain reaction. Elevated genital warts, condyloma acuminata, are due to low-risk human papilloma virus types (usually 6 and 11). They are most common on the vulva, under the prepuce, and on the penile shaft. The vaginal, anal, and oral mucosa may also be affected.

Since current treatments can eradicate only the clinical lesions (the warts) and not the infectious agent (the virus), recurrence is very common (>25%). In addition, treating the sexual partners of infected persons has no effect on the outcome of the treatment in the index case. Warts may spontaneously resolve. It is no longer recommended to search for and treat “subclinical external genital warts”—those seen with acetic acid soaking or identified by special immunologic techniques. The goal of treatment is to provide wart-free intervals with the least discomfort and long-term sequelae to the patient. In monogamous couples the option of no treatment should be discussed. Certain “high-risk” genital human papilloma virus types, usually 16 and 18, cause bowenoid papulosis, flat hyperpigmented papules on the genital skin. Bowenoid papulosis is a high-grade intraepithelial lesion and is associated with penile, vaginal, and cervical dysplasia. Regular gynecologic examination with Pap smears is mandatory. Bowenoid papulosis may be treated like external genital warts.

There are 2 basic forms of therapy: patient applied and health-care worker applied. Patient-applied treatment is either podophyllotoxin or imiquimod. Podophyllotoxin is applied twice daily for 3 days per week for 6–10 weeks. About half of patients clear their warts with 1 course of therapy. It is less irritating and more effective than healthcare worker-applied podophyllum resin. It is contraindicated in pregnancy. Imiquimod is an immune modulator that results in the local production of interferon. It is applied once daily for 3 times per week (usually Monday, Wednesday, and Friday). The response rate is about 40% for men and over 75% for women. Treatment duration is prolonged, with average time to final response being over 2 months. Physician-applied treatments include liquid nitrogen cryotherapy (75% response, 50% durable remission) and electrocautery (100% response, 75% durable remission). Laser therapy and intralesional or systemic immunotherapies are second-line treatments and are rarely indicated for external genital warts.

**Molluscum Contagiosum**

Molluscum contagiosum is a common cutaneous infection that is sexually transmitted in adults. The characteristic lesion is a smooth-surfaced, firm, pearly papule 2–5 mm in diameter with a central umbilication. Most infected persons have 5–15 lesions located on the lower abdomen, upper thighs, or skin of the genitalia. Extensive molluscum contagiosum outside the genital area in adults is rare except...
in immunosuppressed patients, especially those infected with HIV. Treatment involves local destruction of the lesions by cryotherapy or electodesiccation.

**Herpes Simplex**

Genital herpes simplex virus (HSV) is usually caused by HSV 2, but, increasingly, HSV 1 genital herpes due to orogenital sex has been reported. Most infections cause no symptoms initially. Viral culture will confirm the diagnosis. First-episode genital HSV is treated with acyclovir (200 mg) five times daily, valacyclovir (500 mg) twice daily, or famciclovir (250 mg) twice daily, for 7–10 days.

Recurrent disease is frequent, often preceded by tingling at the soon-to-be-affected site (the prodrome). It is virtually always caused by HSV 2. It presents as grouped blisters localized to 1 site and lasting about 1 week. Most patients do not require treatment if the outbreaks are mild. For individual outbreaks, either acyclovir (200 mg) five times daily, or valacyclovir (500 mg) twice daily for 3 days; or famciclovir (1000 mg) twice daily for 1 day only may be used to shorten the duration and severity of the eruption. For frequent recurrences (>6–12/year), suppression may be better than intermittent treatment. Acyclovir (400 mg) twice daily (or 800 mg once daily), valacyclovir (500 mg to 1 g) once daily, or famciclovir (250 mg) twice daily may be used. Suppressive treatment will reduce outbreaks by 85% and reduces the amount of virus shed by 95%. This reduces transmission to sexual partners by 50%.

Herpes simplex is the most common cause of persistent genital ulceration in immunosuppressed patients. In severe cases, intravenous acyclovir may be required. Acyclovir resistance can occur in the setting of immunosuppression and is treated with foscarnet.

**REFERENCES**

**General**


**Inflammatory Dermatoses**


**Common Superficial Infections**


Sexuality is defined as the constitution of an individual in relation to sexual attitudes or activity. What defines our sexuality is a complex interaction between our genetic makeup, environmental stimulus, and cultural influences. The origins of our sexuality occur at the time of conception when the genetic material from 2 sources of the opposite sex coalesces into a new individual. From that moment, sexual differentiation occurs by a highly organized process. Sex chromosomes and autosomes dictate the development of gonads; the gonads in turn produce hormones, which then direct the development of the internal and external genitalia. Disorders of sexual differentiation arise from abnormalities in chromosomes, gonadal development, or hormonal production/activity.

Patients with disorders of sexual differentiation may present (1) during the newborn period as having ambiguous genitalia, (2) as having inappropriate pubertal development, (3) as having delayed pubertal development, or (4) later in life as having infertility.

**NORMAL SEXUAL DIFFERENTIATION**

**Chromosomal Sex**

The genetic material necessary for the development of the male phenotype is normally located on the short arm of the Y chromosome. The critical gene or sex-determining region on the Y chromosome is known as the SRY region. The gene products of the SRY genetic cascade will subsequently direct the development of the testis by interacting with multiple other genes such as SOX-9. Genetic information that is necessary for male and female development beyond gonadal differentiation is located on the X chromosome and on the autosomes.

**Gonadal Differentiation**

The gonads develop from the urogenital ridges (Figure 43–1), which are formed during the 4th week of gestation by the proliferation of the coelomic epithelium and condensation of the underlying mesenchyme along the mesonephros. The germ cells, located in the endoderm of the yolk sac, migrate to the genital ridges. At the early stage of development the gonad is bipotential, capable of forming into either a testis or an ovary. During the 6th–7th week of gestation, at least 4 different genes Wilms’ tumor suppressor gene (WT-1), Fushi-Tarza Factor-1 (FTZ-F1), steroidogenic Factor-1 (SF-1), and LIM-I induce the development of the testis. The primordial germ cells differentiate into the Sertoli cells and associated Leydig cells, which aggregate into spermatogenic cords. Loose mesenchymal tissue condenses into a thick layer, the tunica albuginea, which surrounds the testis and separates its connection with the coelomic epithelium, thereby preventing further migration of mesonephric cells into the testis.

Classic teaching is that the female phenotype is the default developmental pathway in the absence of the SRY cascade. It is now known that at least one gene, dosage-sensitive sex reversal (DAX-1), is essential for ovarian development. DAX-I is located on the short arm of the X chromosome. The gene products of SRY and DAX-I compete to stimulate the steroidogenic acute regulatory protein (StAR). The StAR protein is the first step in steroidogenesis, facilitating the conversion of cholesterol to pregnenolone. In the normal XY male, SRY overwhelms the one functional DAX-I gene, stimulating testicular development and subsequent testosterone production. In the normal XX female, 2 DAX-I genes are present without the competitive SRY; downregulating StAR, hence inhibiting testicular development, which results in ovarian development. In the fetal ovaries, the germ cells differentiate and are arrested in the last phase of meiotic prophase, forming the oocytes. The cells in the genital ridges develop into granulosa cells, which surround the oocytes and complete the formation of the ovaries.

**Hormones**

At 3.5 weeks’ gestation, the Wolffian system appears as 2 longitudinal ducts connecting cranially to the mesonephros and caudally draining into the urogenital sinus (Figure 43–2). At approximately the 6th week of gestation, the Müllerian duct develops as an evagination in the coelomic epithelium just lateral to the Wolffian duct.

During the 8th–9th week of gestation, Sertoli cells of the fetal testis secrete a glycoprotein, Müllerian-inhibiting substance (MIS), or anti-Müllerian hormone. This protein
induces the regression of the Müllerian ducts through the dissolution of the basement membrane and condensation of mesenchymal cells around the Müllerian duct. Because MIS acts locally, Müllerian duct regression occurs only on the ipsilateral side of the fetal testis producing this hormone. MIS also induces the formation of seminiferous tubules and further differentiation of the testis. At the 9th or 10th week of gestation, the Leydig cells appear in the testis and begin to synthesize testosterone. This hormone transforms the Wolffian duct into the male genital tract, which is completed by the end of the 11th week of gestation.

Beginning in the 9th week of gestation, testosterone also induces the development of the external genitalia (Figure 43–3) from the genital tubercle, urogenital sinus, and genital swellings. At the molecular level testosterone is converted to 5α-dihydrotestosterone (DHT) by the microsomal enzyme, type 2 5α-reductase, for complete differentiation of the penis with a male-type urethra and glans. Testosterone dissociates from its carrier proteins in the plasma and enters cells via passive diffusion. Once in the cell, testosterone binds to the androgen receptor (AR) and induces changes in conformation, protecting it from degradation by proteolytic enzymes. This conformational change is also required for AR dimerization, DNA binding, and transcriptional activation, all necessary for testosterone to be expressed. Androgen binding also displaces heat shock proteins, possibly relieving constraints on receptor dimerization or DNA binding. After entering the nucleus, the AR complex then binds androgen response element DNA regulatory sequences within the androgen responsive genes and activates them. DHT also binds the AR, with enhanced androgenic activity, in part because of its slow dissociation rate from the AR.

DHT then binds to nuclear receptors, forming a complex that regulates the transformation of these tissues into the glans penis, penile and cavernous urethra, Cowper’s glands, prostate, and scrotum. Between the 28th and 37th week of gestation, testicular descent into the scrotum begins. While the mechanism of this process is not completely understood, it is clearly androgen dependent.

**Development of the Female Genitalia**

The female internal genitalia develop from the Müllerian ducts. Without the hormones produced by the testis, the Wolffian ducts regress at the 9th week of gestation. At the same time, the Müllerian ducts begin to differentiate; the cranial portions form the fallopian tubes, while the caudal portions fuse to form the uterus, cervix, and the upper portion of the vagina. Concurrently, the external genitalia defined as the lower portion of the vagina, the vestibule, Bartholin and Skene glands, the clitoris, and labia minora and majora develop from the urogenital sinus and genital tubercles. Like the testis, the ovary undergoes a partial transabdominal descent. However, transinguinal descent of the ovary does not occur, leaving the ovaries just below the rim of the true pelvis. The role of estrogen in the differentiation of the female phenotype is unclear.

**Development of the Male External Genitalia**

Formation of the external male genitalia is a complex developmental process involving the SRY genetic programming, cell differentiation, hormonal signaling, enzyme activity, and tissue remodeling. By the end of the 1st month of gestation, the hindgut and future urogenital system reach the ventral surface of the embryo at the cloacal membrane. The urorectal septum divides the cloacal membrane into a posterior, or anal, half and an anterior half, the urogenital membrane. Three protuberances appear around the latter. The most cephalad is the genital tubercle. The other 2, the genital swellings, flank the urogenital membrane on each side. Up to this point, the male and female genitalia are essentially indistinguishable. Under the influence of testosterone in response to a surge of luteinizing hormone from the pituitary, masculinization of the external genitalia takes place. One of the first signs of masculinization is an increase in the distance between the anus and the genital structures, followed by elongation of the phallus, formation of the penile urethra from the urethral groove, and development of the prepuce.

At 8 weeks’ gestation, the external genitalia remain in the indifferent stage. The urethral groove on the ventral surface of the phallus is between the paired urethral folds. The penile urethra forms as a result of fusion of the medial edges of the endodermal urethral folds. As development progresses, the ectodermal edges of the urethral groove begin to fuse to form the median raphe (Figure 43–4A). By
11–12 weeks the coronal sulcus separates the glans from the shaft of the penis. By 16 weeks’ gestation the urethral folds have completely fused in the midline on the ventrum of the penile shaft (Figure 43–4B). Note the normal ventral penile curvature, or chordee, that occurs during development and resolves by the 20th week (Figure 43–4C).

The glandular urethra, which consists of a squamous epithelial-lined tube different from the urothelial-lined anterior urethra, also completes formation during this period. The mechanism of the glandular urethral formation remains controversial. Evidence suggests 2 possible explanations (Figure 43–5): (1) endodermal cellular differentiation where the glandular urethra forms by an extension of urogenital sinus epithelium that undergoes transdifferentiation versus (2) primary intrusion of the ectodermal tissue from the skin of the glans penis. Cross-sectional histologic analysis at 24 weeks’ gestation reveals complete penile development (Figure 43–6A–H). Note the extensive neuronal innervation just above the tunica of the corporeal bodies. Three-dimensional reconstruction of the fetal male penis illustrates the extensive neuronal distribution (Figure 43–7). Note the nerve density in the glans (Figure 43–7E and F).

Anatomical and immunohistochemical studies advocate the new theory of endodermal differentiation, which shows that epithelium of the entire urethra is of urogenital sinus origin. The entire male urethra, including the glandular urethra, is formed by dorsal growth of the urethral plate into the genital tubercle and ventral growth and fusion of the urethral folds. Under proper mesenchymal induction, urothelium has the ability to differentiate into a stratified squamous phenotype with characteristic keratin staining, thereby explaining the cell type of the glans penis.

The future prepuce is forming at the same time as the urethra and is dependent on normal urethral development. At about 8 weeks’ gestation, low preputial folds appear on both sides of the penile shaft, which join dorsally to form a flat ridge at the proximal edge of the corona. The ridge does not entirely encircle the glans because it is blocked on the ventrum by incomplete development of the glandular urethra. Thus, the preputial fold is transported distally by
active growth of the mesenchyme between it and the glandular lamella. The process continues until the preputial fold (foreskin) covers all of the glans, forming a midline seam (Figure 43–4D). The fusion is usually present at birth, but subsequent desquamation of the epithelial fusion allows the prepuce to retract. If the genital folds fail to fuse, the preputial tissues do not form ventrally; consequently, in hypospadias, preputial tissue is absent on the ventrum, and excessive dorsally.

**Disorders of Sexual Differentiation**

Disorders of abnormal sexual differentiation may be divided into the 3 categories.

**Disorders of Chromosomal Sex**

These result from abnormalities in the number or structure of the sex chromosomes. These abnormalities may arise from nondisjunction, deletion, breakage, rearrangement, and translocation of genetic material on these chromosomes. These disorders are summarized in Table 43–1.

**Disorders of Gonadal Sex**

These result from abnormalities in gonadal development. In these disorders, the karyotype is normal (ie, 46XX or 46XY). However, mutations in the sex chromosomes or autosomes, teratogens, or trauma to the gonads interfere with their normal development. These disorders are summarized in Table 43–2.

**Disorders of Phenotypic Sex**

These result from abnormalities in hormonal production or activity. The etiologies include defective synthesis by the gonads, abnormal production by the adrenal glands, presence of exogenous sources, or abnormalities in receptor activity. These disorders are summarized in Table 43–3.
Figure 43–4. Male human fetal external genitalia during gestation. A: 11 weeks. Note the urethra is open and urethral fold (uf) and groove are prominent in the transillumination view of the phallus. B: at 16.5 weeks. Note the normal ventral curvature (vc) is shown as well as the foreskin, which is almost completely formed. C: at 20 weeks’ gestation, penile and urethral development looks complete, with the prepuce covering the glans and the penile curvature resolving. D: At 24 weeks the prepuce covers the whole glans. Note the midline seam (ms). Note the progression of natural curvature to a straight phallus during development.

Figure 43–5. Theories of human penile urethral development. The ectodermal ingrowth theory as described in most textbooks of embryology postulates that the glanular urethra is formed by ingrowth of epidermis. More recent data support the formation of the entire urethra via endodermal differentiation alone.
Clinical Evaluation of Patients with Ambiguous Genitalia

The accurate diagnosis of a patient with ambiguous genitalia is a challenging process. Based on the diagnosis, decisions will be made for gender assignment, which will have a great impact not only on the patient but also on the patient’s family. In most societies the accepted norm is 2 sexes, either male or female. When a new baby arrives and the proclamation as to whether it’s a boy or a girl cannot be made immediately, an anticipated celebration turns into a stressful family dilemma. With prenatal amniocentesis and routine ultrasound, sex determination is often known well before birth. This can compound the emotional trauma when the known and anticipated genotype does not match the newborn’s phenotype. Furthermore, in cases such as severe salt-wasting congenital adrenal hyperplasia (CAH), accurate diagnosis is lifesaving.

History

A detailed history is of great importance. Since many of the disorders such as XX male syndrome and true hermaphrodites are hereditary, a family history should carefully be examined for similarly affected individuals, unexplained death during infancy, infertility, amenorrhea, and hirsutism. Furthermore, drugs ingested during pregnancy (such as progesterone) and virilizing signs in the mother during pregnancy should be ascertained.

Physical Examination

The abdomen and rectum should be carefully palpated for midline structures such as a uterus. These examinations...
will provide information regarding the presence of Müllerian duct derivatives. Other helpful physical findings include dehydration, failure to thrive, pigmentation (in patients with salt-wasting CAH), and the presence of other associated anomalies such as cardiac murmurs or web neck (in patients with Turner’s or Klinefelter syndrome).

It is important to palpate for gonads in the labioscrotal fold or the scrotum. Since ovaries do not descend, it is likely to be a testis and hence unlikely to represent a case of female pseudohermaphroditism. Based on the presence or absence of gonads, an algorithm can be followed to determine the differential diagnosis of patients with ambiguous genitalia (Figure 43–8). It is important to look at the size of phallus (Table 43–4) and the location of the urethral meatus. Any patients with bilateral cryptorchidism or with unilateral cryptorchidism with hypospadias should be suspected of having abnormalities in sexual differentiation. As noted, other helpful physical findings include hyperpigmentation of the areola and labioscrotal fold, common in patients with CAH.

**Chromosomal Evaluation**

Examination of buccal mucosal cells for Barr body (inactivated second X chromosome) cannot be relied on to make an accurate diagnosis in patients with ambiguous genitalia.

A more accurate but more time-consuming method (2–3 days) is the direct assessment of chromosomes from cultured peripheral blood leukocytes. This method provides the exact chromosomal complements, presence of mosaicism, and structural features of the chromosomes. In the case of mosaicism, several different tissue samples may be required to accurately determine the presence of mosaicism.

**Biochemical Evaluation**

In the case of CAH, the specific enzyme defect can be determined based on the presence or absence and the type of steroid excreted in the urine. Figure 43–9 depicts the steroid synthesis pathway from cholesterol to aldosterone, steroids, or DHT. Note the enzymes necessary for conversion from precursors to products (also see Table 43–3).

In other disorders caused by hormonal abnormalities (such as 5α-reductase deficiency and androgen resistance), direct measurement of plasma testosterone is often not helpful, since abnormalities in testosterone levels in these pathologic states have not been consistently characterized. A more useful test is the testosterone response following stimulation by hCG (2000 IU/day for 4 days). If plasma testosterone levels rise more than 2 ng/mL from baseline, the abnormality is consistent with androgen resistance rather than a defect in testosterone synthesis. In addition, this test is also used to diagnose 5α-reductase type 2 deficiency. A post-hCG stimulation ratio of testosterone to DHT >30 establishes this diagnosis.

**Radiographic Evaluation**

In patients with intersex disorders, ultrasonography provides the least invasive and safest means of imaging the abdomen and pelvis. Identification of Müllerian-derived
<table>
<thead>
<tr>
<th>Disorder</th>
<th>Pathology</th>
<th>Chromosomes</th>
<th>Incidence</th>
<th>Gonads</th>
<th>Internal Genitalia</th>
<th>External Genitalia</th>
<th>Other Features</th>
<th>Risk of Cancer</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Klinefelter</td>
<td>Extra X chromosome</td>
<td>47 XXY 46 XY/47XXY</td>
<td>1 in 500</td>
<td>Hyalinized testis</td>
<td>Wolffian</td>
<td>Male</td>
<td>Gynecomastia Tall stature Mild mental retardation</td>
<td>Breast Extragonadal germ cell</td>
<td>Supplemental androgens Surgery for severe gynecomastia</td>
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<td>syndrome</td>
<td></td>
<td></td>
<td></td>
<td>No spermatogenesis</td>
<td></td>
<td></td>
<td>Elevated FSH/LH Low testosterone Elevated estradiol</td>
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<tr>
<td>XX male</td>
<td>No Y chromosome</td>
<td>46 XX</td>
<td>1 in 20,000 to 24,000</td>
<td>Hyalinized testis</td>
<td>Wolffian</td>
<td>Male</td>
<td>Gynecomastia Short stature Inc. incidence of hypospadias Normal mental status May be familial</td>
<td>Rare germ cell</td>
<td>Same as Klinefelter</td>
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Table 43–1. Disorders of Chromosomal Sex.
<table>
<thead>
<tr>
<th>Condition</th>
<th>Karyotype</th>
<th>Side and Position</th>
<th>Gonadal Type</th>
<th>Gonadal Type</th>
<th>Somatic Features</th>
<th>Gender</th>
<th>Surgery</th>
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<tbody>
<tr>
<td>Mixed gonadal dysgenesis</td>
<td>45 X/46 XY (70%)</td>
<td>One testis (usually undescended) and streak gonad</td>
<td>Wolffian and Müllerian</td>
<td>Usually ambiguous 60% reared as female</td>
<td>Somatic features like 45 X</td>
<td>Female</td>
<td>Prophylactic gonadectomy</td>
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<td>Incompl. virilization &amp; Müllerian regression</td>
<td>Undetected mosaic</td>
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<td></td>
<td></td>
<td>Male</td>
<td>Streak gonads removed</td>
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<td></td>
<td></td>
<td>Intra-abd. testis excised unless can be relocated and no ipsilateral Müllerian structure present</td>
</tr>
<tr>
<td>True hermaphrodite</td>
<td>Unknown</td>
<td>Bilateral ovitestedis Ovitestis &amp; ovary or testis (40%) One ovary &amp; testis (40%)</td>
<td>Wolffian and Müllerian</td>
<td>Usually ambiguous 70% reared as male</td>
<td>Gynecomastia at puberty Menstruation at puberty May be familial</td>
<td>Rare germ cell</td>
<td>Reconstructive surgery Poss. remove gonads</td>
</tr>
<tr>
<td></td>
<td>46 XX (70%) 46 XY (10%) Mosaic</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Disorder</td>
<td>Pathology</td>
<td>Chromosomes</td>
<td>Incidence</td>
<td>Gonads</td>
<td>Internal Genitalia</td>
<td>External Genitalia</td>
<td>Other Features</td>
</tr>
<tr>
<td>--------------------------</td>
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<td>----------------------------------</td>
</tr>
<tr>
<td>Pure gonadal dysgenesis</td>
<td>Unknown mutation prevents nl. differentiation of gonads</td>
<td>46 XX</td>
<td>1 in 8000</td>
<td>Bilateral streak gonads</td>
<td>Müllerian</td>
<td>Immature female</td>
<td>Normal to tall stature</td>
</tr>
<tr>
<td></td>
<td>46 XY</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Minimal somatic abn. Female: estrogen def. Male: testosterone def. May be familial</td>
</tr>
<tr>
<td>Absent testes syndrome</td>
<td>Mutation, teratogen or trauma to testis</td>
<td>46 XY</td>
<td>Unknown</td>
<td>Absent/rudiment testis No streak gonads</td>
<td>Wolffian</td>
<td>Var. virilization</td>
<td>Normal</td>
</tr>
<tr>
<td></td>
<td></td>
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<td></td>
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<td></td>
</tr>
</tbody>
</table>
**Table 43–3.** Disorders of Phenotypic Sex.

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Pathology</th>
<th>Chromosomes</th>
<th>Incidence</th>
<th>Gonads</th>
<th>Internal Genitalia</th>
<th>External Genitalia</th>
<th>Other Features</th>
<th>Urinary Steroids</th>
<th>Risk of Cancer</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Female Pseudohermaphrodite</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 β-Hydroxysteroid dehydrogenase def.</td>
<td>Excess androgens</td>
<td>46 XX</td>
<td>Second most common of CAH</td>
<td>Ovary</td>
<td>Müllerian</td>
<td>Mild ambiguous</td>
<td>Severe salt wasting</td>
<td>No cortisol</td>
<td>No aldosterone</td>
<td>DEAS</td>
</tr>
<tr>
<td>11 β-Hydroxylase def.</td>
<td>Excess androgens</td>
<td>46 XX</td>
<td>Rare</td>
<td>Ovary</td>
<td>Müllerian</td>
<td>Ambiguous</td>
<td>Hypertension</td>
<td>Dec. cortisol</td>
<td>Dec. aldosterone</td>
<td>11 DCS</td>
</tr>
<tr>
<td>21 α-Hydroxylase def.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>—Partial</td>
<td>Excess androgens</td>
<td>46 XX</td>
<td>1 in 5000 to 15,000</td>
<td>Ovary</td>
<td>Müllerian</td>
<td>Ambiguous</td>
<td>Normal cortisol</td>
<td>Inc. aldosterone</td>
<td></td>
<td>17 OH-P</td>
</tr>
<tr>
<td>—Severe</td>
<td>Excess androgens</td>
<td>46 XX</td>
<td>Ovary</td>
<td>Müllerian</td>
<td>Ambiguous</td>
<td></td>
<td>Normal cortisol</td>
<td>Inc. aldosterone</td>
<td></td>
<td>17 OH-P</td>
</tr>
<tr>
<td>Excess maternal androgens</td>
<td>Excess androgens</td>
<td>46 XX</td>
<td>Ovary</td>
<td>Müllerian</td>
<td>Ambiguous</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(continued)
Table 43–3. Disorders of Phenotypic Sex. (continued)

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Pathology</th>
<th>Chromosomes</th>
<th>Incidence</th>
<th>Gonads</th>
<th>Internal Genitalia</th>
<th>External Genitalia</th>
<th>Other Features</th>
<th>Urinary Steroids</th>
<th>Risk of Cancer</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male Pseudohermaphrodite</td>
<td>Defect in testosterone synthesis</td>
<td>46 XY</td>
<td>Testis</td>
<td>Wolffian</td>
<td>Ambiguous</td>
<td></td>
<td>Severe salt wasting</td>
<td>None</td>
<td>None</td>
<td>Replacement mineralocorticoids and glucocorticoids</td>
</tr>
<tr>
<td>20,22 Desmolase def.</td>
<td>Defect in testosterone synthesis</td>
<td>46 XY</td>
<td>Testis</td>
<td>Wolffian</td>
<td>Ambiguous</td>
<td></td>
<td>Severe salt wasting</td>
<td>None</td>
<td>None</td>
<td>Replacement mineralocorticoids and glucocorticoids</td>
</tr>
<tr>
<td>3 β-Hydroxysteroid dehydrogenase def.</td>
<td>Defect in testosterone synthesis</td>
<td>46 XY</td>
<td>Second most common in CAH</td>
<td>Testis</td>
<td>Wolffian</td>
<td>Ambiguous</td>
<td>Severe salt wasting</td>
<td>DEAS</td>
<td>None</td>
<td>Replacement mineralocorticoids and glucocorticoids</td>
</tr>
<tr>
<td>17 α-Hydroxylase def.</td>
<td>Defect in testosterone synthesis</td>
<td>46 XY</td>
<td>Testis</td>
<td>Wolffian</td>
<td>Ambiguous</td>
<td></td>
<td>Hypokalemic alkalosis</td>
<td>CS 11 DCS</td>
<td>None</td>
<td>Replacement glucocorticoids</td>
</tr>
<tr>
<td>17,20 Desmolase def.</td>
<td>Defect in testosterone synthesis</td>
<td>46 XY</td>
<td>Rare</td>
<td>Testis</td>
<td>Wolffian</td>
<td>Ambiguous</td>
<td>Hypertension Dec. cortisol Dec. aldosterone one</td>
<td>None</td>
<td>None</td>
<td>Supplemental testosterone</td>
</tr>
<tr>
<td>17 β-Hydroxysteroid dehydrogenase def.</td>
<td>Defect in testosterone synthesis</td>
<td>46 XY</td>
<td>Most common</td>
<td>Testis</td>
<td>Wolffian</td>
<td>Ambiguous</td>
<td>Virilization and puberty</td>
<td>ASD</td>
<td>None</td>
<td>Decision reared as female or male</td>
</tr>
<tr>
<td>5 α-Reductase def.</td>
<td>Defect in androgen action</td>
<td>46 XY</td>
<td>Testis with spermatogenesis</td>
<td>Testis</td>
<td>Female</td>
<td></td>
<td>No gynecomastia NL testosterone one NL virilization</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Condition</td>
<td>Androgen receptor defect</td>
<td>Chromosome</td>
<td>Incidence</td>
<td>Testis</td>
<td>Puberty Status</td>
<td>Estrogen treatment</td>
<td>Androgen treatment</td>
<td>Gonads</td>
<td>Reproductive Status</td>
<td></td>
</tr>
<tr>
<td>------------------------------------------------------</td>
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<td>-------------------</td>
<td>--------</td>
<td>---------------------</td>
<td></td>
</tr>
<tr>
<td>Complete testicular feminization</td>
<td>Androgen receptor defect</td>
<td>46 XY</td>
<td>1 in 20,000 to 64,000</td>
<td>Testis not fertile</td>
<td>Absent</td>
<td>Female reared as female</td>
<td>Inc. testosterone</td>
<td>None</td>
<td>Germ cells</td>
<td>Remove gonads after puberty</td>
</tr>
<tr>
<td>Incomplete testicular feminization</td>
<td>Androgen receptor defect</td>
<td>46 XY</td>
<td>1/10th of complete</td>
<td>Testis not fertile</td>
<td>Wolffian</td>
<td>Female</td>
<td>Inc. testosterone</td>
<td>None</td>
<td>Germ cells</td>
<td>Remove gonads prior to puberty</td>
</tr>
<tr>
<td>Reifenstein syndrome</td>
<td>Androgen receptor defect</td>
<td>46 XY</td>
<td>X linked rec.</td>
<td>Testis not fertile</td>
<td>Wolffian</td>
<td>Hypospadias male</td>
<td>Gynecomastia</td>
<td>None</td>
<td>None</td>
<td>Reconstruction as needed</td>
</tr>
<tr>
<td>Infertile male syndrome</td>
<td>Androgen receptor defect</td>
<td>46 XY</td>
<td>X linked rec.</td>
<td>Testis not fertile</td>
<td>Wolffian</td>
<td>Male</td>
<td>Infertility</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Receptor + resistance</td>
<td>Androgen receptor defect</td>
<td>Unknown</td>
<td>Testis not fertile</td>
<td>Wolffian</td>
<td>Ambiguous</td>
<td>Nl. or inc. testosterone</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td></td>
</tr>
<tr>
<td>Persistent Mullerian duct syndrome</td>
<td>Persistent Mullerian duct syndrome</td>
<td>Unknown</td>
<td>Testis</td>
<td>Wolffian with rudimentary uterus and tubes</td>
<td>Male usually cryptorchid</td>
<td>Nl. testosterone</td>
<td>Nl. estrogen</td>
<td>None</td>
<td>None</td>
<td>Orchipexy</td>
</tr>
</tbody>
</table>

Notes:
- Inc. = Increased
- Nl. = Normal
- Wolffian = Male
- Female = Female
- Males = Males
Figure 43–8. Clinical approach to intersex. Algorithm based on palpating gonads. AS; bil., bilateral; CS/11 DCS, corticosterone/deoxycorticosterone; DEAS, 1,3-bis[4-(diethylamino)-2-hydroxy phenyl]-2,4-dihydroxy cyclobutenediylium dihydroxide, bis(inner salt); DCS/DOC, deoxycorticosterone/11-deoxycorticosterone; def., deficiency; dehyd., dehydration; 17 OH-P, 17 hydroxy progesterone; exp. lap., experimental laparoscopy; resist., resistance; synd., syndrome; T/DHT, testosterone/dihydrotestosterone.
structures such as the uterus and fallopian tubes will be important in determining the diagnosis (Figure 43–10A). The adrenal glands can also be examined for enlargement. While this finding is not diagnostic for CAH, it is suggestive and can direct further evaluation. Magnetic resonance imaging can provide a more detailed examination of the abdomen for internal genital structures. However, in most cases, anesthesia is needed for a good-quality magnetic resonance imaging examination. Injecting radiographic contrast material through the opening in the urogenital sinus is helpful in delineating the internal duct structures. It is most useful in assessing the presence of vagina, cervix, fallopian tube, utricle, and the connection with the urethra (Figure 43–10B and C). Genitography will also provide needed anatomical information for future reconstructive surgery.

### Diagnostic Laparotomy or Laparoscopy

Occasionally, surgery is needed to delineate the internal genitalia and obtain a biopsy specimen of the gonads. It is indicated in patients in whom the biopsy result will influence sex assignment. In addition, surgery may be needed to remove streak or dysgenetic gonads in patients who are at risk for cancer (incomplete testicular feminization, Turner’s Y variant, and mixed gonadal dysgenesis). Laparoscopic surgery has provided an alternative to open surgery in patients with intersex disorders. It can be performed safely in newborns and only requires 1–3 3-mm incisions for placement of the laparoscopic ports. Simple surgical procedures such as hernia repair, orchidopexy, and resection of discordant organs can be readily performed laparoscopically. More complex procedures may require 5-mm ports and larger instruments.

### Sex Assignment

In the past, the baby born with ambiguous genitalia was considered incomplete until either a male or female sex was assigned. Unfortunately, a prompt but inappropriate

---

**Table 43–4. Normal Values for Stretched Penile Length.**

<table>
<thead>
<tr>
<th>Age</th>
<th>Length (cm) (Mean ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Premature 30 wk</td>
<td>2.5 ± 0.4</td>
</tr>
<tr>
<td>Full-term newborn</td>
<td>3.5 ± 0.4</td>
</tr>
<tr>
<td>0–5 months</td>
<td>3.9 ± 0.8</td>
</tr>
<tr>
<td>6–12 months</td>
<td>4.3 ± 0.8</td>
</tr>
<tr>
<td>1–2 y</td>
<td>4.7 ± 0.8</td>
</tr>
<tr>
<td>2–3 y</td>
<td>5.1 ± 0.9</td>
</tr>
<tr>
<td>3–4 y</td>
<td>5.5 ± 0.9</td>
</tr>
<tr>
<td>5–6 y</td>
<td>6.0 ± 0.9</td>
</tr>
<tr>
<td>10–11 y</td>
<td>6.4 ± 1.1</td>
</tr>
<tr>
<td>Adult</td>
<td>12.4 ± 2.7</td>
</tr>
</tbody>
</table>

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**Figure 43–9.** Pathway of steroid hormone biosynthesis and possible enzyme deficiencies. 3β-HSD = 3β-hydroxysteroid dehydrogenase, 21α-H = 21α-hydroxylase, 11β-H = 11β hydroxylase, 17β-HSD = 17β-hydroxysteroid dehydrogenase, 18 HAS = 18 hydroxy-aldosterone synthetase, 18 OAS = 18 oxidase-aldosterone synthetase, 5α-R = 5α reductase, 19A = 19 aromatase, StAR = steroidogenic acute regulatory protein.
Figure 43-10. A: Newborn sonogram revealing a uterus (Ut) behind the bladder (Bl) in a patient with congenital adrenal hyperplasia. Note the dilated vagina (Vag), the cervix (arrow), and the bladder. B: Genitogram showing a high confluence (arrow) of the urethra and vagina with a long common urogenital sinus in a patient with congenital adrenal hyperplasia. C: Schematic of possible genitogram findings: I. and II. Urogenital sinus anomalies note 2 openings on the perineum (common urogenital sinus and rectum). III. and IV. Cloacal anomalies with 1 perineal opening. Note the low and high confluence of the urethra, vagina, and rectum (UVR). (Modified with permission from Dr Hardy Hendron.) Genitogram showing the common urogenital sinus.
assignment, although timely and comforting for family, physicians, nurses, and staff, can lead to more complex problems in the future. The issue of sexual determination remains complex. We reaffirm the teaching of our mentors by advocating an immediate and thorough attempt to make a definitive and accurate diagnosis. Fortunately, for most patients with ambiguous genitalia (ie, CAH) this can be accomplished. In patients where ambiguity remains after initial testing and the diagnosis cannot be made, or when the diagnosis is clear but sex assignment remains difficult, we would now advocate for a more cautious approach. Foremost, this would include a reversible or nonbinding sex assignment. Experience has shown that patients themselves may reassigned their sex. For example, in cases of cloacal exstrophy or iatrogenic penile injuries, past treatment was based on the absence of an “adequate” phallic structure. These patients were converted from genetic males to females with surgical orchiectomy, removal of any excess male genitalia, vaginoplasty, and future hormonal treatment for breast development. Although surgical results can be anatomically successful, these women will not menstruate or have fertility potential, and their sexual function is not known. A number of these patients went through adolescence, have identified as females, and have not had major issues with their assigned discordant, genetic sexual identity. In contrast, some of these patients have subsequently identified with their genetic sex and demanded or reassigned their sex from female to male. In cases where the genotype does not match the phenotype, it is clear that surgical reconstruction from male to female does not guarantee a successful sexual identity.

The clinical experience exemplifies the complexity of sexual determination. It is clear that social factors, or the “nurturing” hypothesis, and biologic factors, or the “genetic” hypothesis, both play a role in determining our sexual identity. The nurturing hypothesis is based on the parent’s perception of their child’s genitalia. This perception will influence interactions such as naming, clothes, play orientation, and social organization. Clearly, how a parent perceives his or her child and the type of environment used to raise the baby is critical to the child’s identity.

In contrast, the genetic hypothesis states that sexual identity is predetermined by the genetic makeup. Increasing laboratory evidence is accumulating to support the genetic hypothesis. For example, animal experimentation supports the concept of steroid or androgen imprinting of the brain. The human evidence supporting masculinization of the brain is supported by (1) women with virilizing CAH, (2) iatrogenic penile ablation in males raised as females, and (3) males with 5α-reductase deficiency who were raised as females. The common theme in these patients is the high level of in utero exposure to androgens theoretically masculinizing the brain and conferring a male identity. Another example of hormonal influence on sexual orientation can be found in women exposed to diethylstilbestrol. Human retrospective studies looking at these women reveal an increase in bisexual and homosexual orientation.

The process of sexual identity in both humans and experimental animals is not an all-or-none process, meaning that male and female characteristics exist as a continuum. For example, although the garbage removal, plumbing, and TV-channel-flipping gene seems to exist almost exclusively on the Y chromosome, these traits may also be found in the female sex.

Two issues must be separated when evaluating patients with intersex or ambiguous genitalia: (1) gender identity (Is the person’s sense of identity male or a female?) and (2) sexual orientation. The incidence of discordant gender identity is approximately 1 in 30,000 males and 1 in 100,000 females. The incidence of same-sex orientation in both males and females is estimated to be approximately 5–10% of the population.

Practical Approach to the Diagnosis of Intersex

In the newborn period, patients with ambiguous genitalia can be approached in a logical fashion (Figure 43–11). As noted above, history, physical examination, laboratory evaluation, and radiographic and in some cases surgical exploration are necessary to make an accurate diagnosis. Once the karyotype is known, along with the gonadal status, an appropriate test can lead to a diagnosis (Figure 43–11 and Table 43–5). Patients may also present at puberty (inappropriate or delayed development) with sexual differentiation abnormalities or later in life with infertility. The differential diagnosis for these disorders is diagrammed in Figure 43–12.

TREATMENT OF SPECIFIC DISORDERS

Female Pseudohermaphrodites

Female pseudohermaphrodites are characterized by a 46 XX genotype, nonpalpable gonads or normal ovaries, and variable degrees of virilization of the external genitalia.

A. CONGENITAL ADRENAL HYPERPLASIA

CAH is the most common cause of female ambiguous genitalia or pseudohermaphroditism and accounts for approximately 70% of all patients with ambiguous genitalia. CAH accounts for >95% of the cases of female pseudohermaphroditism, with exposure to maternal androgens accounting for the remaining 5%. Mutations in 1 of 5 genes result in impaired cortisol secretion, which in turn causes excess secretion of adrenocorticotropic hormone (ACTH) and consequently, adrenal hyperplasia. Four of the 5 genes code for enzymes necessary for steroid hormone synthesis, and
the fifth encodes for an intracellular cholesterol transport protein (StAR) (Figure 43–9). Deficiencies in 21α-hydroxylase and 11β-hydroxylase result in masculinization of the female fetus, while they have no effects on the genitalia of the male fetuses. In contrast, infants with deficiencies in 3β-hydroxysteroid dehydrogenase, 17α-hydroxylase, and StAR have defects in both the synthesis of cortisol and steroid hormones. Affected males have varying degree of ambiguous genitalia due to deficiency in testosterone synthesis, while affected females may or may not be virilized.

21α-hydroxylase deficiency is the most common cause of CAH, accounting for 90% cases. The metabolites 17-hydroxyprogesterone and 17-hydroxypregnenolone, which build up above the 21α-hydroxylase deficiency, are metabolized to androgens, resulting in virilization of the female external genitalia. Three forms of 21α-hydroxylase deficiency exist: classic, simple virilizing, and nonclassic. Each of these disorders is characterized by the activity level of the gene. Patients with the classic disease have both virilization and salt wasting, those with simple virilizing have masculinization without salt losing, and the nonclassic patients present after puberty with virilization.

In general, the classic form of 21α-hydroxylase deficiency exhibits the more severe forms of virilization (Figure 43–13). Impaired cortisol and aldosterone secretion leads to electrolytes and fluid losses, producing hyponatremia, hyperkalemia, acidosis, increased plasma renin, dehydration, and eventual vascular collapse unless recognized and treated. In affected males, deficiency in 21α-hydroxylase does not result in abnormal genitalia and consequently, salt loss may occur unnoticed. Aggressive fluid resuscitation with normal saline should be instituted immediately and repeat serum electrolyte measurement should be obtained to monitor the progress of the resuscitation. Diagnosis is based on an elevated level of 17-hydroxyprogesterone in the urine and blood. After diagnosis and stabilization, replacement therapy should be instituted with glucocorticoids, mineralocorticoids, and salt. Regular measurement of serum electrolytes, renin, and ACTH helps to monitor the adequacy of hormonal replacement. Untreated patients with 21α-hydroxylase deficiency exhibit excessive growth, virilization, advanced bone age, and early closure of epiphyseal growth plates.

11β-hydroxylase deficiency accounts for most of the remaining cases of CAH (approximately 9%). Patients

| Table 43–5. Differential Diagnosis for a Newborn with Ambiguous Genitalia. |
|-------------------------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Female pseudohermaphrodite (CAH) | XX | Ovary | Hypospadias | Present | Elevated |
| Male pseudohermaphrodite | XY | Testes | Hypospadias/micropenis | Absent | Normal |
| Mixed gonadal genesis | XY/XO | Streak dysgenetic | Hypospadias | Variable/rudimentary | Normal |
| True hermaphrodite | XX/mosaic | Ovotestis or ovary and testes | Hypospadias | Variable/rudimentary | Normal |

Figure 43–11. Algorithm for initial workup of intersex based on physical examination and karyotype.
Figure 43–12. Differential diagnosis of patients with inappropriate pubertal development, impaired pubertal development, and infertility. Def., deficiency; devel., development; DOC, 11-deoxycorticosterone; exp. lap., experimental laparoscopy; 17 OH-P, 17 hydroxy progesterone; synd., syndrome.

Figure 43–13. Patient with severe masculinization from congenital adrenal hyperplasia.
with 11β-hydroxylase accumulate 17-hydroxyprogesterone as well as DOC and 11-deoxycortisol, which results in salt accumulation leading to hypertension. Patients with 11β-hydroxylase deficiency are more likely to present with hypertension secondary to the salt-retaining metabolites DOC and 11-deoxycortisol, in contrast to the hypovolemic shock associated with 21α-hydroxylase deficiency. Hypokalemia is also common secondary to an increase in mineralocorticoid activity.

Since CAH is hereditary, it is possible to counsel and offer treatment to families wishing further children. Maternal treatment with dexamethasone prior to the 10th week of gestation can significantly reduce the risk of masculinization of the female fetus. Standard prenatal treatment is 20 (m)g/kg 2 times daily beginning as soon as the pregnancy is confirmed (5th week of gestation) in a family with a positive history of CAH. At 9–10 weeks’ gestation, chorionic villus sampling can confirm karyotype and test for the presence of the gene CYP 21, which is present in 21α-hydroxylase deficiency (90% of CAH cases). If the karyotype is XY or the CAH gene CYP 21 is not present, the maternal dexamethasone treatment is stopped. Statistically, 50% of the fetuses will be male and of the females, only 25% will be affected secondary to the recessive inheritance pattern of 21α-hydroxylase deficiency. Unfortunately, this will result in unnecessary prenatal steroid exposure in 7 of 8 fetuses with unknown long-term health consequences, such as hypertension. Although the short-term success of decreasing female virilization has been documented, long-term follow-up of fetuses exposed to steroids needs to be documented.

B. MATERNAL HORMONAL SOURCES OF VIRILIZATION

Maternal tumors are a rare cause of virilization of the female fetus. The most common type are luteomas of the ovary, which also virilize the mother. Diagnosis can be made by maternal blood samples and imaging studies (sonogram and magnetic resonance imaging). Maternal ingestion of medication is another rare cause of abnormalities in genital development (Table 43–6). Progesterone is a common agent being used early in pregnancy to prevent abortions as well as during in vitro fertilization treatments. The female fetus that is exposed to high concentrations of progesterone can virilize secondary to direct action of progesterone on the AR. In the male fetus, hypospadias can develop by progesterone-inhibiting testosterone synthesis and downregulating the AR. A prenatal history of progesterone exposure should be elicited in the differential diagnosis of patients with abnormalities of the external genitalia.

**Male Pseudohermaphrodites**

Male pseudohermaphrodites are characterized by a 46 XY genotype, normal testes (usual palpable), and partial or complete masculinization of the external genitalia. The differential diagnosis is outlined in Figure 43–14.

Two forms of androgen resistance related to male pseudohermaphrodites are complete androgen insensitivity and partial androgen insensitivity.

A. COMPLETE ANDROGEN INSensitivity

Androgen resistance ranges from partial to complete due to a defect in the AR. Patients with complete androgen resistance or androgen insensitivity syndrome (AIS) (previously called testicular feminization) have a 46 XY karyotype but have unambiguous female external genitalia, hypoplastic labia majora, a blind vaginal pouch, and an absent uterus. Since a functional AR is necessary for the development of auxillary and pubic hair, complete AIS patients have sparse to nonexistent hair growth in these areas. Complete AIS patients either inherit the disease by an X-linked recessive pattern or develop a spontaneous mutation that renders the AR nonfunctional. Patients with complete AIS appear to identify as females. Presumably the functional defect in the AR also exists in the brain, preventing “masculinization.” There is not enough long-term follow-up to assess issues with sexual identity in these patients.

Complete androgen resistance should be suspected in phenotypic females who present with an inguinal hernia that contains a testis (approximately 1% of all prepubertal females undergoing hernia repair). The most common presentation for complete AIS is amenorrhea in adolescent females. Breast development occurs in AIS patients secondary to the peripheral conversion of testosterone to estradiol from aromatase enzyme. After puberty, the testes have approximately a 10% risk of developing cancer, the most common tumor being a seminomatous germ cell. Because of the significantly increased cancer risk, removal of the gonads is recommended after postpubertal breast development. Alternatively, the gonads can be removed at the time of diagnosis, with estrogen replacement therapy initiated in the pubertal time period. Since the vagina may be inadequate in length, some patients may need augmentation procedures. Self-vaginal dilation is the most common technique, followed by vaginal augmentation procedures using skin grafts or bowel.

### Table 43–6. Drugs That May Induce Intersex Disorders if Taken during Pregnancy.

| C21-steroid medroxyprogesterone acetate (progesterone) |
| Finasteride (Proscar) |
| Leuprolide acetate (Lupron) |
| Stilbestrol |
| Danazol |
| Norethynodrel |
| Ethisterone |
| Norethindrone |
B. PARTIAL ANDROGEN INSENSITIVITY

In contrast to complete AIS, patients with partial androgen resistance may have external genitalia ranging from mild to severe hypospadias (with and without cryptorchidism) to micropenis or clitoromegaly with partial labial fusion (Figure 43–15). The testes may be located in the labia, inguinal canal, or abdomen. The testes are histologically normal before puberty. However, after puberty spermatogenesis is usually absent and there is Leydig cell hyperplasia. The testes are predisposed to malignant transformation in 4–9% of the patients.

The defect in partial androgen resistance is typically due to a single base pair mutation in the AR. Inheritance may be X-linked, autosomal recessive, or from a spontaneous mutation. Interestingly, the same genetic defect within a family may have a different phenotypic expression. The variability of phenotypic expression makes counseling difficult in affected families.

In patients with partial androgen resistance, the sex of rearing depends on the degree of androgen resistance and the degree of genital ambiguity. In patients who respond to high-dose androgen therapy (2 mg/kg initially followed by 4 mg/kg) with phallic growth, the sex of rearing as male has been successful. Genital reconstruction repairing the hypospadias and undescended testes is performed at an early age. Patients who have a poor response to androgen stimulation fall into a difficult category of intersex. In the past, patients who were raised as females had feminizing genital surgery and gonadectomy typically in the first year of life. At time of puberty, estrogen replacement is instituted. Presumably in partial androgen insensitivity, sexual identity is influenced by the effects of androgens on central imprinting. A discord may exist between the external genitalia that partially responds to androgen stimulation versus the effects of androgens on determining sexual identity in the brain. The fact that some patients with severe hypospadias and a small phallus have had difficulty with sexual identity in adulthood makes sex assignment difficult. Presently, it seems reasonable to delay irreversible surgery until after the patient has developed a sexual identity and can drive the decision for reconstructive surgery.

5α-REDUCTASE TYPE 2 DEFICIENCY

5α-reductase type 2 deficiency is an autosomal recessive transmitted disorder affecting the formation of the male genitalia. 5α-reductase is responsible for the conversion of the less potent testosterone to the 5–10 times more potent DHT. Type 2 5α-reductase predominates in the tissue of the external genitalia and the prostate, whereas type 1 5α-reductase localizes to the skin and nongenital
tissues. Numerous mutations have been described in the 5α-reductase type 2 gene consistent with the variation in clinical spectrum seen in patients with this defect. Immunohistochemical localization of 5α-reductase type 2 reveals that the enzyme is located in the midline urethral seam (Figure 43–16). The midline seam localization is consistent with the formation of hypospadias in patients with 5α-reductase type 2 gene defects in that the epithelial edges of the urethral seam would fail to fuse, resulting in hypospadias.

Clinically, patients with 5α-reductase type 2 present with a small phallus, severe hypospadias, bifid scrotum, and a residual prostatic utricle or blind-ending vaginal pouch (Figure 43–17). The testes are often undescended in the inguinal canal. Untreated patients will typically virilize during puberty when elevated levels of the less potent testosterone either overwhelm the functioning androgen gene or the functioning 5α-reductase type 1 enzyme cross-reacts with the excess testosterone, converting it to DHT.

Figure 43–15. Partial androgen receptor defect resulting in severe hypospadias with curvature (A) and a small phallus (B).

Figure 43–16. Immunohistochemical localization of 5α-reductase type 2 (A–D) and the androgen receptor (AR) (E–H) in the same human fetal penis at 16.5 weeks of gestation (reduced from 25×). Note strong expression of 5α-reductase type 2 along the urethral seam area (arrows).
Sexual identity appears to be intact for karyotype XY males with 5α-reductase type 2 deficiency, presumably from an intact masculinization of the brain. In specific geographic areas such as the Dominican Republic, where the incidence of 5α-reductase type 2 deficiency is relatively high, it is generally accepted that these children will change from an initial “in-between” sex to a male sexual identity at the time of puberty.

The diagnosis of 5α-reductase type 2 deficiency should be considered in severe phenotypes of hypospadias, especially with associated scrotal anomalies and undescended testes. Diagnosis is based on an increase in ratio of testosterone to DHT. Since these patients have a small phallus, attempts at enlargement with DHT cream are reasonable, although DHT is difficult to obtain in the United States. Reconstructive surgery for the hypospadias and undescended testes is indicated. Fertility has not been reported in patients with 5α-reductase type 2, although sperm production has been documented.

**Persistent Müllerian Duct Syndrome**

Müllerian-inhibiting substance (MIS) or factor (anti-Müllerian duct hormone) causes regression of the structures that would have formed the uterus, fallopian tube, and upper part of the vagina. Defects in the MIS gene or MIS receptor result in retained Müllerian structures typically inherited as an autosomal recessive defect. Male siblings of affected patients, especially with cryptorchidism, should undergo screening; they have a 25% chance of being affected.

Clinically, patients with persistent Müllerian duct syndrome present, unexpectedly, at the time of surgery for cryptorchidism (Figure 43–18). Hence the alternate name

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**Figure 43–17.** A patient with 5α-reductase type 2 deficiency. Note severe hypospadias with a small phallus, bifid scrotum, and visible prostatic utricle or blind-ending vaginal pouch.

**Figure 43–18.** Hernia uterine inguinal, or persistent Müllerian duct syndrome. Note the presence of a fallopian tube (black arrow) and uterus attached (white arrow) to the testicular cord structures.
for persistent Müllerian duct syndrome, hernia uterine inguinal. Within the hernia sac a fallopian tube, uterus, or both are found attached to the testicular cord structures. What makes the treatment difficult is that these structures and hence the diagnosis are found unexpectedly at the time of surgery for cryptorchidism. If persistent Müllerian duct structures are found during orchiopexy, it is reasonable to abort the procedure until a correct diagnosis can be determined. At the initial exploration, a clear description of gonad and surrounding Müllerian structures should be documented, with a biopsy specimen of the gonad taken and a karyotype obtained.

Once a definitive diagnosis is made, reconstructive surgery can then be performed. Separation of inappropriate Müllerian structures from the cord without disturbing the vas deferens, the testicular artery, or both is the goal; however, this may be impossible if the vas runs through the Müllerian structures, which is a common outcome. Fertility is usually impaired in patients with persistent Müllerian duct syndrome even though testosterone levels may be normal. Whether this is a consequence of primary gonadal dysfunction or secondary to the cryptorchid testes is controversial. Efforts should be made to remove the Müllerian structures and deliver the testes into the scrotum or at least a palpable position in the groin for subsequent cancer surveillance. Testes cancer has been reported in 2–10% of patients. In patients where the testes remains in the abdomen or cannot be separated from the Müllerian structures, orchiectomy is indicated.

Abnormal Gonadal Function Syndromes

A. Turner’s Syndrome

Turner’s syndrome is relatively common, occurring in 1 in every 2000 female births. The genotype in patients with Turner’s syndrome is a complete or mosaic X monosomy, 45,X, or 45,X/46,XX. Turner stigmata consist of a web neck, shield chest, aortic valve defects, coartation of the aorta, horseshoe kidney, short stature, and absent puberty. During fetal development in patients with Turner’s syndrome the ovaries develop but subsequently degenerate to streak gonads. The streak gonads are not at risk for cancer because of gonad and surrounding Müllerian structures usually are present secondary to failure of MIS secretion. Laboratory analysis reveals female levels of baseline testosterone with no increase in response to hCG stimulation. Surveillance reveals streak gonads, fallopian tubes, and a uterus. With a 60% chance of tumor, the gonads need to be removed once the diagnosis is confirmed. These patients should be raised as females with estrogen replacement at the time of puberty.

B. Pure Gonadal Dysgenesis

Patients with 46, XX complete gonadal dysgenesis are typically diagnosed following a workup for delayed puberty or primary amenorrhea. Patients have a normal female phenotype without the stigmata of Turner’s syndrome, normal external and internal Müllerian structures, and bilateral streak gonads. Sexual identity is female. Unlike patients with 46, XY gonadal dysgenesis, risk of tumor formation is rare and treatment is directed at hormonal replacement, with removal of the streaks gonads unnecessary.

C. XY Gonadal Dysgenesis

Patients with 46, XY gonadal dysgenesis are characterized by absent testicular function in the presence of a Y chromosome. Classically, patients with 46, XY gonadal dysgenesis have a female phenotype. Patients come to medical attention if the prenatal karyotype (XY) is discordant with the child’s phenotype (female), delayed puberty, amenorrhea, or precocious puberty from a hormonally functional gonadal tumor. The incidence of gonadal tumors is as high as 60%, with gonadoblastoma being the most common, although dysgerminomas, seminomas, and nonseminomatous germ cell tumors have also been reported.

In pure XY gonadal dysgenesis, Müllerian duct structures usually are present secondary to lack of testosterone secretion. Laboratory analysis reveals female levels of baseline testosterone with no increase in response to hCG stimulation. Surgical exploration reveals streak gonads, fallopian tubes, and a uterus. With a 60% chance of tumor, the gonads need to be removed once the diagnosis is confirmed. These patients should be raised as females with estrogen replacement at the time of puberty.

D. Mixed Gonadal Dysgenesis

Patients with mixed gonadal dysgenesis usually have a 45,X/46,XY, 46XY, or other mosaic karyotype. They typically have 1 streak and 1 dysgenetic testis. Most children with mixed gonadal dysgenesis have incomplete virilization resulting in ambiguous genitalia or hypospadias with cryptorchidism. The other classic presentation is a mosaic genotype diagnosed on prenatal amniocentesis. Interestingly, the subsequent phenotype of patients with a prenatal karyotype of 45,X/46,XY is 90% normal male external genitalia. However, with a prenatal genotype of 45,X/46,XY the patient is at risk for progressive gonadal changes leading to fibrosis and decreased fertility and low testosterone levels. The incidence of gonadal tumors does not seem to be increased. Most notably, 20% of these children have mental retardation or autism.

In patients who present with ambiguous genitalia, one gonad is typically palpable in the scrotum or inguinal canal and the other gonad (streak) nonpalpable. The phallus size is typically small with a proximal or more severe hypospadias (Figure 43–19). Testosterone levels are normal with an appropriate response to hCG. MIS levels are usually normal. At surgery, the dysgenetic gonad (streak) may grossly appear normal but have microscopic abnormalities such as hypoplastic tubules surrounded by ovarian or...
fibrotic stroma. Variable Müllerian duct structures, such as fallopian tubes and uterus, are present depending on the degree of gonadal dysgenesis. On biopsy, the contralateral gonad in the scrotum or inguinal canal is either a normal or dysgenetic testis. In patients with mixed gonadal dysgenesis, the risk of gonadoblastoma is 15–30%. Gonadoblastoma is a steroid hormone–secreting gonadal tumor composed of large germ cells, Sertoli cells, and stromal derivatives. The incidence of gonadoblastoma appears to be higher in more undervirilized patients and the most common associated karyotype is 46 XY. Sixty percent of gonadoblastomas arise in an indeterminate gonad, 22% in streak gonads, and 18% in dysgenetic cryptorchid testis. Two cases occurring in a testis located in the scrotum have been reported. One-third of the patients have bilateral disease. Sixty percent of gonadoblastomas are associated with subsequent malignant germ cell tumor (germinoma, seminoma, and dysgerminoma but also embryonal teratoma, embryonal carcinoma, endodermal sinus tumor, or choriocarcinoma). Metastases develop in 10% of patients with germinomas arising within the gonadoblastoma.

In children who are undervirilized, female sex assignment is an option, and the streak and dysgenetic gonads should be removed at time of diagnosis due to increased risk of malignancy. Hormonal replacement with estrogen will be necessary during adolescence. If male gender is assigned, management of the scrotal testis is controversial, ranging from serial observation to surveillance biopsy. In the virilized patients who are raised as males, the testis will inevitably reveal poor hormonal and fertility potential. These patients will require testosterone supplementation in adulthood.

In 5% of patients, mixed gonadal dysgenesis is associated with Wilms’ tumor, ambiguous genitalia, and progressive glomerulopathy known as the Denys-Drash syndrome. Wilms’ tumor occurs in the first 2 years of life and is often bilateral. Classic presentation is an infant with ambiguous genitalia, hypertension, and nephrotic syndrome.

**E. 17β-HYDROXOSTEROID DEHYDROGENASE DEFICIENCY**

Patients with a defect in the enzyme 17β-hydroxysteroid dehydrogenase do not efficiently convert androstenedione to testosterone. 17β-hydroxysteroid dehydrogenase is predominantly located in the testes. The rare disorder of 17β-hydroxysteroid dehydrogenase deficiency is inherited via an autosomal recessive pattern. This disorder is indigenous
to the Arab population of the Gaza strip in the Middle East. Clinical presentation in a patient with XY genotype is mild virilization of the external genitalia, with clitoral hypertrophy, and a blind-ending utricle (vagina). The testes are undescended in the abdomen or inguinal canal or descended into the labioscrotal folds. If the virilization is mild, the diagnosis becomes apparent at puberty, with penile growth and male secondary sexual characteristics. At puberty the increased levels of androstenedione are converted by nongenital, nonmutant 17α-hydroxysteroid dehydrogenase to testosterone. These patients may also present with gynecomastia at puberty by the peripheral conversion of androstenedione to estradiol by aromatase. Diagnosis is based on an increased ratio of androstenedione to testosterone postpubertal or in the prepubertal state in response to an hCG-stimulation test.

If the diagnosis is suspected in infancy, treatment with testosterone, reconstruction of the hypospadias, and male sex assignment is indicated. At puberty in the Gaza strip, gender conversion from female to male is common practice. Long-term outcomes of patients raised as females initially and reassigned to males at puberty await documentation.

**True Hermaphroditism**

True hermaphroditism is defined as the presence of both ovarian and testicular tissue within the same individual (Figure 43–20). The most common karyotype in patients with true hermaphroditism is 46 XX (predominately in African Americans), followed by 46 XY/46 XX mosaicism. The latter karyotype in a patient with ambiguous genitalia strongly suggests the diagnosis of true hermaphroditism. Only 7% of patients with this disorder have a 46 XY karyotype. Interestingly, not all true hermaphrodites express the SRY gene, suggesting that non-SRY genes play a role in the development of the testes in these patients.

In patients with true hermaphroditism, the gonads are a combination of ovotestis, ovaries, or testis. The most common configuration is ovotestis/ovary in 35%, followed by bilateral ovotestis in 25%, ovary/testes in 25%, and ovotestis/testes in the remaining 15%. One or both gonads are palpable in at least 60% of the patients. For unexplained reasons, the testes is more likely to be found on the right side. The testis and ovaries are located in their respective normal position, and the level of descent of the ovotestis is dependent on the amount of testicular tissue. While ovarian histology and function may be normal, testicular histology and function is usually abnormal. Ovotestis can be bilobar in configuration, with the ovarian and testicular tissue relatively separate, or the ovarian and testicular tissue may be intermingled and difficult to surgically separate. At the time of diagnosis, deep biopsies are necessary to determine the histologic status of the gonad. The internal structures tend to correlate with the type of gonad. Approximately 80% of true hermaphrodites will have a functional or rudimentary uterus. The uterus may be found in the abdomen or associated with an inguinal hernia. In patients with normal uterine structures and ovarian histology, fertility and normal pregnancies have been reported.

The external genitalia are usually ambiguous, although 60% of patients are masculinized, with a well-developed

*Figure 43–20.* Finding at the time of surgical exploration in a true hermaphroditic. On the patient’s right side, note the testes, and on the left, note the fallopian tube, uterus, and biopsy-proven ovary.
Hypospadiac phallus. The hypospadias can be severe perineal or penile-scrotal with incomplete fusion of the labioscrotal folds. The degree of masculinization is dependent on the amount of functional testicular tissue present. In childhood, testicular tissue has been documented to have normal spermatogonia. With maturation, however, testicular fibrosis occurs, with fertility in males a rare event. Testicular tumor is uncommon, occurring in 1–2% of the patients.

The diagnosis of true hermaphroditism should be suspected in patients with virilized ambiguous genitalia who have a 46 XX (African American) or mosaic genotype 46 XX/XY associated with the finding of Müllerian structures. Diagnosis is confirmed by gonadal biopsy confirming the presence of both ovarian and testicular tissue. After a decision regarding sex gender assignment has been made, gonadal tissue inappropriate for sex gender assignment should be removed. In patients who are raised as females, removal of all functioning testicular tissue is critical to prevent virilization at puberty. Surgical correction of the urogenital sinus to expose the vagina is necessary. In patients raised as males—who account for approximately 30% of all true hermaphrodites—the hypospadias and undescended testes should be reconstructed. In males, since testicular failure is common at puberty, testosterone supplementation may be required.

Unclassified Forms of Abnormal Sexual Differentiation

A. HYPOSPADIAS

Hypospadias is not a form of intersex (Figure 43–21). The etiology can be defined in less than 5% of patients. This leaves most cases without a defined etiology. The variable expression of the AR in the ventral versus the dorsal urethra may play a role in the etiology of hypospadias (Figure 43–22). Recent theories suggest an abnormality in closure
of the midline urethral seam. Another possible etiology explaining the increase in incidence of hypospadias in Western countries over the last 25 years is an increase in exposure to environmental endocrine disruptors.

In controlled studies, most patients with hypospadias undergo successful surgical reconstruction and have acceptable long-term outcomes. Patients with hypospadias have an unambiguous male sexual identity. In severe forms of hypospadias with perineal or scrotal urethral openings, severe curvature and the phallus buried within the scrotum are the critical issues confirming the correct diagnosis. This is also the case for patients with hypospadias and a nonpalpable or undescended testis. If any doubt exists, patients with severe hypospadias, hypospadias in association with an undescended testis, or both, a karyotype should be checked to document genotype. In severe cases of hypospadias where penile size is difficult to assess secondary to severe chordee, an hCG stimulation will assess the gonadal axis and confirm an intact AR by eliciting penile growth.

### B. MICROPENIS

A small penis defined as less than 2.5 cm in length without hypospadias in a full-term male is defined as micropenis (Figure 43–23 and Table 43–4). Micropenis can be caused by multiple etiologies, the most common being fetal testosterone deficiency followed by partial defects in the AR or 5α-reductase enzyme (Table 43–7). Fetal testosterone synthesis can be divided into 2 categories: (1) primary testicular failure (Leydig cell) and (2) central failure. Central failure can be from congenital hypopituitarism or isolated gonadotropin deficiency. Patients with decreased fetal testosterone production either from Leydig cell failure or lack of Leydig cell stimulation from gonadotropin deficiency respond to treatment with supplementary testosterone enanthate intramuscular injections 25–50 mg each month for 3 consecutive months.

Long-term outcomes of patients with micropenis have documented that final adult penile length is normal for >90% of patients treated with multiple short courses of testosterone enanthate. In addition, patients with micropenis identified with the male gender had normal erections, ejaculation, and orgasm. In the rare patient who does not respond to testosterone stimulation, gender conversion to female had been advocated in the past. Presently, gender conversion would not be considered based solely on the small phallus size.

Reassignment to the female gender with removal of the gonads and feminizing genitoplasty in patients with penile agenesis, iatrogenic penile amputation, or circumcision injury had been standard treatment. In complete penile agenesis, the testicles are normal, corporeal bodies are absent, and the urethra opens into the anterior rectum or

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**Figure 43–22.** Androgen receptor (AR) expression in the human fetal penis at 16.5 weeks. A greater density of AR-positive cells is seen in the ventral portion of the urethral epithelium in the distal glans (A), mid glans (B), and proximal glans (C). In the distal (E), mid (F), and proximal (G) shaft of the penis, all portions of the urethral epithelium show the same density of expression. Three-dimensional reconstruction was performed to demonstrate the urethral AR expression pattern (D). Note the weaker density of AR in the dorsal aspect of the glanular urethra.
ABNORMALITIES OF SEXUAL DETERMINATION & DIFFERENTIATION

perineum. These patients have normal prenatal androgen levels and hence the brain has received signals for male sexual identity. The same is true for the rare patient who has a severe penile injury during circumcision. As in micropenis, gender conversion would now not be considered based solely on the absence or small size of the phallus. Penile reconstruction, although not technically ideal, may provide the best overall outcome.

C. CLOACAL AND EXSTROPHY ANOMALIES

In the past, patients with the most severe and rare form of lower abdominal congenital malformation, cloacal exstrophy (incidence of 1:200,000 live births), were usually left to die. Significant problems associated with surgical reconstruction of cloacal exstrophy include omphalocele; numerous gastrointestinal anomalies such as short gut, malrotation, duplication, duodenal atresia, and Meckel’s diverticulum; and significant genitourinary anomalies such as separate bladder halves, upper tract renal anomalies, and bifid genitalia. Patients with cloacal exstrophy can also have neurologic and orthopedic anomalies such as tethered cord, myelomingingocele, lower extremity paralysis, clubfoot, and hip dislocation.

Historically, newborn males with cloacal exstrophy (Figure 43–24) were often gender-converted to female as a result of inadequate genital development and the poor prognosis for surgically developing a normal male phenotype. In rearing genetic males as females, although the surgical reconstruction can match the assigned female phenotype, a new set of issues was created, such as the need for hormonal replacement with estrogen during adolescence and the issue of a nonmenstruating infertile female. In addition, the fetal and neonatal androgen imprinting on the brain does not seem to be reversible.

Because some of these XY, gender-converted females have self-reassigned their sex during adolescence to coincide with their genetic karyotype, there has been reevaluation of the practice of rearing genetic males as females. With the exact determinates of sexually identity not completely defined, a pragmatic approach is to delay any irreversible surgery such as orchiectomy or phallic removal/reduction in these patients. With modern surgical techniques and a multidisciplinary approach to their care, children with this complex disorder can have a normal sexual identity.

SURGICAL MANAGEMENT OF INTERSEX

The surgical management of patients with intersex is undergoing a reevaluation. The determination of a patient’s sexual identity is strongly influenced by the genetic karyotype and steroid/androgen action on the developing brain. The environmental and social impacts are certainly important but presently seem to have a less dominating influence.
We would advocate surgical management of patients with intersex when the diagnosis is clearly established and the long-term outcome for the diagnosis is favorable. Surgery falls into 5 categories: (1) diagnostic/biopsy, (2) gonadectomy and removal of inappropriate Müllerian structures, (3) clitoral reduction, (4) vaginoplasty, and (5) phallic reconstruction.

Diagnostic techniques have improved with the widespread use of laparoscopy to assess the morphology of the internal genital structures. Laparoscopic techniques allow the gonads and associated structures to be visualized and in some cases have a biopsy specimen taken without the need for a laparotomy open incision. Once the diagnosis is established, it is possible to remove an inappropriate gonad, Müllerian remnant, or both via laparoscopic techniques.

**Clitoroplasty**

Clitoroplasty is presently a controversial topic. No studies exist to clearly document whether androgen stimulation resulting in a large clitoris requires reduction or can be left intact. Clearly, surgery on the clitoral structures can result in nerve damage and removal of erectile tissue.

Historically, the enlarged clitoris/phallic structure has been dealt with by amputation. Subsequently, more refined techniques such as recession clitoroplasty were developed where the entire clitoral organ is preserved by imbricating and burying the proximal corporeal shaft and excess glans clitoris. The disadvantage of the clitoral recession procedures may not become apparent until puberty, when the recessed corporeal bodies become enlarged and painful during sexual stimulation. This leads to the need for a procedure involving subtotal resection of the shaft of the clitoris with preservation of the glans.

In cases where clitoroplasty is performed, the goal is to recreate the normal female anatomy. Presently, more conservative procedures have been employed to preserve both the sensory and cosmetic aspects of the clitoris. An understanding of normal female anatomy has benefited the design of reconstructive surgery in patients with CAH (Figures 43–25 and 43–26). A contemporary reduction clitoroplasty is based on anatomical observations from fetal anatomical dissections. Presently, the tunica of the corporeal body can be preserved to spare as much of the dorsal nerve as possible. The concept of lifting the dorsal nerve off the tunica at the 11 and 1 o’clock positions seems inconsistent with the fact that the nerves fan out extensively around the dorsal and lateral aspects of the clitoral body.

A second issue is the removal of erectile tissue. In severe cases of masculinization of the genitalia (Prader V), consideration may be given to reduce the amount of erectile tissue. Standard treatment was to amputate the erectile body of the clitoris at the pubic arch, leaving each crural body and the neurovascular bundle with a strip of dorsal tunica. The long-term effects of removing this erectile tissue on sexual function are unknown. In contrast, leaving too much erectile tissue has been reported to cause pain in patients at the time of puberty. This, however, may be from fixing the corporeal tissue to the pubic bone, a practice that is no longer advocated. A compromise is to incise

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**Figure 43–24.** A: Male with cloacal exstrophy. B: Female with cloacal exstrophy. In the male note the split scrotal appearance and the small hemi-phallus (arrow). In the female the clitoral bodies/genitalia are not visible.
the corpora cavernosal body on the ventral surface at the 6 o’clock position and remove erectile tissue within the tunica to reduce the size of the erectile body, thereby preserving some erectile tissue and the nerves of the clitoris.

**Vaginoplasty**

The timing of vaginoplasty is also a controversial issue in genital reconstruction. The presence of a vagina is not necessary until puberty and initially only to allow the passage of menstrual fluids. Later the vagina is necessary for vaginal penetration, fertility and, in most females, a healthy female sexual identity. This may not be the case for all females, for example, a women with a small vagina and a female sexual identity with a female sexual preference may not desire a larger vagina. In patients with absent Müllerian structures (specifically a functional uterus) who have a female sexual identity, menstruation is not an issue and timing of vaginoplasty can be driven by the patient’s wishes and motivation. In patients with a common urogenital sinus and hidden vagina, there are advantages and disadvantages of early surgery in the first year of life versus late surgery prior to puberty. The advantage of early vaginoplasty is the closeness of the vagina to the perineum and the reported decreased bleeding in the early years of life. The major disadvantages are the smallness of the structures and that secondary surgery at the time of puberty may be necessary to correct vaginal stenosis. In contrast, delaying surgery has the advantages of operating on larger structures and the possibility that the patient can perform postoperative vaginal dilation to prevent stenosis.

The type of vaginoplasty depends on the level of masculinization. For low urogenital sinus anomalies, a flap vaginoplasty will usually allow for an adequate introitus with separation of the urethra and vagina (Figure 43–27). For high urogenital sinus anomalies, use of the elongated common urogenital sinus as an anterior vaginal flap may be necessary (Figure 43–28). In the case of an absent vagina or a very short vagina, substitution vaginoplasty with bowel or skin grafting maybe required.

**Phallic Reconstruction**

Phallic reconstruction is a formidable task. Nevertheless, it is critical that reconstructive efforts continue in this area, especially for patients with penile agenesis or iatrogenic injuries and a XY genotype and functional AR. Several techniques have been devised, such as free microanastomosis, innervated radial forearm flaps, tubed abdominal flaps with a penile prosthesis, and rectus abdominus myocutaneous flaps. In the free radial forearm flap, the pudendal nerve is anastomosed to the lateral cutaneous nerve of the
Figure 43–26. Normal human fetal clitoris, 24 weeks’ gestation. Four views of a computer-generated three-dimensional reconstruction (A: top; B: bottom; C: back/top; D: bottom.) Note the pathway of the nerves (light gray) with a paucity of nerves on the bottom of the clitoris as well as in the top midline.
Figure 43–27. Female external genitalia reconstruction in patients with a low confluence using a flap vaginoplasty.

A: Surgical schematic of the perineum in patients with common urogenital sinus. B: The anterior flap for the vagina can be created using the phallic skin or the distal portion of the urogenital sinus. 2 openings are created in the midline of the preputial skin flap to accommodate the clitoris and urethra. The preputial skin flap is then brought down and sutured to the anterior wall of the vagina. C: The completed repair. D: Alternatively, the preputial skin can be split in the midline and used for reconstruction of the vaginal introitus and the anterior vaginal wall. E: The completed repair. (Used with permission from Nguyen HT, Baskin LS: A Child with Ambiguous Genitalia. American Urological Association Patient Management Problems, vol. 6:2. Decker Electronic Publishing Inc, 2002.)
Figure 43–28. High Urogenital Sinus Repair. A: The urogenital (UG) sinus is separated from the rectum posteriorly and the pubic bone anteriorly. B: The posterior skin flap (arrows) is assessed for length to reach the vagina. C: The confluence of the vagina and urethra (arrow) is separated. (Used with permission from Nguyen HT, Baskin LS: A Child with Ambiguous Genitalia. American Urological Association Patient Management Problems, vol. 6:2. Decker Electronic Publishing Inc, 2002.)
forearm. The radial artery and vein are anastomosed to the inferior epigastrics, the internal pudendals, or the femoral vessels. The major complications with these procedures are fistula, prosthesis erosion, and poor sensation. The technical nuances of microvascular anastomosis require that the procedures be performed in adolescents and adulthood. The psychological implications of relatively late reconstruction have not been determined. With newer tissue engineering techniques, better phallic reconstruction procedures may be on the horizon.

REFERENCES


Male Infertility

Paul J. Turek, MD

Infertility is defined as the inability to conceive after 1 year of unprotected sexual intercourse. Infertility affects approximately 15% of couples. Roughly 40% of cases involve a male contribution or factor, 40% involve a female factor, and the remainder involve both sexes. The evaluation of male infertility is undertaken methodically to acquire several kinds of information. Before discussing the diagnosis and treatment of male infertility, a review of basic reproductive tract physiology is in order.

■ MALE REPRODUCTIVE PHYSIOLOGY

THE HYPOTHALAMIC–PITUITARY–GONADAL AXIS

The physiology of the hypothalamic-pituitary-gonadal (HPG) axis plays a critical role in each of the following processes, the last 2 of which are relevant for reproduction:

1. Phenotypic gender development during embryogenesis
2. Sexual maturation during puberty
3. Endocrine function of the testis: testosterone
4. Exocrine function of the testis: sperm

Basic Endocrine Concepts

A. HORMONE CLASSES (Figure 44–1)

Two kinds of hormones classically mediate communication in the reproductive axis: peptide and steroid. Peptide hormones are small secretory proteins that act via receptors on the cell surface membrane. Hormone signals are transduced by 1 of 3 second-messenger pathways, as outlined in Figure 44–1. Ultimately, most peptide hormones induce the phosphorylation of various proteins that alter cell function. Examples of peptide hormones are luteinizing hormone (LH) and follicle-stimulating hormone (FSH).

In contrast, steroid hormones are derived from cholesterol and are not stored in secretory granules; consequently, steroid secretion rates directly reflect production rates. In plasma, these hormones are usually bound to carrier proteins. Since they are lipophilic, steroid hormones are generally cell membrane permeable. After binding to an intracellular receptor, steroids are translocated to deoxyribonucleic acid (DNA) recognition sites within the nucleus and regulate the transcription of target genes. Examples of reproductive steroid hormones are testosterone and estradiol.

B. FEEDBACK LOOPS

Normal reproduction depends on the cooperation of numerous hormones, the regulation of which is well controlled. Feedback control is the principal mechanism through which this occurs. With feedback, a hormone can regulate the synthesis and action of itself or of another hormone. Further coordination is provided by hormone action at multiple sites and through multiple responses. In the HPG axis, negative feedback is responsible for minimizing hormonal perturbations and maintaining homeostasis.

Components of the Hypothalamic–Pituitary – Gonadal Axis (Figure 44–2)

A. HYPOTHALAMUS

As the integrative center of the HPG axis, the hypothalamus receives neuronal input from many brain centers, including the amygdala, thalamus, pons, retina, and cortex, and is the pulse generator for the cyclical secretion of pituitary and gonadal hormones. It is anatomically linked to the pituitary gland by both a portal vascular system and neuronal pathways. By avoiding the systemic circulation, the portal vascular system directly delivers hypothalamic hormones to the anterior pituitary. Of the several hypothalamic hormones that act on the pituitary gland, the most important one for reproduction is gonadotropin releasing hormone (GnRH) or luteinizing hormone releasing hormone (LHRH), a 10-amino acid peptide secreted from the neuronal cell bodies in the preoptic and arcuate nuclei. At present, the only known function of GnRH is to stimulate the secretion of LH and FSH from the anterior pituitary. Once secreted into the pituitary portal circulation,
GnRH has a half-life of approximately 5–7 minutes, almost entirely removed on the first pass through the pituitary either by receptor internalization or enzymatic degradation.

GnRH secretion results from integrated input from a variety of influences, including the effects of stress, exercise, and diet from higher brain centers, gonadotropins secreted from the pituitary, and circulating gonadal hormones. Known substances that regulate GnRH secretion are listed in Table 44–1.

GnRH secretion is pulsatile in nature. This secretory pattern governs the concomitant cyclic release of the gonadotropins LH and FSH (to a lesser extent) from the pituitary. The pulse frequency appears to vary from once hourly to as seldom as once or twice in 24 hours. The importance of the pulsatile GnRH secretory pattern in normal reproductive function is aptly demonstrated by the ability of exogenous GnRH agonists Lupron or Zoladex (leuprolide acetate) to halt testosterone production within the testicle by changing the pituitary exposure to GnRH from a cyclic to a constant pattern.

**B. ANTERIOR PITUITARY**

The anterior pituitary gland, located within the bony sella turcica of the cranium, is the site of action of GnRH. GnRH stimulates the production and release of FSH and LH by a calcium flux-dependent mechanism. These peptide hormones were named after their elucidation in the female, but it is recognized that they are equally important in the male. The sensitivity of the pituitary gonadotrophs for GnRH varies with patient age and hormonal status.

LH and FSH are the primary pituitary hormones that regulate testis function. They are both glycoproteins composed of 2 polypeptide chain subunits, termed alpha and beta, each coded by a separate gene. The alpha subunit of each hormone is identical and is similar to that of all other pituitary hormones; biologic and immunologic activity are conferred by the unique beta subunit. Both subunits are required for endocrine activity. Sugars linked to these peptide subunits, consisting of oligosaccharides with sialic acid residues, differ in content between FSH and LH and may account for differences in signal transduction and plasma clearance of these hormones.

Secrertory pulses of LH vary in frequency from 8 to 16 pulses in 24 hours and vary in amplitude by 1- to 3-fold. These pulse patterns generally reflect GnRH release. Both androgens and estrogens regulate LH secretion through negative feedback. On average, FSH pulses occur approximately every 1.5 hours and vary in amplitude by 25%. The FSH response to GnRH is more difficult to measure than that of LH because of a smaller amplitude response.

<table>
<thead>
<tr>
<th>GnRH Modulator</th>
<th>Type of Feedback</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Opioids</td>
<td>Negative/inhibitory</td>
<td>β-endorphin</td>
</tr>
<tr>
<td>Catecholamines</td>
<td>Variable</td>
<td>Dopamine</td>
</tr>
<tr>
<td>Peptide hormones</td>
<td>Negative/inhibitory</td>
<td>FSH, LH</td>
</tr>
<tr>
<td>Sex steroids</td>
<td>Negative/inhibitory</td>
<td>Testosterone</td>
</tr>
<tr>
<td>Prostaglandins</td>
<td>Positive/stimulatory</td>
<td>PGE₂</td>
</tr>
</tbody>
</table>

FSH, follicle-stimulating hormone; LH, luteinizing hormone; PGE₂, prostaglandin E₂.
and a longer serum half-life. The gonadal proteins inhibit and activin may exert significant effects on FSH secretion and are thought to account for the relative secretory independence of FSH from GnRH secretion. They will be discussed in the Testis section.

The only known effects of FSH and LH are in the gonads. They activate adenylate cyclase, which leads to increases in intracellular cyclic adenosine monophosphate (cAMP). In the testis, LH stimulates steroidogenesis within Leydig cells by inducing the mitochondrial conversion of cholesterol to pregnenolone and testosterone. FSH binds to Sertoli cells and spermatogonial membranes within the testis and is the major stimulator of seminiferous tubule growth during development. FSH is essential for the initiation of spermatogenesis at puberty. In the adult, the major physiologic role of FSH is to stimulate quantitatively normal spermatogenesis.

A third anterior pituitary hormone, prolactin, can also affect the HPG axis and fertility. Prolactin is a large, globular protein of 199 amino acids (23 kDa) that is known to affect milk synthesis during pregnancy and lactation in women. The role of prolactin in men is less clear, but it may increase the concentration of LH receptors on the Leydig cell and help sustain normal, high intratesticular testosterone levels. It may also potentiate the effects of androgens on the growth and secretions of male accessory sex glands. Normal prolactin levels may be important in the maintenance of libido. Although low prolactin levels are not necessarily pathologic, evidence suggests that hyperprolactinemia abolishes gonadotropin pulsatility by interfering with episodic GnRH release.

### C. THE TESTIS

Normal male virility and fertility require the collaboration of the exocrine and endocrine testis. Both units are under the direct control of the HPG axis. The interstitial compartment, composed mainly of Leydig cells, is responsible for steroidogenesis. The seminiferous tubules have an exocrine function with spermatogenesis as the product.

#### 1. Endocrine testis

Normal testosterone production in men is approximately 5 g/day, and secretion occurs in a damped, irregular, pulsatile manner. In normal men, approximately 2% of testosterone is “free” or unbound and considered the biologically active fraction. The remainder is almost equally bound to albumin or sex hormone-binding globulin (SHBG) within the blood. Several pathologic conditions can alter SHBG levels within the blood, thus altering the amount of free or bioactive testosterone available for tissues. Elevated estrogens and thyroid hormone decrease plasma SHBG and therefore increase the free testosterone fraction, whereas androgens, growth hormone, and obesity increase SHBG levels and decrease the active androgen fraction. Testosterone is a profound regulator of its own production through negative feedback on the HPG axis.

Testosterone is metabolized into 2 major active metabolites in target tissues: (1) the major androgen dihydrotestosterone (DHT) from the action of 5-alpha-reductase and (2) the estrogen estradiol through the action of aromatases. DHT is a much more potent androgen than testosterone. In most peripheral tissues, testosterone reduction to DHT is required for androgen action, but in the testis and probably skeletal muscle, conversion to DHT is not essential for hormonal activity.

#### 2. Exocrine testis

The primary site of FSH action is on Sertoli cells within the seminiferous tubules. In response to FSH binding, Sertoli cells make a host of secretory products important for germ cell growth, androgens and thyroid hormone decrease plasma SHBG and thus alter the amount of bioactive testosterone available for tissues.

#### 3. Inhibin and activin

Inhibin is a 32-kDa protein derived from Sertoli cells that specifically inhibits FSH release from the pituitary. Within the testis, inhibin production is stimulated by FSH and acts by negative feedback at the pituitary or hypothalamus. Recently, activin, a protein hormone with close structural homology to transforming growth factor-beta, has also been purified and cloned and appears to exert a stimulatory effect on FSH secretion. Activin consists of a combination of 2 of the same beta subunits found in inhibin and is also derived from the testis. Activin receptors are found in a host of extragonadal tissues, suggesting that this hormone may have a variety of growth factors or regulatory roles in the body.

### SPERMATOGENESIS

Spermatogenesis is a complex process by which primitive, multipotent stem cells divide to either renew themselves or produce daughter cells that become spermatozoa. These processes occur within the seminiferous tubules of the testis. In fact, 90% of testis volume is determined by the seminiferous tubules and germ cells at various developmental stages.

#### Sertoli Cells

The seminiferous tubules are lined with Sertoli cells that rest on the tubular basement membrane and extend into its lumen with a complex cytoplasm. Sertoli cells are linked by tight junctions, the strongest intercellular barriers in the body. These junctional complexes divide the seminiferous tubule space into a basal (basement membrane) and adluminal (lumen) compartments. This arrangement forms the basis for the blood-testis barrier, allowing spermatogenesis to occur in an immunologically privileged site. The importance of this sanctuary effect becomes clear when we...
MALE INFERTILITY

remember that spermatozoa are produced at puberty and are considered foreign to an immune system that develops self-recognition during the first year of life.

Sertoli cells serve as “nurse” cells for spermatogenesis, nourishing germ cells as they develop. They also participate in germ cell phagocytosis. High-affinity FSH receptors exist on Sertoli cells and FSH binding induces the production of androgen-binding protein, which is secreted into the tubular luminal fluid. By binding testosterone, androgen-binding protein ensures that high levels of androgen (20–50 times that of serum) exist within the seminiferous tubules. Evidence also suggests that inhibin is Sertoli cell-derived. Ligand-receptor complexes, such as c-kit and kit ligand, may also mediate communication between germinal and Sertoli cells.

Germ Cells

Within the tubule, germ cells are arranged in a highly ordered sequence from the basement membrane to the lumen. Spermatogonia lie directly on the basement membrane, followed by primary spermatocytes, secondary spermatocytes, and spermatids toward the tubule lumen. In all, 13 different germ cell stages have been identified in humans. The tight junction barrier supports spermatogonia and early spermatocytes within the basal compartment; all subsequent germ cells are located within the adluminal compartment. Germ cells are staged by their morphologic appearance; there are dark type A (Ad) and pale type A (Ap) and type B spermatogonia and preleptotene, leptotene, zygotene, and pachytene primary spermatocytes, secondary spermatocytes, and Sa, Sb, Sc, Sd1, and Sd2 spermatids.

Cycles & Waves

A cycle of spermatogenesis involves the division of primitive spermatogonial stem cells into subsequent germ cells. Several cycles of spermatogenesis coexist within the germinal epithelium at any one time. The duration of an entire spermatogenic cycle within the human testis is 60 days. During spermatogenesis, cohorts of developmentally similar germ cells are linked by cytoplasmic bridges and mature together. There is also a specific organization of the steps of the spermatogenic cycle within the tubular space, termed spermatogenic waves. In humans, this is likely a spiral arrangement, which probably exists to ensure that sperm production is a continuous and not a pulsatile process.

MEIOSIS & MITOSIS

Basic Processes

Somatic cells replicate by mitosis, in which genetically identical daughter cells are formed. Germ cells replicate by meiosis, in which the genetic material is halved to allow for reproduction. These differences in cell replication generate genetic diversity through natural selection. The life of a cell is divided into cycles, each of which is associated with different activities. About 5–10% of the cell cycle is spent in the mitotic phase (M), in which DNA and cellular division occurs. Mitosis is a precise, well-orchestrated sequence of events involving duplication of the genetic material (chromosomes), breakdown of the nuclear envelope, and equal division of the chromosomes and cytoplasm into 2 daughter cells (Table 44–2). The essential difference between mitotic and meiotic replication is that a single DNA duplication step is followed by only 1-cell division in mitosis, but 2-cell divisions in meiosis (4 daughter cells). As a consequence, daughter cells contain only half of the chromosome content of the parent cell. Thus, a diploid (2n) parent cell becomes a haploid (n) gamete. Figure 44–3 illustrates how the DNA content of the dividing cell changes with mitosis and meiosis. Other major differences between mitosis and meiosis are outlined in Table 44–3.

Making Sperm

The spermatozoan is an elaborate, specialized cell produced in massive quantity, up to 300 per g of testis per second. Type B spermatogonia divide mitotically to produce diploid primary spermatocytes (2n), which then duplicate their DNA during interphase. After the first meiotic division, each daughter cell contains one partner of the homologous chromosome pair, and they are called secondary spermatocytes (2n). These cells rapidly enter the second meiotic division in which the chromatids then separate at the centromere to yield haploid early round spermatids (n). Thus, each primary spermatocyte theoretically yields 4 spermatids, although fewer actually result, as the complexity of meiosis is associated with germ cell loss.

Table 44–2. Phases of the Cell Cycle and Mitosis.

<table>
<thead>
<tr>
<th>Mitotic Phase</th>
<th>Cell Cycle</th>
<th>Description of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interphase</td>
<td>G1, S, G2</td>
<td>DNA doubling occurs.</td>
</tr>
<tr>
<td>Prophase</td>
<td>M</td>
<td>Nuclear envelope dis-</td>
</tr>
<tr>
<td></td>
<td></td>
<td>solves; spindle forms.</td>
</tr>
<tr>
<td>Metaphase</td>
<td>M</td>
<td>Chromosomes align at cell equator.</td>
</tr>
<tr>
<td>Anaphase</td>
<td>M</td>
<td>Duplicated chromosomes separate.</td>
</tr>
<tr>
<td>Telophase</td>
<td>M</td>
<td>Chromosomes to poles, cytoplasm divides.</td>
</tr>
</tbody>
</table>

DNA, deoxyribonucleic acid.
The process by which spermatids become mature spermatozoa within the Sertoli cell, termed spermiogenesis, can take several weeks and consists of several events:

1. The acrosome is formed from the Golgi apparatus.
2. A flagellum is constructed from the centriole.
3. Mitochondria reorganize around the midpiece.
4. The nucleus is compacted to about 10% of its former size.
5. Residual cell cytoplasm is eliminated.

Many cellular elements contribute to the reshaping process during spermiogenesis, including chromosome structure, associated chromosomal proteins, the perinuclear cytoskeletal theca layer, the manchette of microtubules in the nucleus, subacrosomal actin, and Sertoli cell interactions.

With completion of spermatid elongation, the Sertoli cell cytoplasm retracts around the developing sperm, stripping it of all unnecessary cytoplasm and extruding it into the tubule lumen. The mature sperm has remarkably little cytoplasm.

Sperm Maturation: The Epididymis

Spermatozoa within the testis have very poor or no motility and are incapable of naturally fertilizing an egg. They become functional only after traversing the epididymis and where further maturation occurs. Anatomically, the epididymis is divided into 3 regions: caput or head, corpus or body, and cauda or tail. Passage through the epididymis induces many changes to the newly formed sperm, including alterations in net surface charge, membrane protein composition, immunoreactivity, phospholipid and fatty acid content, and adenylate cyclase activity. These changes improve the membrane structural integrity and increase fertilization ability. The transit time of sperm through the fine tubules of the epididymis is 10–15 days in humans.

Fertilization

Fertilization normally occurs within the ampullary portion of the fallopian tubes. During the middle of the female menstrual cycle the cervical mucus changes, becoming more abundant and watery. These changes facilitate the entry of sperm into the uterus and protect the sperm from highly acidic vaginal secretions. Within the female reproductive tract, sperm undergo physiologic changes, generally referred to as capacitation.

After sperm contact with the egg, a new type of flagellar motion is observed, termed hyperactive motility, characterized by large, lashing motions of the sperm tail. Sperm release lytic enzymes from the acrosome region to help penetrate the egg investments, termed the acrosome reaction. Direct contact between the sperm and egg are mediated by specific receptors on the surface of each gamete.

After penetration of the egg, a “zona reaction” occurs in which the zona pellucida becomes impenetrable to more sperm, providing a block to polyspermy. In addition, the egg resumes its meiosis and forms a metaphase II spindle. The sperm centriole within the midpiece is crucial for early spindle formation within the fertilized egg.

DIAGNOSIS OF MALE INFERTILITY

Given that a male factor can be the cause of infertility in 30–40% of couples and is a contributing factor in 50% of cases, it is important to evaluate both partners in parallel. A
complete urologic evaluation is important because male infertility may be the presenting symptom of otherwise occult but significant systemic disease. The evaluation involves collecting 4 types of information, as outlined in Figure 44–4.

**HISTORY**

The cornerstone of the male partner evaluation is the history. It should note the duration of infertility, earlier pregnancies with present or past partners, and whether there was previous difficulty with conception. A comprehensive list of information relevant to the infertility history is given in Table 44–4.

A sexual history should be addressed. Most men (80%) do not know how to precisely time intercourse to achieve a pregnancy. Since sperm reside within the cervical mucus and crypts for 1–2 days, an appropriate frequency of intercourse is every 2 days. Lubricants can influence sperm motility and should be avoided. Commonly used products such as K-Y Jelly, Surgilube, Lubifax, most skin lotions, and saliva significantly reduce sperm motility in vitro. If needed, acceptable lubricants include vegetable, safflower, and peanut oils.

A general medical and surgical history is also important. Any generalized insult such as a fever, viremia, or other acute infection can decrease testis function and semen quality. The effects of such insults are not noted in the semen until 2 months after the event, because spermatogenesis requires at least 60 days to complete. Surgical procedures on the bladder, retroperitoneum, or pelvis can also lead to infertility, by causing either retrograde ejaculation of sperm into the bladder or anejaculation (aspermia), in which the muscular function within the entire reproductive tract is inhibited. Hernia surgery can also result in vas deferens obstruction in 1% of cases; this incidence may be rising because of the recent increased use of highly inflammatory mesh patches.

Childhood diseases may also affect fertility. A history of mumps can be significant if it occurs postpubertally. After age 11, unilateral orchitis occurs in 30% of mumps infec-

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**Table 44–4.** Components of the Infertility History.

<table>
<thead>
<tr>
<th>Medical history</th>
<th>Surgical history</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fears</td>
<td>Orchidopexy, cryptorchidism</td>
</tr>
<tr>
<td>Systemic illness—diabetes, cancer, infection</td>
<td>Herniorrhaphy</td>
</tr>
<tr>
<td>Genetic diseases—cystic fibrosis, Klinefelter syndrome</td>
<td>Trauma, torsion</td>
</tr>
<tr>
<td></td>
<td>Pelvic, bladder, or retroperitoneal surgery</td>
</tr>
<tr>
<td></td>
<td>Transurethral resection for prostatism</td>
</tr>
<tr>
<td></td>
<td>Pubertal onset</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Fertility history</th>
</tr>
</thead>
<tbody>
<tr>
<td>Previous pregnancies (present and with other partners)</td>
</tr>
<tr>
<td>Duration of infertility</td>
</tr>
<tr>
<td>Previous infertility treatments</td>
</tr>
<tr>
<td>Female evaluation</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sexual history</th>
</tr>
</thead>
<tbody>
<tr>
<td>Erections</td>
</tr>
<tr>
<td>Timing and frequency</td>
</tr>
<tr>
<td>Lubricants</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Family history</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cryptorchidism</td>
</tr>
<tr>
<td>Midline defects (Kartagener syndrome)</td>
</tr>
<tr>
<td>Hypospadias</td>
</tr>
<tr>
<td>Exposure to diethylstilbestrol</td>
</tr>
<tr>
<td>Other rare syndromes—prune belly, etc.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Medication history</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrofurantoin</td>
</tr>
<tr>
<td>Cimetidine</td>
</tr>
<tr>
<td>Sulfasalazine</td>
</tr>
<tr>
<td>Spironolactone</td>
</tr>
<tr>
<td>Alpha blockers</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Social history</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethanol</td>
</tr>
<tr>
<td>Smoking/tobacco</td>
</tr>
<tr>
<td>Cocaine</td>
</tr>
<tr>
<td>Anabolic steroids</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Occupational history</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure to ionizing radiation</td>
</tr>
<tr>
<td>Chronic heat exposure (saunas)</td>
</tr>
<tr>
<td>Aniline dyes</td>
</tr>
<tr>
<td>Pesticides</td>
</tr>
<tr>
<td>Heavy metals (lead)</td>
</tr>
</tbody>
</table>

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**Figure 44–4.** The male infertility evaluation consists of 4 kinds of information: the history, physical examination, semen analysis, and hormone assessment. Several therapeutic directions are possible once this information is collected.
tions and bilateral orchitis in 10%. Mumps orchitis is thought to cause pressure necrosis of testis tissue from viral edema. Marked testis atrophy is usually obvious later in life. Cryptorchidism is also associated with decreased sperm production. This is true for both unilateral and bilateral cases. Longitudinal studies of affected boys have shown that abnormally low sperm counts can be found in 30% of men with unilateral cryptorchidism and 50% of men with bilateral undescended testes. Differences in fertility have not been as easy to demonstrate, but it appears that boys with unilateral cryptorchidism have a slightly higher risk of infertility. However, only 50% of men with a history of bilateral undescended testes are fertile. It is important to remember that orchidopexy performed for this problem does not improve semen quality later in life.

Exposure and medication histories are very relevant to fertility. Decreased sperm counts have been demonstrated in workers exposed to specific pesticides, which may alter normal testosterone/estrogen hormonal balance. Ionizing radiation is also a well-described exposure risk, with temporary reductions in sperm production seen at doses as low as 10 cGy. Several medications (Table 44–5) and ingestants such as tobacco, cocaine, and marijuana have all been implicated as gonadotoxins. The effects of these agents are usually reversible on withdrawal. Androgenic steroids, often taken by bodybuilders to increase muscle mass and development, act as contraceptives with respect to fertility. Excess testosterone inhibits the pituitary-gonadal hormone axis. The routine use of hot tubs or saunas should be discouraged, as these activities can elevate intratesticular temperature and impair sperm production. In general, a healthy body is the best reproductive body.

The family and developmental histories may also provide clues about infertility. A family history of cystic fibrosis (CF), a condition associated with congenital absence of the vas deferens (CAVD), or intersex conditions is important. The existence of siblings with fertility problems may suggest that a Y chromosome microdeletion or a cytogenetic (karyotype) abnormality is present in the family. A history of delayed onset of puberty could suggest Kallmann or Klinefelter syndrome. A history of recurrent respiratory tract infections may suggest a ciliary defect characteristic of the immotile cilia syndromes. It is important to remember that reproductive technologies enable most men afflicted with such conditions to become fathers and therefore allow for the perpetuation of genetic abnormalities that may not be normally sustained.

**PHYSICAL EXAMINATION**

A complete examination of the infertile male is important to identify general health issues associated with infertility. For example, the patient should be adequately virilized; signs of decreased body hair or gynecomastia may suggest androgen deficiency.

The scrotal contents should be carefully palpated with the patient standing. As it is often psychologically uncomfortable for young men to be examined, one helpful hint is to make the examination as efficient and matter of fact as possible. Two features should be noted about the testis: size and consistency. Size is assessed by measuring the long axis and width; as an alternative, an orchidometer can be placed next to the testis for volume determination (Figure 44–5). Standard values of testis size have been reported for normal men and include a mean testis length of 4.6 cm (range 3.6–5.5 cm), a mean width of 2.6 cm (range 2.1–3.2 cm), and a mean volume of 18.6 mL (± 4.6 mL) (Figure 44–6). Consistency is more difficult to assess but can be described as firm (normal) or soft (abnormal). A smaller or softer than normal testis usually indicates impaired spermatogenesis.

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**Table 44–5. Medications Associated with Impaired Ejaculation.**

<table>
<thead>
<tr>
<th>Antihypertensive agents</th>
<th>Alpha-adrenergic blockers (Prazosin, Phentolamine)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thiazides</td>
<td></td>
</tr>
<tr>
<td>Antipsychotic agents</td>
<td>Mellaril (thioridazine)</td>
</tr>
<tr>
<td></td>
<td>Haldol (haloperidol)</td>
</tr>
<tr>
<td></td>
<td>Librium</td>
</tr>
<tr>
<td>Antidepressants</td>
<td>Imipramine</td>
</tr>
<tr>
<td></td>
<td>Amitriptyline</td>
</tr>
</tbody>
</table>

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**Figure 44–5.** Prader orchidometer for measuring testicular volume. (Reproduced, with permission, from McClure RD: Endocrine investigation and therapy. Urol Clin North Am 1987; 14:471.)
The peritesticular area should also be examined. Irregularities of the epididymis, located posterior-lateral to the testis, include induration, tenderness, or cysts. The presence or absence of the scrotal vas deferens is critical to observe, as 2% of infertile men may present with CAVD. Engorgement of the pampiniform plexus of veins in the scrotum is indicative of a varicocele. Asymmetry of the spermatic cords is the usual initial observation, followed by the feeling of a “bag of worms” when retrograde blood flow through the pampiniform veins occurs with a Valsalva maneuver. Varicoceles are usually found on the left side (90%) and are commonly associated with atrophy of the left testis. A discrepancy in testis size between the right and left sides should alert the clinician to this possibility.

Prostate or penile abnormalities should also be noted. Penile abnormalities such as hypospadias, abnormal curvature, or phimosis could result in inadequate delivery of semen to the upper vaginal vault during intercourse. Prostatic infection may be detected by the finding of a boggy, tender prostate on rectal examination. Prostate cancer, often suspected with unusual firmness or a nodule within the prostate, can occasionally be diagnosed in infertile men. Enlarged seminal vesicles, indicative of ejaculatory duct obstruction, may also be palpable on rectal examination.

LABORATORY

Laboratory testing is an important part of the male infertility evaluation.

Urinalysis

A urinalysis is a simple test that can be performed during the initial office visit. It may indicate the presence of infection, hematuria, glucosuria, or renal disease, and as such may suggest anatomic or medical problems within the urinary tract.

Semen Analysis

A carefully performed semen analysis is the primary source of information on sperm production and reproductive tract patency. However, it is not a measure of fertility. An abnormal semen analysis simply suggests the likelihood of decreased fertility. Studies have established that there are certain limits of adequacy below which it may be difficult to initiate a pregnancy. These semen analysis values were identified by the World Health Organization (1999) and are considered the minimum criteria for “normal” semen quality (Table 44–6). It is statistically more difficult to achieve a pregnancy if a semen parameter falls below any of those listed. Of these semen variables, the count and motility appear to correlate best with fertility.

A. SEMEN COLLECTION

Semen quality can vary widely in a normal individual from day to day, and semen analysis results are dependent on collection technique. For example, the period of sexual abstinence before sample collection is a large source of variability. With each day of abstinence (up to 1 week), semen volume can rise by up to 0.4 mL, and sperm concentration can increase by 10–15 million/mL. Sperm motility tends to fall when the abstinence period is longer than 5 days. For this reason, it is recommended that semen be collected after 48–72 hours of sexual abstinence.

To establish a baseline of semen quality, at least 2 semen samples are needed. Semen should be collected by self-stimulation, by coitus interruptus (less ideal), or with a special, nonspermicidal condom into a clean glass or plastic container. Because sperm motility decreases after ejaculation, the specimen should be analyzed within 1 hour of procurement. During transit, the specimen should be kept at body temperature.

B. PHYSICAL CHARACTERISTICS AND MEASURED VARIABLES

Fresh semen is a coagulum that liquefies 15–30 minutes after ejaculation. Ejaculate volume should be at least 1.5 mL, as smaller volumes may not sufficiently buffer against vaginal pH. Table 44–6. Semen Analysis—Minimal Standards of Adequacy.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Adequacy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ejaculate volume</td>
<td>1.5–5.5 mL</td>
</tr>
<tr>
<td>Sperm concentration</td>
<td>&gt;20 ± 10^6 sperm/mL</td>
</tr>
<tr>
<td>Motility</td>
<td>&gt;50%</td>
</tr>
<tr>
<td>Forward progression</td>
<td>2 (scale 1–4)</td>
</tr>
<tr>
<td>Morphology</td>
<td>&gt;30% WHO normal forms (&gt;4% Kruger normal forms)</td>
</tr>
<tr>
<td>No agglutination</td>
<td>(clumping), white cells, or increased viscosity.</td>
</tr>
</tbody>
</table>
nal acidity. Low ejaculate volume may indicate retrograde ejaculation, ejaculatory duct obstruction, incomplete collection, or androgen deficiency. Sperm concentration should be >20 million sperm/mL. Sperm motility is assessed in 2 ways: the fraction of sperm that are moving and the quality of sperm movement (how fast, how straight they swim).

Sperm cytology or morphology is another measure of semen quality. By assessing the exact dimensions and shape characteristics of the sperm head, midpiece, and tail, sperm can be classified as “normal” or not. In the strictest classification system (Kruger morphology), only 14% of sperm in the ejaculate are normal looking. In fact, this number correlates with the success of egg fertilization in vitro and thus is ascribed real clinical significance. In addition, sperm morphology is a sensitive indicator of overall testicular health, because these characteristics are determined during spermatogenesis. The role of sperm morphology in the male infertility evaluation is to complement other information and to better estimate the chances of fertility.

C. COMPUTER-ASSISTED SEMEN ANALYSIS

In an effort to remove the subjective variables inherent in the manually performed semen analysis, computer-aided semen analyses (CASA) couple video technology with digitalization and microchip processing to categorize sperm features by algorithms. Although the technology is promising, when manual semen analyses are compared to CASA on identical specimens, CASA can overestimate sperm counts by 30% with high levels of contaminating cells such as immature sperm or leukocytes. In addition, at high sperm concentrations, motility can be underestimated with CASA. CASA has accepted value in the research setting and in some clinical laboratories.

D. SEMINAL FRUCTOSE AND POSTEJACULATE URINALYSIS

Fructose is a carbohydrate derived from the seminal vesicles and is normally present in the ejaculate. If absent, the condition of seminal vesicle agenesis or obstruction may exist. Seminal fructose testing is indicated in men with low ejaculate volumes and no sperm. A postejaculate urinalysis is the microscopic inspection of the first voided urine after ejaculation for sperm. The presence of sperm in the urine is diagnostic of retrograde ejaculation. This test is indicated in diabetic patients with low semen volume and sperm counts; patients with a history of pelvic, bladder, or retroperitoneal surgery; and patients receiving medical therapy for prostatic enlargement. In general, the semen analyses of infertile men have patterns that may suggest a diagnosis (Table 44–7).

Hormone Assessment

An evaluation of the pituitary-gonadal axis can provide valuable information on the state of sperm production. In turn, it can reveal problems with the pituitary axis that can cause infertility (hyperprolactinemia, gonadotropin deficiency, congenital adrenal hyperplasia). FSH and testosterone should be measured in infertile men with sperm densities of <10 × 10^6 sperm/mL. Testosterone is a measure of overall endocrine balance. FSH reflects more on the state of sperm production rather than endocrine balance. This combination of tests will detect virtually all (99%) endocrine abnormalities. Serum LH and prolactin levels may be obtained if testosterone and FSH are abnormal, to help pinpoint the endocrine defect. Thyroid hormone, liver function, and other organ-specific tests should be obtained if there is clinical evidence of active disease, as uncontrolled systemic illness can affect sperm production. The common patterns of hormonal disorders observed in infertility are given in Table 44–8.

With relatively normal spermatogenesis, low levels of plasma LH and FSH have no clinical meaning; likewise, an isolated low LH with normal testosterone is not significant. The measurement of plasma estradiol should be reserved for those men who appear underandrogenized or have gynecomastia in association with low, normal, or elevated testosterone levels.

In addition to low sperm concentration (<10 million/mL), other indications for hormonal evaluation of the infertile male are evidence of impaired sexual function (impotence, low libido) and findings suggestive of a specific endocrinopathy (eg, thyroid). On initial testing, approximately 10% of infertile men with have an abnormal hormone level, with clinically significant endocrinopathies occurring in 2% of men.

**ADJUNCTIVE TESTS**

Many adjunctive tests are available to help evaluate male-factor infertility if the initial evaluation fails to lead to a diagnosis. One guiding principle in this era of cost containment is to order tests only if they will change patient management.

**Semen Leukocyte Analysis**

White blood cells (leukocytes) are present in all ejaculates and play important roles in immune surveillance and clear-

<table>
<thead>
<tr>
<th>Table 44–7. Frequency of Semen Analysis Findings in Infertile Men.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Findings</strong></td>
</tr>
<tr>
<td>All normal</td>
</tr>
<tr>
<td>Isolated abnormal</td>
</tr>
<tr>
<td>Low motility</td>
</tr>
<tr>
<td>Low count</td>
</tr>
<tr>
<td>Volume</td>
</tr>
<tr>
<td>Morphology</td>
</tr>
<tr>
<td>No sperm</td>
</tr>
</tbody>
</table>
ance of abnormal sperm. Leukocytospermia or pyospermia, an increase in leukocytes in the ejaculate, is defined as \(>1 \times 10^6\) leukocytes/mL semen and is a significant cause of male subfertility. The prevalence of pyospermia ranges from 2.8% to 23% of infertile men. In general, neutrophils predominate among inflammatory cells (Table 44–9). This condition is detected by a variety of diagnostic assays, including differential stains (eg, Papanicolaou), peroxidase stain that detects the peroxidase enzyme in neutrophils, and immunocytology.

**Antisperm Antibody Test**

The testis is a curious organ in that it is an immunologically privileged site, probably owing to the blood-testis barrier. Autoimmune infertility may result when the blood-testis barrier is broken and the body is exposed to sperm antigens. Trauma to the testis and vasectomy are 2 common ways in which this occurs, giving rise to antisperm antibodies (ASA). ASA may be associated with impaired sperm transport through the reproductive tract or impairment in egg fertilization. An assay for ASA should be obtained when

1. The semen analysis shows sperm agglutination or clumping.
2. Low sperm motility exists with history of testis injury or surgery.
3. There is confirmation that increased round cells are leukocytes.
4. There is unexplained infertility.

**Table 44–8.** Characteristic Endocrine Profiles in Infertile Men.

<table>
<thead>
<tr>
<th>Condition</th>
<th>T</th>
<th>FSH</th>
<th>LH</th>
<th>PRL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>NL</td>
<td>NL</td>
<td>NL</td>
<td>NL</td>
</tr>
<tr>
<td>Primary testis failure</td>
<td>Low</td>
<td>High</td>
<td>NL/High</td>
<td>NL</td>
</tr>
<tr>
<td>Hypogonadotropic hypogonadism</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
<td>NL</td>
</tr>
<tr>
<td>Hyperprolactinemia</td>
<td>Low</td>
<td>Low/NL</td>
<td>Low</td>
<td>High</td>
</tr>
<tr>
<td>Androgen resistance</td>
<td>High</td>
<td>High</td>
<td>High</td>
<td>NL</td>
</tr>
</tbody>
</table>

T, testosterone; FSH, follicle-stimulating hormone; LH, luteinizing hormone; PRL, prolactin; NL, normal.

ASAs can be found in 3 locations: serum, seminal plasma, and sperm-bound. Among these, sperm-bound antibodies are the most relevant. The antibody classes that appear to be clinically relevant include immunoglobulin G (IgG) and IgA. IgG antibody is derived from local production and from transudation from the bloodstream (1%), IgA is thought to be purely locally derived.

**Hypoosmotic Swelling Test**

The most clinically useful measure of sperm viability is cell motility. However, a lack of motility does not necessarily signify absent viability. Indeed, there are clinical conditions, such as immotile-cilia syndrome and extracted testicular sperm, in which there may be immotile but otherwise presumably healthy sperm. Such sperm can now be used clinically for micromanipulation and in vitro fertilization (IVF).

Cell viability can be evaluated noninvasively by using the physiologic principle of hypoosmotic swelling. Conceptually, viable cells with functional membranes should swell when placed in a hypoosmotic environment. Since sperm have tails, the swelling response is very obvious in that tail coiling accompanies head swelling. This sperm test is indicated in cases of complete absence of sperm motility.

**Sperm Penetration Assay**

It is possible to measure the ability of human sperm to penetrate a specially prepared hamster egg in the laboratory setting. The hamster egg allows interspecies fertilization but no further development. This form of bioassay can give important information about the ability of sperm to undergo the capacitation process as well as penetrate and fertilize the egg. Infertile sperm would be expected to penetrate and fertilize a lower fraction of eggs than normal sperm. The indications for the diagnostic sperm penetration assay (SPA) are limited to situations in which functional information about sperm are needed, that is, to further evaluate couples with unexplained infertility and to help couples decide whether intrauterine insemination (IUI) (good SPA result) or IVF and micromanipulation (poor SPA result) is the appropriate next treatment.

**Table 44–9.** Cells Involved in Leukocytospermia.

<table>
<thead>
<tr>
<th>Cell Type</th>
<th>Relative Abundance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neutrophils</td>
<td>++++</td>
</tr>
<tr>
<td>Monocyte/macrophage</td>
<td>+</td>
</tr>
<tr>
<td>T-helper lymphocytes</td>
<td>+</td>
</tr>
<tr>
<td>T-suppressor lymphocytes</td>
<td>++</td>
</tr>
<tr>
<td>B lymphocytes</td>
<td>+</td>
</tr>
</tbody>
</table>
Sperm Chromatin Structure

There is now evidence to suggest that the integrity of sperm DNA-chromatin packaging is important for male fertility. The structure of sperm chromatin (the DNA-associated proteins) can be measured by several methods, including the COMET and TUNNEL assays as well as by flow cytometry after acid treatment and staining of sperm with acridine orange. These tests assess the degree of DNA fragmentation that occurs after chemically stressing the sperm DNA-chromatin complex, and can indirectly reflect the quality of sperm DNA-chromatin complex, and can indirectly reflect the quality of sperm DNA integrity. Abnormally fragmented sperm DNA rarely occurs in fertile men, but can be found in 5% of infertile men with normal semen analyses and 25% of infertile men with abnormal semen analyses. This test can detect infertility that is missed on a conventional semen analysis. Often reversible, causes of DNA fragmentation include tobacco use, medical disease, hyperthermia, air pollution, infections, and varicocele.

Chromosomal Studies

Subtle genetic abnormalities can present as male infertility. It is estimated that between 2% and 15% of infertile men with azoospermia (no sperm count) or severe oligospermia (low sperm counts) will harbor a chromosomal abnormality on either the sex chromosomes or autosomes. A blood test for cytogenetic analysis (karyotype) can determine if such a genetic anomaly is present. Patients at risk for abnormal cytogenetic findings include men with small, atrophic testes, elevated FSH values, and azoospermia. Klinefelter syndrome (XXY) is the most frequently detected sex chromosomal abnormality among infertile men (Figure 44–7).

Cystic Fibrosis Mutation Testing

A blood test is indicated for infertile men who present with CF or the much more subtle condition, CAVD. Similar genetic mutations are found in both patients, although the latter group is generally considered to have an atypical form of CF, in which the scrotal vas deferens is nonpalpable. Approximately 80% of men without palpable vasa will harbor a CF gene mutation. Recent data also indicate that azoospermic men with idiopathic obstruction and men with a clinical triad of chronic sinusitis, bronchiectasis, and obstructive azoospermia (Young syndrome) may be at higher risk for CF gene mutations.

Y Chromosome Microdeletion Analysis

As many as 7% of men with oligospermia and 15% of azoospermic men have small, underlying deletions in one or more gene regions on the long arm of the Y chromosome (Yq). Several regions of the Y chromosome have been implicated in spermatogenic failure, identified as AZFa, b, and c (Figure 44–8). Deletion of the DAZ (deleted in azoospermia) gene in the AZFc region is the most commonly observed microdeletion in infertile men. Fertility is possible in most men with these deletions with IVF and micromanipulation of sperm. A polymerase chain reaction-based blood test can examine the Y chromosome from peripheral leukocytes for these gene deletions and is recommended for men with low or no sperm counts and small, atrophic testes.

Radiologic Testing

A. SCROTAL ULTRASOUND

High-frequency (7.5–10 mHz) ultrasound of the scrotum has become a mainstay in the evaluation of testicular and scrotal lesions. Scrotal ultrasound is indicated in men who have a hydrocele within the tunica vaginalis space, such that the testis is nonpalpable, to confirm that it is normal. Any abnormality of the peritesticular region should also undergo a scrotal ultrasound to determine its characteristics or origin.

Recently, scrotal color Doppler ultrasonography has been used to investigate varicoceles (Figure 44–9). By combining measurements of blood-flow patterns and vein size, both physiologic and anatomic information can be obtained to confirm the diagnosis. Although diagnostic criteria that define a varicocele vary widely, a pampiniform venous diameter of >3 mm is considered abnormal. Retrograde blood flow through the veins with a Valsalva maneuver is also an important radiologic feature of a varicocele.

B. VENOGRAPHY

Venography is accepted as the most accurate way to diagnose varicoceles. Although found by palpation in approximately 30–40% of subfertile men, varicoceles can be detected by venography in 70% of patients. Renal and spermatic venography is fairly invasive and is usually performed through percutaneous cannulation of the internal jugular vein or common femoral vein. Venographically, a varicocele is defined by a Valsalva response of a varicocele. Although found by palpation in approximately 30–40% of subfertile men, varicoceles can be detected by venography in 70% of patients. Renal and spermatic venography is fairly invasive and is usually performed through percutaneous cannulation of the internal jugular vein or common femoral vein. Venographically, a varicocele is defined by a Valsalva-induced retrograde flow, of contrast material from the renal vein into the scrotal pampiniform plexus. This test is expensive and technician dependent; at present its main indications are to guide simultaneous percutaneous varicocele embolization or to diagnose recurrent varicoceles after prior treatment.

C. TRANSRECTAL ULTRASOUND

High-frequency (5–7) mHz transrectal ultrasound (TRUS) offers superb imaging of the prostate, seminal vesicles, and ejaculatory ducts. Due to both accuracy and convenience, transrectal ultrasound has replaced surgical vasography in the diagnosis of obstructive lesions that cause infertility.
Demonstration by TRUS of dilated seminal vesicles, (>1.5 cm in width) or dilated ejaculatory ducts, (>2.3 mm) in association with a cyst, calcification, or stones along the duct is highly suggestive of obstruction (Figure 44–10). In addition, prostatic abnormalities such as tumors and congenital anomalies of the vas, seminal vesicle, or ejaculatory ducts are easily defined. The indications for TRUS in infertility include low ejaculate volume, in association with either azoospermia or severe oligospermia and decreased motility.
D. COMPUTED TOMOGRAPHY SCAN OR MAGNETIC RESONANCE IMAGING OF THE PELVIS

The imaging techniques of computed tomography (CT) and magnetic resonance imaging (MRI) can further define reproductive tract anatomy. However, since the advent of TRUS, these studies have relatively few indications. They include evaluation of a patient with a solitary right varicocele, a condition often associated with retroperitoneal pathology, and evaluation of the nonpalpable testis.

Testis Biopsy & Vasography

The testis biopsy is a useful adjunct in the infertility evaluation because it provides direct information regarding the state of spermatogenesis. Most commonly, the technique involves a small, open incision in the scrotal wall and testis tunica albuginea under local anesthesia. A small wedge of testis tissue is removed and examined histologically. Abnormalities of seminiferous tubule architecture and cellular composition are then categorized into several patterns. This procedure is most useful in the azoospermic patient, in which it is often difficult to distinguish between a failure of sperm production and obstruction within the reproductive tract ducts. A testis biopsy allows definitive delineation between these 2 conditions and can guide further treatment options in azoospermic men (Figure 44–11).

In obstructed patients defined by testis biopsy, formal investigation of the reproductive tract is warranted, beginning with a vasogram. A vasogram involves the injection of dye or contrast media into the vas deferens toward the bladder from the scrotum. In plain film radiographs, contrast material can delineate the proximal vas deferens, seminal vesicle, and ejaculatory duct anatomy and determine whether obstruction is present. Sampling of vasal fluid during the same procedure can also determine whether sperm exist within the scrotal vas deferens. Vasal sperm presence implies that there is no obstruction in the testis or epididymis. With this information, the site of obstruction can be accurately determined.
Figure 44–11. Algorithm for evaluation of azoospermia or no sperm in the ejaculate. CBAVD, congenital bilateral absence of the vas deferens; FSH, follicle stimulating-hormone; LH, luteinizing hormone; MRI, magnetic resonance imaging; CF, cystic fibrosis; ACTH, adrenocorticotropic hormone; TSH, thyroid-stimulating hormone; GH, growth hormone; FNA, fine needle aspiration. (Adapted with permission from Turek PJ. Practical approach to the diagnosis and management of male infertility. Nature Clin Pract Urol 2005;2:1.)
Whether biopsy is indicated for oligospermia is controversial. Rare cases of partial reproductive tract obstruction may exist and be diagnosed by biopsy, but the incidence of these disorders is low. While a unilateral testis biopsy is usually sufficient, the finding of 2 asymmetric testes warrants bilateral testis biopsies. This situation may reflect a unilateral unobstructed failing testis paired with a normal obstructed testis. Testis biopsies may also be indicated to identify patients at high risk for intratubular germ cell neoplasia. This premalignant condition exists in 5% of men with a contralateral germ cell tumor of the testis and is more prevalent in infertile than fertile men.

A relatively new indication for the testis biopsy is to determine whether men with atrophic, failing testes and elevated FSH levels actually have mature sperm that may be used for IVF and intracytoplasmic sperm injection (ICSI). A single testis biopsy can detect the presence of sperm in 30% of men with azoospermia, elevated FSH levels, and atrophic testes. Testicular sperm that are harvested by biopsy are now routinely used to help men with severe male-factor infertility to achieve fatherhood.

**Fine-Needle Aspiration “Mapping” of Testes (Figure 44–12)**

Although testicular sperm is used with IVF and ICSI to achieve pregnancies, there is a failure to obtain sperm in 25–50% of men with testis failure. When testis biopsies fail to retrieve sperm, IVF cycles are canceled at great emotional and financial cost. To minimize the chance of failed sperm retrieval, percutaneous fine-needle aspiration and “mapping” of the testis has been described. This technique can detect sperm in 60% of men with azoospermia due to testis failure and has confirmed that spermatogenesis can vary geographically in the failing testis.

Like a testis biopsy, fine-needle aspiration is performed under local anesthesia. Percutaneously aspirated seminiferous tubules from various locations in the testis are smeared on a slide, fixed, stained, and read by a cytologist for the presence of sperm. The information gained from this technique can fully inform patients of their chances of subsequent sperm retrieval for IVF and ICSI.

**Semen Culture**

Seminal fluid that passes through the urethra is routinely contaminated with bacteria. This can make the interpretation of semen culture difficult. Thus, semen cultures should be obtained only in selected situations, given that 83% of all infertile men will have positive semen cultures and that the relationship between bacterial cultures and infertility is at best inconclusive. Semen cultures should be obtained when there are features suggestive of infection, including (1) a history of genital tract infection, (2) abnormal expressed prostatic secretion, (3) the presence of more than 1000 pathogenic bacteria per milliliter of semen, and (4) the presence of \( >1 \times 10^6 \) leukocytes/mL of semen (pyospermia).
The agents most commonly responsible for male genital tract infections are listed in Table 44–10. Gonorrhea is the most common infection. About 10–25% of chlamydial infections may be asymptomatic. *Trichomonas vaginalis* is a protozoan parasite responsible for 1–5% of nongonococcal infections; it is usually symptomatic. *Ureaplasma urealyticum* is a common inhabitant of the urethra in sexually active men (30–50% of normal men) and is responsible for one-fourth of all cases of nongonococcal infections. *Escherichia coli* infections are relatively uncommon in young men and are usually symptomatic. Mycoplasmas are aerobic bacteria that are known to colonize the male reproductive tract. Rarer but possible causes of infection include anaerobic bacteria and tuberculosis.

### Table 44–10. Most Common Organisms in Male Genital Infection.

<table>
<thead>
<tr>
<th>Organism</th>
<th>Infection Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neisseria gonorrhoeae</td>
<td>Cytomegalovirus</td>
</tr>
<tr>
<td>Chlamydia trachomatis</td>
<td>Herpes simplex II</td>
</tr>
<tr>
<td>Trichomonas vaginalis</td>
<td>Human papilloma virus</td>
</tr>
<tr>
<td>Ureaplasma urealyticum</td>
<td>Epstein-Barr virus</td>
</tr>
<tr>
<td><em>Escherichia coli</em> (other</td>
<td>Hepatitis B virus</td>
</tr>
<tr>
<td>gram-negative bacilli)</td>
<td>Human immunodeficiency virus</td>
</tr>
<tr>
<td><em>Mycoplasma hominis</em></td>
<td></td>
</tr>
</tbody>
</table>

The agents most commonly responsible for male genital tract infections are listed in Table 44–10. Gonorrhea is the most common infection. About 10–25% of chlamydial infections may be asymptomatic. *Trichomonas vaginalis* is a protozoan parasite responsible for 1–5% of nongonococcal infections; it is usually symptomatic. *Ureaplasma urealyticum* is a common inhabitant of the urethra in sexually active men (30–50% of normal men) and is responsible for one-fourth of all cases of nongonococcal infections. *Escherichia coli* infections are relatively uncommon in young men and are usually symptomatic. Mycoplasmas are aerobic bacteria that are known to colonize the male reproductive tract. Rarer but possible causes of infection include anaerobic bacteria and tuberculosis.

### CAUSES OF MALE INFERTILITY

The causes underlying male infertility are numerous but are conveniently grouped by effects at one or more of the following levels: pretesticular, testicular, and posttesticular.

**PRETESTICULAR**

Conditions that cause infertility that act at the pretesticular level tend to be hormonal in nature (Table 44–11).

**Hypothalamic Disease**

**A. GONADOTROPIN DEFICIENCY (KALLMANN SYNDROME)**

Kallmann syndrome is a rare (1:50,000 persons) disorder that occurs in familial and sporadic forms. The X-linked form of the disease is a consequence of a single gene deletion (Xp22.3 region, termed KALIG-1). It may also be autosomal transmitted with sex limitation to males. In either case, there is a disturbance of neuronal migration from the olfactory placode during development. This neural region also contains precursors for the LH-releasing cells of the hypothalamus, which explains the 2 most common clinical deficits in the disorder: anosmia and absence of GnRH. Pituitary function is normal. The clinical features include anosmia, facial asymmetry, color blindness, renal anomalies, microphallus, and cryptorchidism. The hallmark of the syndrome is a delay in pubertal development. The differential diagnosis includes delayed puberty. Patients have severely atrophic testes (<2 cm) with biopsies showing germ cell arrest and Leydig cell hypoplasia. Hormone evaluation reveals low testosterone, low LH, and low FSH levels.

Virilization and fertility can be achieved when given FSH and LH are given to stimulate testis function.

**B. ISOLATED LH DEFICIENCY “FERTILE EUNUCH”**

This very rare condition is due to partial gonadotropin deficiency in which there is enough LH produced to stimulate intratesticular testosterone production and spermatogenesis but insufficient testosterone to promote virilization. Affected individuals have eunuchoid body proportions, variable virilization, and often gynecomastia. These men characteristically have normal testis size, but the ejaculate contains reduced numbers of sperm. Plasma FSH levels are normal, but serum LH and testosterone levels are low-normal.

**C. ISOLATED FSH DEFICIENCY**

In this rare condition, there is insufficient FSH production by the pituitary. Patients are normally virilized, as LH is present. Testicular size is normal, and LH and testosterone levels are normal. FSH levels are uniformly low and do not respond to stimulation with GnRH. Sperm counts range from azoospermia to severely low numbers (oligospermia).

**D. CONGENITAL HYPOGONADOTROPIC SYNDROMES**

Several syndromes are associated with secondary hypogonadism. Prader-Willi syndrome (1:20,000 persons) is characterized by genetic obesity, retardation, small hands and feet, and hypogonadism and is caused by a deficiency of hypothalamic GnRH. The single gene deletion associated with this condition is found on chromosome 15. Similar
to Kallmann syndrome, spermatogenesis can be induced with exogenous FSH and LH. Bardet-Biedl syndrome is another autosomal recessive form of hypogonadotropic hypogonadism that results from GnRH deficiency. It is characterized by retardation, retinitis pigmentosa, polydactyly, and hypogonadism. The presentation is similar to Kallmann syndrome except it includes genetic obesity. The hypogonadism can be treated with FSH and LH. Cerebellar ataxia can be associated with hypogonadotropic hypogonadism. This rare condition can result from consanguineous unions. Cerebellar involvement includes abnormalities of speech and gait. These patients can be eunuchoid-looking with atrophic testes. Hypothalamic-pituitary dysfunction due to pathologic changes in cerebral white matter is thought to be the reason for infertility.

**Pituitary Disease**

**A. PITUITARY INSUFFICIENCY**

Pituitary insufficiency may result from tumors, infarcts, surgery, radiation, or infiltrative and granulomatous processes. In sickle cell anemia, pituitary and testicular microinfarcts from sickling of red blood cells are suspected of causing infertility. Men with sickle cell anemia have decreased testosterone and variable LH and FSH levels. Beta Thalassemia patients have mutations in the beta-globin gene that lead to an imbalance in alpha and beta globin composition of hemoglobin; these patients are mainly of Mediterranean or African origin. Infertility is also believed to result from the deposition of iron in the pituitary gland and testes. Similarly, hemochromatosis results in iron deposition within the liver, testis, and pituitary and is associated with testicular dysfunction in 80% of cases.

**B. HYPERPROLACTINEMIA**

Another form of hypogonadotropic hypogonadism is due to elevated circulating prolactin. If hyperprolactinemia occurs, secondary causes such as stress during the blood draw, systemic diseases, and medications should be ruled out. With these causes excluded, the most common and important cause of hyperprolactinemia is a prolactin-secreting pituitary adenoma. High-resolution CT scanning or MRI of the sella turcica has classically been used to distinguish between microadenoma (<10 mm) and macroadenoma (>10 mm) forms of tumor.

Stratification of disease based on radiologic diagnosis alone is misleading, as surgery for hyperprolactinemia almost always reveals a pituitary tumor. Elevated prolactin usually results in decreased FSH, LH, and testosterone levels and causes infertility. Associated symptoms include loss of libido, impotence, galactorrhea, and gynecomastia. Signs and symptoms of other pituitary hormone derangements (adrenocorticotrophic hormone, thyroid-stimulating hormone) should also be investigated.

**C. EXOGENOUS OR ENDOGENOUS HORMONES**

1. **Estrogens**—An excess of sex steroids, either estrogens or androgens, can cause male infertility due to an imbalance in the testosterone-estrogen ratio. Hepatic cirrhosis increases endogenous estrogens because of augmented aromatase activity within the diseased liver. Likewise, excessive obesity may be associated with testosterone-estrogen imbalance owing to increased peripheral aromatase activity. Less commonly, adrenocortical tumors, Sertoli cell tumors, and interstitial testis tumors may produce estrogens. Excess estrogens mediate infertility by decreasing pituitary gonadotropin secretion and inducing secondary testis failure. Exposure to exogenous estrogens has been implicated as a reason for the controversial finding of decreased sperm concentrations in men over the last 50 years. Supporters of this claim suggest that men are overexposed to estrogenic compounds during fetal life, which results in compromised semen quality later. Postulated sources of exposure include anabolic estrogens in livestock, consumed plant estrogens, and environmental estrogenic chemicals like pesticides. This xenoestrogen exposure theory, however, remains unproved as a cause of impaired fertility.

2. **Androgens**—An excess of androgens can suppress pituitary gonadotropin secretion and lead to secondary testis failure. The use of exogenous androgenic steroids (anabolic steroids) by as many as 15% of high school athletes, 30% of college athletes, and 70% of professional athletes may result in temporary sterility due to this effect. Initial treatment is to discontinue the steroids and reevaluate semen quality every 3–6 months until spermatogenesis returns. The most common reason for excess endogenous androgens is congenital adrenal hyperplasia, in which the enzyme 21-hydroxylase is most commonly deficient. As a result, there is defective cortisol synthesis and excessive adrenocorticotrophic hormone production, leading to abnormally high production of androgenic steroids by the adrenal cortex. High androgen levels in prepubertal boys results in precocious puberty, with premature development of secondary sex characteristics and abnormal enlargement of the phallus. The testes are characteristically small because of central gonadotropin inhibition by androgens. In young girls, virilization and clitoral enlargement may be obvious. In cases of the classic 21-hydroxylase-deficient congenital adrenal hyperplasia that presents in childhood, normal sperm counts and fertility have been reported, even without glucocorticoid treatment. This disorder is one of the few intersex conditions associated with fertility. Other sources of endogenous androgens include hormonally active adrenocortical tumors or Leydig cell tumors of the testis.

3. **Glucocorticoids**—Exposure to excess glucocorticoids either endogenously or exogenously can result in decreased spermatogenesis. Elevated plasma cortisol levels depress LH secretion and induce secondary testis failure.
Source of exogenous glucocorticoids include chronic therapy for ulcerative colitis, asthma, or rheumatoid arthritis. Cushing’s syndrome is a common reason for excess endogenous glucocorticoids. Correction of the problem usually improves spermatogenesis.

4. Hyper- and hypothyroidism—Abnormally high or low levels of serum thyroid hormone affect spermatogenesis at the level of both the pituitary and testis. Thyroid balance is important for normal hypothalamic hormone secretion and for normal sex hormone-binding protein levels that govern the testosterone-estrogen ratio. Thyroid abnormalities are a rare cause (0.5%) of male infertility.

5. Growth hormone—There is emerging evidence that growth hormone may play a role in male infertility. Some infertile men have deficient responses to growth hormone challenge tests and may respond to growth hormone treatment with improvements in semen quality. Growth hormone is an anterior pituitary hormone that has receptors in the testis. It induces insulin-like growth factor-1, a growth factor important for spermatogenesis. The routine measurement of serum growth hormone is presently not indicated in the infertility evaluation.

TESTICULAR

Conditions that cause infertility that act at the testicular level are listed in Table 44–12. Unlike most pretesticular conditions, which are treatable with hormone manipulation, testicular effects are, at present, largely irreversible. If sperm are observed, however, assisted reproductive technology can provide biological children for affected men.

Chromosomal Causes

Abnormalities in chromosomal constitution are well-recognized causes of male infertility. In a study of 1263 infertile couples, a 6.2% overall incidence of chromosomal abnormalities was detected. Among men whose sperm count was <10 million/mL, the incidence was 11%. In azoospermic men, 21% had significant chromosomal abnormalities. For this reason, cytogenetic analysis (karyotype) of autosomal and sex chromosomal anomalies should be considered in men with severe oligospermia and azoospermia.

A. KLINEFELTER SYNDROME (FIGURE 44–7)

Klinefelter syndrome is the most common genetic reason for azoospermia, accounting for 14% of cases (overall incidence 1:500 males). It has a classic triad: small, firm testes; gynecomastia; and azoospermia. This syndrome may present with delayed sexual maturation, increased height, decreased intelligence, varicosities, obesity, diabetes, leukemia, increased likelihood of extragonadal germ cell tumors, and breast cancer (20-fold higher than in normal males). In this abnormality of chromosomal number, 90% of men carry an extra X chromosome (47, XXY) and 10% are mosaic, with a combination of XXY/XY chromosomes. Paternity with this syndrome is rare but more likely in the mosaic or milder form of the disease. The testes are usually <2 cm in length and always <3.5 cm; biopsies show sclerosis and hyalinization of the seminiferous tubules with normal numbers of Leydig cells. Hormones usually demonstrate decreased testosterone and frankly elevated LH and FSH levels. Serum estradiol levels are commonly elevated. Since testosterone tends to decrease with age, these men will require androgen replacement therapy both for virilization and for normal sexual function.

B. XX MALE SYNDROME

XX male syndrome is a structural and numerical chromosomal condition, a variant of Klinefelter syndrome, that presents as gynecomastia at puberty or as azoospermia in adults. Average height is below normal, and hypospadias is common. Male external and internal genitalia are otherwise normal. The incidence of mental deficiency is not increased. Hormone evaluation shows elevated FSH and LH and low or normal testosterone levels. Testis biopsy reveals absent spermatogenesis with fibrosis and Leydig cell clumping. The most obvious explanation is that sex determining ratio (SRY), or the testis-determining region, is translocated from the Y to the X chromosome. Thus, testis differentiation is present; however, the genes that control spermatogenesis on the Y chromosome are not similarly translocated, resulting in azoospermia.

C. XYY SYNDROME

The incidence of XYY syndrome is similar to that of Klinefelter, but the clinical presentation is more variable. Typically, men with 47, XYY are tall, and 2% exhibit aggressive or antisocial behavior. Hormone evaluation reveals elevated FSH and normal testosterone and LH levels. Semen analyses show either oligospermia or
Other Syndromes

A. NOONAN SYNDROME

Also called male Turner syndrome, Noonan syndrome is associated with clinical features similar to Turner syndrome (45, X). However, the karyotype is either normal (46, XY) or mosaic (X/XY). Typically, patients have dysmorphic features like webbed neck, short stature, low-set ears, wide-set eyes, and cardiovascular abnormalities. At birth, 75% have cryptorchidism that limits fertility in adulthood. If testes are fully descended, then fertility is possible and likely. Associated FSH and LH levels depend on the degree of testicular function.

B. MYOTONIC DYSTROPHY

Myotonic dystrophy is the most common reason for adult-onset muscular dystrophy. In addition to having myotonia, or delayed relaxation after muscle contraction, patients usually present with cataracts, muscle atrophy, and various endocrinopathies. Most men have testis atrophy, but fertility has been reported. Infertile men may have elevated FSH and LH with low or normal testosterone, and testis biopsies show seminiferous tubule damage in 75% of cases. Pubertal development is normal; testis damage seems to occur later in life.

C. VANISHING TESTIS SYNDROME

Also called bilateral anorchia, vanishing testis syndrome is rare, occurring in 1:20,000 males. Patients present with bilateral nonpalpable testes and sexual immaturity due to the lack of testicular androgens. The testes are lost due to fetal torsion, trauma, vascular injury, or infection. In general, functioning testis tissue must have been present during weeks 14–16 of fetal life, since Wolffian duct growth and Müllerian duct inhibition occur along with appropriate growth of male external genitalia. Patients have eunuchoid body proportions but no gynecomastia. The karyotype is normal. Serum LH and FSH levels are elevated, and serum testosterone levels are extremely low. There is no treatment for this form of infertility; patients receive lifelong testosterone for normal virilization and sexual function.

D. SERTOLI-CELL-ONLY SYNDROME

Also referred to as germ cell aplasia, the hallmarks of Sertoli-cell-only syndrome are an azoospermic male with testis biopsies that show the presence of all testis cell types except for germinal epithelium. Several causes have been proposed, including genetic defects, congenital absence of germ cells, and androgen resistance. Clinically, these men have normal virilization with small testes of normal consistency. There is no gynecomastia. Testosterone and LH levels are normal, but FSH levels are usually (90%) elevated. The use of the word “syndrome” implies that no recognized insult has occurred, since gonadotoxins like ionizing radiation, chemotherapy, and mumps orchitis can also render the testes aplastic of germ cells. There is no known treatment for this condition. In some patients, extensive testis sampling with fine-needle aspiration mapping or multiple biopsies can reveal sperm that can be used for pregnancy with assisted reproductive technologies.

E. Y CHROMOSOME MICRODELETIONS

Approximately 7% of men with low sperm counts and 13% with azoospermia have a structural alteration in the long arm of the Y chromosome (Yq). The testis-determining region genes that control testis differentiation are intact, but there may be gross deletions in other regions that may lead to defective spermatogenesis. The recent explosion in molecular genetics has allowed for sophisticated analysis of the Y chromosome. At present, 3 gene sites are being investigated as putative AZF (azoospermia factor) candidates: AZFa, b, and c. The most promising site is AZFc, which contains the DAZ gene region. The gene, of which there are at least 6 copies in this region, appears to encode a ribonucleic acid (RNA)-binding protein that regulate the meiotic pathway during germ cell production. Homologs of the DAZ gene are found in many other animals, including mouse and Drosophila. A quantitative polymerase chain reaction-based assay is used to test blood for these deletions. In the future, sperm DNA may also be tested as part of a semen analysis. Since men with these microdeletions can have sperm in the ejaculate, they are likely to pass them on to offspring if assisted reproductive technology is used.

Gonadotoxins

A. RADIATION

The effects of radiotherapy on sperm production are well described. They are derived mainly from a series of remarkable experiments performed during the “atomic age” but only recently published. In a study of healthy prisoners in Oregon and Washington in the 1960s, Clifton and Brenner (1983) examined the effects of ionizing irradiation on semen quality and spermatogenesis. Before a vasectomy, each of 111 volunteers was exposed to different levels of radiation. There was a distinct dose-dependent, inverse relationship between irradiation and sperm count. A significant reduction in sperm count was observed at 15 cGy, and sperm counts were temporarily abolished at 50 cGy. Azoospermia was induced at 400 cGy, this persisted for at least 40 weeks. Despite these profound effects, sperm counts rebounded to preirradiation levels in most patients during recovery.

From examination of testis tissue after irradiation, it is observed that spermatogonia are the germ cells most sensi-
tive to irradiation. Given the dramatic sensitivity of testis tissue to irradiation, recent studies have focused on the "scatter" to testes of men undergoing radiation therapy for cancer. In cases of abdominal radiation with gonadal shielding, the estimated mean unintended gonadal exposure is approximately 75 cGy. There does not appear to be an increase in congenital birth defects in offspring of irradiated men.

**B. DRUGS**

Medications are usually tested for their potential as reproductive hazards before marketing. Despite this, it is wise to discontinue unnecessary medications that can be safely stopped during attempts to conceive. A list of gonadotoxic medications can be found in Table 44–13. These can result in infertility by various mechanisms. Ketoconazole, spironolactone, and alcohol inhibit testosterone synthesis, whereas cimetidine is an androgen antagonist. Recreational drugs such as marijuana, heroin, and methadone are associated with lower testosterone levels. Certain pesticides, like dibromochloropropane, are likely to have estrogen-like activity.

Cancer chemotherapy is designed to kill rapidly dividing cells; an undesired outcome is the cytotoxic effect on normal tissues. Differentiating spermatogonia are the germinal cells most sensitive to cytotoxic chemotherapy. Alkylating agents such as cyclophosphamide, chlorambucil, and nitrogen mustard are the most toxic agents. The toxic effects of chemotherapeutic drugs vary according to dose and duration of treatment, type and stage of disease, age and health of the patient, and baseline testis function. Despite this toxicity, the mutagenic effects of chemotherapy agents do not appear to be significant enough to increase the chance of birth defects or genetic diseases among offspring of treated men. However, patients should wait at least 6 months after chemotherapy ends before attempting to conceive.

**Systemic Disease**

**A. RENAL FAILURE**

Uremia is associated with infertility, decreased libido, erectile dysfunction, and gynecomastia. The cause of hypogonadism is controversial and probably multifactorial. Testosterone levels are decreased, and FSH and LH levels can be elevated. Serum prolactin levels are elevated in 25% of patients. It is likely that estrogen excess plays a role in hormone axis derangement. Medications and uremic neuropathy may play a role in uremic-related impotence and changes in libido. After successful renal transplantation, the hypogonadism usually improves.

**B. LIVER CIRRHOSIS**

Hypogonadism related to liver failure may have various contributing factors. The reason for organ failure is important. Hepatitis is associated with viremia, and associated fevers can affect spermatogenesis. Excessive alcohol intake inhibits testicular testosterone synthesis, independent of its liver effects. Liver failure and cirrhosis are associated with testicular atrophy, impotence, and gynecomastia. Levels of testosterone and its metabolic clearance are decreased; estrogen levels are increased owing to augmented conversion of androgens to estrogens by aromatases. Decreased testosterone levels are not accompanied by proportionate elevations in LH and FSH levels, suggesting that a central inhibition of the HPG axis may accompany liver failure.

**C. SICKLE CELL DISEASE**

As mentioned earlier, sickle cell disease can cause pituitary dysfunction, likely due to the sludging of erythrocytes and associated microinfarcts. This same mechanism may also occur in testis tissue and contribute to primary hypogonadism. As a result, spermatogenesis is decreased, accompanied by lower serum testosterone levels.

**Defective Androgen Activity**

Peripheral resistance to androgens occurs with 2 basic defects: (1) a deficiency of androgen production through the absence of 5-alpha-reductase or (2) a deficiency in the androgen receptor. In general, these conditions are a consequence of single gene deletions. Figure 44–13 shows the algorithm of normal male development. Androgen insensitivity syndromes stem from aberrations in this pathway.

**A. 5-ALPHA-REDUCTASE DEFICIENCY**

5-Alpha-reductase deficiency results in normal development of the testes and Wolffian duct structures (internal genitalia) but ambiguous external genitalia. The ambiguity results from an inborn deficiency of the 5-alpha-reductase enzyme that converts testosterone to DHT in androgen-sensitive tissues like the prostate, seminal vesicle, and external genitalia. Thus far, 29 mutations have been described in the culprit enzyme. The diagnosis is made by measuring the ratio of testosterone metabolites in urine and confirmed by finding decreased 5-alpha-reductase in genital skin fibroblasts. Spermatogenesis has been described in descended testes; however, fertility has not been reported

<table>
<thead>
<tr>
<th>Table 44–13. Medications Associated with Infertility.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium channel blockers</td>
</tr>
<tr>
<td>Cimetidine</td>
</tr>
<tr>
<td>Sulfasalazine</td>
</tr>
<tr>
<td>Valproic acid</td>
</tr>
<tr>
<td>Spironolactone</td>
</tr>
<tr>
<td>Colchicine</td>
</tr>
</tbody>
</table>
in these patients. The lack of fertility may be due largely to functional abnormalities of the external genitalia.

B. ANDROGEN RECEPTOR DEFICIENCY

Androgen receptor deficiency is an X-linked genetic condition marked by resistance to androgens. The androgen receptor, a nuclear protein, is absent or functionally altered such that testosterone or DHT cannot bind to it and activate target cell genes. Since androgens have no effect on tissues, both internal and external genitalia are affected. Fertility effects depend on the specific receptor abnormality. Some patients are 46, XY males with complete end-organ resistance to androgens. They have female external genitalia with intra-abdominal testes. Testes show immature tubules and the risk of testis cancer is elevated: Tumors will develop in 10–30% of patients without orchiectomy. Fertility is absent. Patients with mild receptor defects may present as normal-appearing infertile men. Spermatogenesis may be present, although impaired. It is unclear exactly how common this occurs in infertile men.

Testis Injury

A. ORCHITIS

Inflammation of testis tissue is most commonly due to bacterial infection, termed epididymo-orchitis. Viral infections also occur in the testis in the form of mumps orchitis. Orchitis is observed in approximately 30% of postpubertal males who contract parotitis. Testis atrophy is a significant and frequent result of viral orchitis but is less common with bacterial infections.

B. TORSION

Ischemic injury to the testis secondary to twisting of the testis on the spermatic cord pedicle is common in prepubertal and early postpubertal boys. When diagnosed and corrected surgically within 6 hours of occurrence, the testis can usually be saved. Torsion may result in inoculation of the immune system with testis antigens that may predispose to later immunological infertility. It recognized that the “normal” contralateral mate of a torsed testis could also exhibit histologic abnormalities. It has not been clearly demonstrated whether this is related to the actual torsion or to an underlying abnormality in testes predisposed to torsion.

C. TRAUMA

Because of the peculiar immunologic status of the testis in the body (ie, it is an immunologically privileged site), trauma to the testis can invoke an abnormal immune response in addition to atrophy resulting from injury. Both may contribute to infertility. Trauma to the testis that results in fracture of the testis tunica albugineaal layer should be surgically explored and repaired to minimize exposure of testis tissue to the body.

Cryptorchidism

The undescended testis is a common urologic problem, observed in 0.8% of boys at 1 year of age. It is considered a developmental defect and places the affected testis at higher risk of developing cancer. Although the newborn undescended testis is morphologically fairly normal, deterioration in germ cell numbers is often seen by 2 years of age. The contralateral, normally descended testis is also at increased risk of harboring germ cell abnormalities. Thus, males with either unilaterally or bilaterally undescended testes are at risk for infertility later in life. Prophylactic orchidopexy is performed by 2 years of age to allow the testis to be palpated for cancer detection. It is unclear whether orchidopexy alters fertility potential in cryptorchidism.

Varicocele

A varicocele is defined as dilated and tortuous veins within the pampiniform plexus of scrotal veins. It is the most surgically correctable cause of male subfertility. The varicocele is a disease of puberty and is only rarely detected in boys <10 years of age. A left-sided varicocele is found in 15% of healthy young men. In contrast, the incidence of a left varicocele in subfertile men approaches 40%. Bilateral varicoceles are uncommon in healthy men (<10%) but are palpated in up to 20% of subfertile men. In general, varicoceles do not spontaneously regress. The cornerstone of varicocele diagnosis rests on an accurate physical examination.

Several anatomic features contribute to the predominance of left-sided varicoceles. The left internal spermatic
vein is longer than the right; in addition, it usually joins the left renal vein at right angles. The right internal spermatic vein has a more oblique insertion into the inferior vena cava. This particular anatomy in the standing man may cause higher venous pressures to be transmitted to the left scrotal veins and result in retrograde reflux of blood into the pampiniform plexus.

Varicoceles are associated with testicular atrophy and varicocele correction can reverse atrophy in adolescents. There is indisputable evidence that the varicocele affects semen quality. In fact, a classic semen analysis pattern has been attributed to varicoceles in which low sperm count and motility is found in conjunction with abnormal sperm morphology. The finding of semen abnormalities constitutes the main indication for varicocele surgery in infertile men.

Precisely how a varicocele exerts an effect on the testicle remains unclear. Several theories have been postulated; it is likely that a combination of effects results in infertility. Pituitary-gonadal hormonal dysfunction, internal spermatic vein reflux of renal or adrenal metabolites, and an increase in hydrostatic pressure associated with venous reflux are also postulated effects of a varicocele. The most intriguing theory of how varicoceles affect testis function invokes an inhibition of spermatogenesis through the reflux of warm corporeal blood around the testis, with disruption of the normal countercurrent heat exchange balance and elevation of intratesticular temperature.

**Idiopathic**

It has been estimated that at least 25–50% of male infertility has no identifiable cause. As our knowledge expands, it is likely that genetic and environmental factors will explain many of these cases. For example, based on findings from animal models, it is likely that X-chromosome gene mutations will play a significant role in human male infertility.

**POSTTESTICULAR (TABLE 44–14)**

**Reproductive Tract Obstruction**

The posttesticular portion of the reproductive tract includes the epididymis, vas deferens, seminal vesicles, and associated ejaculatory apparatus.

**A. CONGENITAL BLOCKAGES**

1. **Cystic fibrosis**—CF is the most common autosomal recessive genetic disorder in the United States and is fatal. It is associated with fluid and electrolyte abnormalities (abnormal chloride–sweat test) and presents with chronic lung obstruction and infections, pancreatic insufficiency, and infertility. Interestingly, 99% of men with CF are missing parts of the epididymis. In addition, the vas deferens, seminal vesicles, and ejaculatory ducts are usually atrophic or absent, causing obstruction. Spermatogenesis is usually normal. CAVD accounts for 1–2% of infertility cases. On physical examination, no palpable vas deferens is observed on one or both sides. As in CF, the rest of the reproductive tract ducts may also be abnormal and unreconstructable. This disease is related to CF. Even though most of these men demonstrate no symptoms of CF, up to 80% of patients will harbor a detectable CF mutation. In addition, 15% of these men will have renal malformations, most commonly unilateral agenesis.

2. **Young syndrome**—Young syndrome presents with a triad of chronic sinusitis, bronchiectasis, and obstructive azoospermia. The obstruction is in the epididymis. The pathophysiology of the condition is unclear but may involve abnormal ciliary function or abnormal mucus quality. Reconstructive surgery is associated with lower success rates than that observed with other obstructed conditions.

3. **Idiopathic epididymal obstruction**—Idiopathic epididymal obstruction is a relatively uncommon condition found in otherwise healthy men. There is recent evidence linking this condition to CF in that one-third of men so obstructed may harbor CF gene mutations.

4. **Adult polycystic kidney disease**—Adult polycystic kidney disease is an autosomal dominant disorder associated with numerous cysts of the kidney, liver, spleen, pancreas, epididymis, seminal vesicle, and testis. Disease onset usually occurs in the twenties or thirties with symptoms of
abdominal pain, hypertension, and renal failure. Infertility with this disease is usually secondary to obstructing cysts in the epididymis or seminal vesicle.

5. Blockage of the ejaculatory ducts—Blockage of the ejaculatory ducts, the delicate, paired, collagenous tubes that connect the vas deferens and seminal vesicles to the urethra, is termed ejaculatory duct obstruction. It is the cause of infertility in 5% of azoospermic men. Obstruction can be congenital and result from Müllerian duct (utricular) cysts, Wolffian duct (diverticular) cysts, or congenital atresia or is acquired from seminal vesicle calculi or postsurgical or inflammatory scar tissue. It presents as hematospermia, painful ejaculation, or infertility. The diagnosis is confirmed by finding a low-volume ejaculate and TRUS showing dilated seminal vesicles or dilated ejaculatory ducts.

B. ACQUIRED BLOCKAGES

1. Vasectomy—Vasectomy is performed on 800,000 men per year in the United States for contraception. Subsequently, 5% of these men have the vasectomy reversed, most commonly because of remarriage.

2. Groin and hernia surgery—Groin and hernia surgery can result in inguinal vas deferens obstruction in 1% of cases. There has been concern that Marlex mesh used for hernia repairs may add to perivasal inflammation and increase the likelihood of vassal obstruction.

3. Bacterial infections—Bacterial infections (E. coli in men age, >35) or Chlamydia trachomatis in young men) may involve the epididymis, with scarring and obstruction.

C. FUNCTIONAL BLOCKAGES

Besides physical obstruction, functional obstruction of the seminal vesicles may exist. Functional blockages may result from nerve injury or medications that impair the contractility of seminal vesicle or vasa musculature. A classic example of nerve injury affecting ejaculation is after retroperitoneal lymph node dissection for testis cancer. This can cause either retrograde ejaculation or complete anejaculation, depending on the degree of injury to postganglionic sympathetic fibers arising from the thoracolumbar spinal cord. These autonomic nerves overlie the inferior aorta and coalesce as the hypogastric plexus within the pelvis and control seminal emission. Multiple sclerosis and diabetes are other conditions that result in disordered ejaculation.

Evidence from animal models indicates that the seminal vesicles possess contractile properties similar to those of the urinary bladder, suggesting that seminal vesicle organ dysfunction may underlie some cases of ejaculatory duct “obstruction.” Medications implicated in this functional problem are those classically associated with ejaculatory impairment. Table 44–5 lists these medications.

Disorders of Sperm Function or Motility

A. IMMOTILE CILIA SYNDROMES

Immotile cilia syndromes are a heterogeneous group of disorders (1:20,000 males) in which sperm motility is reduced or absent. The sperm defects are due to abnormalities in the motor apparatus or axoneme of sperm and other ciliated cells. Normally, 9 pairs of microtubules are organized around a central pair within the sperm tail and are connected by dynein arms (ATPase) that regulate microtubule and therefore sperm tail motion. Various defects in the dynein arms cause deficits in ciliary and sperm activity. Kartagener syndrome is a subset of this disorder (1:40,000 males) that presents with the triad of chronic sinusitis, bronchiectasis, and situs inversus. Most immotile cilia cases are diagnosed in childhood with respiratory and sinus difficulties. Cilia present in the retina and ear may also be defective and lead to retinitis pigmentosa and deafness in Usher’s syndrome. Men with immotile cilia characteristically have nonmotile but viable sperm in normal numbers. Sperm nuclear material is thought to be unaffected. The diagnosis is made with electron microscopy of sperm.

B. MATURATION DEFECTS

After vasectomy reversal, normal sperm counts but low motility is often observed. This is thought to be due to elevated epididymal intratubular pressure and epididymal dysfunction, a consequence of time after vasectomy-induced blockage. As a result, sperm may not gain the usual maturation and motility capacities during transit through the epididymis.

C. IMMUNOLOGIC INFERTILITY

Autoimmune infertility has been implicated as a cause of infertility in 10% of infertile couples. The tests is a curious organ in that sperm are highly antigenic, yet normally coexist within the host; it is an immunologically privileged site, probably owing to the blood-testis barrier, which consists of Sertoli cell tight junctions and locally down regulated cellular immunity. Autoimmune infertility may result from an abnormal exposure to sperm antigens after, for example, vasectomy, testis torsion, or biopsy, which then incites a pathologic immune response. Antibodies may disturb sperm transport or disrupt normal sperm-egg interaction. Antibodies may cause clumping or agglutination of sperm, which inhibits passage, or may block normal sperm binding to the oocyte. Many assays are available to detect (ASAs), but assays that detect sperm-bound, and not serum, antibodies are the most accurate.

D. INFECTION

The agents most commonly responsible for male genital tract infections are listed in Table 44–10. Various products of activated leukocytes can exist in infected semen. A correlation exists between leukocytes in semen and the gener-
ation of superoxide anions, hydrogen peroxide, and hydroxyl radicals (reactive oxygen species), all of which can damage sperm membranes. Sperm are highly susceptible to the effects of oxidative stress because they possess little cytoplasm and therefore little antioxidant activity. Damage to sperm from oxidative stress has been correlated to loss of function and damaged DNA. Although genital tract infection has been linked to infertility in epidemiologic studies, the correlation between individual organisms and infertility is unclear. Uncontrolled studies suggest that pregnancy rates may improve after treatment, but controlled studies do not confirm these findings.

Disorders of Coitus

A. IMPOTENCE

Sexual dysfunction stemming from low libido or impotence is a frequent cause of infertility. The male hormonal evaluation can detect organic reasons for such problems. Most cases of situational impotence, in which the stress of attempting to conceive results in poor erections, are treated with sexual counseling and oral phosphodiesterase inhibitors.

B. HYPOSPADIAS

Anatomic problems like hypospadias can cause inappropriate placement of the seminal coagulum too distant from the cervix and result in infertility.

C. TIMING AND FREQUENCY

Simple problems of coital timing and frequency can be corrected by a review of the couple’s sexual habits. An appropriate frequency of intercourse is every 2 days, performed within the periovulatory period, the window of time surrounding ovulation when egg fertilization is possible. Charting of basal body temperature by the female partner allows for the calculation of that period for the next ovulatory cycle. Home kits that detect the LH surge in the urine before ovulation are also helpful. Couples should be counseled to avoid lubricants if at all possible. It is also wise to discontinue any unnecessary medications during attempts to conceive. Other coital toxins include heat exposure from regular saunas, hot saunas, hot tubs, or Jacuzzis and the use of cigarettes, cocaine, marijuana, and excessive alcohol.

TREATMENT OF MALE INFERTILITY

SURGICAL TREATMENTS

The role of surgery in the treatment of male infertility is well established and cost effective when compared to high-technology approaches. Surgery also attempts to reverse specific pathophysiologic effects and may allow for conception at home rather than in the laboratory.

Microsurgery in Urology

The rise of microsurgery as a surgical discipline followed 3 advances. The first was refinements in optical magnification; the second, the development of more precise microsuture and microneedles; and the third, the ability to manufacture smaller and more refined surgical instruments. In urology, microsurgical techniques were first applied to renal transplantation and vasectomy reversal. Microsurgery in urology is one of the most challenging disciplines in the field.

Varicocele

Although most men with varicoceles are fertile, the association of varicoceles with infertility is well established. Several treatment modalities, both surgical and nonsurgical, are available for varicoceles. These include incisional ligation of the veins through the retroperitoneal, inguinal, or subinguinal approaches; percutaneous embolization; and laparoscopy. The common goal of all treatments is to eliminate the retrograde reflux of venous blood through the internal spermatic veins. Treatments can be compared in terms of expected success rates (semen improvement and pregnancy), cost, and outcomes (pain pills, return to work or other activity), and their relative merits can be analyzed. A basic comparison of 3 treatment options is outlined in Table 44–15. Remember that if watchful waiting is chosen, a pregnancy rate of 16% can be expected. If IVF is chosen, a pregnancy rate of 35% can be expected. An overall complication rate of 1% is associated with the incisional approach, compared with a 4% complication rate for laparoscopy and 10–15% for radiologic occlusion. A significant problem with the radiologic approach is technical failure, meaning the inability to access and occlude the spermatic vein.

Vasovasostomy

About 35,000 men per year undergo vasectomy reversal in the United States. The most common reason is remarriage and the desire for more children. Occasionally, an unfortunate individual will have lost a child and desire another. Infection, deformities, trauma, and previous surgery are less frequent indications for vasovasostomy or epididymovasostomy. A problem with duct obstruction is suspected in men with normal hormones and normal testis size and no sperm in the ejaculate.

There are several methods for performing a vasovasostomy. None has been proved superior to any other, except that magnification with an operating microscope results in better success rates. Generally, either a single-layer anasto-
mosis or a strict, 2 layer anastomosis is performed (Figure 44–14). Although these procedures are technically different, the experience of the surgeon is the most important factor for success. Depending on these factors, 95% or more of patients may have a return of sperm after a vasovasostomy. If the vas fluid contains no sperm below the vasectomy site, a second problem may exist in the delicate tubules of the epididymis. The longer the time since vasectomy, the greater the “back-pressure” behind the blocked vas deferens. This may cause a blowout at some point in the single, 18-feet-long epididymal tubule, the weakest point in the system. A blowout results in blockage of the tubule as it heals. In this case, the vas must be connected to the epididymis above the blowout to allow sperm to travel through the reproductive tract. This is called an epididymovasostomy. After epididymovasostomy, approximately 60–65% of men will have sperm in the ejaculate. These rates, however, have improved remarkably during the last several years, with the evolution of surgical techniques and equipment.

Table 44–15. Varicocele Treatments: Comparison of Outcomes.

<table>
<thead>
<tr>
<th>Outcome Parameter</th>
<th>Incisional</th>
<th>Laparoscopic</th>
<th>Radiologic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Semen improvement</td>
<td>66%</td>
<td>50–70%</td>
<td>60%</td>
</tr>
<tr>
<td>Pregnancy rate</td>
<td>35%</td>
<td>12–32%</td>
<td>10–50%</td>
</tr>
<tr>
<td>Recurrence</td>
<td>0–15%</td>
<td>5–25%</td>
<td>0–10%</td>
</tr>
<tr>
<td>Technical failure</td>
<td>Negligible</td>
<td>Small</td>
<td>10–15%</td>
</tr>
<tr>
<td>Pain pills</td>
<td>9.4</td>
<td>11</td>
<td>Minimal</td>
</tr>
<tr>
<td>Days to work</td>
<td>5.0</td>
<td>5.3</td>
<td>1</td>
</tr>
</tbody>
</table>

Figure 44–14. Two-layer microsurgical vasovasostomy. A: Mucosal stitches of 10–0 nylon are placed in the “back wall” of the vas lumen, incorporating mucosa and a small amount of submucosal tissue. B: The “front wall” mucosal sutures are then placed. C: Finally, serosal sutures of 9–0 nylon are placed in the outside wall of the vas deferens to complete the anastomosis. (Reproduced, with permission, from McClure RD: Microsurgery of the male reproductive system. World J Urol 1986;4:105.)
The achievement of sperm in the ejaculate after vasovasostomy depends on the surgeon but pregnancy after surgery obviously involves a third party. It is rare that >67% of men who have normal sperm counts after vasectomy reversal will impregnate a woman. Therefore, it is critical to understand the reproductive health of the female partner before embarking on the procedure. Other reasons that reproductive tract microsurgery fails are (1) the quality of preblockage semen may not have been normal; (2) ASAs develop in roughly 30% of men who have had vasectomies (high antibody levels may impair fertility); (3) postsurgical scar tissue can develop at the anastomotic site, causing another blockage; (4) when the vas deferens has been blocked for a long time, the epididymis is adversely affected and sperm maturation may be compromised.

Ejaculatory Duct Obstruction

For over 20 years, transurethral resection of the ejaculatory ducts (TURED) has been used to relieve pain due to ejaculatory duct obstruction. Ejaculatory duct obstruction is suspected when the ejaculate volume is <2 mL and no sperm or fructose is present. Clinical suspicion can be confirmed by TRUS demonstration of dilated seminal vesicles or dilated ejaculatory ducts. Patients with ejaculatory duct obstruction sufficient to cause coital discomfort, recurrent hematospermia, or infertility should be considered for treatment.

Transurethral resection of the ejaculatory ducts is performed cystoscopically (Figure 44–15). A small resectoscope is inserted, and the verumontanum is resected in the midline. Since the area of resection is at the prostatic apex, near the external urethral sphincter and the rectum, careful positioning of the resectoscope is essential. Long-term relief of postcoital pain after TURED can be expected in 60% of patients.

Hematospermia has also been effectively treated with TURED, but this literature is anecdotal. There is convincing evidence from several large studies of infertility patients that 65–70% of men show significant improvement in semen quality after TURED and that a 30% pregnancy rate can be expected. The complication rate from TURED is approximately 20%. Most complications are self-limited and include hematospermia, hematuria, urinary tract infection, epididymitis, and a watery ejaculate. Rarely reported complications include retrograde ejaculation, rectal perforation, and urinary incontinence.

Electroejaculation

A complete failure of emission and ejaculation occurs most commonly from spinal cord injury (10,000 cases/year in the United States) and as a result of deep pelvic or retroperitoneal surgery that injured the pelvic sympathetic nerves. With rectal probe electroejaculation, the pelvic sympathetic nerves undergo controlled stimulation, with contraction of the vas deferens, seminal vesicle, and prostate, such that a reflex ejaculation is induced. The semen is collected from the penis and the bladder as retrograde ejaculation is often associated with electroejaculation. Semen acquired in this way generally requires assisted reproductive technology for success.

In men with anejaculation after retroperitoneal surgery or spinal trauma, successful recovery of sperm with electroejaculation is possible in the vast majority of patients. Sperm motility tends to be lower than normal when obtained in this way, an effect independent of electrical or heat effects inherent to the procedure. In men with spinal cord injuries...
above the T5 level, it is often possible to induce a reflex ejaculation with high-frequency penile vibration, termed vibratory stimulation. With the use of handheld vibrators set to a frequency of 110 cycles/s at an amplitude of 3 mm, patients may be taught to perform the procedure and attempt to conceive at home with cervical insemination.

**Sperm Aspiration**

Sperm aspiration techniques are indicated for men in whom the transport of sperm is not possible because the ductal system is absent or surgically unreconstructable. An example of this is vasal agenesis. Acquired forms of obstruction may also exist, the most common of which is failed vasectomy reversal. Aspiration procedures can involve microsurgery to collect sperm from the sperm reservoirs within the genital tract. At present, sperm are routinely aspirated from the vas deferens, epididymis, or testicle. It is important to realize that IVF is required to achieve a pregnancy with these procedures. Thus, success rates are intimately tied to a complex program of assisted reproduction for both partners (Table 44–16). In cases of sperm aspiration from the testicle and epididymis, IVF along with ICSI is required. An obvious prerequisite for these procedures is ongoing sperm production. Although evaluated indirectly by hormone levels and testis volume, the most direct way to verify sperm production is with a testis biopsy.

**A. VASAL ASPIRATION**

After a scrotal incision and with an operating microscope, a vasotomy is made, and leaking sperm are aspirated into culture medium. Once enough sperm are obtained (>10–20 million), the vasotomy is closed with microscopic sutures. Vasal aspiration provides the most mature or fertilizable sperm, as they have already passed through the epididymis, where sperm maturation is completed.

**B. EPIDIDYMAL SPERM ASPIRATION**

Epididymal sperm aspiration is performed when the vas is not present or is scarred and unusable. Sperm are directly collected from a single, isolated epididymal tubule (Figure 44–16). After sperm are obtained, the epididymal tubule is closed with microscopic suture, and the sperm are processed. Epididymal sperm are not as mature as vasal sperm; as a consequence, epididymal sperm require ICSI to fertilize the egg. Egg fertilization rates of 65% and pregnancy rates of 50% are possible with epididymal sperm, but results vary among individuals because of differences in sperm and egg quality.

**C. TESTIS SPERM RETRIEVAL**

The most recently developed aspiration technique is testicular sperm retrieval, begun in 1995. It is a breakthrough in that it demonstrates that sperm do not have to pass through the epididymis to fertilize the egg. Testicular sperm extraction is indicated for patients in whom there is an unreconstructable blockage in the epididymis, or in cases of severe testis failure, in which so few sperm are produced that they cannot reach the ejaculate. In this proce-
dure, a small piece of testis tissue is taken in a manner similar to that of a regular testis biopsy. The testis tissue is specially treated in the laboratory to separate sperm from other cells. High egg fertilization rates (60–75%) and pregnancy rates (40–50%) are possible with testis sperm.

**Orchidopexy**

An undescended testis occurs in 0.8% of male infants at 1 year of age. Although the most important reason for orchidopexy is to make testicles with a higher risk of cancer palpable, preservation of fertility is another debatable reason. Histologic studies of undescended testis show that significant decreases in spermatogonial numbers occur between birth and 2 years of age. Orchidopexy has been recommended within 2 years of age to potentially prevent this germ cell degeneration, although proof of this is lacking. Given that sperm can be retrieved from very atrophic testes and used with assisted reproduction, orchidopexy and not orchiectomy should be the primary goal in these cases.

Torsion of the testis is a urologic emergency. There are significant data from animal (but not human) studies to suggest that the unaffected, contralateral testis can become infertile after torsion of its mate. This has been termed sympathetic orchidopathia and is assumed to be immunologic in nature. It is the basis for the recommendation that the nonviable torsed testicle be removed at diagnosis. However, given the advances in assisted reproductive technologies, such recommendations should be reconsidered.

**Pituitary Ablation**

Elevated serum prolactin levels stemming from a pituitary adenoma can be treated medically and surgically. If the adenoma is radiologically visible (macroadenoma), then transsphenoidal surgical ablation of the lesion is possible. If the adenoma is not visible (microadenoma), then medical therapy with the dopamine agonist bromocriptine or a derivative is indicated.

**NONSURGICAL TREATMENTS**

**Specific Therapy**

Specific therapy seeks to reverse known pathophysiologic effects to improve fertility. For the most part, they are cost-effective treatments.

**A. PYOSPERMIA**

The presence of elevated numbers of leukocytes in semen is termed pyospermia and has been associated with (1) subclinical genital tract infection, (2) elevated reactive oxygen species, and (3) poor sperm function and infertility. The treatment of pyospermia is controversial in the absence of overt bacteriologic infection. It is important to evaluate the patient for sexually transmitted diseases, penile discharge, prostatitis, or epididymitis. An expressed prostatic secretion is examined for leukocytes, and urethral cultures are obtained for chlamydia and mycoplasma. The use of broad-spectrum antibiotics such as doxycycline and trimethoprimsulfamethoxazole has been shown to reduce seminal leukocyte concentrations, improve sperm function, and increase conception. Generally, the female partner is also treated.

In pyospermia with a documented prostatic source (>20 leukocytes per high-power field in expressed prostatic secretion), frequent ejaculation (more than every 3 days) and doxycycline may result in a more durable resolution of pyospermia than either treatment alone. There is increasing evidence that the antioxidant vitamins (A, C, and E) as well as glutathione and other antioxidants may help scavenge reactive oxygen species within semen and improve sperm motility in pyospermic men.

**B. COITAL THERAPY**

Simple counseling on issues of coital timing, frequency, and gonadotoxin avoidance can improve fertility. It is important to review the essentials of basal body temperature charting or home kits that detect the LH surge in the urine immediately, (<24 hours) before ovulation. Since sperm reside in the cervical mucus for 48 hours and are released continuously, it is not necessary that coitus and ovulation occur at the exact same time, a fact that can reduce the stress associated with infertility. Coitus every other day around ovulation is the best recommendation. Coital lubricants should be avoided if possible. If necessary, vegetable oils, olive oil, and petroleum jelly are the safest.

Retrograde ejaculation results from a failure of the bladder neck to close during ejaculation. Diagnosed by the finding of sperm within the postejaculate bladder urine, it can be treated with a trial of sympathomimetic medications. Approximately 30% of men will respond to treatment with some degree of antegrade ejaculation. Begun several days before ejaculation, imipramine (25–50 mg twice a day), or Sudafed Plus (60 mg three times a day) have all been used with success. The side effects associated with these medications usually limit the efficacy of therapy. For medication failure, sperm harvesting techniques can be used with IUI to achieve a pregnancy. Premature ejaculation occurs when men ejaculate before the partner is ready. Sexual counseling combined with tricyclic antidepressants or serotoninergic uptake inhibitors can be very effective.

**C. IMMUNOLOGIC INFERTILITY**

ASA’s are a complex problem underlying male infertility. Available treatment options include corticosteroid suppression (Table 44–17), sperm washing, IUI, IVF, and ICSI. Steroid suppression is based on the concept that an overactive immune system can be weakened to reduce antibodies on sperm. Intruterine insemination places more sperm nearer the ovulated egg to optimize the sperm-egg environment. Pregnancy rates with this technique generally fall in
the 10–15%/cycle range. Assisted reproductive technology with IVF and ICSI is very effective in this scenario. In general, if >50% of sperm are bound with antibodies, then treatment should be offered. In addition, head-directed or midpiece-directed sperm antibodies appear more relevant than tail-directed antibodies. Since the presence of ASA is associated with obstruction in the genital tract, such lesions should be sought and corrected. There is renewed interest in the causes and possible treatments of this interesting problem, as several animal models exist that mimic the condition in humans.

D. MEDICAL THERAPY

Effective hormonal therapy can be offered to patients with diseases that predispose to infertility. Hormone therapy is effective when it is used as specific and not empiric treatment. Specific replacement therapy seeks to reverse well-established, pathophysiologic states. Empiric treatments attempt to overcome pathologic conditions that are ill-defined or have no proven treatment.

1. Hyperprolactinemia—Normal levels of prolactin in men help sustain high intratesticular testosterone levels and affect the growth and secretions of the accessory sex glands. Hyperprolactinemia abolishes gonadotropin pulsatility by interfering with episodic GnRH release. Visible lesions are generally treated with transsphenoidal surgery, and nonvisible lesions are treated with bromocriptine, 5–10 mg daily, to restore normal pituitary balance.

2. Hypothyroidism—Both elevated and depressed levels of thyroid hormone alter spermatogenesis. Replacement or removal of low or excessive thyroid hormone is effective treatment for infertility. As these diseases are clinically evident, routine thyroid screening is not recommended for infertility patients.

3. Congenital adrenal hyperplasia—Most commonly, the 21-hydroxylase enzyme is deficient, and defective cortisol production results. The testes fail to mature because of gonadotropin inhibition due to excessive androgens. The diagnosis is rare and classically presents as precocious puberty; careful laboratory evaluation is essential. In both sexes, the condition and the infertility associated with it are treated with corticosteroids.

4. Testosterone excess/deficiency—Patients with Kallmann syndrome lack GnRH that stimulates normal pituitary function. Infertility associated with this condition can be very effectively treated with hCG, 1000–2000 U three times weekly, and recombinant FSH 75 IU twice weekly, to replace LH and FSH. It is also possible to give GnRH replacement in a pulsatile manner, 25–50 ng/kg every 2 hours, by a portable infusion pump. Individuals with fertile eunuch syndrome or isolated LH deficiency respond well to hCG therapy alone. One can expect to find sperm in the ejaculate beginning 9–12 months after therapy is started. Since injectable drug regimens are long, complex, and costly, it is good practice for men to cryopreserve motile sperm once achieved in the ejaculate. Anabolic steroids are a common and underdiagnosed reason for testicular failure in which excess exogenous testosterone and metabolites depress the pituitary-gonadal axis and spermatogenesis. Initially, the patient should discontinue the offending hormones to allow the return of normal homeostatic balance. Second-line therapy generally consists of “jump-starting” the testis with hCG and FSH as with Kallmann syndrome.

Empiric Medical Therapy

In at least 25% of infertile men, no identifiable cause can be attributed to the problem. Because the pathophysiology is ill-defined, this is termed idiopathic infertility. There is a second group of men in whom a cause of infertility may be identified but no specific therapy is available. Both groups of men are candidates for empiric medical therapy. This form of therapy seeks to overcome pathologic conditions that are ill-defined or have no proven treatment. As a rule, it is important to establish a timeline of therapy and decide with the patient when empiric treatment is to be discontinued and other avenues pursued.

A. CLOMIPHENE CITRATE

Clomiphene citrate is a synthetic nonsteroidal drug that acts as an antiestrogen and competitively binds to estrogen.
gen receptors in the hypothalamus and pituitary. This blocks the action of the normally low levels of estrogen on the male hormone axis and results in increased secretion of GnRH, FSH, and LH. The enhanced output of these hormones increases testosterone production and sperm production. Its use in male infertility treatment is “off-label,” as it is only FDA-approved for the treatment of female infertility. Clomiphene therapy is given for idiopathic low sperm count in the setting of low-normal LH, FSH, and testosterone levels. It is less effective as a treatment for low motility. The dose is 12.5–50 mg/day either continuously or with a 5-day rest period each month. Serum gonadotropins and testosterone should be monitored at 3 weeks and the dose adjusted to keep the testosterone level within the normal range. Higher than normal testosterone levels may result in decreased semen quality. Therapy should be discontinued if no semen quality response is observed in 6 months. Although there have been over 30 published trials on clomiphene since 1964, only a few include control arms. In general, there are as many trials showing that clomiphene is equivalent to placebo as there are showing that it improves sperm density and pregnancy rates. Decreased sperm densities have also been observed on this therapy.

B. ANTIOXIDANT THERAPY

There is evidence that up to 40% of infertile men have increased levels of reactive oxygen species in the reproductive tract. These species (OH, O₂ radicals, and hydrogen peroxide) can cause lipid peroxidation damage to sperm membranes. Treatment with scavengers of these radicals may protect sperm from oxidative damage: glutathione, 600 mg daily for 3–6 months, or vitamin E, 400–1200 U/day. These agents may be useful in a subgroup of infertile men with elevated levels of seminal reactive oxygen species. Non-FDA approved vitamin supplements abound as treatments for male infertility, but well-controlled trials demonstrating their efficacy are scarce.

C. GROWTH HORMONE

There is emerging evidence that growth hormone-induced insulin-like growth factor-1 may be important for spermatogenesis. In recent European trials of growth hormone in infertile men, individuals with maturation arrest and azoospermia developed sperm counts. The use of growth hormone or its releasing factor may become a new and effective treatment for oligospermia.

ASSISTED REPRODUCTIVE TECHNOLOGIES

If neither surgery nor medical therapy is appropriate for male infertility treatment, assisted reproductive techniques can be used to achieve a pregnancy.

Intrauterine Insemination

IUI involves the placement of a washed pellet of ejaculated sperm within the female uterus, beyond the cervical barrier. The principal indication for IUI is for a cervical factor; if the cervix is bypassed, then pregnancies may ensue. IUI is also used for low sperm quality, for immunologic infertility, and in men with mechanical problems of sperm delivery (eg, hypospadias). There should be at least 5–40 million motile sperm in the ejaculate (volume × concentration × motility) to make this procedure worthwhile. Success rates vary widely and are directly related to female reproductive potential; given this, pregnancy rates of 8–16% per cycle have been reported with IUI as a treatment for male infertility. Success rates are improved if ultrasound is used to document that follicles are enlarging and if urine testing is used to predict ovulation precisely.

In Vitro Fertilization and ICSI (Figure 44–17)

In vitro fertilization is a more complex technique than IUI and removes even more of the formidable obstacles to

Figure 44–17. The intracytoplasmic sperm injection procedure. (Top) A mature oocyte (left) is readied for injection with a sperm (arrow) in a micropipet under the microscope. (Bottom) The micropipet is placed directly into the oocyte, and the sperm is deposited into the cytoplasm.
Preimplantation Genetic Diagnosis

Preimplantation genetic diagnosis is a specialized technique that enables the laboratory to precisely define the genetic normality of embryos. In patients with heritable, possibly life-threatening diseases, it is possible that offspring conceived with IVF and ICSI may have these diseases transmitted to them. This complex technique involves the removal of single cells from the early embryo while it is grown in petri dishes before transfer to the uterus. The genetic material from these “biopsied” cells can then be examined to determine whether the embryo carries an abnormal chromosome or gene. Through preimplantation genetic diagnosis, early human embryos that result from IVF and ICSI can be individually examined as they develop for the presence or absence of suspected genetic traits. Because of the real-time nature of the technique, decisions regarding embryo transfer are made within 24 hours and help ensure that lethal diseases are not transmitted to offspring. Remarkably, the removal of a few cells from the embryo is not detrimental to the survival and normal development of most embryos.

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Causes of Male Infertility—Pretesticular


Belker AM et al: Results of 1,469 microsurgical vasectomy reversals by the vasovasostomy study group. J Urol 1991;145:505. [PMID: 1997700]


Causes of Male Infertility—Posttesticular


Causes of Male Infertility—Pretesticular

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INTRODUCTION

Is it not strange that desire should so many years outlive performance?

W. Shakespeare

The average age of individuals in the United States is projected to rise significantly over the next 25 years, the largest increase occurring in individuals >65 years old. According to census data, the proportion of Americans >65 years or older will rise 80% from 35 million today to 62 million in 2025. As a consequence, medicine will experience a dramatic increase in age-related health problems, including cancer, cerebrovascular and ischemic heart disease, and hormone deficiency. Among these, the health risks associated with age-related hormonal decline has been addressed mainly in women. However, there is now firm evidence to suggest that hormonal changes in the aging male may be associated with significant health problems. This chapter will review the alterations in testis biology that occur with age and the effects that these changes may have on semen quality, fertility, birth defects in offspring, and on the overall health of older men.

CHANGES IN TESTIS BIOLOGY WITH AGE

The Endocrine Testis

A. LEYDIG CELLS

The clinical observation that the aging male experiences a gradual decline in testosterone production has led to histologic investigation of Leydig cell populations in human testes. Located in the interstitial space between seminiferous tubules, Leydig cells are responsible for the production of 95% of adult male testosterone. In an early quantitative study of testes retrieved at autopsy after sudden death in men aged 18–87, Kaler and Neaves (1978) noted that total Leydig cell volume declines significantly with age and that this change is driven mainly by a fall in the absolute number of Leydig cells. From such studies, it is estimated that a pair of young testes (20-year old) is endowed with 700 million Leydig cells and undergoes an attrition of about 80 million cells per decade of life. Other autopsy studies have also shown that serum luteinizing hormone (LH) levels are significantly higher in older men compared to younger men, providing physiologic corroboration of the Leydig cell studies.

B. TESTOSTERONE

There is a progressive decline in androgen production associated with aging in men. This phenomenon has been variously termed male menopause, male climacteric, andropause, or more appropriately, partial androgen deficiency in the aging male (PADAM). Serum testosterone levels in men fall progressively from the third decade to the end of life, mainly due to a decline in testicular Leydig cell mass. This decline may be associated with changes in the circadian rhythm and hypothalamic-pituitary homeostatic control of LH secretion, which regulates testosterone production. Thus, mechanisms exist both at the testicular and hypothalamic-pituitary levels that may result in decreased testosterone with age.

Another feature of testosterone physiology that complicates matters is the fact that it exists in several different forms in plasma, with each form having different bioactivity (Figure 45–1). Free or unbound testosterone is fully bioavailable, but protein-bound testosterone is only partly bioavailable. Among protein bound forms, albumin-bound testosterone is more freely bioavailable than sex hormone-binding globulin (SHBG)-bound testosterone, which is considered an inactive form of testosterone (Figure 45–2). Aging is associated with an increase in SHBG that, by binding to and inactivating testosterone, further lowers the levels of bioavailable androgens, as illustrated in Figure 45–3. These changes in testosterone bioactivity are pronounced with age, such that 50% of men over age 60 have below normal levels of non-SHBG-bound testosterone. The age-related onset, rate and degree of change in testosterone production are all variable, such that no single factor can predict the course of age-related hypogonadism. However, a general rule of thumb is that mean testosterone levels decrease approximately 1% annually after the age of 50. Indeed, the concentration of bioavailable androgens decreases by as much as 50% between the ages of 25 and 75 years.

Interestingly, the age-related decline in testosterone is observed both in the general population and in individuals. Serum estradiol levels fall less dramatically than serum testosterone levels, and dihydrotes-
testosterone levels (a primary metabolite of testosterone and potent androgen) show even less decline with time. Thus, the complex physiology of testosterone balance in both young and older individuals often clouds the interpretation of age-related hypoandrogenism as will be discussed later.

The Exocrine Testis

A. SERTOLI CELLS

Anatomical studies of Sertoli cell populations reveal that the young adult male testis is endowed with about 500 million Sertoli cells. Similar to Leydig cell numbers, an age-related decline in Sertoli cell numbers is thought to occur in the human testis. Features of the relationship between Sertoli cells and germ cells with age are given in Table 45–1.

![Figure 45–1](image1.png)  
*Figure 45–1.* Relative amounts of various forms of testosterone in the blood. Albumin-bound testosterone is considered “bioavailable” and has physiologic activity but the SHBG-bound testosterone is chemically unavailable. Note: SHBG, sex hormone-binding globulin.

Semen Quality

Although an age-related decrease in semen quality might be expected from calculated changes in testis biology with age, this has not been obvious to demonstrate clinically. Cross-sectional studies have observed both decreased and

### Table 45–1. Comparison of Sertoli Cells and Germ Cells in Younger and Older Men.

<table>
<thead>
<tr>
<th>Testis Parameter</th>
<th>20–48 yrs</th>
<th>50–85 yrs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average testis weight</td>
<td>19 g</td>
<td>16 g</td>
</tr>
<tr>
<td>No. Sertoli cells/testis</td>
<td>503 million</td>
<td>312 million</td>
</tr>
<tr>
<td>No. round spermatids/g testis</td>
<td>55 million/g</td>
<td>41 million/g</td>
</tr>
<tr>
<td>No. spermatids/Sertoli cell</td>
<td>4.0</td>
<td>4.3</td>
</tr>
</tbody>
</table>

Data from Johnson et al, 1984.

Figure 45–2. Diagram outlining the various forms of testosterone in the blood. Total testosterone includes all forms of the hormone, both free and bound. The affinity of sex hormone-binding globulin (SHBG) for testosterone is much higher (thick arrow) than that of albumin. Available forms of testosterone that exert physiologic activity include free and albumin-bound fractions. Note: T, testosterone.

Figure 45–3. Changes in sex hormone-binding globulin (SHBG) with age. Although total testosterone levels may be similar in younger and older men, there is less “available” testosterone due to an increase in SHBG with age. Note: T, testosterone.

### SPERMATOGENESIS

#### Production

The changes that occur in the seminiferous tubules with age include a decrease in total volume of the testis occupied by seminiferous tubules and a decrease in actual tubule length. Calculations of mature sperm production in testis tissue homogenates also suggests that daily sperm production decreases significantly with age, as outlined in Table 45–2. The age-related decrease in sperm production in older testes appears to stem from a decrease in primary spermatocytes or a decrease in spermatogonial proliferation rather than cellular degeneration. Correspondingly, follicle-stimulating hormone (FSH) levels increase significantly with age, with mean values 3-fold higher in older than younger men.

### Table 45–2. Comparison of Sertoli Cells and Germ Cells in Younger and Older Men.

<table>
<thead>
<tr>
<th>Testis Parameter</th>
<th>Age Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average testis weight</td>
<td>19 g</td>
</tr>
<tr>
<td>No. Sertoli cells/testis</td>
<td>503 million</td>
</tr>
<tr>
<td>No. round spermatids/g testis</td>
<td>55 million/g</td>
</tr>
<tr>
<td>No. spermatids/Sertoli cell</td>
<td>4.0</td>
</tr>
</tbody>
</table>

Data from Johnson et al, 1984.
unchanged ejaculated sperm concentrations in older versus younger men. However, most studies show that sperm motility is consistently lower in older compared to younger men, with a decrease of approximately 0.7% motility per year after the third decade.

Fertility
The effect of paternal age on fertility is controversial. Studies that address this issue are confounded by the variables of the aging female partner and decreased coital frequency that occurs with age. Advancing paternal age has been implicated in increasing conception times in couples with secondary infertility and also in populations of fertile couples.

Genetics

A. SPERM CHROMOSOMAL ANOMALIES
The chromosomal status of sperm was first investigated because of the concern that advanced paternal age was associated with increased cases of trisomy, especially trisomy 21 or Down syndrome. Early sperm cytogenetic studies in fertile men stratified by age showed a 10% overall incidence of sperm chromosomal abnormalities, but no relationship between paternal age and the frequency of numerical abnormalities (aneuploidy) in sperm chromosomes. However, recent studies using more sensitive fluorescence in situ hybridization (FISH) technology have shown more subtle paternal-age effects on sperm aneuploidy. Interestingly, paternal age appears to increase the fraction of sperm with sex chromosomal aneuploidies. Even more pronounced is the highly significant positive, linear relationship demonstrated between paternal age and the frequency of structural anomalies in sperm ($r=0.63$, Figure 45–4).

An explanation for these age-related genetic changes may be that the continued cell divisions that characterize spermatogenesis place the germ cells at risk for chromosomal injury, especially given the extended exposure to clastogens that occur with age. However, it is important to realize that there is no evidence that these associations lead to an increased frequency of offspring with de novo structural chromosomal anomalies as assessed by studies of live newborns or prenatally diagnosed fetuses, with the exception of inherited reciprocal translocations.

B. SPERM GENETIC MUTATIONS
Single gene defects in sperm result from errors in the DNA replication process. To date, it has been difficult to assess the presence or absence of such defects in sperm. However, the effect of advanced paternal age on new cases of conditions associated with single gene deletions has been extensively studied. These disorders are listed in Table 45–3 and discussed in next section. One mechanism for the development of new single gene mutations with advanced paternal age implicates the characteristic and continuous process of spermatogonial cell division in spermatogenesis. By puberty, 30 cell divisions of spermatogonia have occurred, resulting in a large pool of undifferentiated cells. After puberty, 23 divisions per year occur; in a 35-year old man, these cells

Table 45–2. Age-Related Changes in Seminiferous Tubules and Sperm Production.

<table>
<thead>
<tr>
<th>Testis Parameter</th>
<th>20–48 yrs</th>
<th>50–90 yrs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paired testis weight</td>
<td>41 g</td>
<td>31 g</td>
</tr>
<tr>
<td>Seminiferous tube volume</td>
<td>24 mL/person</td>
<td>18 mL/person</td>
</tr>
<tr>
<td>Daily sperm production</td>
<td>250 million</td>
<td>121 million</td>
</tr>
</tbody>
</table>

Data from Johnson et al, 1986.

Figure 45–4. Incidence of sperm chromosomal structural anomalies by paternal age (data from Martin and Rademaker, 1987).

Table 45–3. Selected Genetic Disorders Associated with Advanced Paternal Age.

<table>
<thead>
<tr>
<th>Achondroplasias</th>
<th>Aniridia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apert syndrome</td>
<td>Bilateral retinoblastoma</td>
</tr>
<tr>
<td>Crouzon syndrome</td>
<td>Fibrodyplasia ossificans</td>
</tr>
<tr>
<td>Hemophilia A</td>
<td>Lesch-Nyhan syndrome</td>
</tr>
<tr>
<td>Marfan syndrome</td>
<td>Neurofibromatosis</td>
</tr>
<tr>
<td>Oculodentodigital syndrome</td>
<td>Polycystic kidney disease</td>
</tr>
<tr>
<td>Polyposis coli</td>
<td>Progeria</td>
</tr>
<tr>
<td>Treacher-Collins syndrome</td>
<td>Tuberous sclerosis</td>
</tr>
<tr>
<td>Waardenburg syndrome</td>
<td></td>
</tr>
</tbody>
</table>
will have undergone 540 divisions. The simple fact that the spermatogonial stem cells of older men have undergone numerous cell divisions may make it them more likely to contain errors in DNA transcription, the source of single gene defects.

C. PATERNAL AGE AND BIRTH DEFECTS AND DISEASE IN OFFSPRING

Although the incidence of chromosomal disorders in children does not appear to exhibit a paternal age-related increase, there is no question of the association between advanced paternal age and an increase likelihood of autosomal dominant diseases in offspring. A list of implicated disorders in given in Table 45–3. They are termed “sentinel phenotypes,” as they are disorders of significant frequency and low fitness, and occur sporadically due to highly penetrant mutations. Several investigators have established formal risk estimates for the contribution of advanced paternal age to autosomal dominant mutations. In men <29 years old, the risk of a mutation occurring in offspring is 0.22 per 1000 births. At paternal ages 40–44, this risk peaks at 4.5 per 1000 births and the shape of the risk curve is similar to that observed for aneuploid conceptions with female age. Interestingly, this genetic mechanism has also been postulated to explain the strong correlation of paternal age and schizophrenia among offspring in several countries around the world.

Studies of birth defects (anatomic and genetic) have also been assessed with respect to paternal age. One study estimated that fathers over the age of 40 years had a 20% greater incidence of having a baby born with a serious birth defect, raising the possibility that anatomic birth defects such as ventricular septal defects, atrial septal defects, and situs inversus may increase with paternal age. Overall, these risks form the basis for general guidelines that sperm donors be less than 50 years old.

THE DIAGNOSIS OF ANDROGEN DEFICIENCY

The Role of Testosterone

Testosterone has anabolic effects on a number of target organs (Figure 45–5). In the brain, it influences libido, male aggression, mood, and some aspects of cognition. Androgens have been shown to enhance verbal memory and visual-spatial skills and memory in older men. It has also been shown to improve fatigue and depression in hypogonadal men. Testosterone is responsible for an increase in muscle strength and growth. In the kidney and

![Figure 45–5. Anabolic effects of testosterone on normal male physiology.](image)
Bone marrow, testosterone stimulates production of erythropoietin and stem cells, and in bone, testosterone causes accelerated linear growth and closure of epiphyses. Testosterone is responsible for growth and maturation of male sexual organs, specifically penile growth and erections, spermatogenesis, and prostatic growth and function. It helps the liver to produce serum proteins and influences the male external appearance, for example, body hair growth, temporal balding, and other secondary characteristics. Thus, normal testosterone balance is critical for normal male growth, maturation, and reproductive function (Table 45–4).

The potential benefits of testosterone replacement therapy in PADAM include an increase in bone mineral density and reduction in fractures. The magnitude of these responses in men is roughly equivalent to that seen in postmenopausal women receiving estrogen replacement. This is an important consideration given that hip fractures are 2–3 times as lethal in older men than in women of equivalent age, with 40% of older male fracture patients dying within 1 year. Testosterone therapy also results in increases in lean body mass, possibly strength and can decrease fat mass and potentially improve insulin sensitivity. By stimulating erythropoietin production, testosterone replacement increases the red blood cell production (hematocrit) in aging men. Its effect on lipid profiles appears favorable and epidemiological data indicate that, in general, hypogonadal men have a greater incidence of myocardial infarction than eugonadal men. However, there is no data that demonstrates that testosterone replacement for PADAM reduces the risk of cardiovascular events. A positive effect of testosterone on cardiovascular risk is suggested by the lowering of body and visceral fat, and by the relatively favorable effects of androgens on clotting factors and coronary vasodilation. In contrast, androgens are also known to lower high-density lipoprotein (HDL)-cholesterol without altering low-density lipoproteins (LDL) or total cholesterol, which would suggest that androgen supplementation might in fact increase cardiovascular risk. Based on current evidence, the use of androgen replacement for the purpose of reducing cardiovascular risk is unjustifiable.

Sexual function also improves with testosterone replacement. Although a complex symptom to assess, most studies agree that sexual drive is improved by testosterone. Penile erections can be improved with testosterone supplements in hypogonadal men, but remember that isolated hypogonadism is a rare (6%) cause of erectile dysfunction in older men. Erectile dysfunction in older men is usually the net effect of decreased libido and impaired penile vascular smooth muscle dilation. The latter is best treated more specifically, such as with phosphodiesterase inhibitors.

### Table 45–4. Testosterone Effects in the Normal Male.

<table>
<thead>
<tr>
<th>Target Organ</th>
<th>Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male sex organs</td>
<td>Growth, development, maintenance of secondary sex characteristics, sperm production erections, and prostatic function</td>
</tr>
<tr>
<td>Behavior</td>
<td>Improved libido, mood, memory, and energy</td>
</tr>
<tr>
<td>Bone</td>
<td>Linear growth, closure of the epiphyses, increases bone mineral density</td>
</tr>
<tr>
<td>Fat tissue</td>
<td>Body and visceral fat reduction</td>
</tr>
<tr>
<td>Muscle</td>
<td>Anabolic; increases muscle mass and possibly strength</td>
</tr>
<tr>
<td>Liver</td>
<td>Stimulates production of serum proteins</td>
</tr>
<tr>
<td>Kidney</td>
<td>Stimulates erythropoietin production</td>
</tr>
<tr>
<td>Heart</td>
<td>Coronary vasodilation</td>
</tr>
<tr>
<td>Blood</td>
<td>Suppression of clotting factors (II, V, VII), low HDL-cholesterol</td>
</tr>
<tr>
<td>Bone marrow</td>
<td>Stimulates stem cell production</td>
</tr>
<tr>
<td>Hair</td>
<td>Influences body hair growth, especially facial hair</td>
</tr>
</tbody>
</table>

### Patient History and Examination

The PADAM syndrome is characterized by a variety of physical and intellectual alterations, most of which are not specific for hypogonadism. This complicates the diagnosis of androgen deficiency in many cases. Most clinicians agree that symptoms or physical findings need to accompany the laboratory demonstration of low testosterone to further consider treatment for PADAM. Symptoms associated with androgen deficiency include decreased sexual desire and erectile dysfunction, changes in mood associated with fatigue, depression and anger, and decreases in memory and spatial orientation ability. On physical examination, there may be a decrease in lean body mass with associated reduced muscle volume and strength, and increases in visceral fat (abdominal girth). Decreased or thinning facial and chest hair and skin alterations such as increases in facial wrinkling and pale-appearing skin suggestive of anemia may also be noted. On genital examination, testicular atrophy with either reduced testis volume or soft consistency may exist. Finally, decreased bone mineral density with osteopenia or osteoporosis may also suggest hypogonadism.

Not all of these clinical manifestations need to be present simultaneously to diagnose the syndrome. In fact, many of these symptoms are multifactorial and could be attributed simply to the natural and unavoidable consequence of aging. For example, frailty with aging is a major socioeconomic and health-care problem as it reduces independent living. Frailty may be due to many causes, some
of which include loss of muscle strength, bone fractures, decreased mood, and impaired cognition, symptoms typical of testosterone deficiency. To complicate matters further, some studies suggest that testosterone levels may be completely unrelated to the symptoms classically ascribed to androgen deficiency. The association of such symptoms with documentation of testosterone deficiency certainly implicates PADAM as a possible underlying etiology. By these criteria, it is estimated that only 10% of men with true hypogonadism are currently being diagnosed.

Laboratory

A. TOTAL AND FREE TESTOSTERONE

There is considerable debate as to what laboratory test should be ordered to diagnose androgen deficiency in aging men. In general, a total serum testosterone be ordered first and if <200 ng/dL, the diagnosis is confirmed; if >350 ng/dL androgen deficiency is ruled out; and if between 200 and 349 ng/dL, a bioactive testosterone, free testosterone by dialysis, or a total serum testosterone, SHBG, and albumin be measured together and used to calculate a free or bioavailable testosterone level (www.issam.ch/freetesto.htm). In other words, it is helpful to quantify all of the non-SHBG-bound testosterone. Currently, free testosterone levels assessed by other methods are not recommended or reflective of true free testosterone concentrations. If the measured or calculated active or bioavailable testosterone levels are normal in this situation, then androgen deficiency is unlikely. A working management algorithm for the laboratory diagnosis of androgen deficiency is outlined in Figure 45–6. Presently, the assessment of levels of testosterone metabolites such as dihydrotestosterone (DHT), estradiol, and dihydroandrosterone (DHEA) are not helpful in making the diagnosis of androgen deficiency.

B. ADJUNCTIVE TESTS

Although not indicated in all situations, an assessment of red blood cell mass with a hemoglobin or hematocrit may be helpful to confirm hypogonadal anemia. Although plain-films x-rays have been used in the past to assess bone mineral density, the DEXA scan is currently the most accurate method. This is an enhanced form of x-ray technology, termed dual-energy x-ray absorptiometry (DXA or DEXA) that is a quick and painless procedure for measuring bone loss. The DEXA machine sends low-dose x-rays with 2 distinct energy peaks through the body. One peak...
is absorbed mainly by soft tissue and the other by bone. Soft tissue absorption is subtracted from the total absorption, resulting in calculated levels of bone absorption that are converted into bone density equivalents. The amount of radiation used is extremely small—less than $\frac{1}{10}$ the dose of a standard chest x-ray. Measurement of the lower spine and hips are most often done. Portable devices are also available that measure the wrist, fingers, or heel. These are sometimes used for screening, including some that use ultrasound waves rather than x-rays. The test is scored in two ways: (1) the T score is the amount of bone present compared with a young adult of the same gender with normal bone mass. A score above –1 is considered normal. A score between –1 and –2.5 is classified as osteopenia, the first stage of bone loss. A score below –2.5 is defined as osteoporosis and can be used to estimate the risk of developing a fracture; (2) the Z score reflects the amount of bone present compared with others of similar age, size, and gender. Bone density assessments can also be performed periodically during testosterone replacement to assess the bone response to treatment.

**TREATMENT OF ANDROGEN DEFICIENCY**

**Formulations**

Ideal testosterone replacement therapy maintains physiologic serum concentrations of the hormone and its active metabolites without inducing significant side effects. Several types of hormone replacement are currently available, including oral, injectables, transdermal, and buccal mucosal systems as outlined in Table 45–5. The most popular formulations are the transdermal gels, injectables, and then transdermal patches, in that order. These 3 types of replacement therapies account for over 95% of prescribed testosterone medication. The formula-specific risks of each type of treatment are also given in Table 45–6. General risks of testosterone replacement are described below.

**Risks**

**Fluid and Electrolyte Disturbances.** Water retention is known to occur with androgen therapy. In aging men, it can lead to hypertension, peripheral edema, or exacerbation of congestive heart failure. Moreover, retention of sodium, chloride, potassium, calcium, and inorganic phosphates can occur. Weight and blood pressure monitoring are important for at-risk patients.

**Hematologic Reactions.** Polycythemia is a predictable consequence of testosterone therapy due to its stimulatory effect on erythropoietin. Indeed, polycythemia was the most commonly observed side effect in a meta-analysis of placebo-controlled clinical trials of testosterone replacement, occurring 4 times as often in treated patients (odds ratio 3.69). Hematocrit levels above 50 have been associated with an increased risk of stroke in several studies. Monitoring of hematocrit levels is important for patients on testosterone replacement. In addition, treatment-related suppression of clotting factors II, V, and VII, and bleeding in patients on concomitant anticoagulant therapy can occur.

**Hepatotoxicity.** Liver toxicity has been reported with oral administration of methyl testosterone and fluoxymesterone, however, it is only rarely observed with parenteral, transdermal and transbuccal formulations.

**Spermatogenesis and Infertility.** Exogenous testosterone of any type generally leads to spermatogenic arrest through negative feedback inhibition of both pituitary LH and FSH secretion. Azoospermia occurs in >90% of patients within 10 weeks. With cessation, sperm levels rebound usually within 18 months, but some patients may remain azoospermic. Patients receiving hormone replace-

<table>
<thead>
<tr>
<th>Available Preparation</th>
<th>Formula</th>
<th>Dose</th>
<th>Specific Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oral</td>
<td>Methyltestosterone&lt;br&gt;Fluoxymesterone</td>
<td>10–50 mg/day</td>
<td>Multiple daily doses. High first pass inactivation, with risk of hepatotoxicity</td>
</tr>
<tr>
<td>Parenteral</td>
<td>Testosterone cypionate&lt;br&gt;Testosterone enanthate</td>
<td>200–250 mg IM every 2–3 weeks</td>
<td>Deep intramuscular injection. Gives supraphysiologic peaks and low trough hormone levels; mood fluctuations</td>
</tr>
<tr>
<td>Transdermal patch</td>
<td>Testoderm (scrotal)&lt;br&gt;Testoderm TTS&lt;br&gt;Androderm</td>
<td>4–6 mg/day&lt;br&gt;5 mg/day</td>
<td>Scrotal patch: requires daily shaving, variable absorption Non-scrotal patch: high incidence of skin welts at placement site</td>
</tr>
<tr>
<td>Transdermal gel</td>
<td>AndroGel&lt;br&gt;Testim</td>
<td>2.5–5 mg/day&lt;br&gt;5 mg/day</td>
<td>Transference of hormone to partner or children. Odor</td>
</tr>
<tr>
<td>Transbuccal</td>
<td>Striant</td>
<td>30 mg buccal tablet</td>
<td>Twice daily dosing. Gum or mouth irritation. Taste perversion</td>
</tr>
</tbody>
</table>
Baseline (Pretreatment) Hgb, HCT, and PSA level
Digital rectal exam
Ascertain voiding symptoms
Consider DEXA scan for bone mineral density

1–2 months of treatment
Assess treatment efficacy: testosterone level and symptom relief. Consider adjusting dose for either variable

3–6 month intervals during 1st year
Assess symptomatic response to treatment, voiding symptoms, and sleep apnea.
Perform physical exam with DRE and obtain testosterone, LFTs, lipid profile, PSA, and Hgb levels

Anually after 1st year
Assess symptomatic response to treatment, voiding symptoms, and sleep apnea
Perform physical exam with DRE and obtain testosterone, LFTs, lipid profile, PSA, and Hgb levels

Altered Cholesterol Balance. Testosterone replacement generally does not affect total cholesterol or LDL, but can lower HDL levels. However, the extent to which cholesterol balance is altered varies widely among individuals and among studies. It is reasonable to follow lipid levels during treatment.

Exacerbation of Sleep Apnea. Testosterone therapy can worsen preexisting sleep apnea. Although not a cause of sleep apnea, testosterone can exacerbate the problem in patients predisposed to the condition (ie, elderly men, obese men, and patients with chronic obstructive pulmonary disease). All potential patients should be questioned for sleep-related breathing disorders before androgen replacement is considered.

Gynecomastia or Breast Tenderness. Painful breast enlargement due to elevated levels of estrogen (a metabolite of testosterone) frequently develops, and may occasionally persist, in patients on testosterone therapy. Estrogen receptor blockers can be used to treat this side effect.

Effects on the Prostate Gland. One of the most concerning risks of androgen replacement is the potential to exacerbate preexisting (detected or undetected) prostate cancer. The concern arises partly because of the known sensitivity of prostate cancer to androgen deprivation, an established treatment for this cancer for 50 years. Given this, it is important to realize that there is no obvious association between testosterone replacement and the development of prostate cancer. Several testosterone replacement studies in aging men have monitored the effect on prostate-specific antigen (PSA) levels, prostate size, volume, and prostatic symptoms. Although only a few studies noted a significant change in PSA levels or an increase in the rate of diagnosis of prostate cancer with therapy, all were of relatively short duration, which limits their ability to detect real changes. Reflecting the overall clinical concern for prostate cancer in testosterone-treated patients, a recent meta-analysis of 19 placebo-controlled clinical trials revealed that rates of prostate cancer, PSA >4 ng/mL and prostate biopsies were numerically higher in the testosterone group than in the placebo group. Importantly, however, these differences were not individually statistically significant. Additionally, there is evidence to suggest that hypogonadal men (ie, those who might need testosterone therapy) are more likely to harbor prostate cancer than eugonadal men. What is required to properly answer questions about risk with testosterone therapy are long-term studies with large numbers of patient. Until these studies are completed, there is still significant controversy regarding the relationship between testosterone replacement and prostate cancer.

A second prostate-related concern is whether testosterone replacement worsens symptoms of bladder outlet obstruction in patients with underlying benign prostatic hypertrophy. For this reason, voiding symptoms should be monitored in treated patients.

Contraindications and Precautions

For the reasons mentioned above, testosterone therapy is contraindicated in men with known or suspected carcinoma of the prostate and breast because it is unclear how the hormone will affect the growth of these tumors. Furthermore, it may also be inappropriate in men with bladder outlet obstruction related to severe benign prostatic hypertrophy and severe sleep apnea.

Just as importantly, there is insufficient evidence to justify the use of testosterone supplementation (not replacement) in men of any age with low-normal, but not truly “androgen-deficient” testosterone levels. In this scenario, the risk profile associated with treatment may outweigh the benefits. This recommendation is supported by a consensus statement made by The Institute of Medicine in 2003 after a multidisciplinary panel reviewed the published literature on the topic. They concluded that there was insufficient evidence to justify the use of testosterone therapy for widespread, generalized use for preventing age-related disease or to enhance strength or mood in otherwise healthy older men (www.nia.nih.gov/NewsAndEvents/PressReleases/FrequentlyAskedQuestionsAboutTestosteroneandtheOMReport.htm).
Monitoring Treatment

Testosterone replacement for PADAM is normally given for life. Monitoring of the patient during treatment is also a lifetime commitment on the part of the clinician. As outlined in Table 45–6, prior to starting therapy, a digital rectal examination and serum PSA determination are mandatory. Within a month or two after treatment is started, symptoms and testosterone levels should be assessed. As testosterone levels can fluctuate, particularly after intramuscular administration, clinical indicators may be a better guide for adjusting the dose of testosterone. During the first year of therapy, patients should be followed quarterly to assess clinical and biochemical response, with serial digital rectal examinations and PSA levels if they are >40 years old. After the first year, patients who remain stable may subsequently be followed annually. Annual evaluations should include testosterone, hemoglobin, liver function tests, lipid profile, and PSA measurements. Bone density and psychological evaluations should be performed depending on the initial indications for androgen supplementation.

FUTURE TREATMENTS FOR ANDROGEN DEFICIENCY

Dihydrotestosterone

The natural androgen DHT is a metabolite of testosterone, resulting from the action of 5-alpha reductase on testosterone (see Chapter 44). It is a selective androgen because, unlike testosterone, it cannot be converted to estrogens. DHT is also a very potent androgen, as it binds to androgen receptors more avidly than testosterone. Based on data from men with DHT deficiency, it is apparent that DHT has an effect on several target tissues, including external genitalia, prostate, and skin. DHT-deficient men have normal muscle mass and are not osteoporotic. In normal men, DHT supplements suppress pituitary FSH and LH secretion, likely causing infertility. In hypogonadal men, DHT has strong androgenic effects as predicted, but, paradoxically, is relatively prostate sparing. This may be due to the fact that estrogens may work synergistically with androgens to cause prostatic growth with age and DHT, as a selective androgen, may result in lower estrogen levels during treatment. Because of its potency and potential, significant research is being conducted with DHT supplements for androgen replacement.

Dehydroepiandrosterone

DHEA is available in over-the-counter formulations in the United States. DHEA is a steroid hormone made by the adrenal gland and exhibits a progressive decline in serum levels beginning the third decade of life and beyond. As a consequence of this decline, studies have attempted to correlate levels of DHEA and DHEA sulfate with a myriad of health conditions. Clinical trials that have investigated the DHEA for multiple conditions have shown largely inconsistent findings. Placebo-controlled studies suggest that doses of 30–50 mg of oral DHEA may produce physiologic androgen levels. Clinical trials suggest that 50 mg of oral DHEA, but not <30 mg, can increase serum androgen levels to within the physiologic range for young adults with primary and secondary adrenal insufficiency, and improve sexual function, mood and self-esteem, and decrease fatigue/exhaustion. Whereas DHEA replacement therapy may be effective in patients with adrenal insufficiency, its efficacy in aging males is not well established.

Growth Hormone

It is well established that there are decreases in growth hormone and insulin-like growth factor-I with age in both men and women. In addition, treatment of young GH-deficient adults with growth hormone improves body composition, muscle strength, physical function, and bone density, and reduces blood cholesterol and cardiovascular disease risk. Some of these improvements occur in health domains that are similar to those affected by androgen deficiency. However, growth hormone treatment is often accompanied by carpal tunnel syndrome, peripheral edema, joint pain and swelling, gynecomastia, glucose intolerance, and possibly increased cancer risk. In aged individuals, growth hormone treatment augments lean body mass and reduces body fat. However, clinically significant functional benefits, prolongation of youth, and life extension have not been demonstrated. Until more research better defines these risk/benefit relationships, treatment of elderly individuals with growth hormone is recommended only as part of controlled research studies.

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Changes in Testis Physiology with Age


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**Androgen Deficiency**


Appendix: Normal Laboratory Values*

Marcus A. Krupp, MD, FACP

Values may vary with method of measurement and population.

HEMATOLOGY

Antithrombin III: [P] 86–120%

Bleeding time: Template method, 3–9 minutes (180–540 seconds)

Cellular measurements of red blood cells (erythrocytes):

Average diameter = 7.3 µm (5.5–8.8 µm)
Mean corpuscular volume (MCV): Men, 80–94 fl; women, 81–99 fl (by Coulter counter)
Mean corpuscular hemoglobin (MCH): 27–32 pg
Mean corpuscular hemoglobin concentration (MCHC): 32–36 g/dL; erythrocytes (32–36%)
Color, saturation, and volume indices: 1 (0.9–1.1)

Clot retraction: Begins in 1–3 hours; complete in 6–24 hours. No clot lysis in 24 hours.

Fibrin D-dimer: [P] 0–250 ng/mL

Fibrinogen split products: <10 µg

Fragility of erythrocytes: Begins at 0.45–0.38% NaCl; complete at 0.36–0.3% NaCl

Glucose-6-phosphate dehydrogenase (G6PD): [B] 4–8 µg/g Hb

Hematocrit (PCV): Men, 40–52%; women, 37–47%

Hemoglobin: [B] Men, 14–18 g/dL (2.09–2.79 mmol/L as Hb tetramer); women, 12–16 g/dL (1.86–2.48 mmol/L). [S] 2–3 mg/dL

Partial thromboplastin time: Activated, 25–37 seconds

Platelets: 150,000–400,000/mL (0.15–0.4 × 10¹²/L)

Prothrombin: International normalized ratio (INR), 1–1.4

Erythrocyte count: Men, 4.5–6.2 million/µL (4.5–6.2 × 10¹²/L); women, 4.5–5.5 million/µL (4–5.5 × 10¹²/L)

Reticulocytes: 0.6–1.8% of erythrocytes

Sedimentation rate: <20 mm/h (Westergren)

White blood count (leukocytes) and differential:

5,000–10,000/µL (5–10 × 10⁹/L)

Segmented neutrophils 40–70%
Myelocytes 0%
Juvenile neutrophils 0%
Band neutrophils 0–15%
Lymphocytes 15–45%
Eosinophils 1–3%
Basophils 0–5%
Monocytes 0–7%

Lymphocytes: Total, 1500–4000/µL

B cell 5–25%

T cell 60–88%

Suppressor 10–43%

Helper 32–66%

H:S >1

BLOOD, PLASMA, OR SERUM CHEMICAL CONSTITUENTS (Values vary with method used)

Acetone and acetoacetate: [S] 0.3–2 mg/dL (3–20 mg/L)

Aldolase: [S] Values vary with method used.

α-Amino acid nitrogen: [S, fasting] 3–5.5 mg/dL (2.2–3.9 mmol/L)

Aminotransferases:

Aspartate aminotransferase (AST; SGOT): [S] 15–55 IU/L

Alanine aminotransferase (ALT; SGPT): [S] 10–70 IU/L. Values vary with method used.

Ammonia: [B] 9–33 µmol/L

Amylase: [S] 80–180 units/dL (Somogyi). Values vary with method used.

α₁-Antitrypsin: [S] >180 mg/dL

Ascorbic acid: [P] 0.4–1.5 mg/dL (23–85 33 µmol/L)

Base, total serum: [S] 145–160 mEq/L (145–160 mmol/L)

Bicarbonate: [S] 24–28 mEq/L (24–28 mmol/L)

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*Blood [B], Plasma [P], Serum [S], Urine [U].
Bilirubin: [S] Total, 0.2–1.2 mg/dL (3.5–20.5 µmol/L). Direct conjugated, 0.1–0.4 mg/dL (<7 µmol/L). Indirect, 0.2–0.7 mg/dL (<12 µmol/L).

Calcium: [S] 8.5–10.3 mg/dL (2.1–2.6 mmol/L). Values vary with albumin concentration.

Calcium, ionized: [S] 4.25–5.25 mg/dL; 2.1–2.6 mEq/L (1.05–1.3 mmol/L)

β-Carotene: [S, fasting] 50–300 µg/dL (0.9–5.58 µmol/L)

Ceruloplasmin: [S] 25–43 mg/dL (1.7–2.9 µmol/L)

Chloride: [S or P] 96–106 mEq/L (96–106 mmol/L)

Cholesterol: [S or P] 150–240 mg/dL (3.9–6.2 mmol/L). (See Lipid fractions.) Values vary with age.

Cholesteryl esters: [S] 65–75% of total cholesterol

CO2 content: [S or P] 24–29 mEq/L (24–29 mmol/L)
Transferrin: [S] 200–400 mg/dL (23–45 µmol/L)
Triglycerides: [S] <165 mg/dL (1.9 mmol/L). (See Lipid fractions.)
Troponin: [S] <0.5 ng/mL
Urea nitrogen: [S or P] 8–25 mg/dL (2.9–8.9 mmol/L). Do not use anticoagulant-containing ammonium oxalate.
Uric acid: [S or P] Men, 3–9 mg/dL (0.18–0.54 mmol/L); women, 2.5–7.5 mg/dL (0.15–0.45 mmol/L)
Vitamin A: [S] 15–60 µg/dL (0.53–2.1 µmol/L)
Vitamin B12: [S] >200 pg/mL (>148 pmol/L)
Vitamin D: [S] Cholecalciferol (D₃): 25-hydroxycholecalciferol, 8–55 ng/mL (19.4–137 nmol/L); 1,25-dihydroxycholecalciferol, 26–65 pg/mL (62–155 pmol/L); 24,25-dihydroxycholecalciferol, 1–5 ng/mL (2.4–12 nmol/L)
Volume, blood (Evans blue dye method): Adults, 2990–6980 mL. Women, 46.3–85.5 mL/kg; men, 66.2–97.7 mL/kg.
Zinc: [S] 50–150 µg/dL (7.65–22.95 µmol/L)

HORMONES, SERUM, OR PLASMA

Adrenal:
Aldosterone: [P] Supine, normal salt intake, 2–9 ng/dL (56–250 pmol/L); increased when upright.
Cortisol: [S] 8:00 AM, <5–20 µg/dL (0.14–0.55 µmol/L); 8:00 PM, <10 µg/dL (0.28 µmol/L)
Deoxycortisol: [S] After metyrapone, >7 µg/dL (>0.2 µmol/L)
Dopamine: [P] <135 pg/mL
Epinephrine: [P] <0.1 pg/mL (<0.55 nmol/L)
Norepinephrine: [P] <0.5 µg/L (<3 nmol/L). See also Miscellaneous Normal Values.
Gonad:
Testosterone, free: [S] Men, 10–30 ng/dL; women, 0.3–2 ng/dL. (1 ng/dL = 0.035 nmol/L)
Testosterone, total: [S] Prepubertal, <100 ng/dL; adult men, 300–1000 ng/dL; adult women, 20–80 ng/dL; luteal phase, up to 120 ng/dL
Estradiol (E₂): [S, special handling] Men, 12–34 pg/mL; women, menstrual cycle 1–10 days, 24–68 pg/mL; 11–20 days, 50–300 pg/mL; 21–30 days, 73–149 pg/mL (by radioimmunoassay [RIA]). (1 pg/mL = 3.6 pmol/L)
Progesterone: [S] Follicular phase, 0.2–1.5 ng/mL; luteal phase, 6–32 ng/mL; pregnancy, >24 ng/mL; men, <1 ng/mL (by RIA). (1 ng/mL = 3.2 nmol/L)

Islets:
Insulin: [S] 4–25 µU/mL (29–181 pmol/L)
C-peptide: [S] 0.9–4.2 ng/mL
Glucagon: [S, fasting] 20–100 pg/mL
Kidney:
Renin activity: [P, special handling] Normal sodium intake: Supine, 1–3 ng/mL/h; standing, 3–6 ng/mL/h. Sodium depleted: Supine, 2–6 ng/mL/h; standing, 3–20 ng/mL/h
Parathyroid: Parathyroid hormone levels vary with method and antibody. Correlate with serum calcium.
Pituitary:
Growth hormone (GH): [S] Adults, 1–10 ng/mL (46–465 pmol/L) (by RIA)
Thyroid-stimulating hormone (TSH): [S] <10 µU/mL
Follicle-stimulating hormone (FSH): [S] Prepubertal, 2–12 mIU/mL; adult men, 1–15 mIU/mL; adult women, 1–30 mIU/mL; castrate or postmenopausal, 30–200 mIU/mL. (by RIA)
Luteinizing hormone (LH): [S] Prepubertal, 2–12 mIU/mL; adult men, 1–15 mIU/mL; adult women, <30 mIU/mL; castrate or postmenopausal, >30 mIU/mL
Corticotropin (ACTH): [P] 8:00–10:00 AM, up to 100 pg/mL (22 pmol/L)
Prolactin: [S] 1–25 ng/mL (0.4–10 nmol/L)
Somatomedin C: [P] 0.4–2 U/mL
Antidiuretic hormone (ADH; vasopressin): [P] Serum osmolality 285 mOsm/kg, 0–2 pg/mL; >290 mOsm/kg, 2–12+ pg/mL
Placenta:
Estriol (E₃): [S] Men and nonpregnant women, <0.2 µg/dL (<7 nmol/L) (by RIA)
Chorionic gonadotropin: [S] β-Subunit: Men, <9 mIU/mL; pregnant women after implantation, >10 mIU/mL
Stomach:
Gastrin: [S, special handling] Up to 100 pg/mL (47 pmol/L). Elevated, >200 pg/mL
Pepsinogen I: [S] 25–100 ng/mL
Thyroid:
Thyroxine, free (FT₄): [S] 0.8–2.4 ng/dL (10–30 pmol/L)
Thyroxine, total (TT₄): [S] 5–12 µg/dL (65–156 nmol/L) (by RIA)
Thyroxine-binding globulin capacity (T₄): [S] 12–28 µg/dL (150–360 nmol/L)
Triiodothyronine (T₃): [S] 80–220 ng/dL (1.2–3.3 nmol/L). Reverse triiodothyronine (rT₃): [S] 30–80 ng/dL (0.45–1.2 nmol/L).
Triiodothyronine uptake (rT3U): [S] 25–36%; as TBG assessment (rT3U ratio), 0.85–1.15.
Calcitonin: [S] <100 pg/mL (<29.2 pmol/L)

NORMAL CEREBROSPINAL FLUID VALUES

**Appearance:** Clear and colorless.
**Cells:** Adults, 0–5 mononuclear cells/µL; infants, 0–20 mononuclear cells/µL.
**Glucose:** 50–85 mg/dL (2.8–4.7 mmol/L). (Draw serum glucose at same time.)
**Pressure (reclining):** Newborns, 30–88 mm water; children, 50–100 mm water; adults, 70–200 mm water (avg = 125)
**Proteins:** Total, 20–45 mg/dL (200–450 mg/L) in lumbar cerebrospinal fluid. IgG, 2–4 mg/dL (0.02–0.04 g/L)
**Specific gravity:** 1.003–1.008

RENAL FUNCTION TESTS

ρ-Aminohippurate (PAH) clearance (RPF): Men, 560–830 mL/min; women, 490–700 mL/min
**Creatinine clearance:** Calculation from serum creatinine:

\[
\text{Men: } \left( \frac{140 - \text{age}}{72} \right) \times (\text{wt in kg}) = \text{creatinine clearance} \text{ mL/min}
\]

Women: calculated mg/dL × 0.85

**Creatinine clearance, endogenous (GFR):** Approximates insulin clearance (see below).
**Insulin clearance (GFR):** Men, 110–150 mL/min; women, 105–132 mL/min (corrected to 1.73 m² surface area)
**Maximal glucose reabsorptive capacity (TmG):** Men, 300–450 mg/min; women, 250–350 mg/min
**Maximal PAH excretory capacity (TmPAH):** 80–90 mg/min

**Osmolality:** On normal diet and fluid intake: Range 500–850 mOsm/kg water. Achievable range, normal kidney: Dilution 40–80 mOsm; concentration (dehydration) up to 1400 mOsm/kg water (at least three to four times plasma osmolality).

**Specific gravity of urine:** 1.003–1.030

MISCELLANEOUS NORMAL VALUES

**Adrenal hormones and metabolites:**
**Aldosterone:** [U] 2–26 µg/24 h (5.5–72 nmol). Values vary with sodium and potassium intake.
**Catecholamines:** [U] Total, <100 µg/24 h. Epinephrine, <10 µg/24 h (<55 nmol); norepinephrine, <100 µg/24 h (<591 nmol). Values vary with method used.
**Cortisol, free:** [U] 20–100 µg/24 h (0.55–2.76 mmol)

11,17-Hydroxycorticoids: [U] Men, 4–12 mg/24 h; women, 4–8 mg/24 h. Values vary with method used.

17-Ketosteroids: [U] <8 years, 0–2 mg/24 h; adolescents, 2–20 mg/24 h. Men, 10–20 mg/24 h; women, 5–15 mg/24 h. Values vary with method used. (1 mg = 3.5 mmol)

Metanephrine: [U] <1.3 mg/24 h (<6.6 µmol) or <2.2 µg/mg creatinine. Values vary with method used.

Vanillylmandelic acid (VMA): [U] Up to 7 mg/24 h (<35 µmol).

Fecal fat: <30% dry weight

**Fecal Lead:** [U] <80 µg/24 h (<0.4 µmol/d)

**Porphyrains:**
δ-Aminolevulinic acid: [U] 1.5–7.5 mg/24 h (11.4–57.2 µmol)
Coproprphyrin: [U] <230 µg/24 h (<345 nmol)
Uroporphyrin: [U] <50 µg/24 h (<60 nmol)
Porphobilinogen: [U] <2 mg/24 h (<8.8 µmol)
Urobilinogen: [U] 0–2.5 mg/24 h (<4.23 µmol)
Urobilinogen, fecal: 40–280 mg/24 h (68–474 µmol)
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