To all our students
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Preface

In the several years since publication of the previous edition of this book, the creative work of the research community has penetrated ever more deeply into the intricacies of signaling systems that regulate events in different regions of the kidney. This progress not only gives us a more complete picture of how the kidney accomplishes its various tasks, but also presents a still bigger challenge to the student to learn the principles of renal function in an integrated manner that is not split into isolated fragments. Our approach in this text is to distill the essence of renal processes and their regulation into a core that is accurate, yet not so burdened with detail that it overwhelms the reader. Rather than add complexity, we have purposely limited detail in order to keep the reader focused on the logic of renal processes. Accordingly, our goals in the preparation of this edition were (1) to incorporate the results of new literature, (2) point out the logical connection between events at the cellular level and their consequences for the well being of the body, and (3) to revise the text to improve its clarity. Consistent with prior editions we allot considerable coverage to body systems other than the kidneys, particularly the cardiovascular system, in order to bring out how the kidneys influence those systems, and in turn are influenced by those systems.

To help orient the reader and to prevent students from being mired in details, we have added key concepts at the end of each chapter. This feature also highlights the major points throughout the text and allows easy and quick review. We hope that these key concepts, together with the learning objectives, will be effective tools in learning renal physiology.
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OBJECTIVES

The student understands the disparate role of the kidneys in maintaining health.

- States the 7 major functions of the kidneys.
- Defines the balance concept.

The student understands the structural makeup of the kidneys, their blood supply, and the relation between major functional components.

- Defines the gross structures and their interrelationships: renal pelvis, calyces, renal pyramids, renal medulla (inner and outer zones), renal cortex, and papilla.
- Defines the components of the nephron and their interrelationships: renal corpuscle, glomerulus, nephron, and collecting-duct system.
- Draws the relationship between glomerulus, Bowman’s capsule, and the proximal tubule.
- Describes the 3 layers separating the lumen of the glomerular capillaries and Bowman’s space; defines podocytes, foot processes, and slit diaphragms.
- Defines glomerular mesangial cells and states their functions and location within the glomerulus.
- Lists the individual tubular segments in order; states the segments that comprise the proximal tubule, Henle’s loop, and the collecting-duct system; defines principal cells and intercalated cells.
- Lists, in order, the vessels through which blood flows from the renal artery to the renal vein; contrasts the blood supply to the cortex and the medulla; defines vasa recta and vascular bundles.
- Describes, in general terms, the differences among superficial cortical, midcortical, and juxtamedullary nephrons.
- Defines juxtaglomerular apparatus and describes its 3 cell types; states the function of the granular cells.

The student understands how the 2 kidneys handle substances in order to achieve balance of each.

- Defines the basic renal processes: glomerular filtration, tubular reabsorption, and tubular secretion.
- Defines renal metabolism of a substance and gives examples.
The kidneys perform an incredibly wide array of functions for the body, most of which are essential for life. Some renal functions have obvious logical and necessary connections to each other. Others seem to be totally independent. Most involve matching renal excretion of substances out of the body to inputs into the body (ie, providing a balance between input and output).

FUNCTIONS

A popular view considers the kidney to be an organ primarily responsible for the removal of metabolic waste from the body. Although this is certainly a major function of the kidneys, there are other functions that are arguably more important.

Function 1: Regulation of Water and Electrolyte Balance

The balance concept states that our bodies are in balance for any substance when the inputs and outputs of that substance are matched. Any difference between input and output leads to an increase or decrease in the amount of a substance within the body. Our input of water and electrolytes is enormously variable and is only sometimes driven in response to body needs. For example, we drink water when thirsty but we drink much more because it is a component of beverages that we consume for reasons other than hydration. We also consume food to provide energy, but food often contains large amounts of water. The kidneys respond by varying the output of water in the urine, thereby maintaining balance for water (ie, constant total body water content). Minerals like sodium, potassium, magnesium, and so on are components of foods and generally present far in excess of body needs. As with water, the kidneys excrete minerals at a highly variable rate that, in the aggregate, matches input. One of the amazing feats of the kidneys is their ability to regulate each of these minerals independently (ie, we can be on a high-sodium, low-potassium diet or low-sodium, high-potassium diet, and the kidneys will adjust excretion of each of these substances appropriately).

Function 2: Excretion of Metabolic Waste

Our bodies continuously form end products of metabolic processes. In most cases, those end products serve no function and are harmful at high concentrations. Some of these waste products include urea (from protein), uric acid (from nucleic acids), creatinine (from muscle creatine), the end products of hemoglobin breakdown (which give urine much of its color), and the metabolites of various hormones, among many others.

\footnote{Being in balance does not by itself imply a normal state or good health. A person may have an excess or deficit of a substance, yet still be in balance as long as output matches input. This is often the case in chronic disorders of renal function or metabolism.}
Function 3: Excretion of Bioactive Substances (Hormones and Many Foreign Substances, Specifically Drugs) That Affect Body Function

Drugs and hormones in the blood are removed in many ways, mostly in the liver, but a number of them are removed in parallel by renal processes. Physicians have to be mindful of how fast the drugs are excreted in order to prescribe a dose that achieves the appropriate body levels.

Function 4: Regulation of Arterial Blood Pressure

Although many people appreciate at least vaguely that the kidneys excrete waste substances like urea (hence the name urine) and salts, few realize the kidneys’ crucial role in controlling blood pressure. Blood pressure ultimately depends on blood volume, and the kidneys’ maintenance of sodium and water balance achieves regulation of blood volume. Thus, through volume control, the kidneys participate in blood pressure control. They also participate in regulation of blood pressure via the generation of vasoactive substances that regulate smooth muscle in the peripheral vasculature.

Function 5: Regulation of Red Blood Cell Production

Erythropoietin is a peptide hormone that is involved in the control of erythrocyte (red blood cell) production by the bone marrow. Its major source is the kidneys, although the liver also secretes small amounts. The renal cells that secrete it are a particular group of cells in the interstitium. The stimulus for its secretion is a reduction in the partial pressure of oxygen in the kidneys, as occurs, eg, in anemia, arterial hypoxia, and inadequate renal blood flow. Erythropoietin stimulates the bone marrow to increase its production of erythrocytes. Renal disease may result in diminished erythropoietin secretion, and the ensuing decrease in bone marrow activity is one important causal factor of the anemia of chronic renal disease.

Function 6: Regulation of Vitamin D Production

When we think of vitamin D, we often think of sunlight or additives to milk. In vivo vitamin D synthesis involves a series of biochemical transformations, the last of which occurs in the kidneys. The active form of vitamin D (1,25-dihydroxyvitamin D₃) is actually made in the kidneys, and its rate of synthesis is regulated by hormones that control calcium and phosphate balance.

Function 7: Gluconeogenesis

Our central nervous system is an obligate user of blood glucose regardless of whether we have just eaten sugary doughnuts or gone without food for a week. Whenever the intake of carbohydrate is stopped for much more than half a day, our body begins to synthesize new glucose (the process of gluconeogenesis) from noncarbohydrate sources (amino acids from protein and glycerol from triglycerides). Most gluconeogenesis occurs in the liver, but a substantial fraction occurs in the kidneys, particularly during a prolonged fast.
Most of what the kidneys actually do to perform the functions just mentioned involves transporting water and solutes between the blood flowing through the kidneys and the lumina of tubules (nephrons and collecting tubules that comprise the working mass of the kidneys). The lumen of a nephron is topologically outside the body, and any substance in the lumen that is not transported back into the blood is eventually excreted in the urine. As we explore renal function in more detail, we will constantly refer to tubular structure and the surrounding vasculature. Therefore, in the following section, we present the essential aspects of renal anatomy that are necessary to understand function.

ANATOMY OF THE KIDNEYS AND URINARY SYSTEM

The 2 kidneys lie outside the peritoneal cavity close to the posterior abdominal wall, 1 on each side of the vertebral column. Each of the 2 kidneys is a bean-shaped structure. The rounded, outer convex surface of each kidney faces the side of the body, and the indented surface, called the hilum, is medial. Each hilum is penetrated by a renal artery, renal vein, nerves, and a ureter, which carries urine out of the kidney to the bladder. Each ureter within a kidney is formed from funnel-like structures called major calyces, which, in turn, are formed from minor calyces. The minor calyces fit over underlying cone-shaped renal tissue called pyramids. The tip of each pyramid is called a papilla and projects into a minor calyx. The calyces act as collecting cups for the urine formed by the renal tissue in the pyramids. The pyramids are arranged radially around the hilum, with the papillae pointing toward the hilum and the broad bases of the pyramids facing the outside, top, and bottom of the kidney (from the 12-o’clock to the 6-o’clock position). The pyramids constitute the medulla of the kidney. Overlying the medullary tissue is a cortex, and covering the cortical tissue on the very external surface of the kidney is a thin connective tissue capsule (Figure 1–1).

The working tissue mass of both the cortex and medulla is constructed almost entirely of tubules (nephrons and collecting tubules) and blood vessels (capillaries and capillary-like vessels). Tubules and blood vessels are intertwined (something like a plateful of spaghetti) or arranged in parallel arrays (like bundles of soda straws) and, in either case, are always close to each other. Between the tubules and blood vessels lies an interstitium, which comprises less than 10% of the renal volume. The interstitium contains fluid and scattered interstitial cells (fibroblasts and others) that synthesize an extracellular matrix of collagen, proteoglycans, and glycoproteins.

The cortex and medulla have very different properties both structurally and functionally. On closer examination, we see that (1) the cortex has a highly granular appearance, absent in the medulla, and (2) each medullary pyramid is divisible into an outer zone (adjacent to the cortex) and an inner zone, which includes the papilla. All these distinctions reflect the arrangement of the various tubules and blood vessels.
The urinary system. The urine formed by a kidney collects in the renal pelvis and then flows through the ureter into the bladder, from which it is eliminated via the urethra. B, Section of a human kidney. Half the kidney has been sliced away. Note that the structure shows regional differences. The outer portion (cortex) contains all the glomeruli. The collecting ducts form a large portion of the inner kidney (medulla), giving it a striped, pyramid-like appearance, and these drain into the renal pelvis. The papilla is in the inner portion of the medulla.

**THE NEPHRON**

Each kidney contains approximately 1 million *nephrons*, one of which is shown diagrammatically in Figure 1–2. Each nephron consists of a spherical filtering component, called the *renal corpuscle*, and a tubule extending from the renal corpuscle. Let us begin with the renal corpuscle, which is responsible for the initial step in urine formation: the separation of a protein-free filtrate from plasma.
The Renal Corpuscle

The renal corpuscle consists of a compact tuft of interconnected capillary loops, the glomerulus (pl. glomeruli) or glomerular capillaries, surrounded by a balloon-like hollow capsule: Bowman’s capsule (Figure 1–3). Blood enters and leaves Bowman’s capsule through arterioles that penetrate the surface of the capsule at the vascular pole. A fluid-filled space (the urinary space or Bowman’s space) exists within the capsule (labeled US in Figure 1–3), and it is into this space that fluid filters. Opposite the vascular pole, Bowman’s capsule has an opening that leads into the first portion of the tubule (see Figure 1–3, bottom).

Figure 1–2. Relationships of component parts of a long-looped nephron, which has been “uncoiled” for clarity (relative lengths of the different segments are not drawn to scale). The combination of glomerulus and Bowman’s capsule is the renal corpuscle.
Figure 1–3. Diagram of a longitudinal section through a glomerulus and its juxtaglomerular (JG) apparatus. The JG apparatus consists of the granular cells (GC), which secrete renin, the macula densa (MD), and the extraglomerular mesangial cells (EGM). E, endothelium of the capillaries; EA, efferent arteriole; AA, afferent arteriole; PE, parietal (outer) epithelium of Bowman’s space; PO, podocytes of Bowman’s capsule; GBM, glomerular basement membrane; US, “urinary” (Bowman’s) space. (Reproduced with permission from Kriz W et al. In: Davidson AM, ed. Proceedings of the 10th International Congress on Nephrology, Vol 1. London: Balliere Tindall; 1987.)
The filtration barrier in the renal corpuscle through which all filtered substances must pass consists of 3 layers: the capillary endothelium of the glomerular capillaries, a rather thick basement membrane, and a single-celled layer of epithelial cells (Figure 1–4). The first layer, the endothelial cells of the capillaries, is perforated by many large fenestrae (“windows”), like a slice of Swiss cheese, and

Figure 1–4.  A, Anatomy of the glomerulus.  B, Cross-section of glomerular membranes. US, “urinary” (Bowman’s) space; E, epithelial foot processes; GBM, glomerular basement membranes; End, capillary endothelium; Cap, lumen of capillary. (Courtesy HG Rennke. Originally published in Fed Proc 1977;36:2019; reprinted with permission.)  C, Scanning electron micrograph of podocytes covering glomerular capillary loops; the view is from inside Bowman’s space. The large mass is a cell body. Note the remarkable interdigitation of the foot processes from adjacent podocytes and the slits between them. (Courtesy of C. Tisher.)
is freely permeable to everything in the blood except red blood cells and platelets. The middle layer, the capillary basement membrane, is not a membrane in the sense of a lipid bilayer membrane but is a gel-like acellular meshwork of glycoproteins and proteoglycans, with a structure like a kitchen sponge. The third layer consists of epithelial cells that rest on the basement membrane and face Bowman’s space. These cells are called podocytes. They are quite different from the relatively simple, flattened epithelial cells that line the outside of Bowman’s capsule. The podocytes have an unusual octopus-like structure. Small “fingers,” called pedicels (or foot processes), extend from each arm of the podocyte and are embedded in the basement membrane. Pedicels from a given podocyte interdigitate with the pedicels from adjacent podocytes. Spaces between adjacent pedicels constitute the path through which the filtrate, once through the endothelial cells and basement membrane, travels to enter Bowman’s space. The foot processes are coated by a thick layer of extracellular material, which partially occludes the slits, and extremely thin processes called slit diaphragms bridge the slits between the pedicels. Slit diaphragms are widened versions of the tight junctions and adhering junctions that link all contiguous epithelial cells together. These are like miniature ladders. The pedicels form the sides of the ladder, and the slit diaphragms are the rungs.

The functional significance of this anatomic arrangement is that it permits the filtration of large volumes of fluid from the capillaries into Bowman’s space but restricts filtration of large plasma proteins such as albumin.

Another cell type—the mesangial cell—is found in the central part of the glomerulus between and within capillary loops (see Figure 1–3). Glomerular mesangial cells act as phagocytes and remove trapped material from the basement membrane. They also contain large numbers of myofilaments and can contract in response to a variety of stimuli in a manner similar to vascular smooth muscle cells. The role of such contraction in influencing filtration by the renal corpuscles is discussed in Chapters 2 and 7.

The Tubule

Throughout its course, the tubule, which begins at and leads out of Bowman’s capsule, is made up of a single layer of epithelial cells resting on a basement membrane. The structural and immunocytochemical characteristics of these epithelial cells vary from segment to segment of the tubule. A common feature is the presence of tight junctions between adjacent cells that physically link them together (like the plastic form that holds a 6-pack of soft drinks together).

Table 1–1 lists the names and sequence of the various tubular segments, as illustrated in Figures 1–2 and 1–5. Physiologists and anatomists have traditionally grouped 2 or more contiguous tubular segments for purposes of reference, but the terminologies have varied considerably. Table 1–1 also gives the combination terms used in this text.

The proximal tubule, which drains Bowman’s capsule, consists of a coiled segment—the proximal convoluted tubule—followed by a straight segment—the proximal straight tubule—which descends toward the medulla, perpendicular to the cortical surface of the kidney (2 and 3 in Figure 1–5).
The next segment, into which the proximal straight tubule drains, is the descending thin limb of Henle’s loop (or simply the descending thin limb). The descending thin limb is in the medulla and is surrounded by an interstitial environment that is quite different from that in the cortex. The descending thin limb ends at a hairpin loop, and the tubule then begins to ascend parallel to the descending limb. The loops penetrate to varying depths within the medulla. In long loops (see later discussion), the epithelium of the first portion of this ascending limb remains thin, although different from that of the descending limb. This segment is called the ascending thin limb of Henle’s loop (or simply the ascending thin limb) (see Figure 1–5). Beyond this segment, in these long loops, the epithelium thickens, and this next segment is called the thick ascending limb of Henle’s loop (or simply the thick ascending limb). In short loops (see later discussion), there is no ascending thin limb, and the thick ascending limb begins right at the hairpin loop (see Figure 1–5). The thick ascending limb rises back into the cortex. Near the end of every thick ascending limb, the tubule returns to Bowman’s capsule, from which it originated, and passes directly between the afferent and efferent arterioles, as they enter and exit that renal corpuscle at its vascular pole (see Figure 1–3). The cells in the thick ascending limb closest to Bowman’s capsule (between the afferent and efferent arterioles) are specialized cells known as the macula densa. The macula densa marks the end of the thick ascending limb and the beginning of the distal convoluted tubule. This is followed by the connecting tubule, which leads to the cortical collecting tubule, the first portion of which is called the initial collecting tubule.

From Bowman’s capsule through the loop of Henle to the initial collecting tubules, each of the 1 million nephrons in each kidney is completely separate from the others. However, connecting tubules from several nephrons merge to form cortical collecting tubules, and a number of initial collecting tubules then join end
Figure 1–5. Standard nomenclature for structures of the kidney (1988 Commission of the International Union of Physiological Sciences). Shown are a short-looped and a long-looped (juxtamedullary) nephron, together with the collecting system (not drawn to scale). A cortical medullary ray—the part of the cortex that contains the straight proximal tubules, cortical thick ascending limbs, and cortical collecting ducts—is delineated by a dashed line. 1, renal corpuscle (Bowman's capsule and the glomerulus); 2, proximal convoluted tubule; 3, proximal straight tubule; 4, descending thin limb; 5, ascending thin limb; 6, thick ascending limb; 7, macula densa (located within the final portion of the thick ascending limb); 8, distal convoluted tubule; 9, connecting tubule; 9*, connecting tubule of a juxtamedullary nephron that arches upward to form a so-called arcade (there are only a few of these in the human kidney); 10, cortical collecting duct; 11, outer medullary collecting duct; 12, inner medullary collecting duct. (Reproduced with permission from Kriz W, Bankir L. *Am J Physiol* 1988;254LF:F1–F8.)
to end or side to side to form larger cortical collecting ducts. All the cortical collecting ducts then run downward to enter the medulla and become outer medullary collecting ducts and then inner medullary collecting ducts. The latter merge to form several hundred large ducts, the last portions of which are called papillary collecting ducts, each of which empties into a calyx of the renal pelvis.

The pathway taken by fluids flowing within a nephron always begins in the cortex (in Bowman’s capsule), descends into the medulla (descending limb of the loop of Henle), returns to the cortex (thick ascending limb of the loop of Henle), passes down into the medulla once more (medullary collecting tubule), and ends up in a renal calyx. Each renal calyx is continuous with the ureter, which empties into the urinary bladder, where urine is temporarily stored and from which it is intermittently eliminated. The urine is not altered after it enters a calyx. From this point on, the remainder of the urinary system serves only to maintain the fluid composition established by the kidney.

As noted earlier, the tubular epithelium has a one-cell thickness throughout. Before the distal convoluted tubule, the cells in any given segment are homogeneous and distinct for that segment. Thus, e.g., the thick ascending limb contains only thick ascending limb cells. However, beginning in the second half of the distal convoluted tubule, 2 cell types are found intermingled in most of the remaining segments. One type constitutes the majority of cells in the particular segment, is considered specific for that segment, and is named accordingly: distal convoluted tubule cells, connecting tubule cells, and collecting-duct cells, the latter known more commonly as principal cells. Interspersed among the segment-specific cells in each of these 3 segments are individual cells of the second type, called intercalated cells. There are actually several types of intercalated cells; 2 of them are called type A and type B. (The last portion of the medullary collecting duct contains neither principal cells nor intercalated cells but is composed entirely of a distinct cell type called the inner medullary collecting-duct cells.)

**BLOOD SUPPLY TO THE NEPHRONS**

The kidneys receive an enormous amount of blood relative to their mass. Blood enters each kidney via a renal artery, which then divides into progressively smaller branches: interlobar, arcuate, and finally interlobular arteries (usually called cortical radial arteries because they radiate outward toward the kidney surface). As each of the interlobular arteries projects toward the outer kidney surface, a series of parallel arterioles branch off at right angles (Figure 1–6), each of which leads to a glomerulus. These are called afferent arterioles. Note that these arteries and glomeruli are found only in the cortex, never in the medulla.

Normally about 20% of the plasma (and none of the erythrocytes) entering the glomerulus is filtered from the glomerulus into Bowman’s capsule, leaving the remaining 80% to flow on to the next vascular segment. In most organs, capillaries recombine to form the beginnings of the venous system, but the glomerular
capillaries instead recombine to form another set of arterioles called the *eff erent arterioles*. Thus, blood enters each glomerulus through a single afferent arteriole and leaves via a single efferent arteriole at the vascular pole of Bowman’s capsule (see Figure 1–3). The afferent and efferent arterioles both penetrate Bowman’s capsule on the same side, with the thick ascending limb of the nephron that originated from that capsule passing between and in contact with each arteriole. The efferent arterioles soon subdivide into a second set of capillaries. These are usually the peritubular capillaries, which are profusely distributed throughout the cortex. The peritubular capillaries then rejoin to form the veins by which blood ultimately leaves the kidney.

The medulla receives much less blood than does the cortex, and in a quite different manner. There are no glomeruli in the medulla. In contrast to most efferent arterioles in the cortex, those from juxtamedullary glomeruli do not branch into peritubular capillaries, but rather descend downward into the outer medulla, where they divide many times to form bundles of parallel vessels that penetrate deep into the medulla (see Figure 1–6). These are called descending *vasa recta* (Latin *recta* for “straight” and *vasa* for “vessels”). Although it is still uncertain, a small fraction of the descending vasa recta may branch off from the cortical radial arteries before the glomeruli, not after. The vasa recta on the outside of the vascular bundles “peel off” and give rise to interbundle plexi of capillaries that surround Henle’s loops and the collecting ducts in the outer medulla. Only the center-most vasa recta supply capillaries in the inner medulla; thus, limited blood flows into the papilla. The capillaries from the inner medulla re-form into ascending vasa recta that run in close association with the descending vasa recta within the vascular bundles. The structural and functional properties of the vasa recta are rather complex. The beginnings of the descending vasa recta are like arterioles, with pericytes containing smooth muscle in their walls, but become more capillary like as they descend (see Figure 1–6). The ascending vasa recta have a fenestrated endothelium like that found in the glomerular capillaries. Therefore, the vasa recta, in addition to being conduits for blood, also participate in exchanging water and solutes between plasma and interstitium. The whole arrangement of descending and ascending blood flowing in parallel has major significance for the formation of both concentrated and diluted urine (described in Chapter 6) because plasma and medullary interstitial constituents exchange between descending and ascending vessels.

**Categories of Nephrons**

There are important regional differences in the various tubular segments of the nephron. All the renal corpuscles are in the cortex (accounting for its granular appearance) as well as the convoluted portions of the proximal tubule, cortical portions of Henle’s loops, distal convoluted tubules, connecting tubules, and cortical collecting ducts. The medulla contains the medullary portions of Henle’s loops and the medullary collecting ducts.
Figure 1–6. The renal microcirculation. The kidney is divided into a cortex and a medulla. The cortex contains an arterial network, glomeruli, a dense peritubular capillary plexus, and a venous drainage system. In the cortex, arcuate arteries, which run parallel to the surface, give rise to cortical radial (interlobular) arteries radiating toward the surface. Afferent arterioles originate from the cortical radial arteries at an angle that varies with cortical location. Blood is supplied to the peritubular capillaries of the cortex from the efferent flow out of superficial glomeruli. Blood is supplied to the medulla from the efferent flow out of juxtamedullary glomeruli. Efferent arterioles of juxtamedullary glomeruli give rise to bundles of descending vasa recta in the outer stripe of the outer medulla. In the inner stripe of the outer medulla, descending vasa recta and ascending vasa recta returning from the inner medulla run side by side in the vascular bundles, allowing exchange of solutes and water as described in Chapter 6. The descending vasa recta from the bundle periphery supply the interbundle capillary plexus of the inner stripe, whereas those in the center supply blood to the capillaries of the inner medulla. Contractile pericytes in the walls of the descending vasa recta regulate flow. DVR, descending vasa recta. AVR, ascending vasa recta. (Used with permission from Pallone TL, Zhang Z, Rhinehart K. Am J Physiol Renal Physiol 2003;284:F253–F266.)
Nephrons are categorized according to the locations of their renal corpuscles in the cortex (see Figure 1–5): (1) In superficial cortical nephrons, renal corpuscles are located within 1 mm of the capsular surface of the kidneys; (2) in midcortical nephrons, renal corpuscles are located, as their name implies, in the midcortex, deep relative to the superficial cortical nephrons but above (3) the juxtamedullary nephrons, which, as mentioned previously, have renal corpuscles located just above the junction between cortex and medulla. One major distinction among these 3 categories of nephrons is the length of Henle’s loop. All superficial cortical nephrons have short loops, which make their hairpin turn above the junction of outer and inner medulla. All juxtamedullary nephrons have long loops, which extend into the inner medulla, often to the tip of a papilla. Midcortical nephrons may be either short looped or long looped. The additional length of Henle’s loop in long-looped nephrons is due to a longer descending thin limb and the presence of an ascending thin limb. Finally, the beginning of the thick ascending limb marks the border between the outer and inner medulla; in other words, the thick ascending limbs are found only in the cortex and outer medulla.

**Nephron Heterogeneity**

As stated earlier, there are more than 2 million nephrons in the 2 human kidneys. These nephrons manifest significant differences in anatomic, biochemical, and functional characteristics beyond those described in the previous section. For simplicity, however, we generally ignore these complexities, many of which currently are not fully understood.

**The Juxtaglomerular Apparatus**

Reference was made earlier to the macula densa, a portion of the late thick ascending limb at the point where, in all nephrons, this segment comes between the afferent and efferent arterioles at the vascular pole of the renal corpuscle from which the tubule arose. This entire area is known as the juxtaglomerular (JG) apparatus (see Figure 1–3). (Do not confuse the term juxtaglomerular apparatus with juxtamedullary nephron.) Each JG apparatus is made up of 3 cell types: (1) granular cells, also called juxtaglomerular cells (GC in Figure 1–3), which are differentiated smooth muscle cells in the walls of the afferent arterioles; (2) extraglomerular mesangial cells (EGM in Figure 1–3); and (3) macula densa cells (MD in Figure 1–3), which are specialized thick ascending limb epithelial cells.

The granular cells (so called because they contain secretory vesicles that appear granular in light micrographs) are the cells that secrete the hormone renin. As we will describe in Chapter 7, renin is a crucial substance for the control of renal function and systemic blood pressure. The extraglomerular mesangial cells are morphologically similar to and continuous with the glomerular mesangial cells but lie outside Bowman’s capsule. The macula densa cells are detectors of the luminal content of the nephron at the very end of the thick ascending limb and contribute to the control of glomerular filtration rate (GFR) and to the control of renin secretion.
Renal Innervation
The kidneys receive a rich supply of sympathetic neurons. These are distributed to the afferent and efferent arterioles, the JG apparatus, and many portions of the tubule. There is no significant parasympathetic innervation. The sensory endings of many afferent neurons are distributed throughout the kidneys as well, which feedback to the neural centers that regulate sympathetic outflow.

BASIC RENAL PROCESSES
The working structures of the kidney are the nephrons and collecting tubules into which the nephrons drain. Figure 1–7 illustrates the meaning of several key words that we use to describe how the kidneys function. It is essential that any student of the kidney grasp their meaning.

Filtration is the process by which water and solutes in the blood leave the vascular system through the filtration barrier and enter Bowman’s space (a space that is topologically outside the body). Secretion is the process of moving substances into the tubular lumen from the cytosol of epithelial cells that

Figure 1–7. The 3 basic renal processes. Only the directions of reabsorption and secretion, not specific sites or order of occurrence, are shown. Depending on the specific substance, reabsorption and secretion can occur at various sites along the tubule.
form the walls of the nephron. Secreted substances may originate by synthesis within the epithelial cells or, more often, by crossing the epithelial layer from the surrounding renal interstitium. **Reabsorption** is the process of moving substances from the lumen across the epithelial layer into the surrounding interstitium. In most cases, reabsorbed substances then move from the interstitium into surrounding blood vessels, so that the term **reabsorption** implies a 2-step process of removal from the lumen followed by movement into the blood. **Excretion** means exit of the substance from the body (ie, the substance is present in the final urine produced by the kidneys). **Synthesis** means that a substance is constructed from molecular precursors, and **catabolism** means the substance is broken down into smaller component molecules.

The renal handling of any substance consists of some combination of the just-mentioned processes. If we can answer the following questions, we can know what the kidney does with a given substance. Is it filtered? Is it secreted? Is it reabsorbed? Is it synthesized? Is it catabolized?

**Glomerular Filtration**

Urine formation begins with glomerular filtration, the bulk flow of fluid from the glomerular capillaries into Bowman’s capsule. The glomerular filtrate (ie, the fluid within Bowman’s capsule) is very much like blood plasma. However, it contains very little total protein. The large plasma proteins like albumin and globulins are virtually excluded from moving through the filtration barrier. Smaller proteins, such as many of the peptide hormones, are present in the filtrate, but their mass in total is miniscule compared with the mass of large plasma proteins in the blood. The filtrate contains most inorganic ions and low-molecular-weight organic solutes in virtually the same concentrations as in the plasma. Substances that are present in the filtrate at the same concentration as found in the plasma are said to be **freely filtered**. (Note that *freely* filtered does not mean *all* filtered. The amount filtered is in exact proportion to the fraction of plasma volume that is filtered.) Many low-molecular-weight components of blood are freely filtered. Among the most common substances included in the freely filtered category are the ions sodium, potassium, chloride, and bicarbonate; the neutral organics glucose and urea; amino acids; and peptides like insulin and antidiuretic hormone (ADH).

The volume of filtrate formed per unit time is known as the GFR. In a normal young adult male, the GFR is an incredible 180 L/day (125 mL/min)! Contrast this value with the net filtration of fluid across all the other capillaries in the body: approximately 4 L/day. The implications of this huge GFR are extremely important. When we recall that the average total volume of plasma in humans is approximately 3 L, it follows that the entire plasma volume is filtered by the kidneys some 60 times a day. The opportunity to filter such huge volumes of plasma enables the kidneys to excrete large quantities of waste products and to regulate the constituents of the internal environment very precisely. One of the general consequences of aging and of many renal pathologies is a reduction in the GFR.

The forces that determine the GFR and their physiological control are described in Chapters 2 and 7.
Tubular Reabsorption and Tubular Secretion

The volume and solute contents of the final urine that enters the renal pelvis are quite different from those of the glomerular filtrate. Clearly, almost all the filtered volume must be reabsorbed; otherwise, with a filtration rate of 180 L/day, we would urinate ourselves into dehydration very quickly. As the filtrate flows from Bowman’s capsule through the various portions of the tubule, its composition is altered, mostly by removing material (tubular reabsorption) but also by adding material (tubular secretion). As described earlier, the tubule is, at all points, intimately associated with peritubular capillaries in the cortex or capillary-like vessels in the medulla, a relationship that permits rapid transfer of materials between the capillary plasma and the lumen of the tubule via the interstitial space.

The most common relationships among these basic renal processes, glomerular filtration, tubular reabsorption, and tubular secretion, are shown in the hypothetical examples of Figure 1–8. Plasma, containing 3 low-molecular-weight substances (X, Y, and Z), enters the glomerular capillaries, and approximately 20% of the plasma is filtered into Bowman’s capsule. The filtrate contains substances X, Y, and Z in the same concentrations as the plasma (ie, each one is freely filtered). The filtrate enters the proximal convoluted tubule and begins its flow through the rest of the tubule. Simultaneously, the remaining 80% of the plasma, with its substances X, Y, and Z in the same concentrations as they had when entering the kidney, leaves the glomerular capillaries via the efferent arterioles and enters the peritubular capillaries.

Suppose the cells of the tubular epithelium can secrete all the peritubular-capillary substance X into the tubular lumen but cannot reabsorb substance X. Thus, by the combination of filtration and tubular secretion, all the plasma that originally entered the renal artery is cleared of substance X, which leaves the body via the urine. Now suppose the tubule can reabsorb some of substance Y. The amount of substance Y reabsorbed is small, so most of the filtered substance Y escapes from the body in the urine. In contrast, let substance Z be reabsorbed fully. Therefore, no substance Z is lost from the body. Hence, the processes of filtration and reabsorption have canceled each other, and the net result is as though substance Z had never entered the kidney at all.

As we will see, most of the tubular transport consists of reabsorption rather than tubular secretion. An idea of the magnitude and importance of tubular reabsorption can be gained from Table 1–2, which summarizes data for a few plasma components that undergo reabsorption. The values in Table 1–2 are typical for a normal person on an average diet. There are at least 3 important generalizations to be drawn from this table:

1. Because of the huge GFR, the quantities filtered per day are enormous, generally larger than the amounts of the substances in the body. For example, the body contains about 40 L of water, but the volume of water filtered each day may be as large as 180 L. If reabsorption of water ceased but filtration continued, the total plasma water would be urinated within 30 min.
Figure 1–8. Renal manipulation of 3 hypothetical substances, X, Y, and Z. Substance X is filtered and secreted but not reabsorbed. Substance Z is filtered but is completely reabsorbed.
2. Reabsorption of waste products, such as urea, is incomplete, so that large fractions of their filtered amounts are excreted in the urine, like substance Y in our hypothetical example.

3. Reabsorption of most “useful” plasma components (eg, water, electrolytes, and glucose) varies from essentially complete, so that urine concentrations should normally be undetectable (eg, glucose), to almost complete (eg, water and most electrolytes), so that the amounts excreted in the urine represent only very small fractions of the filtered amounts.

For each plasma substance, a particular combination of filtration, reabsorption, and secretion applies. The relative proportions of these processes then determine the amount excreted. A critical point is that the rates at which the relevant processes proceed for many of these substances are subject to physiological control. By triggering changes in the rates of filtration, reabsorption, or secretion when the body content of a substance goes above or below normal, these mechanisms can regulate excretion to keep the body in balance. For example, consider what happens when a person drinks a large quantity of water: Within 1–2 h, all the excess water has been excreted in the urine, partly as the result of an increase in GFR but mainly as the result of decreased tubular reabsorption of water. The body is kept in balance for water by increasing excretion. By keeping the body in balance, the kidney is the effector organ of a reflex that maintains body water concentration within very narrow limits.

### Metabolism by the Tubules

Although renal physiologists traditionally list glomerular filtration, tubular reabsorption, and tubular secretion as the 3 basic renal processes, we cannot overlook metabolism by the tubular cells. For example, the tubular cells may extract organic nutrients from the glomerular filtrate or peritubular capillaries and metabolize them as dictated by the cells’ own nutrient requirements. In doing so, the renal cells are behaving no differently from any other cells in the body. In contrast, other metabolic transformations performed by the kidney are not directed toward its own nutritional requirements but rather toward altering the composition of the urine and plasma. The most important of these are the synthesis of ammonium from glutamine and the production of bicarbonate, both described in Chapter 9.

### Table 1–2. Average values for several substances handled by filtration and reabsorption

<table>
<thead>
<tr>
<th>Substance</th>
<th>Amount filtered per day</th>
<th>Amount excreted</th>
<th>% Reabsorbed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water, L</td>
<td>180</td>
<td>1.8</td>
<td>99.0</td>
</tr>
<tr>
<td>Sodium, g</td>
<td>630</td>
<td>3.2</td>
<td>99.5</td>
</tr>
<tr>
<td>Glucose, g</td>
<td>180</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>Urea, g</td>
<td>56</td>
<td>28</td>
<td>50</td>
</tr>
</tbody>
</table>
Regulation of Renal Function

By far, the most difficult aspect of renal physiology for students (and authors alike) is regulation of renal function. Neural signals, hormonal signals, and intrarenal chemical messengers combine to regulate the basic renal processes presented previously in a manner to help the kidneys meet the needs of the body. Unfortunately, our collective knowledge on much of this is, as yet, incomplete. Of necessity, much of the coverage in this textbook will attempt to draw an overview of renal function without an emphasis on nuance and detail that is more appropriate for advanced texts.

As with many organs, signals regulating the kidney arise from both neural and hormonal input. Neural signals originate in the sympathetic celiac plexus. Sympathetic signals exert major control over renal blood flow, glomerular filtration, and the release of vasoactive substances (the renin-angiotensin system, described later). Hormonal signals originate in the adrenal gland, pituitary gland, and heart. The adrenal cortex secretes the steroid hormones aldosterone and cortisol, and the adrenal medulla secretes the catecholamines epinephrine and norepinephrine. All of these hormones, but mainly aldosterone, are regulators of sodium and potassium excretion by the kidney. The pituitary gland secretes the hormone arginine vasopressin (also called ADH). ADH is a major regulator of water excretion, and via its influence on the renal vasculature and possible collecting-duct principal cells, probably sodium excretion as well. The heart secretes hormones, natriuretic peptides, that contribute to signaling increased excretion of sodium by the kidneys. The most difficult aspect of regulation lies in the realm of intrarenal chemical messengers (ie, messengers that originate in one part of the kidney and act in another part). It is clear that an array of substances (eg, nitric oxide, purinergic agonists, superoxide, various eicosanoids) influence basic renal processes, but, for the most part, the role of these substances is beyond the scope of this text.

Overview of Regional Function

We conclude this chapter with a broad overview of the tasks performed by the various nephron segments. Later, we examine renal function substance by substance and see how tasks performed in the various regions combine to produce an overall result that is useful for the body.

The glomerulus is the site of filtration—about 180 L/day of volume and proportional amounts of solutes that are freely filtered, which is the case for most solutes (large plasma proteins are an exception). The glomerulus is where the greatest mass of excreted substances enter the nephron. The proximal tubule (convoluted and straight portions) reabsorbs about two thirds of the filtered water, sodium, and chloride. The proximal convoluted tubule reabsors all of the useful organic molecules that the body wishes to conserve (eg, glucose, amino acids). It reabsors significant fractions, but by no means all, of many important ions, such as potassium, phosphate, calcium, and bicarbonate. It is the site of secretion of a number of organic substances that are either metabolic waste products (eg, urate, creatinine) or drugs (eg, penicillin) that physicians must replace to make up for renal excretion.
The loop of Henle contains different segments that perform different functions, but the key functions occur in the thick ascending limb (a region that begins in the outer medulla for all nephrons and continues outward into the renal cortex until it reaches the renal corpuscle from which the tubule arose (which can, depending on the nephron, be near the corticomedullary border or close to the cortical surface)). As a whole, the loop of Henle reabsorbs about 20% of the filtered sodium and chloride and 10% of the filtered water. A crucial consequence of these different proportions is that, by reabsorbing relatively more salt than water, the luminal fluid becomes diluted relative to normal plasma and the surrounding interstitium. During periods when the kidneys excrete dilute final urine, the role of the loop of Henle in diluting the luminal fluid is crucial.

The end of the loop of Henle contains cells of the macula densa, which senses or assays the sodium and chloride content of the lumen and generates signals that influence other aspects of renal function, specifically the renin-angiotensin system (discussed in Chapter 7).

The distal tubule and connecting tubule together reabsorb some additional salt and water, perhaps 5% of each.

The cortical collecting tubule is where several (6–10) connecting tubules join to form 1 tubule. Cells of the cortical collecting tubule are strongly responsive to and are regulated by the hormones aldosterone and ADH. Aldosterone enhances sodium reabsorption and potassium secretion by this segment, and ADH enhances water reabsorption. The degree to which these processes are stimulated or not stimulated plays a major role in regulating the amount of solutes and water present in the final urine. With large amounts of ADH present, most of the water remaining in the lumen is reabsorbed, leading to concentrated, low-volume urine. With little ADH present, most of the water passes on to the final urine, producing dilute, high-volume urine.

The medullary collecting tubule continues the functions of the cortical collecting tubule in salt and water reabsorption. In addition, it plays a major role in regulating urea reabsorption and in acid-base balance (secretion of protons or bicarbonate).

**KEY CONCEPTS**

1. One major function of the kidneys is to regulate excretion of substances at a rate that, on average, exactly balances their input into the body and, thereby, maintain total body homeostatic balance for many substances.

2. A second major function of the kidneys is to regulate blood volume, blood osmolality, and total body sodium content in a way that determines average blood pressure.

3. The working tissues of the kidney are divided into an outer cortex and inner medulla.
Each functional renal unit is composed of a filtering component (glomerulus) and a transporting tubular component (the nephron and collecting duct).

The cortex receives an enormous volume of blood that flows in series through glomerular capillaries and then peritubular capillaries, whereas blood flow to the medulla is highly restricted.

The renal handling of any substance is defined by its rate of filtration, reabsorption, secretion, and, in some cases, metabolism.

**STUDY QUESTIONS**

1–1. Is the following statement true or false? The difference between superficial and juxtamedullary nephrons is that superficial nephrons have their glomeruli in the cortex whereas the glomeruli of juxtamedullary nephrons are in the medulla.

1–2. What percentage of the blood entering the kidney flows directly into the medulla without passing through the cortex?

1–3. Substance T is present in the urine. Does this prove that it entered the renal tubule by filtration at the glomerulus?

1–4. Substance V is not normally present in the urine. Does this mean that it does not enter the kidney at all (in the blood), or is neither filtered nor secreted?

1–5. A substance is filtered into Bowman’s space and excreted in the urine. How many cell plasma membrane barriers must it cross in order to move from the blood to outside the body?

1–6. A substance is freely filtered. Does this mean that it is all filtered?

1–7. If you immunologically labeled cells of the macula densa with label X, and labeled cells of the thick ascending limb with label Y, would you find labels X and Y in the cortex, medulla, or both?

1–8. Given the generalizations about transport events in the medulla (secretion, reabsorption), can you say that blood flow into the medulla is in any way different in volume from blood flow out of the medulla?
Renal Blood Flow and Glomerular Filtration

OBJECTIVES

The student understands the hemodynamics of renal blood flow.

- Defines renal blood flow, renal plasma flow, glomerular filtration rate, and filtration fraction and gives normal values.
- States the formula relating flow, pressure, and resistance in an organ.
- Describes the relative resistances of the afferent arterioles and efferent arterioles.
- Describes the effects of changes in afferent and efferent arteriolar resistances on renal blood flow.

The student understands how glomerular filtrate is formed and the forces that determine its rate of formation.

- Describes how molecular size and electrical charge determine filterability of plasma solutes; states how protein binding of a low-molecular-weight substance influences its filterability.
- States the formula for the determinants of glomerular filtration rate, and states, in qualitative terms, why the net filtration pressure is positive.
- Defines filtration coefficient and states how mesangial cells might alter the filtration coefficient; states the reason why glomerular filtration rate is so large relative to filtration across other capillaries in the body.
- Describes how arterial pressure, afferent arteriolar resistance, and efferent arteriolar resistance influence glomerular capillary pressure.
- Describes how changes in renal plasma flow influence average glomerular capillary oncotic pressure.

The student understands the normal controls of renal blood flow and glomerular filtration rate.

- States the Starling forces involved in capillary filtration.
- States how changes in each Starling force affect glomerular filtration rate.
- Defines autoregulation of renal blood flow and glomerular filtration rate.
- Describes the myogenic and tubuloglomerular feedback mechanisms of autoregulation.
GLOMERULAR FILTRATION AND RENAL BLOOD FLOW

The kidneys receive an enormous blood flow: more than 1 L/min, or about 20% of the cardiac output. This blood flow is far in excess of the kidney’s metabolic need and provides the kidneys with the flexibility to alter their blood flow in response to physiological demand. All of this blood flows through glomeruli in the cortex. The vast majority continues on (via efferent arterioles) to peritubular capillaries in the cortex and then into the renal venous system. A much smaller fraction, about 5–10%, flows from efferent arterioles down into the medulla. This medullary blood derives from juxtamedullary glomeruli that are situated near the corticomedullary border. Consider some typical numbers. A normal hematocrit is 0.45, i.e., 45% of the blood volume is composed of red blood cells and the remaining 55% is almost entirely plasma. Typical renal blood flow (RBF) is 1.1 L/min. The renal plasma flow (RPF) = 0.55 × 1.1 L/min = 605 mL/min. As stated in Chapter 1, a typical glomerular filtration rate (GFR) is about 125 mL/min. Thus, of the 605 mL of plasma that enters the glomeruli via the afferent arterioles, 125 mL, or 20%, filters into Bowman’s space. The remaining 480 mL passes via the efferent arterioles into the peritubular capillaries. This ratio—GFR/RPF—is known as the filtration fraction. Because freely filtered substances are passing into Bowman’s space along with the water, about 20% of all freely filtered substances (e.g., sodium) that enter the kidney also move into Bowman’s space.

FLOW, RESISTANCE, AND PRESSURE IN THE KIDNEYS

The basic equation for blood flow through any organ is as follows:

\[ Q = \frac{\Delta P}{R} \]

where \( Q \) is organ blood flow, \( \Delta P \) is mean pressure in the artery supplying the organ minus mean pressure in the vein draining that organ, and \( R \) is the total vascular resistance in that organ. Resistance is determined by the blood viscosity and the lengths and radii of the organ’s blood vessels, the arteriolar radii being overwhelmingly the major contributor. As described by Poiseuille’s law, resistance of a cylindrical vessel varies inversely with the fourth power of vessel radius. It takes only a 19% decrease or increase in vessel radius to double or halve vessel resistance. The radii of arterioles are regulated by the state of contraction of the arteriolar smooth muscle.

The presence of 2 two sets of arterioles (afferent and efferent) and 2 sets of capillaries (glomerular and peritubular) makes the vasculature of the cortex unusual. (The vasculature of the medulla is even more unusual, but we concentrate on the cortex for now.) Normally, the resistances of the afferent and efferent arterioles are approximately equal and account for most of the total renal vascular resistance. Resistance in arteries preceding afferent arterioles (i.e., cortical radial arteries) plays some role also, but we concentrate on the arterioles. Vascular pressures (i.e,

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1There is some evidence that a small fraction of blood approaching Bowman’s capsule in afferent arterioles is diverted directly to the medulla without going through glomeruli. This fraction, if it exists at all, is small.
hydrostatic or hydraulic pressure) in the 2 capillary beds are quite different. The peritubular capillaries are downstream from the efferent arteriole and have a lower hydraulic pressure. Typical glomerular pressures are near 60 mm Hg in a normal unstressed individual, whereas peritubular pressures are closer to 20 mm Hg. The high glomerular pressure is crucial for glomerular filtration, whereas the low peritubular capillary pressure is equally crucial for the tubular reabsorption of fluid.

To repeat, total RBF is determined mainly by the mean pressure in the renal artery and the contractile state of the smooth muscle of the renal arterioles of the cortex. Now for a simple but very important point: A change in arteriolar resistance produces the same effect on RBF regardless of whether it occurs in the afferent arteriole or efferent arteriole. Because these vessels are in series, a change in either one has the same effect on the total. When the 2 resistances both change in the same direction, the most common state of affairs, their effects on RBF will be additive. When they change in different directions—one resistance increasing and the other decreasing—they exert opposing effects on RBF. We see in the next section that the story is totally different for GFR.

GLOMERULAR FILTRATION

Formation of Glomerular Filtrate

As stated in Chapter 1, the glomerular filtrate is nearly protein-free and contains most inorganic ions and low-molecular-weight organic solutes in virtually the same concentrations as in the plasma.

In order to form a glomerular filtrate, filtered fluid must pass through the glomerular filtration barrier. As described anatomically in Chapter 1 (see Figure 1–4), the filtration barrier separates the blood from the urinary space that topologically connects to the outside world via the renal tubules, ureters, bladder, and urethra. The route that filtered substances take from the blood through the filtration barrier of a renal corpuscle into Bowman’s space is a 3-step process: through fenestrae in the glomerular-capillary endothelial layer, through the basement membrane, and finally through slit diaphragms between podocyte foot processes. The fraction of endothelial surface area occupied by fenestrae is about 10%. Both the slit diaphragm and basement membrane are composed of an array of proteins, and while the basement membrane may contribute to selectivity of the filtration barrier, integrity of the slit diaphragms is essential to prevent excessive leak of plasma protein (albumin). Some protein-wasting diseases are associated with abnormal slit diaphragm structure. Selectivity of the barrier to filtered solute is based on both molecular size and electrical charge. Let us look first at size.

The filtration barrier of the renal corpuscle provides no hindrance to the movement of molecules with molecular weights less than 7000 Da (ie, solutes this small are all freely filtered). This includes all small ions, glucose, urea, amino acids, and many hormones. The filtration barrier almost totally excludes plasma albumin.

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2It is protein-free in the sense that the total concentration of protein is very low. However, many small proteins that have low plasma concentrations, such as many peptide hormones, are freely or nearly freely filtered.
(molecular weight of approximately 66,000 Da). (We are, for simplicity, using molecular weight as our reference for size; in reality, it is molecular radius and shape that is critical.) The hindrance to plasma albumin is not 100%, however, and so the glomerular filtrate does contain extremely small quantities of albumin, on the order of 10 mg/L or less. This is only about 0.02% of the concentration of albumin in plasma and is the reason for the use of the phrase “nearly protein-free” earlier. (Note: Some small substances are partly or mostly bound to large plasma proteins and are thus not free to be filtered, even though, when not bound to plasma proteins, they can easily move through the filtration barrier. This includes hydrophobic hormones of the steroid and thyroid categories and about 40% of the calcium in the blood.)

For molecules with a molecular weight ranging from 7000 and 70,000 Da, the amount filtered becomes progressively smaller as the molecule becomes larger. Thus, many normally occurring plasma peptides and small proteins are filtered to a significant degree. Moreover, when certain small proteins not normally present in the plasma appear because of disease (eg, hemoglobin released from damaged erythrocytes or myoglobin released from damaged muscles), considerable filtration of these may occur.

Electrical charge is the second variable determining filterability of macromolecules. For any given size, negatively charged macromolecules are filtered to a lesser extent, and positively charged macromolecules to a greater extent, than neutral molecules. This is because the surfaces of all the components of the filtration barrier (the cell coats of the endothelium, the basement membrane, and the cell coats of the podocytes) contain fixed polyanions, which repel negatively charged macromolecules during filtration. Because almost all plasma proteins bear net negative charges, this electrical repulsion plays a very important restrictive role, enhancing that of purely size hindrance. (For example, when neutral dextrans, the same size as plasma albumin, are administered to experimental animals, they are found to be 5–10% filterable rather than albumin’s 0.02%.) In other words, if albumin were not charged or the filtration barrier were not charged, even albumin would be filtered to a considerable degree. Certain diseases that cause glomerular capillaries to become “leaky” to protein do so by eliminating negative charges in the membranes.

It must be emphasized that the negative charges in the filtration membranes act as a hindrance only to macromolecules, not to mineral ions or low-molecular-weight organic solutes. Thus, chloride and bicarbonate ions, despite their negative charge, are freely filtered.

**Direct Determinants of GFR**

Variation in GFR is a crucial determinant of renal function. Everything else being equal, a higher GFR means greater excretion of salt and water. Regulation of the GFR is straightforward in terms of physical principles but very complex functionally because there are so many regulated variables. The rate of filtration in any of the body’s capillaries, including the glomeruli, is determined by the hydraulic permeability of the capillaries, their surface area, and the net filtration pressure (NFP) acting across them.
Rate of filtration = hydraulic permeability × surface area × NFP

Because it is difficult to estimate the area of a capillary bed, a parameter called the filtration coefficient \(K_f\) is used to denote the product of the hydraulic permeability and the area.

The net filtration pressure is the algebraic sum of the hydrostatic pressures and the osmotic pressures resulting from protein—the oncotic or colloid osmotic pressures (for additional description of the importance of oncotic pressure, see Chapter 4)—on the 2 sides of the capillary wall. There are 4 pressures to contend with: 2 hydrostatic pressures and 2 oncotic pressures. These are referred to as *Starling forces*, named after the physiologist who first described them. Applying this to the glomerular capillaries:

\[
NFP = (P_{GC} - \pi_{GC}) - (P_{BC} - \pi_{BC}),
\]

where \(P_{GC}\) is glomerular capillary hydraulic pressure, \(\pi_{BC}\) is oncotic pressure of fluid in Bowman’s capsule, \(P_{BC}\) is hydraulic pressure in Bowman’s capsule, and \(\pi_{GC}\) is oncotic pressure in glomerular capillary plasma, shown schematically in Figure 2–1.

Because there is normally little protein in Bowman’s capsule, \(\pi_{BC}\) may be taken as zero and not considered in our analysis. Accordingly, the overall equation for GFR becomes

\[
GFR = K_f (P_{GC} - \pi_{GC}).
\]

---

*Figure 2–1.* Net filtration pressure in the renal corpuscle equals glomerular-capillary hydraulic pressure \(P_{GC}\) minus Bowman’s capsule hydraulic pressure \(P_{BC}\) minus glomerular-capillary oncotic pressure \(\pi_{GC}\).
The hydrostatic pressures in glomerular capillaries and Bowman’s capsule have not been directly measured in humans. However, several lines of indirect evidence suggest that the human values are probably similar to those for the dog, and these values are shown in Table 2–1 and Figure 2–2 along with glomerular capillary oncotic pressure.

**Table 2–1.** Estimated forces involved in glomerular filtration in humans

<table>
<thead>
<tr>
<th>Forces</th>
<th>Afferent end of glomerular capillary (mm Hg)</th>
<th>Efferent end of glomerular capillary (mm Hg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Favoring filtration</td>
<td>60</td>
<td>58</td>
</tr>
<tr>
<td>Glomerular-capillary hydraulic pressure, $P_{GC}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 Opposing filtration</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>a Hydraulic pressure in Bowman’s capsule, $P_{BC}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>b Oncotic pressure in glomerular capillary, $\pi_{GC}$</td>
<td>21</td>
<td>33</td>
</tr>
<tr>
<td>3 Net filtration pressure (1 − 2)</td>
<td>24</td>
<td>10</td>
</tr>
<tr>
<td>$P_{GC}$ − $\pi_{GC}$ − $P_{BC}$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The estimated forces involved in glomerular filtration in humans (these are the same values shown in Table 2–1). Net filtration pressure (NFP) $= P_{GC} − \pi_{GC} − P_{BC}.$
Note that the hydraulic pressure changes only slightly along the glomeruli; this is because the very large total cross-sectional area of the glomeruli collectively provides only a small resistance to flow. Importantly, note that the oncotic pressure in the glomerular capillaries does change substantially along the length of the glomeruli. Water is moving out of the vascular space and leaving protein behind, thereby raising protein concentration and, hence, the oncotic pressure of the unfiltered plasma remaining in the glomerular capillaries. Mainly because of this large increase in oncotic pressure, the net filtration pressure decreases from the beginning of the glomerular capillaries to the end. The average net filtration pressure over the whole length of the glomerulus is about 17 mm Hg. This average net filtration pressure is higher than found in most nonrenal capillary beds. Along with a high value for $K_f$, it accounts for the enormous filtration of 180 L of fluid/day (compared with 3 L/day or so in all other capillary beds combined).

As we have noted, the GFR is not fixed but shows marked fluctuations in differing physiological states and in disease. If all other factors remain constant, any change in $K_f$, $P_{GC}$, $P_{BC}$, or $\pi_{GC}$ will alter GFR. However, “all other factors” do not always remain constant, and so other simultaneous events may oppose the effect of any one factor. To grasp this situation, it is essential to see how a change in any one factor affects GFR under the assumption that all other factors are held constant.

Table 2–2 presents a summary of these factors. It provides, in essence, a checklist to review when trying to understand how diseases or vasoactive chemical messengers and drugs change GFR. In this regard, it should be noted that the major

Table 2–2. Summary of direct GFR determinants and factors that influence them

<table>
<thead>
<tr>
<th>Direct determinants of GFR: $GFR = K_f(P_{GC} - P_{BC} - \pi_{GC})$</th>
<th>Major factors that tend to increase the magnitude of the direct determinant</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_f$</td>
<td>1. ↑ Glomerular surface area (because of relaxation of glomerular mesangial cells)</td>
</tr>
<tr>
<td></td>
<td>Result: ↑ GFR</td>
</tr>
<tr>
<td>$P_{GC}$</td>
<td>1. ↑ Renal arterial pressure</td>
</tr>
<tr>
<td></td>
<td>2. ↓ Afferent-arteriolar resistance (afferent dilation)</td>
</tr>
<tr>
<td></td>
<td>3. ↑ Efferent-arteriolar resistance (efferent constriction)</td>
</tr>
<tr>
<td></td>
<td>Result: ↑ GFR</td>
</tr>
<tr>
<td>$P_{BC}$</td>
<td>1. ↑ Intratubular pressure because of obstruction of tubule or extrarenal urinary system</td>
</tr>
<tr>
<td></td>
<td>Result: ↓ GFR</td>
</tr>
<tr>
<td>$\pi_{GC}$</td>
<td>1. ↑ Systemic-plasma oncotic pressure (sets $\pi_{GC}$ at beginning of glomerular capillaries)</td>
</tr>
<tr>
<td></td>
<td>2. ↓ Renal plasma flow (causes increased rise of $\pi_{GC}$ along glomerular capillaries)</td>
</tr>
<tr>
<td></td>
<td>Result: ↓ GFR</td>
</tr>
</tbody>
</table>

GFR, glomerular filtration rate; $K_f$, filtration coefficient; $P_{GC}$, glomerular-capillary hydraulic pressure; $P_{BC}$, Bowman’s capsule hydraulic pressure; $\pi_{GC}$, glomerular-capillary oncotic pressure. A reversal of all arrows in the table will cause a decrease in the magnitudes of $K_f$, $P_{GC}$, $P_{BC}$, and $\pi_{GC}$. 
cause of decreased GFR in renal disease is not any change in these parameters within individual nephrons but rather simply a decrease in the number of functioning nephrons.

$K_f$
Changes in $K_f$ can be caused by glomerular disease and drugs, but this variable is also subject to normal physiological control by a variety of chemical messengers. The details are still not completely clear, but these messengers cause contraction of glomerular mesangial cells. Such contraction may restrict flow through some of the capillary loops, effectively reducing the area available for filtration and, hence, $K_f$. This decrease in $K_f$ will tend to lower GFR.

$P_{GC}$
Hydrostatic pressure in the glomerular capillaries ($P_{GC}$) is the most complex of the variables in the basic filtration equation because it is itself influenced by so many factors. We can help depict the situation by using the analogy of a leaking garden hose. If pressure feeding the hose (pressure in the pipes leading to the faucet) goes up or down, this directly affects pressure in the hose and, hence, the rate of leak. Resistances in the hose also affect the leak. If we kink the hose upstream from the leak, pressure at the region of leak falls, and less water leaks out. However, if we kink the hose beyond the leak, this raises pressure at the region of leak and increases leak rate. These same principles apply to $P_{GC}$ and GFR. First, a change in renal arterial pressure will cause a change in $P_{GC}$ in the same direction. If resistances remain constant, $P_{GC}$ will rise and fall as renal artery pressure rises and falls. This is a crucial point because a major regulator of renal function is arterial blood pressure. Second, changes in the resistance of the afferent and efferent arterioles have opposite effects on $P_{GC}$. An increase in resistance upstream from the glomerulus in the afferent arteriole (like kinking the hose above the leak) will lower $P_{GC}$, whereas an increase in resistance downstream from the glomerulus in the efferent arteriole (like kinking the hose beyond the leak) will increase $P_{GC}$. In contrast, a decrease in afferent resistance ($R_A$) (resulting from afferent arteriolar dilation) will tend to raise $P_{GC}$. Similarly, a decrease in efferent resistance ($R_E$) (caused by efferent arteriolar dilation) tends to lower $P_{GC}$. It should also be clear that when $R_A$ and $R_E$ both change simultaneously in the same direction (ie, both increase or decrease), they exert opposing effects on $P_{GC}$.

It is possible for both resistances to rise by the same fraction, with the result that there is no effect on $P_{GC}$ (even though, in this case, RBF would fall). In contrast, when they change in different directions, they cause additive effects on $P_{GC}$ (and can have no effect on RBF). The real significance of this is that the kidney can regulate $P_{GC}$ and, hence, GFR independently of RBF. The effect of changes in $R_A$ and $R_E$ are summarized in Figure 2–3.

$P_{BC}$
Changes in this variable generally are of very minor physiological importance. The major pathological cause of increased hydraulic pressure in Bowman’s capsule is obstruction anywhere along the tubule or in the external portions of the urinary
Figure 2-3. Effects of afferent- and/or efferent-arteriolar constriction on glomerular capillary pressure ($P_{GC}$) and renal blood flow (RBF). The RBF changes reflect changes in total renal arteriolar resistance, the location of the change being irrelevant. In contrast, the changes in $P_{GC}$ are reflected in which set of arterioles the altered resistance occurs. Pure afferent constriction lowers both $P_{GC}$ and RBF, whereas pure efferent constriction raises $P_{GC}$ and lowers RBF. Simultaneous constriction of both afferent and efferent arterioles has counteracting effects on $P_{GC}$ but additive effects on RBF; the effect on $P_{GC}$ may be a small increase, small decrease, or no change. Vasodilation of only 1 set of arterioles would have effects on $P_{GC}$ and RBF opposite those shown in parts B and C. Vasodilation of both sets would cause little or no change in $P_{GC}$, the same result as constriction of both sets but would cause a large increase in RBF. Constriction of 1 set of arterioles and dilation of the other would have maximal effects on $P_{GC}$ but little effect on RBF.
Oncotic pressure in the plasma at the very beginning of the glomerular capillaries is, of course, simply the oncotic pressure of systemic arterial plasma. Accordingly, a decrease in arterial plasma protein concentration, as occurs, eg, in liver disease, will lower arterial oncotic pressure and tend to increase GFR, whereas increased arterial oncotic pressure will tend to reduce GFR.

However, now recall that $\pi_{GC}$ is identical to arterial oncotic pressure only at the very beginning of the glomerular capillaries; $\pi_{GC}$ then progressively increases along the glomerular capillaries as protein-free fluid filters out of the capillary, concentrating the protein left behind. This means that net filtration pressure and, hence, filtration progressively decrease along the capillary length. Accordingly, anything that causes a steeper rise in $\pi_{GC}$ will tend to lower average net filtration pressure and hence GFR.

This steep increase in oncotic pressure tends to occur when RPF is low. It should not be hard to visualize that the filtration of a given volume of fluid from a small total volume of plasma flowing through the glomeruli will cause the protein left behind to become more concentrated than if the total volume of plasma were large. In other words, a low RPF, all other factors remaining constant, will cause the $\pi_{GC}$ to rise more steeply and reach a final value at the end of the glomerular capillaries that is higher than normal. This increase in average $\pi_{GC}$ along the capillaries lowers average net filtration pressure and, hence, GFR. Conversely, a high RPF, all other factors remaining constant, will cause $\pi_{GC}$ to rise less steeply and reach a final value at the end of the capillaries that is less than normal, which will increase the GFR.

Another way of thinking about this is in terms of filtration fraction: the ratio GFR/RPF. The increase in $\pi_{GC}$ along the glomerular capillaries is directly proportional to the filtration fraction (ie, the more volume that is filtered from plasma, the higher is the rise in $\pi_{GC}$). Therefore, if you know that filtration fraction has changed, you can be certain that there has also been a proportional change in $\pi_{GC}$ and that this has played a role in altering GFR.

**Filtered Load**

A term we use in other chapters is *filtered load*. It is the amount of substance that is filtered per unit time. For freely filtered substances, the filtered load is just the product of GFR and plasma concentration. Consider sodium. Its normal plasma concentration is 140 mEq/L, or 0.14 mEq/mL. (Note: 1 mEq of sodium is 1 mmol.) A normal GFR is 125 mL/min, so the filtered load of sodium is 0.14 mEq/mL × 125 mL/min = 17.5 mEq/min. We can do the same calculation for any other substance, being careful in each case to be aware of the unit of measure in which concentration is expressed. The filtered load is what is presented to the rest of the nephron to handle. A high filtered load means a substantial amount of
material to be reabsorbed. The filtered load varies with plasma concentration and GFR. A rise in GFR, at constant plasma concentration, increases the filtered load, as does a rise in plasma concentration at constant GFR.

**AUTOREGULATION**

It is extremely important for the kidneys to keep the GFR at a level appropriate for the body because, as we have emphasized, the excretion of salt and water is strongly influenced by the GFR. We have also emphasized that the GFR is strongly influenced by renal arterial pressure. A rise in blood pressure causes an increased excretion of salt and water, a process called *pressure natriuresis* (see Chapter 7), whereas a fall in blood pressure diminishes excretion. These changes in excretion are mediated partly via changes in GFR. The effect is so strong that urinary excretion would tend to vary widely with the ordinary daily excursions of arterial pressure. Also, vascular pressure in the thin-walled glomerular capillaries is higher than in capillaries elsewhere in the body and hypertensive damage ensues if this pressure is too high.

Therefore, both to protect the glomerular capillaries from hypertensive damage and to preserve a healthy GFR, changes in GFR and RBF are severely blunted by mechanisms that we collectively call *autoregulation*. Consider first a situation in which mean arterial pressure rises 20%. Such a modest rise occurs many times throughout the day in association with changes in excitement level and activity. Pretend, for the moment, that all renal vascular resistances remain constant. By the basic flow equation \( Q = \Delta P/R \), RBF would rise 20% also (actually slightly more if pressure in the renal vein is unaffected). What would this do to GFR? It would rise much more than 20%, in fact almost 50%. This is because net filtration pressure would rise almost 50%. In effect, fractional changes in upstream pressure (in the renal artery) are *magnified* in terms of net filtration pressure. Why is this? At the beginning of the glomerulus, capillary hydrostatic pressure is about 60 mm Hg and the pressures opposing filtration sum to 36 mm Hg, yielding a net filtration pressure of about 24 mm Hg (Table 2–1). With an increase in arterial pressure to 120 mm Hg, capillary pressure would rise to about 71 mm Hg, but there would be no increase in the pressures that oppose filtration—plasma oncotic pressure and Bowman’s capsule pressure. Therefore, net filtration pressure would rise to 71 – 36 = 35 mm Hg (an increase of almost 50%). The higher net filtration pressure would cause a parallel increase in GFR. (In turn, this would raise plasma oncotic pressure at the *distal* end of the glomerulus, tending to reduce filtration somewhat, but the total effect is still a major rise in GFR.) This emphasizes the crucial role of glomerular capillary pressure in glomerular filtration.

Now, what actually happens in the face of changes in mean arterial pressure? As is the case in many organs, blood flow does not change in proportion to changes in arterial pressure. The changes are blunted. A rise in driving pressure is counteracted by a rise in vascular resistance that *almost* offsets the rise in pressure. The word “almost” is crucial here. Higher driving pressures do indeed lead to higher flow but not proportionally. Consider Figure 2–4. Within the range of mean arterial pressures commonly found in the human body (between the dashed
vertical lines in Figure 2–4), RBF varies only modestly when mean arterial pressure changes. This is partly a result of a direct reaction of the vascular smooth muscle to stretch or relaxation—or the myogenic response—and partly the result of intrarenal signals that we describe shortly. The myogenic response is very fast-acting and protects the glomeruli from short-term fluctuations in blood pressure. In addition to keeping changes in RBF fairly small, autoregulatory processes also keep changes in GFR fairly small. Again, GFR does rise with an increase in arterial pressure, just not substantially.

How do the intrarenal processes work? Much is by way of a process with the clumsy name of tubuloglomerular feedback. Tubuloglomerular feedback is feedback from the tubules back to the glomerulus (ie, an influence of events in the tubules that is exerted on events in the glomeruli). We will return to the mechanism in Chapter 7, but for now the essence of tubuloglomerular feedback can be summarized as follows: As the filtration rate in an individual nephron increases or decreases, the amount of sodium that escapes reabsorption in the proximal tubule and the loop of Henle also increases or decreases. More sodium filtered means more sodium remaining in the lumen of the nephron and more sodium flowing from the thick ascending limb into the distal tubule. Recall that at the division between these nephron segments lies the macula densa, a special group of cells in the nephron wall where the nephron passes between the afferent and efferent arterioles (see Figure 1–7). The macula densa cells sense the amount of sodium and chloride in the lumen. They act, in part, as salt detectors. One result of changing levels of luminal sodium chloride is to increase or decrease the

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*Figure 2–4.* Autoregulation of renal blood flow (RBF). A similar pattern holds for glomerular filtration rate.
secretion of transmitter agents into the interstitial space that affect the filtration in the nearby glomerulus. High levels of sodium flowing past the macula densa cause a decrease in filtration rate; low levels of sodium flowing past allow a higher filtration rate. It is as though each nephron adjusts its filtration so that the right amount of sodium remains in the lumen to flow past the macula densa. How can it adjust its filtration? The transmitter agents released by the salt-sensing macula densa cells produce vasoconstriction of the afferent arteriole, thereby reducing hydrostatic pressure in the glomerular capillaries. These same agents also produce contraction of glomerular mesangial cells, thereby reducing the effective filtration coefficient. Both processes reduce the single-nephron filtration rate and keep it at a level appropriate for the rest of the nephron.

In conclusion, we emphasize that autoregulation blunts or lowers the RBF and GFR responses to changes in arterial pressure but does not totally prevent those changes.

**KEY CONCEPTS**

1. **RBF** is much higher than required for metabolic needs and is regulated for functional reasons, not metabolic demand.

2. **RBF** varies with driving pressure and varies inversely with the sum of resistances in the renal vasculature.

3. **GFR** varies with the net filtration pressure and capillary filtration coefficient.

4. Net filtration pressure is determined by renal artery pressure, resistances in the afferent and efferent arterioles, and plasma oncotic pressure.

5. The kidney has autoregulatory mechanisms that blunt changes in blood flow and GFR in response to changes in renal artery pressure.

**STUDY QUESTIONS**

2–1. A substance \( X \) is known to be freely filtered. Therefore, all of the \( X \) that enters the glomerular capillaries is filtered. True or false?

2–2. The concentration of glucose in plasma is 100 mg/dL, and the GFR is 125 mL/min. How much glucose is filtered per minute?

2–3. The concentration of calcium in Bowman’s capsule is 3 mEq/L, whereas its plasma concentration is 5 mEq/L. Why are the concentrations different?
2–4. A protein has a molecular weight of 30,000 and a plasma concentration of 100 mg/L. The GFR is 100 L/day. How much of this protein is filtered per day?

2–5. A drug is noted to cause a decrease in GFR. Identify 4 possible actions of the drug that might decrease GFR.

2–6. A drug is noted to cause an increase in GFR with no change in net filtration pressure. What might the drug be doing?

2–7. A person is given a drug that dilates the afferent arteriole and constricts the efferent arteriole by the same amounts. Assuming no other actions of the drug, what happens to this person’s GFR, RBF, and filtration fraction?

2–8. A clamp around the renal artery is partially tightened to reduce renal arterial pressure from a mean of 120 to 80 mm Hg. How much do you predict RBF will change?
A. 33% decrease
B. No change
C. 5–10% decrease
D. 33% increase
Clearance

OBJECTIVES

The student understands the principles and applications of clearance technique.

- Defines the terms clearance and metabolic clearance rate, and differentiates between general clearance and renal clearance.
- Lists the information required to calculate clearance.
- States the criteria that must be met for a substance so that its clearance can be used as a measure of glomerular filtration rate; states which substances are used to measure glomerular filtration rate and effective renal plasma flow.
- Given data, calculates $C_{\text{in}}$, $C_{\text{PAH}}$, $C_{\text{urea}}$, $C_{\text{glucose}}$, $C_{\text{Na}}$.
- Predicts whether a substance undergoes net reabsorption or net secretion by comparing its clearance with that of inulin or by comparing its rate of filtration with its rate of excretion.
- Given data, calculates net rate of reabsorption or secretion for any substance.
- Given data, calculates fractional excretion of any substance.
- Describes how to estimate glomerular filtration rate from $C_{\text{Cr}}$ and describes the limitations.
- Describes how to use plasma concentrations of urea and creatinine as indicators of changes in glomerular filtration rate.

One of the crucial functions of the kidneys is to remove metabolic wastes and excess amounts of ingested substances from the blood. Nitrogenous wastes such as urea and creatinine are generated by metabolism and removed from the body by the kidneys. The average excretion rate normally reflects the rate at which these substances are added to the blood by metabolic processes, thus keeping the body in balance for these substances. Other substances, those that are crucial for normal function, including sodium and potassium, enter the body by ingestion and are also removed from the body by the kidneys. These excretion rates parallel ingestion rate in the long term but are altered transiently to reflect other needs, such as the regulation of blood pressure or plasma osmolality.

Ridding the body of a substance is often called clearance. This term in a biomedical context has both a general meaning and a specific renal meaning. The general meaning of clearance is simply that a substance is removed from the blood by any of several mechanisms. For instance, a drug may be cleared...
by excretion in the urine or the feces, or it may be transformed by the liver and other peripheral tissues to an inactive form. Renal clearance, on the other hand, means that the substance is removed from the blood and excreted in the urine.

The rate at which something is cleared can be expressed in several different ways. The most obvious is excretion rate. If a substance leaves the body at a certain amount per hour, this is one way to quantify its clearance. The units of excretion rate are the amount leaving the body per time. Another way to quantify the clearance of a substance is by the substance’s plasma half-life. The time it takes for the plasma concentration to fall to half of its current value is the plasma half-life, or $t_{1/2}$. The units are time. Still another way, and the one we develop here, places a more specific meaning on the word clearance. Both general clearance and specific renal clearance can be expressed as the volume of plasma per unit time from which all of a substance is removed.

**CLEARANCE UNITS**

The units of clearance are often confusing to the first-time reader, so let us be sure of the meaning. First, the units are volume per time (not amount per time). The easiest way to think of this is to ask what volume of plasma contains the amount excreted in a given time. Suppose 5 mg of a substance is excreted per hour and 200 mL of plasma contains 5 mg. Then the clearance of the substance is 200 mL/h, i.e., 200 mL of volume has been completely cleared of the substance. Again, note the units: volume per time (i.e., the volume of plasma from which a substance is completely removed [cleared]).

The general meaning and specific renal meaning of clearance can be illustrated by comparing how the body handles 2 substances with similar-sounding names but very different properties: insulin and inulin. **Insulin** is the familiar pancreatic hormone involved in regulating blood sugar (glucose). It is a protein with a molecular weight of 5.8 kDa and is small enough to be freely filtered by the glomerulus. Once in Bowman’s space, it moves along with every other filtered substance into the proximal convoluted tubule, where it is largely taken up by endocytosis and degraded into its constituent amino acids. Very little insulin escapes this uptake, and very little of the filtered insulin survives to be excreted in the urine. Thus, the kidney takes part in clearing insulin from the blood; however, because so little appears in the urine, the specific renal clearance is very low (<1 mL/min). However, the body has additional mechanisms for clearing insulin, and its metabolic clearance rate is quite high (half-life less than 10 min). Let us contrast this with inulin. **Inulin** is a polysaccharide starch of about 5 kDa molecular weight. Like insulin, it is freely filtered by the glomerulus, but it is not reabsorbed or secreted by the nephron. All the inulin that is filtered flows through the nephron and appears in the urine. Thus, inulin’s renal clearance is relatively large. Inulin in the blood is not taken up by other tissues, and the kidneys are the only excretion route. As we will see, this makes inulin a very special substance with respect to assessing renal function.
Quantification of Clearance

Consider again a substance X that is excreted in the urine. How do we actually calculate clearance in the proper units? The amount excreted exists in some volume of urine, and that amount must have been contained in some volume of plasma. This is depicted in Figure 3–1. The large boxes on the left show the plasma before, during, and after clearance of substance X (dots), while the smaller boxes on the right show the urine gaining X as it is removed from the plasma. The *amount* cleared per unit time is the product of the volume of plasma cleared per unit time (C_x) and the plasma concentration (P_x). Amount cleared = C_x \cdot P_x. The amount appearing in the urine during this time is the product of the urine flow rate (V) and the urine concentration of X (U_x). Amount in urine = V \cdot U_x. The amount removed from the plasma must equal the amount appearing in the urine. This equality is shown in Equations 3-1 and 3-2. Finally, by rearrangement we solve for clearance (C_x) as shown in Equation 3-3. Thus we have equated amounts per unit time, but by rearrangement we end up with clearance in its proper units—volume per time.

\[
\begin{align*}
\text{amount in plasma} &= C_x \cdot P_x \\
\text{amount in urine} &= V \cdot U_x
\end{align*}
\]

\[
\begin{align*}
C_x \cdot P_x &= V \cdot U_x \\
C_x &= \frac{V \cdot U_x}{P_x}
\end{align*}
\]

*Figure 3–1. Derivation of the basic clearance formula. Substance X (dots) is removed from the plasma (large boxes) and excreted in the urine (small boxes). The amount removed from the plasma in a given time is matched to the amount excreted in the urine in that same time. By rearrangement (Equation 3-3) we solve for the volume of plasma containing the amount of X excreted per unit time, that is, the renal clearance of X. C_x = clearance of X; P_x = plasma concentration of X; V = urine flow rate; U_x = urine concentration of X.*
While we are addressing the quantification of clearance, note that the product of urine flow rate and urine concentration (numerator on the right-hand side of Equation 3-3) is excretion rate. Therefore, we can also state that the clearance of substance X is the excretion rate divided by the plasma concentration.

Let us now examine the clearance of several substances that are important for the quantification of renal function, starting with inulin. Inulin, as described previously, is a polysaccharide that is freely filtered and neither reabsorbed nor secreted. All that is filtered is excreted. Therefore, the volume of plasma cleared per unit time is the same as the glomerular filtration rate (GFR) (Figure 3–2). Inulin clearance is indeed the hallmark method for measuring GFR. Why is inulin so good in this regard? The first property is its filterability. It moves into Bowman’s space in the same proportion as the volume filtered. Second, it cannot move in either direction by the paracellular route around the tubular epithelium. The tight junctions are too restrictive to permit saccharides of any sort to move through them. Third, there are no transport mechanisms either on the apical or basolateral surface of the tubular epithelium to take up inulin. Finally, there are no enzymes (amylases) present in the tubular lumen to break down inulin. Thus, it is freely filtered, and all that is filtered moves through the nephron into the urine.

Can something have a clearance greater than the GFR? Indeed, yes. One such substance is para-aminohippurate (PAH).

This is a small (molecular weight of 194 Da) water-soluble organic anion that is freely filtered and also avidly secreted by the proximal tubule epithelium (via the transcellular route). The secretion rate is saturable. (That is, there is a maximum rate of PAH secretion into the tubule. Such tubular
maximum, or \( T_m \), transport systems are common; see Chapter 4.) However, at low
plasma concentrations, about 90% of the PAH entering the kidney is removed
from the plasma and excreted in the urine. Its clearance, therefore, is nearly as
great as the renal plasma flow. In fact, the PAH clearance is used as a measure of
renal plasma flow, usually called the **effective renal plasma flow** to indicate that its
value is slightly less than the true renal plasma flow.

So far, the freely filtered substances we have discussed have clearance values
lying between the GFR and the renal plasma flow. Can a freely filtered substance
have a clearance value less than the GFR? Yes. In fact, many freely filtered sub-
stances have clearance values of zero! If a filtered substance is completely removed
from the nephron by reabsorption or degradation, then none appears in the urine,
and the clearance is zero. We have already seen one example of this: insulin, which
is all degraded. Another is glucose, which is normally all reabsorbed.

Many freely filtered substances have clearance values less than the GFR but
greater than zero (eg, sodium, chloride, and urea). On occasion, some freely fil-
tered substances have a clearance greater than the GFR (eg, PAH and potassium).
What can the clearance of a substance tell us? If we know the GFR (as assessed
from inulin clearance) and the clearance of a given substance, then any difference
between clearance and GFR represents net secretion or reabsorption (or, in a few
rare cases, renal synthesis). If the clearance of a substance exactly equals the GFR,
then there has been no **net** reabsorption or secretion. If the clearance is greater
than the GFR, there must have been net reabsorption. Finally, if the clearance is
less than the GFR, there must have been net reabsorption. The word **net** in this
description is important. As we will see, a number of substances are reabsorbed
in certain regions of the nephron and secreted in other regions. The net result of
these processes is the sum of everything that happens along the nephron.

To illustrate these concepts, consider potassium clearance. Potassium is a small,
freely filtered ion. About 70% of the filtered load is reabsorbed in the proximal
tubule, and another 20% or so is reabsorbed in the loop of Henle. The rest of the
nephron has mechanisms to both secrete and reabsorb potassium. Under some con-
ditions (eg, when potassium ingestion is low), reabsorption dominates, reducing the
excretion of potassium and thus resulting in a clearance less than the GFR. In other
conditions (eg, after ingesting large amounts of potassium in citrus fruit), secretion
dominates. In this case, not only is the previously reabsorbed amount secreted back
into the tubule, but additional potassium is secreted. The result is that more potas-
sium is excreted than is filtered, and potassium clearance is greater than the GFR.

**A Practical Method of Measuring GFR: Creatinine Clearance**

The gold standard for measuring GFR is the inulin clearance, as described
previously, and this method is used in research studies and some clinical
situations when a very accurate value is needed. The method is cumber-
some, however, because inulin must be infused, and it must be infused at a rate
sufficient to keep its plasma concentration constant during the period of urine for-
mation and collection. If GFR is normal, between 2.5% and 3.5% of plasma inulin
is removed every minute and must be replaced by infusion if GFR is to be
accurately determined. For routine assessment of GFR in hospitalized patients, there is an easier method: creatinine clearance. Creatinine is an end product of creatine metabolism and is exported into our blood continuously by skeletal muscle. The rate is proportional to skeletal muscle mass, and to the extent that muscle mass is constant in a given individual, the creatinine production is constant. Creatinine is freely filtered and not reabsorbed. A small amount, however, is secreted by the proximal tubule. Therefore, the creatinine appearing in the urine represents both a filtered component and a secreted component. Because of the secretion, creatinine clearance is slightly higher than the GFR. The secreted fraction is normally about 10–20% of what is excreted, so the measured creatinine clearance overestimates GFR by the same percentage. For routine assessment of GFR, this degree of error is acceptable. How does one measure creatinine clearance? Usually, a patient’s urine is collected for 24 h, and a blood sample is taken sometime during the collection period. Blood and urine are assayed for creatinine concentration, and we apply the clearance formula (Figure 3–1, Equation 3-3) to yield creatinine clearance. Additional errors cloud the issue (eg, errors in the assays for plasma and urine creatinine concentration or drug-induced alteration of creatinine secretion), so the method is not perfect. For a patient with a very low GFR, the secreted component is a relatively larger fraction of the total amount excreted; therefore, the creatinine clearance more severely overestimates GFR in patients with a very low GFR than in those with normal GFR values. Nevertheless, because of low cost and convenience, creatinine clearance continues to be the most common method for routine assessment of patient GFR and the integrity of renal filtration.

PLASMA CREATININE AND UREA CONCENTRATIONS AS INDICATORS OF GFR CHANGES

Although creatinine clearance is a valuable clinical measure of GFR, in practice it is far more common to measure plasma creatinine alone and to use this as an indicator of GFR. This approach is justified by the fact that most excreted creatinine gains entry to the tubule by filtration. If we ignore the small amount secreted, there should be an excellent inverse correlation between plasma creatinine concentration and GFR (Figure 3–3).

A normal person’s plasma creatinine concentration is about 1 mg/dL. It remains stable because each day the amount of creatinine excreted is equal to the amount of creatinine produced. Suppose one day an individual’s GFR suddenly decreases by 50% because of a blood clot in the renal artery. On that day the person filters only 50% as much creatinine as normal so that creatinine excretion is also reduced by 50%. (We are ignoring the small contribution of secreted creatinine.) Therefore, assuming no change in creatinine production, the person transiently goes into positive creatinine balance, and the plasma creatinine rises. However, despite the persistent 50% GFR reduction, the plasma creatinine does not continue to rise indefinitely; rather, it stabilizes at 2 mg/dL (ie, after it has doubled). At this point, the person once again is able to excrete creatinine at the normal rate and so goes back in balance with a stable plasma level. The reason is that the 50% GFR reduction has been just offset by the doubling of plasma creatinine
concentration, restoring the filtered load of creatinine to normal. To understand this point, assume an original daily filtration volume of 180 L (1800 dL).

Original normal state:
Filtered creatinine = 1 mg/dL × 1800 dL/day = 1800 mg/day

New steady-steady state:
Filtered creatinine = 2 mg/dL × 900 dL/day = 1800 mg/day

This is a very important point: In the new steady state, creatinine excretion is normal (the person is in balance) because of the doubling of plasma creatinine concentration. In other words, creatinine excretion is below normal only transiently until plasma creatinine has increased as much proportionally as the GFR has fallen.

What if the GFR then fell to 300 dL/day? Again, creatinine retention would occur until a new steady state had been established (ie, until the person was again filtering 1800 mg/day). What would the new plasma creatinine be?

\[ 1800 \text{ mg/day} = P_{Cr} \times 300 \text{ dL/day} \]
\[ P_{Cr} = 6 \text{ mg/dL} \]

The rise in plasma creatinine results directly from the fall in GFR. Therefore, a single plasma creatinine measurement is a reasonable indicator of GFR. It is

**Figure 3–3.** Steady-state relationship between GFR and plasma creatine, assuming no creatinine is secreted.
not completely accurate, however, for several reasons: (1) As before, some creatinine is secreted; (2) there is no way of knowing exactly what the person’s original creatinine was when GFR was normal; (3) creatinine production may not remain completely unchanged. However, a rising plasma creatinine is a red flag that there may be a renal problem.

Because urea is also handled by filtration, the same type of analysis suggests that the measurement of plasma urea concentration could also serve as an indicator of GFR. However, it is a much less accurate indicator than plasma creatinine because the range of normal plasma urea concentration varies widely, depending on protein intake and changes in tissue catabolism, and because urea excretion is under partial hormonal regulation.

**KEY CONCEPTS**

1. Clearance has both a general meaning, describing loss of material from the body, and a specific renal meaning involving the kidney’s ability to remove substances from the blood. Renal clearance is always expressed in units of volume per time.

2. Renal clearance of any substance X is quantified by a general clearance formula relating urine flow to urine and plasma concentrations:

   \[ C_x = \frac{U_x}{P_x} \cdot V \]

   *Inulin clearance is used to measure GFR because inulin is freely filtered and neither secreted nor reabsorbed.*

3. Para-aminohippurate clearance is used as a practical estimate of renal blood flow.

4. Creatinine clearance is used as a practical estimate of GFR.

**STUDY QUESTIONS**

3–1. The hospital lab reports that your patient’s renal creatinine clearance is 120 g/day. This value is

   A. Normal
   
   B. Significantly above normal
   
   C. Not an interpretable number as presented
3–2. A substance is known to be cleared from the body both by renal excretion and by nonrenal mechanisms. Is the renal clearance higher, lower, or the same as the metabolic clearance rate?

3–3. Inulin clearance is measured twice; the first time at a low inulin infusion rate, and the second time at a higher infusion rate that results in a higher plasma inulin concentration during the test. Assuming the kidneys behave the same in both cases, which measurement will yield a higher inulin clearance?
   
   A. The first
   B. The second
   C. Both measurements are the same.

3–4. The clearance of substance A is less than inulin clearance. Give 3 possible explanations.
OBJECTIVES

The student understands the basic mechanisms of tubular reabsorption and secretion:

- Defines and states the major characteristics of diffusion, uniport (facilitated diffusion), primary active transport, secondary active transport (including symport and antiport), and endocytosis.
- Defines channel gating and states three common means of gating channels.
- Describes the major morphological components of an epithelial tissue, including lumen, interstitium, apical and basolateral membranes, tight junctions, and lateral spaces.
- States how transport mechanisms can be combined to achieve active transcellular reabsorption in epithelial tissues.
- Defines paracellular transport and differentiates between transcellular and paracellular transport.
- Defines osmolality and osmolarity, and states why osmolarity is commonly used to approximate osmolality.
- Describes what is meant by the expression “water follows the osmoles.”
- Describes qualitatively the forces that determine movement of reabsorbed fluid from the interstitium into peritubular capillaries.
- Compares the Starling forces governing glomerular filtration with those governing peritubular capillary absorption.
- Compares and contrasts the concepts of Tm and gradient-limited transport.
- States the consequences of pump-leak systems.
- Contrasts “tight” and “leaky” epithelia.

CROSSING THE EPITHELIAL BARRIERS

The basic processes of moving substances between blood and tubular lumen (secretion and reabsorption) require that solutes and water cross two cell layers: the tubular epithelium and the vascular endothelium, plus a thin region of interstitial fluid between them. In the cortex, where the fluxes of many filtered substances are enormous, the vascular endothelium (peritubular capillaries) is fenestrated. The fenestrae and the loose underlying basement membrane offer virtually no resistance to the passive movement of water and small solutes. This facile permeation has two consequences. First, overall transport is governed
almost exclusively by events in the tubular epithelium rather than the vascular endo-
thelium; second, the cortical interstitium, which is the medium faced by the
basolateral membranes of the tubular epithelia, has an osmolality and concentra-
tion of small solutes close to those in plasma. The interstitial composition changes
when plasma composition changes. In the medulla, where both blood flow and
transport events are quantitatively lower, things are far more complicated. Only
some regions of the vasculature are fenestrated, so that (1) the overall transport
depends on both the properties of the vascular endothelium and tubular epithe-
lium, and (2) the medullary interstitium is most definitely not plasma-like in its
composition. We will address the importance of these medullary properties later.
For now, we direct our attention in the rest of this chapter to epithelial transport.

Crossing the tubular epithelium can be performed in a single step or two
steps. The paracellular route (single step) is when the substance goes around
the cells (ie, through the matrix of the tight junctions that link each epi-
thelial cell to its neighbor). More often, however, a substance goes through the cells,
a two-step process: across the apical membrane facing the tubular lumen and across
the basolateral membrane facing the interstitium. This is called the transcellular
route. These structures and pathways are depicted in Figures 4–1A and B.

An array of mechanisms exists by which substances cross the various barriers.
The general classes of mechanisms are no different from those used elsewhere
in the body to transport substances across cell membranes. We can view these
mechanisms as a physiological tool box. Renal cells use whichever set of tools is
most suitable for the task. The general classes of mechanisms for traversing the
barriers are depicted in Figure 4–2.

Movement by Diffusion

Diffusion is the frenzied random movement of free molecules in solution (like
the Ping-Pong balls in a lottery drawing). Net diffusion occurs across a barrier (ie,
more molecules moving one way than the other) if there is driving force (a concen-
tration gradient or, for charged molecules, a potential gradient) and if the barrier
is permeable. This applies to almost all substances crossing the endothelial barrier
lining the peritubular capillaries. It applies to substances taking the paracellular
route around the tubular epithelium and to some substances taking the transcellu-
lar route through membranes. Substances that are lipid soluble, such as the blood
gases or steroids, can diffuse directly through the lipid bilayer.

Movement Through Channels

Most substances that are biologically important cannot penetrate lipid
membranes. To cross a membrane, they must move through specific integ-
ral membrane proteins, which are divided into categories of channels and
transporters (see Figure 4–2). Channels are small pores (proteins with a “channel”
or pathway through the interior of the protein) that permit, depending on their
structure, water or specific solutes to diffuse through them. Thus, we use the terms
Figure 4–1. A, Transcellular and paracellular reabsorption. Transcellular reabsorption involves separate influx and efflux steps, in most cases utilizing transporters or channels. Paracellular transport is always a passive process through the tight junctions, driven by electrochemical gradients between the lumen and interstitium. B, Diagrammatic representation of tubular epithelium. The tight junctions can be visualized three dimensionally as the sheet of plastic holding together a 6-pack of soda, each cell being one of the cans.
sodium channel and potassium channel to designate channels that permit diffusion of these molecular species. Aquaporins are channels selectively permeable to water. Channels typically flicker open and close like camera shutters so that the permeability of a membrane containing many channels is proportional to the probability of their being open. Movement through channels is passive (ie, no external energy is required). The energy to drive the diffusion is inherent in the concentration gradient or, strictly speaking, the electrochemical gradient, because charged ions are driven through channels and around cells via the paracellular route not only by gradients of concentration but also by gradients of voltage. Channels represent a mechanism for rapidly moving across membranes large amounts of specific substances, which would otherwise diffuse slowly or not at all.

A characteristic of channels critical for renal function is the regulation of their permeability by a number of environmental factors and signaling cascades (Figure 4–3). First, many channel types can be gated, meaning that the probability that the channel is open can be increased or decreased. The topic of channel gating

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\[ \text{ADP} + \text{Pi} \rightarrow \text{ATP} \]

**Figure 4–2.** Basic mechanisms of transmembrane solute transport. See text for details.

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\[ ^1 \text{In tissues where movement of dissolved gas (CO}_2\text{ and O}_2\text{) is quite rapid, a substantial amount of these gases may move through aquaporins.} \]
is a whole story by itself, but several ways of gating channels include (1) reversible binding of small molecules that are components of signaling cascades (ligand-gated channels), (2) changes in membrane potential (voltage-gated channels), and (3) mechanical distortion (stretch-gated channels). Second, many channel types have phosphorylation sites such that phosphorylation either locks the channel shut or allows it to be gated by one of the mechanisms above. Also, some channel species can be moved back and forth between the surface membrane and intracellular vesicles, thereby regulating how many of the existing channels are actually functioning as permeability pathways. Third, and on a slower time scale, the genomic expression of channels is regulated so that the total number of channels, whether in the membrane or sequestered in vesicles, is altered up or down.

**Movement by Transporters**

Our genome codes for a large array of proteins that function as transporters, all with names and acronyms that suffuse the physiological literature. Transporters, like channels, permit the transmembrane flux of a solute that is otherwise impermeable in the lipid bilayer. Channels can move large amounts of materials across membranes in a short period of time, but many transporters have a lower rate of transport because the transported solutes bind much more strongly to the transport protein. Furthermore, the protein must undergo a more elaborate cycle of conformational change to move the solute from one side of the membrane to the other. We can group transporters into categories according to basic functional properties. As is the case for channels, the amount of substance moved via transporters is highly regulated. The regulation includes changes in phosphorylation of

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**Figure 4–3.** Regulation of channel permeability. (1) Existing channels may be gated to an open configuration by (a) ligand binding, (b) changes in membrane potential, or by (c) mechanical deformation. (2) Existing channels can be phosphorylated, which either locks them closed or permits them to be gated. (3) Channels may be shuttled back and forth between the surface membrane and intracellular storage vesicles.
the transporter (thereby turning its activity on or off), sequestration into vesicles, and of course changes in genomic expression.

**Uniporters**

Uniporters permit the movement of a single solute species through the membrane. The basic difference between a channel and a uniporter is that a channel is a tiny hole, whereas a uniporter requires the solute to bind to a site that is alternately available to one side and then the other side of the membrane (like entering a vestibule through an outside door and then leaving the vestibule to enter a hallway through an inside door). Movement through a uniporter is often called facilitated diffusion because, like diffusion, it is driven by concentration gradients, but the transported material moves through the uniporter protein rather than the membrane. A set of uniporters crucial for all cells includes those that facilitate the movement of glucose across cell membranes. These are members of the GLUT family of proteins that permit, in the kidney’s proximal tubule epithelial cells, glucose to move from the cytosol across the basolateral membrane into the interstitium.

**Symporters and Antiporters**

Symporters and antiporters move 2 or more solute species in the same direction across a membrane (symporters) or in opposite directions across a membrane (antiporters). In the literature, transport by a symporter is sometimes called cotransport, whereas transport by an antiporter is called exchange or counter transport. Thus, there are symporters that move sodium and glucose together into cells (members of the SGLT protein family) and symporters that move sodium, potassium, and chloride all together into a cell. In the case of the SGLT family, each transport cycle moves 1 glucose molecule and either 1 or 2 sodium ions depending on the particular SGLT. There are antiporters that move sodium into a cell and protons out of a cell (often called sodium-hydrogen exchangers, members of the NHE protein family). Another key antiporter in many cells, including the kidney, moves chloride in one direction and bicarbonate in the opposite direction.

All molecular transport requires energy. In the case of diffusion through a channel or movement via a uniporter, the energy is inherent in the electrochemical gradient for the solute. With symporters and antiporters, at least 1 of the solutes moves down its electrochemical gradient and provides the energy to move 1 or more of the other solutes up its electrochemical gradient. Movement of any solute up its electrochemical gradient is called active transport. In the case of symporters and antiporters that do not hydrolyze adenosine triphosphate (ATP), the active transport is called secondary active transport because the energy is provided indirectly from the transport of another solute rather than directly from a chemical reaction. In a large number of cases, sodium is one of the solutes moved by a symporter or antiporter to provide energy. The energetics of sodium distribution always favors entrance (eg, if a transmembrane sodium-permeable pathway exists, sodium will enter, not leave, the cell). When sodium movement is coupled with that of another solute, as in sodium-proton antiport (exchange), sodium will enter passively, and the other solute will move in the opposite direction actively if the energy obtained from moving sodium down its electrochemical gradient is greater than that required.
to move the other solute up its gradient. The stochiometry is important here. The energy available from a gradient is multiplied by the number of molecules that move per transport cycle. For example, some SGLT proteins move 2 sodium ions per transport cycle; others move only 1. There is more energy available to actively transport glucose through an SGLT protein that moves 2 sodium ions than one that transports 1 sodium ion per glucose. Another example is the coupled transport of bicarbonate and sodium. An important symporter in the proximal tubule is a so-called NBCe transporter, which moves 3 bicarbonate ions and 1 sodium ion per transport cycle. The electrochemical gradient for bicarbonate is directly outward, and the energy gained from moving 3 bicarbonate ions outward is greater than the energy it takes to move 1 sodium ion outward. Therefore, this transporter moves these solutes outward, up the electrochemical gradient for sodium.

**Primary Active Transporters**

Primary active transporters are membrane proteins that are capable of moving 1 or more solutes up their electrochemical gradients, using the energy obtained from the hydrolysis of ATP. All transporters that move solutes in this manner are ATPases (ie, their structure is both that of an enzyme that splits ATP and a transporter that has binding sites that alternately are open to one side and then the other side of the membrane). Among the key primary active transporters in the kidney is the ubiquitous Na-K-ATPase (often called the “sodium pump”), some isoform of which is present in virtually all cells of the body. This transporter simultaneously moves sodium against its electrochemical gradient out of a cell and potassium against its gradient into a cell. The stochiometry is 3 sodium ions out and 2 potassium ions for every ATP molecule hydrolyzed. Other crucial primary active transport systems are a set of H-ATPases, which move protons out of cells, and Ca-ATPases, which move calcium out of cells. All of these ATPases belong to a large family of homologous transporter proteins.

**Receptor-Mediated Endocytosis and Transcytosis**

Almost all the secretion and reabsorption of solutes discussed throughout this textbook use some combination of the above-mentioned set of membrane permeability mechanisms. One other solute transport process of some importance is receptor-mediated endocytosis. In this case, a solute, usually a protein, binds to a site on the apical surface of an epithelial cell, and then a patch of membrane with the solute bound to it is internalized as a vesicle in the cytoplasm. Subsequent processes then degrade the protein into its constituent amino acids, which are transported across the basolateral membrane and into the blood.

For a few proteins, particularly immunoglobulins, endocytosis can occur at either the apical or basolateral membranes, after which the endocytic vesicles remain intact and are transported to the opposite cellular membrane, where they undergo exocytosis to release the protein intact. Such transcytosis is very important in the host defense mechanisms of the kidney and in the prevention of urinary tract infections.

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2An exception is the macula densa cell, which removes sodium via a different species of ATPase.
Hydrostatic Flow, Osmosis, and Osmotic Pressure

Hydrostatic pressure (hydraulic pressure) drives the volume flux of filtration across the endothelial walls of glomerular capillaries (the filtration process described in Chapter 2). We described the role of plasma proteins in generating an oncotic pressure that opposed filtration. We now discuss their role a little further, along with the general role of solutes in controlling volume flux across a barrier.

As long as solutes are equally permeable as water, they will move along with filtered water or reabsorbed water and have the same concentration in the filtrate as in the plasma. Solutes play a different role when a barrier (epithelial layer or cell membrane) is less permeable to solutes than to water.

Solute dissolved in water reduce the concentration of water and, therefore, reduce the tendency of water to diffuse out of a solution. Solutions that are highly concentrated with solutes are lower in water concentration. Therefore, when solutions of different solute concentration are separated by a barrier, water will move from the more dilute solution to the more concentrated solution (ie, from where the water is more concentrated to where the water is less concentrated). This process is called osmosis. The ability of solutes to lower the concentration of water is called osmolality. It is a function of both the concentration of solutes and the kind of solutes. For example, proteins are better than sugars, and sugars are better than small ions, at lowering the concentration of water.

Osmolality is expressed in units of osmoles per kilogram of water (or, more commonly, milliosmoles per kilogram). Osmolality is often called osmotic pressure.

Osmolality and osmotic pressure have the same meaning; they are just expressed in different units (1 mOsm/kg = 19.3 mm Hg of osmotic pressure). To say that a solution has a high osmotic pressure means that it has a high osmolality. Given a cell membrane or epithelial layer in which the solutions on the 2 sides have different osmolalities, water will move by osmosis toward the side with the higher osmolality. A convenient way to express this is to say that “water follows the osmoles.”

Now for a crucial point: Osmolality (osmotic pressure) is only effective in driving osmosis when the barrier is less permeable to solutes than to water. (Imagine a barrier made up of chicken wire. Regardless of solute concentrations, there would be no osmosis because there would be no restriction on the diffusion of solute.) In the fenestrated endothelial barriers of glomerular capillaries and peritubular capillaries, most of the solutes are as permeable through the fenestrae as water and thus do not influence water movement. However, the large plasma proteins are not permeable, and they do indeed influence water movement. The osmotic pressure resulting from the proteins only (ignoring everything else) is called the colloid osmotic pressure or oncotic pressure. Colloid osmotic pressure is a component of the Starling forces governing filtration and absorption across endothelial layers. In other barriers, specifically the epithelial lining of the renal tubules, permeabilities of all solutes are generally lower than water permeability. Therefore, all solutes contribute to driving a water flux. Here, all of the osmolality, not just the component resulting from proteins, is important.
Knowing the osmolality of a solution is impossible without measuring it (it cannot be calculated even if we know the concentrations of everything in the solution. For some solutes, we can consult tables and then can interpolate between table values. For most solutes, no such tables exist.) However, we can get a rough idea of the osmolality or “guessimate” it from a related quantity called the \textit{osmolarity}. If we assume all solutes behave as ideal solutes, the osmolarity is simply the sum of the molar concentrations of all solutes without regard to kind.\textsuperscript{3} Osmolarity is expressed in units of osmoles per liter (or usually milliosmoles per liter). A solution containing 140 mEq/L of sodium, 140 mEq/L of chloride, and 10 mmol/L glucose has an osmolarity of 190 mOsm/L (140 + 140 + 10 = 190).

Fortunately, when osmolality is measured (milliosmoles per kilogram of water) and osmolarity is calculated from solution ingredients (milliosmoles per liter) and again assuming solutes to be ideal, the results are usually within 10% of each other. For convenience, physiologists often calculate osmolarity and then call it osmolality, and accept the error in this calculation as the price paid for convenience.

The difference between osmolality and osmolarity is illustrated in the case of physiological saline (0.9% NaCl, or 154 mmol/L NaCl). This solution is commonly used as a hospital infusion solution because it matches the normal osmolality of human plasma (280–290 mOsm/kg). The osmolarity of this solution is 154 + 154 = 308 mOsm/L, but when measured, the osmolality is 287 mOsm/kg.

**TRANSPORT MECHANISMS IN REABSORPTION**

Quantitatively, most of the transport in the kidney consists of reabsorption. Virtually all of the 180 L of water and several pounds of salt that are filtered each day into Bowman’s space in the glomeruli must be reabsorbed, along with large amounts of many other substances. Much of the reabsorption is very nearly \textit{iso-osmotic}, meaning that water and solutes are reabsorbed in equal proportions. Recall that filtration in the glomerulus is iso-osmotic. Almost all solutes (except large plasma proteins) move from plasma into the filtrate in the same proportion as water; thus, their concentration in the glomerular filtrate is the same as in the plasma. In the proximal tubule, where the majority of reabsorption occurs, the process is virtually iso-osmotic, ie, water and solutes are absorbed in equal proportions. In the later portions of the nephron, reabsorption is generally not iso-osmotic (a point that is crucial for our ability to separately regulate solute and water balance).

Most of the solute reabsorbed in the proximal tubule consists of sodium and the anions that must accompany the sodium to maintain electroneutrality: mostly chloride and bicarbonate. These solutes are removed from the tubular lumen and put into the interstitium by a combination of the processes described previously. Thus, a large amount of solute is transferred from lumen to interstitium, setting

\textsuperscript{3}Sometimes, the distinction between real and ideal solutions confuses students. If all solutions were ideal, then a mole of any dissolved solute at any concentration would always equal one osmole. In real solutions, however, the number of osmoles per mole of dissolved solute depends both on the species of solute and the volume in which it is dissolved (ie, its concentration). Fortunately, these differences are not great in physiological solutions.
up an osmotic gradient that favors the parallel movement of water. The proximal tubule epithelium is very permeable to water, and water—a substantial amount—indeed follows solute from the lumen to the interstitium. The water moves in equal proportions as solute, so that both the fluid removed from the lumen and that remaining behind are essentially iso-osmotic with the original filtrate (i.e., have the same osmolality). We say “essentially” because there must be some difference in osmolality to induce water movement, but for an epithelial barrier (like the proximal tubule) that is very permeable to water, a difference of less than 1 mOsm/kg is sufficient to drive reabsorption of water.

Tubular hydrostatic pressure is several millimeters of mercury greater than interstitial hydrostatic pressure, and this pressure gradient also favors reabsorption. However, under normal circumstances, this is a small influence. It requires a hydrostatic pressure gradient of 19.3 mm Hg to act as a driving force equivalent to an osmotic gradient of 1 mOsm/kg, and the hydrostatic pressure difference is usually not more than 5–8 mm Hg.

Once in the interstitium, the solutes and water move from interstitium into the peritubular capillaries and are returned to the systemic circulation. Otherwise, the kidney would swell without limit.

Fortunately, the Starling forces across the peritubular capillaries favor reabsorption. The capillary hydraulic pressure, which was about 60 mm Hg in the glomerular capillaries, and opposes uptake of interstitial fluids has fallen to about 15–20 mm Hg within the peritubular capillaries, but the plasma oncotic pressure, resulting from filtration in the glomerular capillaries, has risen to more than 30 mm Hg. There is a small but significant, interstitial pressure (Table 4–1). Therefore, the net filtration pressure is now a net absorptive pressure, and net fluid movement is into the peritubular capillaries. The alert student can appreciate the fact that if cortical Starling forces are abnormal (e.g., low plasma oncotic pressure as when liver disease prevents normal production of serum albumin), absorption of fluid from the cortical interstitium can be slowed, causing a backup of fluid that inhibits fluid movement from tubular lumen to interstitium. Ultimately, this can lead to increased excretion of water and electrolytes from the body.

As blood flows through the peritubular capillaries, there is a rapid diffusion of individual molecules back and forth between capillary plasma and cortical interstitial fluid. The total volume of interstitial space is only 4% of the total cortical volume, and the vascular volume is a little higher. Given the very high renal blood flow, the solute concentrations in the interstitial fluid are essentially clamped to those in the blood perfusing the cortex. The cortical interstitium remains quite plasma-like (minus the proteins) in its composition, even though large amounts of solute continuously cross through the interstitium from tubule to blood.

With all of these factors as a background, let us more closely examine how things are reabsorbed. We can take an anthropomorphic view and ask, “If I’m a molecule in the lumen, what induces me to go across the epithelial layer instead of just staying where I am?”
Generalized Epithelial Transport: The Transcellular Route

Epithelial transport requires that epithelial cells are polarized (ie, the proteins present in the apical and the basolateral membranes are not the same). This polarization can promote net flux of sodium from lumen to interstitium, which is the linchpin around which the transport of virtually every other substance depends. Figure 4–4 shows the morphology of a generalized renal epithelium in which salt and water transport can be viewed as a 4-step process. Step 1 is the active extrusion of sodium via Na-K-ATPase from the cell to the interstitium. This creates a low concentration of sodium within the cell so that sodium moves downhill from the lumen to the cell interior via a variety of symporters, antiporters, and channels. A key player in the proximal tubule is the sodium-proton antiporter (NHE-3 isoform) that we discuss further later. The consequence of this transcellular sodium movement is the separation of charge (excess Na\(^+\) on the interstitial side) that promotes Step 2, the movement of anions through anion-specific transcellular and paracellular pathways to balance the positive charge. The accumulation of sodium and anions in the interstitial space produces an osmotic gradient from lumen to interstitium that promotes water movement (Step 3). Finally, the accumulation of salt and water in the interstitium promotes the bulk flow of solute and water into the peritubular capillaries (Step 4) driven by Starling forces (see Table 4–1). In Step 1, any reabsorbed substance that enters the epithelial cells with sodium across the apical membrane must exit across the basolateral membrane. The particular mechanism by which this occurs depends on the

Table 4–1. Estimated forces involved in the movement of fluid from interstitium into peritubular capillaries

<table>
<thead>
<tr>
<th>Forces</th>
<th>mm Hg</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Favoring uptake</td>
<td></td>
</tr>
<tr>
<td>a Interstitial hydraulic pressure, (P_{\text{int}})</td>
<td>3</td>
</tr>
<tr>
<td>b Oncotic pressure in peritubular capillaries, (\pi_{\text{pc}})</td>
<td>33</td>
</tr>
<tr>
<td>2 Opposing uptake</td>
<td></td>
</tr>
<tr>
<td>a Hydraulic pressure in peritubular capillaries, (P_{\text{pc}})</td>
<td>20</td>
</tr>
<tr>
<td>b Interstitial oncotic pressure, (\pi_{\text{int}})</td>
<td>6</td>
</tr>
<tr>
<td>3 Net pressure for uptake (1–2)</td>
<td>10</td>
</tr>
</tbody>
</table>

*The values for peritubular-capillary hydraulic and oncotic pressures are for the early portions of the capillary. The oncotic pressure, of course, decreases as protein-free fluid enters it (ie, as absorption occurs) but would not go below 25 mm Hg (the value of arterial plasma) even if all fluid originally filtered at the glomerulus were absorbed.
Chapter 4

substance. For example, we know that glucose crosses the apical membrane via the Na/Glucose symporter and exits across the basolateral membrane via a GLUT uniporter. Many other substances also move via their own uniporters. We examine some of these more closely later as we discuss transport mechanisms in individual tubular segments, but, in general, the transcellular route involves crossing 2 membranes—apical and basolateral—and the transporters in the 2 membranes are different and, thereby, promote unidirectional movement of molecules.

The Paracellular Route

As water follows sodium and its anions across the epithelium, the volume remaining in the lumen decreases. Therefore, any solute that has not been specifically transported via the transcellular route will become more concentrated. If two thirds of the water is removed, a nontransported solute will increase in concentration to 3 times its original value. As the luminal concentration rises, this generates a concentration gradient across the tight junctions between the lumen and the interstitium. If the tight junctions are permeable to the substance in question (“leaky”), the substance will diffuse from the lumen to the interstitium. This is precisely what happens to many solutes (eg, urea, potassium, chloride, calcium,

Figure 4–4. Steps involved in transporting solute and water from the tubular lumen to the peritubular capillary. Everything depends on and follows logically from Step 1, which is the active extrusion of sodium into the interstitium. This induces a parallel transport of anions (Step 2). The movement of sodium and anions generates an osmotic drive that causes reabsorption of water (Step 3). Finally, the increased volume in the interstitium alters peritubular Starling forces and induces the bulk flow of water and solute from interstitium into the peritubular capillary (Step 4).
and magnesium) in the proximal tubule. The exact fractions that are reabsorbed depend on the permeability of the tight junctions but are generally in the range of one half to two thirds. Because ions are driven not only by concentration gradients but also by voltage gradients, the transepithelial voltage plays a role here also. Early in the proximal tubule, the lumen is slightly negative relative to the interstitium (a few millivolts), whereas later it is slightly positive. This voltage enhances paracellular anion reabsorption early and reduces it later. To keep things simple, we can account for most paracellular reabsorption purely on the basis of the rise in luminal concentration that occurs when water is reabsorbed. One substance that does not get reabsorbed by the paracellular route is glucose. First, it is transported by the transcellular route. Second, the tight junctions are not permeable to saccharides. Thus, it cannot diffuse no matter how large the concentration gradient might be.

**Limits on Rate of Transport: T_m and Gradient-Limited Systems**

Even though the transport capacity of the renal cortex is huge, it is not infinite. There are upper limits to the speed with which any given solute can be reabsorbed from the tubular lumen to capillary blood. In certain situations, these limits are reached, with the consequence that more than the usual amount of solute is not reabsorbed (ie, left in the lumen to be passed on to the next nephron segment). In general, transport mechanisms can be classified by the properties of these limits as either (1) tubular maximum-limited (T_m) systems or (2) gradient-limited systems. T_m systems reach an upper limit because the transporters moving the substance become saturated; any further increase in solute concentration does not increase the rate at which the substance binds to the transporter and thereafter moves through the membrane. Gradient-limited systems reach an upper limit because the tight junctions are leaky, and any significant lowering of luminal concentration relative to the interstitium results in a leak back into the lumen as fast as the substance is transported out. Thus, the upper rate for the T_m-limited system is a property of the transporter, whereas the upper rate of a gradient-limited system is a property of the permeability of the epithelial monolayer regardless of the maximal rate of the transport protein.

Let us explain T_m-limited systems using glucose as an example. Glucose is present in plasma at a concentration of about 5 mmol/L (90 mg/dL) and is freely filtered. It is reabsorbed by the transcellular route. Glucose enters the epithelial cells across the apical membrane via a symporter with sodium (a member of the SGLT protein family) and exits across the basolateral membrane into the interstitium via a uniporter (a GLUT protein family member). Normally, all the filtered glucose is reabsorbed in the proximal tubule, with none remaining in the lumen to be passed on to the loop of Henle. However, if the filtered load of glucose is abnormally high, the SGLT proteins’ upper limit for reabsorption is reached. That upper limit is the tubular maximum, or T_m, for glucose. It is the maximum rate at which the substance (glucose in this case) can be reabsorbed regardless of the luminal concentration. Any increase in filtered load above the T_m, which for glucose
represents a pathological situation, results in glucose being passed on to the loop of Henle. $T_m$ systems are perhaps like snow shovelers keeping a driveway clear during a snowstorm. With an ordinary snowfall, they can remove snow as it falls and keep a driveway clear. However, during a blizzard, snow falls faster than it can be shoveled and accumulates. (Although we are discussing $T_m$ systems in terms of reabsorption, the same concept applies to secretion. For some secreted substances, there is a $T_m$ limit to how fast they are secreted, and a rise in plasma concentration does not increase the secretion rate above the $T_m$.)

Gradient-limited systems are more complicated than $T_m$ systems. The key to gradient-limited systems is that the epithelium has a significant passive permeability to the substance, usually through the tight junctions, such that the establishment of a large concentration gradient between the interstitium and lumen results in a large passive back-leak. Consider the case of sodium. In the proximal tubule, the tight junctions are quite permeable to sodium. Any significant lowering of the concentration in the lumen as sodium is reabsorbed results in a large passive flux from the interstitium back into the lumen. Sodium is freely filtered and present in the luminal fluid at a concentration of about 140 mEq/L, the same as in plasma. It is reabsorbed by a variety of pathways described previously (ie, symporters and antiporters in the apical membrane and the Na-K-ATPase across the basolateral membrane into the interstitium). As sodium is transported into the interstitium, the interstitial concentration begins to rise and the luminal concentration falls. The rise in interstitial concentration not only drives a flux of sodium into the peritubular capillaries, but also back through the tight junctions into the lumen. Most of the sodium moves into the blood, accomplishing the goal of reabsorption, but some does leak back into the lumen. When the concentration of sodium reaches a sufficiently low level in the lumen, the concentration gradient between the interstitium and the lumen drives sodium back across the tight junctions as fast as it can be transported through the transcellular pathway from lumen to interstitium. At this point, transport through paracellular and transcellular pathways is large, but net transport is zero: The system has established the largest gradient possible, its gradient limit. The leakier the epithelium, the lower is the gradient limit (in the proximal tubule, the gradient limit is about 2 millimolar sodium).

In normal conditions the reabsorption of sodium is accompanied by a proportional reabsorption of water, so that the luminal sodium concentration actually falls very little, ie, it does not reach its limiting gradient. The main factor governing sodium reabsorption is the activity of its transporters (particularly, the NHE-3 sodium-hydrogen antiporters). However, if there is an unusually large amount of poorly reabsorbed solute in the lumen (eg, infused mannitol), this restrains reabsorption of water because nonreabsorbed osmoles remain in the lumen. In turn, less water accompanies reabsorbed sodium. Then as sodium is reabsorbed, its luminal sodium concentration falls and reaches the gradient limit. This reduces the amount of sodium that is reabsorbed and leads to an osmotic diuresis (see Chapter 6).

Technically, back-leak of sodium is a secretion, and net sodium reabsorption is the difference between reabsorption from lumen to interstitium and secretion from interstitium to lumen. Because the net transport is indeed reabsorption, we
will say simply that sodium is reabsorbed, with a limit placed by the amount of back-leak.

The functional reasons for differentiating between $T_m$ and gradient-limited systems is that solutes handled by $T_m$ systems may, if the filtered load is below the $T_m$, be reabsorbed essentially completely, whereas solutes handled by gradient-limited systems are never reabsorbed completely, ie, a substantial amount always remains in the tubule to be passed on to the next nephron segment.

**KEY CONCEPTS**

1. Reabsorption is a 2-step process: lumen to interstitium, and interstitium to peritubular capillary.

2. Flux from lumen to interstitium can be transcellular, using separate transport steps in the apical and basolateral membranes, or paracellular, around the cells through tight junctions.

3. Channels and transporters promote the transmembrane flux of solutes that cannot permeate lipid bilayers.

4. Osmotic gradients drive a volume flux across membranes and epithelia.

5. Osmotic pressure and osmolality mean the same thing and represent the power of dissolved solute to drive an osmotic flux of water.

6. For convenience, osmolality is approximated by the easier concept of osmolarity.

7. Water and solutes, which are reabsorbed from lumen to interstitium, then move from interstitium to peritubular capillaries by bulk flow, driven by Starling forces.

8. The reabsorption of water and almost all solutes is linked, directly or indirectly, to the active reabsorption of sodium.

9. All reabsorptive processes have a limit on how fast they can occur, either because the transporters saturate ($T_m$ systems) or because the substance leaks back into the lumen (gradient-limited systems).
STUDY QUESTIONS

4–1. Flux of a solute out of a cell, whether via a uniporter, symporter, or an ATPase, is always a process of active transport (primary or secondary). True or false?

4–2. Reabsorption in the proximal tubule is described as being iso-osmotic, leaving the luminal fluid isosmotic with plasma. Yet we already know from earlier chapters that the excreted urine usually is quite different osmotically from the surrounding interstitium. Why is the final urine not always iso-osmotic?

4–3. In the proximal tubule, the tubular epithelium is far less permeable to small solutes than is the endothelium of the surrounding peritubular capillaries. True or false?

4–4. Low plasma oncotic pressure inhibits volume reabsorption from tubular lumen to interstitium. Because this is plasma oncotic pressure, how can it affect transepithelial transport?

4–5. Even though values of osmolality and osmolarity differ numerically, any 2 solutions of equal osmolarity will have equal osmolality. True or false?

4–6. Given the high volume of fluid normally moving from interstitium to blood in the renal cortex, how can secreted substances move from blood to epithelium? Are they not going the wrong way?

4–7. The T_m for glucose is set at what level?
   A. Close to the normal filtered load
   B. Well above the normal filtered load
   C. Well below the normal filtered load

4–8. Define channel “gating,” and state whether changing interstitial osmolality is a way of gating channels.
OBJECTIVES

The student understands the renal handling of certain organic substances, specifically, including urea:

- States the physiological utility of either excreting or saving organic solutes.
- States the general characteristics of the proximal tubular systems for active reabsorption or secretion of organic nutrients.
- Describes the renal handling of glucose and states the conditions under which glucosuria is likely to occur.
- Describes the renal handling of proteins and small peptides.
- Describes the secretion of para-aminohippurate.
- Outlines the handling of urate.
- Describes the secretion of organic cations.
- Describes, in general terms, the renal handling of weak acids and bases and how tubular pH affects reabsorption.
- Describes the renal handling of urea, including the medullary recycling of urea from the collecting duct to the loop of Henle.

Subsequent chapters of this textbook deal almost exclusively with the renal handling of salt and water and the other major physiological ions in blood because homeostatic regulation of their excretion is of such crucial importance. However, as pointed out in Chapter 1, another major renal function is the excretion of organic waste products, foreign chemicals, and their metabolites. Furthermore, the kidneys filter large amounts of substances that they do not excrete; therefore, reabsorptive processes must exist to prevent inappropriate loss of useful organic nutrients. Because the blood contains many small, filterable molecular species, the kidney has to handle all of them. An analysis of the renal transport pathways for all these organic substances is an overwhelming task, so this chapter briefly describes some of the major pathways. In short, however, the kidney (1) keeps or saves (reabsorbs) organic metabolites that should not be lost and (2) excretes by not reabsorbing or actually secreting waste products and foreign organic substances to prevent their accumulation in the blood.
One organic substance, urea, is unique in this regard. It is a waste product that must be excreted to prevent accumulation. However, it also plays a key role in renal regulation of water balance. We briefly present the renal handling of urea later in this chapter and again in Chapter 6 in the discussion of renal handling of water.

ACTIVE PROXIMAL REABSORPTION OF ORGANIC NUTRIENTS (EG, GLUCOSE, AMINO ACIDS)

Most major cellular nutrients are freely filterable, including glucose, amino acids, acetate, Krebs cycle intermediates, certain water-soluble vitamins, lactate, acetoacetate, β-hydroxybutyrate, and many others. The proximal tubule is the major site for reabsorption of the large quantities of organic nutrients filtered each day by the renal corpuscles. The characteristics of glucose reabsorption described in Chapter 4 are typical of the transport processes for most nutrients. In general,

1. They are actively transported (ie, can be reabsorbed up their respective electrochemical gradients). Indeed, the luminal concentration of the substances in many cases can be reduced virtually to zero (ie, reabsorption can be very close to 100% complete).
2. The “uphill” step is across the luminal membrane, usually via a symporter with sodium.
3. Most are characterized as Tm systems (have an upper limit to the speed at which they can transport). These limits are usually well above the amounts normally filtered. Accordingly, the kidneys return these filtered substances back to the plasma; however, because there is no opportunity to vary the amount excreted (there is none), the kidneys do not help regulate their levels in the body. It is also true, however, that under abnormal conditions the plasma concentration of these substances may increase so much that the filtered load exceeds the reabsorptive Tm and large quantities are excreted in the urine. Examples are glucose, acetoacetate, and β-hydroxybutyrate in patients with severe uncontrolled diabetes.
4. They manifest specificity. This means that a given transporter selectively takes up one or a few substrates and ignores all others. However, there is not a separate transporter for every solute in the body. Two or more closely related substances may use the same transporter. For example, the amino acid transporters are distinct from those for glucose and other monosaccharides, but there are not 20 separate transporters, one for each amino acid. Rather there is one for arginine, lysine, and ornithine; another for glutamate and aspartate; and so on. Shared pathways imply that there is competition among those substances using the same transporter. Practically, this means that an excess of one substance, ornithine for example, in the blood may lead to not
only excess ornithine excretion but also inappropriate excretion of arginine and lysine.

5. They are inhibitable by a variety of drugs, and several monogenetic diseases are associated with loss of function in one or more of these proximal re-absorptive systems. In some cases, the deficit may be highly specific (eg, involving only one amino acid), whereas in others multiple systems may be involved (eg, glucose and many amino acids). This range of defects is also seen when the deficit is due to an ingested toxin (eg, heavy metal toxicity) rather than a genetic abnormality.

Glucose

Because of the importance of glucose as the basic coin of cellular energy exchange and the prevalence of diabetes, which manifests itself as renal disease along with other pathologies, we review the normal renal handling of glucose. The normal plasma glucose level is about 90 mg/dL (5 mmol/L). It rises transiently to well over 100 mg/dL during meals and can reach levels of over 1000 mg/dL (over 55 mmol/L) in severe diabetes. Normally, all the filtered glucose is reabsorbed in the proximal tubule. This involves removing glucose from the tubular lumen along with sodium via a sodium-dependent glucose symporter (SGLUT) across the apical membrane of proximal convoluted tubule epithelial cells, followed by its exit across the basolateral membrane into the interstitium via a GLUT uniporter. Unlike the case for sodium and many other solutes discussed later, the tight junctions are not significantly permeable to glucose. Therefore, as glucose is removed from the lumen and the luminal concentration falls, there is no back-leak. The transport of a solute with no back-leak depends only on the characteristics of the rate-limiting transporter, in this case the SGLT symporter, and is a Tₘ-limited system.

Because glucose reabsorption is a Tₘ system, abnormally high filtered loads overwhelm the reabsorptive capacity (exceed the Tₘ; Figure 5–1). This occurs when plasma glucose rises above roughly 300 mg/dL. Again, this is a pathological situation but one that is relatively common. Assume that the glucose Tₘ is 375 mg/min (a typical value). The normal filtered load is well below this level. With a glomerular filtration rate (GFR) of 125 mL/min (1.25 dL/min) and plasma glucose of 90 mg/dL, the filtered load is 1.25 dL/min × 90 mg/dL = 112.5 mg/min, much less than the Tₘ of 375 mg/min. When plasma glucose reaches 300 mg/dL, the filtered load is now 1.25 dL/min × 300 mg/dL = 375 mg/min. At this point, the proximal convoluted tubule fails to reabsorb all the filtered glucose, and a little glucose begins to spill into the urine. Further increases in plasma glucose above 300 mg/dL lead to progressively greater renal losses. We discuss the various reasons for the diuresis (high urine volume) associated with diabetes later, but one can appreciate that any glucose not reabsorbed is an osmole in the tubule that has consequences for water reabsorption.
PROTEINS AND PEPTIDES

Although we sometimes say that the glomerular filtrate is protein-free, it is not truly free of all protein. First, small and medium-size proteins (e.g., angiotensin, insulin) are filtered in considerable quantities. Second, although the movement of large plasma proteins across glomerular filtration barrier is extremely limited, a small amount does make it through into Bowman’s space. For albumin, the plasma protein of highest concentration in the blood, the concentration in the filtrate is normally about 10 mg/L, or roughly 0.02% of the plasma albumin concentration (50 g/L). Yet because of the huge volume of fluid filtered per day, the total filtered amount of protein is not negligible. However, the proximal tubule is capable of taking up filtered albumin and other proteins, and we treat this protein uptake separately here to emphasize its importance. We use the word uptake rather than reabsorption because the proteins, although they are transported intact out of the lumen into the epithelial cells, are degraded into their constituent amino acids before being transported into the cortical interstitium. Thus, the term reabsorption in the context of proteins and peptides, although widely used, is actually a misnomer.

The initial step for the uptake of larger proteins is endocytosis at the luminal membrane. This energy-requiring process is triggered by the binding of filtered
protein molecules to specific receptors on the luminal membrane. Therefore, the rate of endocytosis is increased in proportion to the concentration of protein in the glomerular filtrate until a maximal rate of vesicle formation, and thus the $T_m$ for protein uptake, is reached. The pinched-off intracellular vesicles resulting from endocytosis merge with lysosomes, whose enzymes degrade the protein to low-molecular-weight fragments, mainly individual amino acids. These end products then exit the cells across the basolateral membrane into the interstitial fluid, from which they gain entry to the peritubular capillaries.

To understand the potential problem associated with a failure to take up filtered protein, remember that

\[
\text{Total filtered protein} = \text{GFR} \times \text{concentration of protein in filtrate} \\
= 180 \text{ L/day} \times 10 \text{ mg/L} = 1.8 \text{ g/day}
\]

If none of this protein was removed from the lumen, the entire 1.8 g would be lost in the urine. In fact, almost all the filtered protein is taken up, so that the excretion of protein in the urine is normally only 100 mg/day. The endocytic mechanism by which protein is taken up is easily saturated; however, any large increase in filtered protein resulting from increased glomerular permeability can cause the excretion of large quantities of protein.

Discussions of the renal handling of protein logically tend to focus on albumin because it is by far the most abundant plasma protein. There are, of course, many other plasma proteins, and it should be emphasized that many of these proteins, being smaller than albumin, are more easily filtered than albumin. For example, growth hormone (molecular weight 22,000 kDa) is approximately 60% filterable, and insulin is 100% filterable. The total mass of these filtered hormones is insignificant; however, because even tiny levels in the plasma have important signaling functions in the body, renal filtration becomes an important influence on levels in the blood. Relatively large fractions of these smaller plasma proteins are filtered and then degraded in tubular cells. Accordingly, the kidneys are major sites of catabolism of many plasma proteins, specifically including polypeptide hormones. Decreased rates of degradation occurring in renal disease may result in elevated plasma hormone concentrations.

Very small peptides, such as angiotensin II, are handled differently from larger proteins, although the end result is the same: the catabolism of the peptide and preservation of its amino acids. The very small peptides are completely filterable at the renal corpuscles and are then catabolized mainly into amino acids within the proximal tubular lumen by peptidases located on the luminal surface of the plasma membrane. The amino acids (as well as any di- and tripeptides generated by this process) are then reabsorbed by the same transporters that normally reabsorb filtered amino acids.

Finally, it should be noted that, in certain types of renal damage, proteins released from damaged tubular cells rather than filtered at the renal corpuscles may appear in the urine and provide important diagnostic information.
The proximal tubule actively secretes a large number of different organic anions, both endogenously produced and foreign (see Table 5–1 for a partial listing). Many of the organic anions handled by this system are also filterable at the renal corpuscles, and so the amount secreted proximally adds to that which gains entry to the tubule via glomerular filtration. Others, however, are extensively bound to plasma proteins and undergo glomerular filtration only to a limited extent; accordingly, proximal tubular secretion constitutes the only significant mechanism for their excretion.

The active secretory pathway for organic anions in the proximal tubule in some sense is the reverse of reabsorption of organic solutes: There are active transporters for the anions at the basolateral membrane of tubular epithelial cells that are the rate-limiting step in overall transport. Transport out of the cell across the apical membrane into the lumen is via facilitated diffusion on a variety of uniporters or more specific sodium-dependent antiporters. Several different organic anion transporters (members of the OAT family of proteins) have been cloned. They are interesting in their similarity to amino acid transporters; they are specific for classes of organic anions (eg, OAT3 transports most tricarboxylic organic anions like citrate) but do not distinguish well among members of the class. Because the basolateral membrane of proximal convoluted tubule epithelial cells contains all of these different transporters, the proximal tubule has the capacity to secrete all the organic anions listed in Table 5–1 and many more. Like glucose, organic anions are not significantly permeable through tight junctions or membranes, so that their transport is also characterized by a tubular maximum. If the blood concentration of an organic anion is too high, it will not be efficiently removed from the blood by the kidneys (one aim of the dose regimen for many prescribed drugs).

Table 5–1. Some organic anions actively secreted by the proximal tubule

<table>
<thead>
<tr>
<th>Endogenous substances</th>
<th>Drugs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bile salts</td>
<td>Acetazolamide</td>
</tr>
<tr>
<td>Fatty acids</td>
<td>Chlorothiazide</td>
</tr>
<tr>
<td>Hippurates</td>
<td>Ethacrynate</td>
</tr>
<tr>
<td>Hydroxybenzoates</td>
<td>Furosemide</td>
</tr>
<tr>
<td>Oxalate</td>
<td>Penicillin</td>
</tr>
<tr>
<td>Prostaglandins</td>
<td>Probenecid</td>
</tr>
<tr>
<td>Urate</td>
<td>Saccharin</td>
</tr>
<tr>
<td></td>
<td>Salicylates</td>
</tr>
<tr>
<td></td>
<td>Sulfonamides</td>
</tr>
</tbody>
</table>
The relatively nondiscriminating nature of this collection of transporters accounts for their ability to eliminate from the body so many drugs and other foreign environmental chemicals. In this regard, the liver’s metabolic transformations are very important. In the liver, many foreign (and endogenous) substances are conjugated with either glucuronate or sulfate. The addition of these groups then renders the parent molecule far more water soluble. These 2 types of conjugates are actively transported by the organic-anion secretory pathway.

The most intensively studied organic anion secreted by this pathway is para-aminohippurate (PAH), the substance used for the measurement of effective renal plasma flow (see Chapter 3). PAH secretion involves a pair of antiporters, one at each membrane. At the basolateral membrane, PAH is taken up in exchange for the anion (base) form of a dicarboxylic acid. PAH is extruded into the lumen across the apical membrane via another antiporter.

As the plasma concentration of an anion secreted by this system increases, so does the rate of secretion (until the $T_m$ for that substance is reached). This provides a mechanism for homeostatically regulating the endogenous organic anions handled by the system and for speeding the excretion of foreign organic anions.

PAH is typical, in yet another way, of many of the organic anions secreted proximally: It undergoes no significant additional transport anywhere along the nephron. In contrast, some of the other organic anions secreted by the proximal tubule can also undergo other forms of transport in both the proximal tubule and more distal segments. The most important of these is passive tubular reabsorption or secretion, which is described later.

**Urate**

Urate (the base form of uric acid) provides a fascinating example of the renal handling of organic anions that is particularly important for clinical medicine and is illustrative of renal pathology. An elevated plasma concentration of urate can cause gout; therefore, its removal from the blood is important. However, it is as though the kidney cannot make up its mind what to do with urate. Urate is not protein bound and so is freely filterable. Almost all the filtered urate is reabsorbed early in the proximal tubule; however, further on in the proximal tubule, urate undergoes active tubular secretion. Then, in the straight portion, urate is once again reabsorbed. The total rate of tubular reabsorption is normally much greater than the rate of tubular secretion, and so the mass of urate excreted per unit time is only a small fraction of the mass filtered. We will not discuss the specific transport steps required to accomplish all of this, but most involve antiporters that exchange urate for another organic anion.

Although urate reabsorption is greater than secretion, the secretory process is homeostatically controlled to maintain relative constancy of plasma urate. In other words, if plasma urate begins to increase because of increased urate production, the active proximal secretion of urate is stimulated, thereby increasing urate excretion.

Given these mechanisms of renal urate handling, the reader should be able to deduce the 3 ways by which altered renal function can lead to decreased urate
excretion and hence increased plasma urate, as in gout: (1) decreased filtration of urate secondary to decreased GFR, (2) excessive reabsorption of urate, and (3) diminished secretion of urate.

ACTIVE PROXIMAL SECRETION OF ORGANIC CATIONS

Proximal tubules possess several closely related transport systems for organic cations that are analogous to those for organic anions: Because of the large number of different transporters, a substantial amount of foreign and endogenous substances are transported (Table 5–2), the cations compete with one another for transport, and the transporters manifest a $T_m$ limitation. Organic cations enter across the basolateral membrane via one of several uniporters, members of the OCT family (organic cation transporter), and exit into the lumen via an antiporter, which exchanges a proton for the organic cation.

The proximal secretion of organic cations, as for organic anions, is particularly critical for the excretion of those cations extensively bound to plasma proteins and not filterable at the renal corpuscle. However, again similar to organic anions, many of the organic cations secreted by the proximal tubules are not protein bound and, therefore, also undergo glomerular filtration and tubular secretion; creatinine is a good example.

Finally, and again analogous to organic anions, some organic cations are not only actively secreted by the proximal tubules but also may undergo other forms of tubular handling, mainly passive reabsorption or secretion.

<table>
<thead>
<tr>
<th>Endogenous substances</th>
<th>Drugs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetylcholine</td>
<td>Atropine</td>
</tr>
<tr>
<td>Choline</td>
<td>Isoproterenol</td>
</tr>
<tr>
<td>Creatinine</td>
<td>Cimetidine</td>
</tr>
<tr>
<td>Dopamine</td>
<td>Meperidine</td>
</tr>
<tr>
<td>Epinephrine</td>
<td>Morphine</td>
</tr>
<tr>
<td>Guanidine</td>
<td>Procaine</td>
</tr>
<tr>
<td>Histamine</td>
<td>Quinine</td>
</tr>
<tr>
<td>Serotonin</td>
<td>Tetraethyl ammonium</td>
</tr>
<tr>
<td>Norepinephrine</td>
<td></td>
</tr>
<tr>
<td>Thiamine</td>
<td></td>
</tr>
</tbody>
</table>
**pH DEPENDENCE OF PASSIVE REABSORPTION OR SECRETION**

Many substances handled by the kidney are weak acids or bases. At a given pH, the total amount of each one is split between a neutral form and an ionized form. Many weak acids are neutral at low pH (acid form) and are dissociated into an anion and a proton at higher pH. The lower the pH, the more of the total there is in the neutral acid form, whereas the higher the pH, the greater is the fraction in the dissociated anionic form. In general, neutral forms of organic acids and bases are more permeable in lipid membranes than ionized forms, and the neutral forms can diffuse either into or out of the tubular lumen down the concentration gradient of the neutral form. In contrast, the ionized forms, once in the lumen, cannot diffuse; they are effectively trapped there. Imagine the case in which the tubular fluid becomes acidified relative to the plasma, which it does on a normal diet. For a weak acid in the tubular fluid, relatively more will be converted to the neutral free acid form and, therefore, become more permeable. This favors diffusion out of the lumen (reabsorption). Therefore, a highly acidic urine (low pH) tends to increase passive reabsorption of weak acids (and promote less excretion). For many weak bases, the pH dependence is just the opposite. At low pH they are protonated cations (trapped in the lumen), whereas at high pH they are converted to neutral free base. As the urine becomes acidified, more is converted to the impermeable charged form and is trapped in the lumen. Less is reabsorbed passively, and more is excreted.

Having said this, what difference does it make? Because so many medically useful drugs are weak organic acids and bases, all these factors have important clinical implications. For example, if one wishes to enhance the excretion of a drug that is a weak acid, one attempts to alkalinize the urine (because this traps the ionic form in the lumen). In contrast, acidification of the urine is desirable if one wishes to prevent excretion of the drug. Of course, exactly the opposite applies to weak organic bases. At any pH, increasing the urine flow increases the excretion of both weak acids and bases (Figure 5–2). Finally, excretion can be reduced by administering another drug that interferes with any active proximal secretory pathway for the drug.

**UREA**

Urea is a very special substance for the kidney. It is an end product of protein metabolism, simply waste to be excreted, and also a useful tool for the regulation of water excretion. Proteins, from which urea is derived, are the action molecules in cells (e.g., transporters and enzymes) and the structural substance of the connective tissue (e.g., collagen). In addition, proteins are the source of metabolic fuel. Excess dietary protein not needed for tissue synthesis is either oxidized right away or converted to fat and stored for later oxidation. During fasting, the body breaks down its own protein for fuel, in essence consuming itself. When oxidized for fuel, protein is first split into its constituent amino acids. These are then separated into a nitrogen moiety (ammonium) and a carbohydrate moiety. The carbohydrate goes on to further metabolic processing, but the ammonium cannot be further
oxidized and is a waste product. The problem is, ammonium is rather toxic to most tissues (except the medullary interstitium) and the liver immediately converts it to urea (mostly) and a smaller, but a crucial, amount of glutamine. (We will take up the fate of this glutamine in Chapter 9 on acid-base balance.) Urea production proceeds continuously and constitutes about half of the normal solute content of urine.

The normal level in the blood is quite variable (3 mmol/L–9 mmol/L), reflecting variations in both protein intake and renal handling of urea. Over the long term (days to weeks), renal urea excretion must match hepatic production; otherwise, plasma levels would rise into the pathological range, producing a condition

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1 Plasma urea concentration is usually expressed as blood urea nitrogen (BUN) in units of milligrams per deciliter. Each molecule of urea contains 2 atoms of nitrogen, so 1 mmol of urea contains 2 mmol of nitrogen, with a combined weight of 28 mg. Thus, the normal levels of plasma urea are expressed as BUN values ranging from 8.4 mg/dL to 25.2 mg/dL. We use units of millimoles per liter because we can then directly convert to osmolality.
called *uremia*. On a shorter term basis (hours to days), urea excretion rate may not exactly match production rate because urea excretion is also regulated for purposes other than keeping a stable plasma level. As discussed in Chapter 6, urea is a key solute involved in regulating the excretion of water. To summarize renal handling of urea, urea is freely filtered. About half is reabsorbed in the proximal tubule. An amount equal to that reabsorbed is then secreted back into the loop of Henle. Finally, about half is reabsorbed again in the medullary collecting duct. The net result is that about half the filtered load is excreted (Figure 5–3).

*Figure 5–3.* Urea handling by the kidney. The arrows indicate the regions where transport occurs and the direction of transport. The boxes show (in the top half) the percentage of the filtered load remaining in the tubule and, in the bottom half, the luminal concentration relative to the concentration in plasma. The numbers are subject to considerable variability, particularly for regions beyond the proximal tubule, because they are so dependent on hydration status.
As a molecule, urea is small (molecular weight, 60 Da), is water soluble, and is freely filtered. Because of its highly polar nature, it does not permeate lipid bilayers, but a set of uniporters (the UT family) transport urea in various places in the kidney and in other sites within the body (particularly red blood cells). Because urea is freely filtered, the filtrate contains urea at a concentration identical to that in plasma. Let us assume a normal plasma level (5 mmol/L). Roughly half the filtered load is reabsorbed in the proximal tubule. This occurs primarily by the paracellular route. As water is reabsorbed (about two thirds of the filtered water is reabsorbed in the proximal tubule), solutes in the lumen that are not reabsorbed by the transcellular route become concentrated. Urea is prominent among these solutes. As urea becomes concentrated, it is driven passively through the leaky tight junctions. By the time the tubular fluid enters the loop of Henle, about half the filtered urea has been reabsorbed, and the urea concentration has increased to a little more than its value in the filtrate (because proportionally more water than urea was reabsorbed). At this point, the process becomes fairly complicated. First, conditions in the medulla depend highly on the individual’s state of hydration. Second, there is a difference between superficial nephrons, with short loops of Henle that only penetrate the outer medulla, and juxtamedullary nephrons, with long loops of Henle that reach all the way down to the papilla. We simplify the issue by considering all nephrons together.

The interstitium of the medulla has a considerably higher urea concentration than plasma. The concentration increases from the outer to the inner medulla (part of the so-called medullary osmotic gradient), and its peak value in the inner medulla depends on hydration status and levels of antidiuretic hormone. We explain these variations in Chapter 6 in connection with regulation of water excretion. For now, we note that the medullary urea concentration is greater than in the tubular fluid entering the loop of Henle, so there is a concentration gradient favoring secretion into the lumen. The tight junctions in the loop of Henle are no longer permeable, but the epithelial membranes of the thin regions of the Henle loops express urea uniporters, members of the UT family. This permits secretion of urea. In fact, the urea secreted from the medullary interstitium into the thin regions of the loop of Henle replaces the urea previously reabsorbed in the proximal tubule. Thus, when tubular fluid enters the thick ascending limb, the amount in the lumen is at least as large as the filtered load. Because about 80% of the filtered water has now been reabsorbed, the luminal urea concentration is now several times greater than in the plasma. Beginning with the thick ascending limb and continuing all the way to the medullary collecting ducts (through the distal tubule and cortical collecting ducts), the luminal membrane urea permeability (and the tight junction permeability) is essentially zero. Therefore, a large amount (roughly the filtered load or more) of urea is still within the tubular lumen and flowing from the cortical into the medullary collecting ducts. The concentration is now much greater than in the plasma. Just how much greater depends on hydration status, because, as is discussed in Chapter 6, a variable fraction of the remaining water is reabsorbed in the cortical collecting ducts.
As tubular fluid flows in the collecting-duct system from cortex to medulla, additional water is reabsorbed. Thus, luminal urea concentration rises even more and can easily reach 50 times greater than in plasma. We indicated earlier that the urea concentration in the medullary interstitium is also greater than in plasma, but the luminal concentration is a little higher, so the gradient favors reabsorption in the inner medulla. Therefore, urea is reabsorbed for the second time. In fact, this reabsorbed urea increases medullary interstitial urea concentration and is the source of urea that is secreted into the loop of Henle. Finally, the result is that half the original amount of filtered urea passes into the final urine, an amount that, over the long term, must match hepatic production of urea if the body is to remain in balance for urea. These processes are summarized in Figure 5–3.

**KEY CONCEPTS**

1. The myriad array of organic solutes in the plasma is handled by the kidney; important metabolites are almost completely reabsorbed (saved), whereas waste products are, for the most part, excreted.
2. Most organic solutes are transported transcellularly by a large number of different saturable multiporters ($T_m$ systems).
3. Normal filtered loads of glucose are completely reabsorbed by a sodium-glucose symporter that saturates in conditions of pathological hyperglycemia, leading to the appearance of glucose in the urine.
4. Urea is reabsorbed proximally and recycled between the collecting ducts and loop of Henle in the medulla, resulting in a net excretion of about half the filtered load.

**STUDY QUESTIONS**

5–1. If 50% of a person’s nephrons were destroyed, which of the following compounds would be likely to show increased blood concentration?

A. Urea
B. Creatinine
C. Uric acid
D. Most amino acids
E. Glucose
5–2. The concentration of urea in urine is always much higher than the concentration in plasma. Is this because the overall tubular handling of urea is secretion?

5–3. If the concentration of protein in the glomerular filtrate was 0.005 g/100 mL and none was reabsorbed, how much protein would be excreted per day (assuming a normal GFR)?

5–4. Suppose there is an excessively high filtered load of glucose, and only half is reabsorbed in the proximal tubule. How much of the remaining half now flowing into the loop of Henle is reabsorbed from this point on?
   A. None
   B. About half, leaving one quarter of the filtered load to be excreted
   C. The amount is variable, depending on hydration status

5–5. If you wished to increase your patient’s excretion of quinine, a weak organic base, what change in urinary pH would you try to induce?

5–6. How much of the filtered load of urea remains at the following sites: (a) beginning of the loop of Henle, (b) end of the loop of Henle, and (c) end of the cortical collecting duct?
OBJECTIVES

The student understands the role of different tubular segments in the reabsorption of salt and water.

- Lists approximate percentages of sodium reabsorbed in major tubular segments.
- Lists approximate percentages of water reabsorbed in major tubular segments.

The student understands the role of the proximal tubule in reabsorbing the large filtered load of salt and water.

- Defines the term iso-osmotic volume reabsorption.
- Describes proximal tubule sodium reabsorption, including the functions of the apical membrane sodium entry mechanisms and the basolateral sodium-potassium-adenosine triphosphatase.
- Explains why chloride reabsorption is coupled with sodium reabsorption, and lists the major pathways of proximal tubule chloride reabsorption.

The student understands how the kidney can proliferate either a concentrated or dilute urine.

- States the maximum and minimum values of urine osmolality.
- Defines osmotic diuresis and water diuresis.
- Explains why there is an obligatory water loss.
- Describes the handling of sodium by the descending and ascending limbs, distal tubule, and collecting-duct system.
- Describes the role of sodium-potassium-2 chloride symporters in the thick ascending limb.
- Describes the handling of water by descending and ascending limbs, distal tubule, and collecting-duct system.
- Describes the process of “separating salt from water” and how this permits excretion of either concentrated or dilute urine.
- Describes how antidiuretic hormone affects water reabsorption.
- Describes the characteristics of the medullary osmotic gradient.
- Explains the role of the thick ascending limb, urea recycling, and medullary blood flow in generating the medullary osmotic gradient.
- States why the medullary osmotic gradient is partially “washed out” during a water diuresis.
OVERVIEW

This chapter is devoted to the renal handling of sodium (Na), chloride (Cl), and water. Sodium and chloride are crucial for the body because they are the two solutes of highest concentration in the extracellular fluids, while water is the solvent for all of the body’s dissolved substances and constitutes the majority of the body’s volume. In terms of transport, the transport of water is the simplest. As we pointed out in Chapter 4, “water follows the osmoles.” Thus, much of the description for water transport really amounts to describing solute transport, taking into account the fact that in some regions of the kidney low-water permeability limits the amount of water that follows the osmoles. Transport of chloride is a little more complicated, but is for the most part passive and, because of the constraints of electroneutrality, tied to the transport of sodium. Sodium transport is clearly the most complicated, mostly because it is linked to the transport of so many other substances. However, if we keep in mind the generalized model of epithelial transport developed in Chapter 4 (Figure 4–4) it is not difficult to grasp the key features of sodium transport.

The excretory rates of sodium, chloride, and water can vary over an extremely wide range. For example, some persons may ingest 20–25 g of sodium chloride/day, whereas a person on a low-salt diet may ingest only 0.05 g. The normal kidney can readily alter its excretion of salt over this range. Similarly, urinary water excretion can be varied physiologically from approximately 0.4 L/day to 25 L/day, depending on whether one is lost in the desert or participating in a beer-drinking contest.

Sodium, chloride, and water are all freely filterable at the renal corpuscle. They all undergo considerable tubular reabsorption, usually more than 99%, but normally no tubular secretion. Most of the considerable renal ATP energy expended every day is used to accomplish this enormous reabsorptive task. The major tubular mechanisms for reabsorption of these substances can be summarized by 3 generalizations:

1. The reabsorption of sodium is mainly an active, transcellular process driven mostly by sodium-potassium-adenosine triphosphatase (Na-K-ATPase).
2. The reabsorption of chloride is both passive (paracellular diffusion) and active (transcellular), but it is directly or indirectly coupled with the reabsorption of sodium, thus explaining why the reabsorption of the 2 ions usually occurs in parallel. When describing the reabsorption of sodium, a parallel reabsorption of chloride is usually implied.
3. The reabsorption of water is by osmosis and is secondary to reabsorption of solutes, particularly sodium and substances whose reabsorption is dependent on sodium reabsorption (mostly chloride).

**Sodium Reabsorption**

Table 6–1 is a balance sheet for sodium chloride. Clearly, the major route of salt excretion from the body under normal circumstances is via the kidneys. The large amount excreted should not obscure the fact that nearly all the filtered sodium
and chloride is reabsorbed. Table 6–2 summarizes the approximate quantitative contribution of each tubular segment to sodium reabsorption. In an individual with an average salt intake, the proximal tubule reabsorbs 65% of the filtered sodium, the thin and thick ascending limbs of Henle’s loop 25%, and the distal convoluted tubule and collecting-duct system most of the remaining 10%, so that the final urine contains less than 1% of the total filtered sodium. As discussed in Chapter 7, reabsorption at several of these tubular sites is under physiological control by neural, hormonal, and paracrine signals, so that the exact amount of sodium excreted is homeostatically regulated. Because so much sodium is filtered, even a small percentage change in reabsorption results in a relatively large change in excretion.

An absolutely crucial generalization is this: In all nephron segments, the essential event for active transcellular sodium reabsorption is the primary active transport of sodium from cell to interstitial fluid by the Na-K-ATPase pumps in the basolateral membrane. These pumps keep the intracellular sodium concentration lower than in the surrounding media. Because the inside of the cell is negatively charged with respect to the lumen, luminal sodium ions enter the cell passively, down their electrochemical gradient. (In the proximal tubule, working in parallel with the Na-K-ATPase is a sodium-bicarbonate antiporter that actively extrudes one sodium ion in symport with three bicarbonate ions. But as described in Chapter 9, this process ultimately depends on the activity of the Na-K-ATPase pumps.)

On examination of the luminal membrane depicted in Figure 6–1, note that there are various types of entry processes for sodium: Na-hydrogen antiporters (mostly NHE-3 isoform), Na-nutrient, Na-phosphate, or Na-sulfate symporters; and sodium channels. Quantitatively, the Na-H antiporters bring in the majority of the sodium (and serve as a major site for regulating sodium reabsorption in the proximal tubule). In other chapters, you will learn other roles for particular segments besides reabsorption of sodium. For example, as discussed in Chapter 5, the proximal tubule reabsorbs nutrients, and the active step in this process is often by symport with sodium across the luminal membrane.
Chloride Reabsorption

Because chloride reabsorption is dependent on sodium reabsorption, the tubular locations that reabsorb chloride and the percentages of filtered chloride reabsorbed by these segments are similar to those for sodium (see Table 6–1). When examining chloride reabsorption, it is helpful to keep in mind the absolute constraint of electroneutrality: any finite volume of fluid reabsorbed must contain equal amounts of anion and cation equivalents. Let us do a “guesstimate” calculation.

One liter of normal filtrate contains 140 mEq of sodium, and thus must contain about 140 mEq of anions, mainly chloride (110 mEq) and bicarbonate (24 mEq). (We say “about” because there are other cations [eg, potassium and calcium] and anions [eg, sulfate and phosphate] that must factor into the calculation to achieve an exact balance, but their contributions are much smaller than sodium, chloride, and bicarbonate.) If 65% of the filtered sodium is reabsorbed in the proximal tubule, 0.65 X 140 = 91 mEq of sodium in each liter of filtrate are reabsorbed. Therefore, about 91 mEq of some combination of chloride and bicarbonate must also be reabsorbed to accompany this sodium. As described in Chapter 9, about 90% of the filtered bicarbonate is reabsorbed in the proximal tubule (0.9 X 24 = 22). This leaves 91 – 22 = 69 mEq of chloride that must be reabsorbed in the proximal tubule. This is more than 60% of the filtered chloride and almost as much as the fractional reabsorption of sodium and water.

To understand active transcellular chloride reabsorption, it is necessary to recognize that the critical transport step for chloride is from lumen to cell. The chloride transport process in the luminal membrane must achieve a high enough intracellular chloride concentration to cause downhill chloride movement out of the cell across the basolateral membrane (of course, the movement of chloride across the basolateral membrane is also promoted by the negative potential within the cell). Thus, luminal membrane chloride transporters serve essentially the same function for chloride that the basolateral membrane Na-K-ATPase pumps do for sodium: They use energy to move chloride uphill from lumen to cell against its electrochemical gradient.

Table 6–2. Comparison of sodium and water reabsorption along the tubule

<table>
<thead>
<tr>
<th>Tubular segment</th>
<th>Percent of filtered load reabsorbed (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sodium</td>
</tr>
<tr>
<td>Proximal tubule</td>
<td>65</td>
</tr>
<tr>
<td>Descending thin limb of Henle’s loop</td>
<td>—</td>
</tr>
<tr>
<td>Ascending thin limb and thick</td>
<td>25</td>
</tr>
<tr>
<td>ascending limb of Henle’s loop</td>
<td></td>
</tr>
<tr>
<td>Distal convoluted tubule</td>
<td>5</td>
</tr>
<tr>
<td>Collecting-duct system</td>
<td>4–5</td>
</tr>
</tbody>
</table>
According to the luminal membrane depicted in Figure 6–1, the major routes are (1) paracellular absorption and (2) a complicated parallel set of Na-H and Cl-base antiporters (described later). These mechanisms are dependent on sodium movement across the membrane and are, therefore, linked to sodium reabsorption.

**Water Reabsorption**

With a large water load, the renal response is to produce a large-volume, very dilute urine (osmolality much lower than in blood plasma). In contrast, during a state of dehydration, the urine volume is low and very concentrated (ie, the urine osmolality is much greater than in blood plasma). That the urine osmolality is so variable brings us to a crucial aspect of renal function. Terrestrial animals must be able to independently control excretion of salt and water, because their ingestion and loss is not always linked (see Tables 6–1 and 6–3). To excrete water in excess of salt and vice versa (ie, produce a range of urine osmolalities), the kidneys must be able to separate the reabsorption of solute from the reabsorption of water, ie, to “separate salt from water.” How do they do this?
Regardless of hydration state the collective actions of renal tubular segments before the cortical collecting tubule reabsorb more solute than water. This leaves a large volume of dilute tubular fluid (~110 mOsm/kg H₂O) entering the limited segment of the cortical collecting tubule. If an individual is overhydrated and, therefore, requires maximum water excretion, most of this water simply passes through the collecting-duct system to appear in the urine, with only limited further reabsorption. In contrast, when an individual is dehydrated, water excretion should be low. The vast majority of this dilute water is reabsorbed, leaving a low volume of concentrated final urine.

A balance sheet for total body water is given in Table 6–3. These are average values, which are subject to considerable variation. The two sources of body water are metabolically produced water, resulting largely from the oxidation of carbohydrates, and ingested water, obtained from liquids and so-called solid food (eg, a rare steak is approximately 70% water). There are several sites from which water is always lost to the external environment: skin, lungs, gastrointestinal tract, and kidneys. Menstrual flow and, in lactating women, breast milk constitute two other potential sources of water loss in women.

The loss of water by evaporation from the cells of the skin and the lining of respiratory passageways is a continuous process, often referred to as insensible loss because people are unaware of its occurrence. Additional water evaporates from the skin during the production of sweat. Fecal water loss is normally quite small but can be severe in diarrhea. Gastrointestinal loss can also be large during severe bouts of vomiting.

Water reabsorption always occurs in the proximal tubule (65% of the filtered water), descending thin limb of Henle’s loop (10%), and collecting-duct system

### Table 6–3. Normal routes of water gain and loss in adults

<table>
<thead>
<tr>
<th>Route</th>
<th>mL/day</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Intake</strong></td>
<td></td>
</tr>
<tr>
<td>Beverage</td>
<td>1200</td>
</tr>
<tr>
<td>Food</td>
<td>1000</td>
</tr>
<tr>
<td>Metabolically produced</td>
<td>350</td>
</tr>
<tr>
<td>Total</td>
<td>2550</td>
</tr>
<tr>
<td><strong>Output</strong></td>
<td></td>
</tr>
<tr>
<td>Insensible loss (skin and lungs)</td>
<td>900</td>
</tr>
<tr>
<td>Sweat</td>
<td>50</td>
</tr>
<tr>
<td>In feces</td>
<td>100</td>
</tr>
<tr>
<td>Urine</td>
<td>1500</td>
</tr>
<tr>
<td>Total</td>
<td>2550</td>
</tr>
</tbody>
</table>
Basic Renal Processes for Sodium, Chloride, and Water (where the fractional reabsorption is the most variable). A comparison of water and sodium reabsorption (see Table 6–2) reveals several important points. First, sodium and water reabsorption occur in the proximal tubule to the same extent. Second, both are also reabsorbed in Henle’s loop, but not in equal proportions. The part of the loop involved in water reabsorption is different from that for sodium reabsorption, and the fraction of sodium reabsorbed by the loop as a whole is always greater than that of water (i.e., the loop overall is a site where salt is reabsorbed and excess water is left in the lumen of the nephron: “separating salt from water”). Third, sodium reabsorption, but not water reabsorption, occurs in the distal convoluted tubule. Fourth, both occur in the collecting-duct system, but the percentages of sodium and water reabsorbed in the collecting-duct system vary enormously depending on a number of factors.

The movement of water down an osmotic gradient can occur by several routes: simple net diffusion through the lipid bilayer, through aquaporins in plasma membranes of the tubular cells, and through the tight junctions between the cells. The amount of water that moves for a given osmotic gradient and its route depends on the water permeability of the different cellular components. The basolateral membranes of all renal cells are quite permeable to water due to the presence of protein aquaporins that act as water pores: As a result, the cytosolic osmolality is always close to that of the surrounding interstitium. It is the luminal membrane and tight junctions where most of the variability lies. The segments of the renal tubule fall into 3 general categories with regard to water permeability: (1) The luminal membranes of the proximal tubule and descending thin limb of Henle’s loop always have a high water permeability; (2) the luminal membrane of the ascending limbs of Henle’s loop (both thin and thick; recall from Chapter 1 that only long loops have ascending thin limbs) and the luminal membranes of distal convoluted tubule are always relatively water impermeable, as are the tight junctions; and (3) the water permeability of the luminal membrane of the collecting-duct system is intrinsically low but can be regulated so that its water permeability increases substantially. These differences in water permeability account for the sites of water reabsorption as well as the large range of water reabsorptions given for the collecting-duct system in Table 6–2.

The ability of the kidneys to produce low-volume hyperosmotic urine is a major determinant of one’s ability to survive without water. The human kidney can produce a maximal urinary concentration of 1400 mOsm/kg in extreme dehydration. This is almost 5 times the osmolality of plasma. The sum of the urea, sulfate, phosphate, other waste products, and a small number of nonwaste ions excreted each day normally averages approximately 600 mOsm/day. Therefore, the minimal volume of water in which this mass of solute can be dissolved is roughly 600 mmol/1400 mOsm/L = 0.43 L/day.

This volume of urine is known as the obligatory water loss. It is not a strictly fixed volume but changes with different physiological states. For example,

\[^{1}\text{In this calculation, osmolarity is used as an approximation to osmolality to simplify the math.}\]
increased tissue catabolism, as during fasting or trauma, releases excess solute and so increases obligatory water loss.

The obligatory water loss contributes to dehydration when a person is deprived of water intake. For example, if we could produce urine with an osmolarity of 6000 mOsm/L, the obligatory water loss would only be 100 mL of water, and survival time would be greatly increased. A desert rodent, the kangaroo rat, does just that. This animal does not need to drink water because the water content of its food and the water produced by metabolism of the foods is sufficient to meet its needs.²

INDIVIDUAL TUBULAR SEGMENTS

Because we know that water reabsorption is driven by osmolality differences across the epithelium of water-permeable tubular segments, our major task in reviewing the individual tubular segments is to describe how these transtubular osmolality differences arise. We also explain how the kidneys generate the osmolality differences by separating salt from water and form a hypo-osmotic or hyperosmotic urine.

The important principles to be understood regarding individual tubular segments are how the reabsorption of sodium, chloride, and water are related to one another and how the amount of reabsorption quantitatively varies from one segment to another.

Proximal Tubule

As shown in Figure 6–1, several luminal entry steps are involved in the active transepithelial reabsorption of sodium in the proximal tubule. In the early portion (the proximal convoluted tubule), a large fraction of the filtered sodium enters the cell across the luminal membrane via antiport with protons. As described in Chapter 9, these protons, which are supplied by carbon dioxide and water, cause the secondary active reabsorption of filtered bicarbonate. Therefore, in the early proximal tubule, bicarbonate is a major anion reabsorbed with sodium, and the luminal bicarbonate concentration decreases markedly (Figure 6–2). Organic nutrients and phosphate are also absorbed with sodium, and their luminal concentrations decrease rapidly. A major percentage of chloride reabsorption in the proximal tubule occurs via paracellular diffusion. The concentration of chloride in Bowman’s capsule is, of course, essentially the same as in plasma (about 110 mEq/L). Along the early proximal tubule, however, the reabsorption of water, driven by the osmotic gradient created by the reabsorption of sodium plus its cotransported solutes and bicarbonate, causes the chloride concentration in the tubular lumen to increase somewhat above that in the peritubular capillaries (see Figure 6–2). Then, as the fluid flows through the middle and late proximal tubule, this concentration

²The obligatory solute excretion explains why a thirsty sailor cannot drink sea water, even if the urine osmolality is slightly greater than that of the sea water. To excrete all the salt in 1 L of sea water (to prevent a net gain of salt) plus the obligatory organic solutes produced by the body, the volume of urine would have to be much greater than 1 L.
gradient, maintained by continued water reabsorption, provides the driving force for paracellular chloride reabsorption by diffusion.

There is also an important component of active chloride transport from lumen to cell in the later proximal tubule. As illustrated in Figure 6–1, it uses parallel Na-H and Cl-base antiporters. Chloride transport into the cell is powered by the downhill antiport of organic bases (including formate and oxalate), which are continuously generated in the cell by dissociation of their respective acids into a proton and the base. Simultaneously, the protons generated within the cell by the dissociation of the acids are actively transported into the lumen by Na-H antiporters. In the lumen, the protons and organic bases recombine to form the acid, which is a neutral molecule. This nonpolar neutral acid then diffuses across the luminal membrane back into the cell, where the entire process is repeated. Thus, the overall achievement of the parallel Na-H and Cl-base antiporters is the same as though the Cl and Na were simply cotransported into the cell together.

Figure 6–2. Changes in tubular fluid composition along the proximal convoluted tubule. Values below 1.0 indicate that relatively more of the substance than water has been reabsorbed. Values above 1.0 indicate that relatively less of the substance than water has been reabsorbed. The concentrations of inorganic phosphate, bicarbonate, glucose, and lactate all rapidly decrease in the proximal tubule because these substances are actively reabsorbed much more rapidly than water. This is because these substances are preferentially reabsorbed with sodium in the early proximal tubule. In contrast, the concentration of chloride increases because chloride reabsorption lags behind sodium and, hence, water reabsorption in the early proximal tubule. TF, concentration of the substance in tubular fluid; P, its concentration in arterial plasma. (Modified from Rector FC, Am J Physiol 1983;249:F461; Maddox DA, Gennari JF, Am J Physiol 1987;252:F573.)
Importantly, the recycling of protons and base means that most of the protons are not acidifying the lumen but are simply combining with the base and moving back into the cells. It should also be recognized that everything is ultimately dependent on the basolateral membrane Na-K-ATPases to establish the gradient for sodium that powers the luminal Na-H antiporter.

Regarding water reabsorption, the proximal tubule, as mentioned, has a very high permeability to water. This means that very small differences in osmolality (less than 1 mOsm/L) will suffice to drive the reabsorption of very large quantities of water, normally about 65% of the filtered water. This osmolality difference is created by the reabsorption of solute. The osmolality of the freshly filtered tubular fluid at the very beginning of the proximal tubule is, of course, essentially the same as that of plasma and interstitial fluid. Then, as solute is reabsorbed from the proximal tubule, the movement of this solute out of the lumen lowers luminal osmolality (ie, raises water concentration) compared with interstitial fluid. Simultaneously, it also tends to raise the interstitial fluid osmolality. (Interstitial osmolality rises only a small amount because the high perfusion through peritubular capillaries keeps the interstitial osmolality close to the plasma value.) The osmotic gradient from lumen to interstitial fluid causes osmosis of water from the lumen across the plasma membranes via aquaporins and tight junctions into the interstitial fluid. The Starling forces across the peritubular capillaries in the interstitium favor reabsorption, as explained in Chapter 4, and so the water and solutes then move into the peritubular capillaries and are returned to the general circulation.

The term solute was used to describe how reabsorption creates an osmolality difference between lumen and interstitial fluid. It should be clear by now, however, that we could just as well have referred simply to “sodium” because the reabsorption of virtually all solutes by the proximal tubule is dependent directly or indirectly on the reabsorption of sodium (Table 6–4). In other words, sodium and solutes whose reabsorption is coupled in one way or another with sodium

### Table 6–4. Summary of mechanisms by which reabsorption of sodium drives reabsorption of other substances in the proximal tubule

<table>
<thead>
<tr>
<th>Reabsorption of sodium</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Creates transtubular osmolality difference, which favors reabsorption of water by osmosis; in turn, water reabsorption concentrates many luminal solutes (eg, chloride and urea), thereby favoring their reabsorption by diffusion.</td>
</tr>
<tr>
<td>2 Achieves reabsorption of many organic nutrients, phosphate, and sulfate by cotransport across the luminal membrane.</td>
</tr>
<tr>
<td>3 Achieves secretion of hydrogen ion by countertransport across the luminal membrane; these hydrogen ions are required for reabsorption of bicarbonate (as described in Chapter 9).</td>
</tr>
<tr>
<td>4 Achieves reabsorption of chloride by indirect cotransport across the luminal membrane (the parallel Na/H and Cl/base countertransporters).</td>
</tr>
</tbody>
</table>
reabsorption constitute the overwhelming majority of all solutes reabsorbed. Thus, the terms sodium reabsorption and total solute reabsorption are almost interchangeable when considering the proximal tubule.

Given the tremendous amount of sodium reabsorbed, how can the luminal sodium concentration and osmolality not progressively decrease along the proximal tubule? As shown in Figure 6–2, both remain almost equal to their values in plasma. Actually, the luminal values are slightly lower than the plasma values, but the difference is usually too small to detect. Remember that we are dealing here with concentrations of sodium and total solute (osmolality). Whereas 65% of the mass of filtered sodium and total solute has been reabsorbed by the end of the proximal tubule, so has virtually the same percentage of filtered water. This is because the water permeability of the proximal tubule is so great that passive water reabsorption always keeps pace with total solute reabsorption. Therefore, the concentrations of sodium and total solute (osmolality), as opposed to their masses, remain virtually unchanged during fluid passage through the proximal tubule. This process, therefore, is called iso-osmotic volume reabsorption.

A good example of what happens when tight coupling between proximal sodium and water reabsorption is disrupted is the phenomenon known as osmotic diuresis. The term diuresis simply means increased urine flow, and osmotic diuresis denotes the situation in which the increased urine flow is due to an abnormally high amount of any substance in the glomerular filtrate that is reabsorbed incompletely or not at all by the proximal tubule. As water reabsorption begins in this segment, secondary to sodium reabsorption, the concentration of any unreabsorbed solute rises, and its osmotic presence retards the further reabsorption of water here (and downstream as well). Moreover, the failure of water to follow sodium causes the sodium concentration in the proximal tubular lumen to fall slightly below that in the interstitial fluid. This concentration difference, even though small, drives a net passive diffusion of sodium across the epithelium (mostly the tight junctions) back into the lumen (remember that the proximal tubule is a “leaky” epithelium and sodium transport is a gradient-limited system), resulting in more sodium than usual in the lumen and passing on to the loop of Henle. Thus, osmotic diuretics inhibit the reabsorption of both water and sodium (as well as other ions). Osmotic diuresis can occur in persons with uncontrolled diabetes mellitus; the filtered load of glucose exceeds the tubular maximum \( T_m^G \) for this substance, and the unreabsorbed glucose then acts as an osmotic diuretic.

**Henle’s Loop**

As stated earlier (see also Table 6–2), Henle’s loop, taken as a whole, always reabsorbs proportionally more sodium and chloride (about 25% of the filtered loads) than water (10% of the filtered water). This is a key difference from the proximal tubule, which always reabsorbs water and sodium in essentially equal proportions.

Also as shown in Table 6–2, the reabsorption of sodium chloride and water occur in different places. The descending limb does not reabsorb sodium or chloride significantly, but it is quite permeable to water and reabsorbs it. In contrast,
the ascending limbs (both thin and thick) reabsorb sodium and chloride but little water (because they are quite impermeable to water).

What are the mechanisms of sodium and chloride reabsorption by the ascending limbs? These are mainly passive in the thin ascending limb and active in the thick ascending limb. Water reabsorption in the descending limb (see later discussion) concentrates luminal sodium and creates a favorable gradient for passive sodium reabsorption. The epithelium of the thin ascending limb permits this gradient to drive reabsorption, probably by the paracellular route. As tubular fluid enters the thick ascending limb, the transport properties of the epithelium change again, and active processes become dominant. As shown in Figure 6–3, the major luminal entry step for the sodium and chloride in this segment is via the Na-K-2Cl symporter (NKCC2 isoform). This symporter is the target for a major class of diuretics collectively known as the loop diuretics, which include the drugs furosemide (Lasix) and bumetanide. The luminal membrane of this segment also has a Na-H antiporter isoform, which, like the isoform in the proximal tubule, provides another mechanism for sodium movement into the cell.

Figure 6–3. Major transport pathways for sodium and chloride in thick ascending limb cells within the loop of Henle. The major transporter in the thick ascending limb is the Na-K-2Cl symporter (NKCC), which is the target for inhibition by loop diuretics like furosemide and bumetanide. In addition to NKCC, the cells contain potassium channels that recycle potassium from the cell interior to the lumen and to the interstitium (see Chapter 8). Besides transcellular routes, some sodium also moves paracellularly in response to the lumen positive potential. The apical membranes and tight junctions have a very low water permeability. Because the cells reabsorb salt, but not water, the thick ascending limb is the point in the nephron at which salt is separated from water. This ultimately allows water excretion and salt excretion to be controlled independently. Defects in NKCC, the recycling potassium channel, and the basolateral chloride channel lead, respectively, to the 3 different types of Bartter’s syndrome. ATP, adenosine triphosphate.
The Na-K-2Cl symporter requires that equal amounts of potassium and sodium be transported. However, there is far less potassium in the lumen than sodium, and it seems that the lumen would be depleted of potassium long before very much sodium was reabsorbed. Interestingly, the luminal membrane has a large number of potassium channels that allow much of the potassium transported into the cell on the Na-K-2Cl symporter to leak back (as described in Chapter 8, potassium recycles between the cytosol and lumen in order to be available for symport with sodium and chloride). Thus, under normal circumstances, luminal potassium does not limit sodium and chloride reabsorption through Na-K-2Cl symporters.

In addition to the active transcellular reabsorption of sodium, a large percentage (perhaps as much as 50%) of total sodium reabsorption in this segment occurs by paracellular diffusion. There is a high paracellular conductance for sodium in the thick ascending limb, and the luminal potential in this segment is positive, a significant driving force for cations. (We see in later chapters that this paracellular pathway also allows substantial reabsorption of potassium and calcium as well.) However, none of this would work without the continuous operation of the Na-K-ATPase in the basolateral membrane.

To summarize the most important feature of the loop of Henle, the descending limb reabsorbs water but not sodium chloride, whereas the ascending limb reabsorbs sodium chloride but not water, with the net of the loop as a whole being reabsorption of more salt than water. The ascending limb is called a diluting segment, and because Henle's loop as a whole has reabsorbed more solute than water, the fluid leaving the loop to enter the distal convoluted tubule is hypo-osmotic (more dilute) compared with plasma.

**Distal Convoluted Tubule**

The major luminal entry step in the active reabsorption of sodium and chloride by the distal convoluted tubule is via the Na-Cl symporter (Figure 6–4). The characteristics of this transporter differ significantly from the TAL Na-K-2Cl symporter and are sensitive to different drugs. In particular, the Na-Cl symporter is blocked by the thiazide diuretics, including hydrochlorothiazide. (Sodium channels, like those in the collecting tubule principal cells, also exist in the distal convoluted tubule.)

**Collecting-Duct System**

In the collecting ducts, there is a division of labor among several different cell types. Reabsorption of sodium and water is associated with principal cells (so called because they make up approximately 70% of the cells; Figure 6–5). Principal cells also play a major role in maintaining potassium homeostasis (see Chapter 8). Reabsorption of chloride can occur partially via paracellular pathways, but active

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3Besides sodium and chloride reabsorption, distal convoluted tubule cells are a major site for the control of calcium homeostasis because they have apical calcium channels that are regulated by parathyroid hormone (see Chapter 10).
Figure 6–4. Major transport pathways for sodium and chloride in the distal convoluted tubule. The apical membrane contains the Na-Cl symporter (NCC), which is the target for inhibition by thiazide diuretics. There is also some sodium reabsorption via apical sodium channels (ENaCs). The apical membranes and tight junctions have a very low water permeability. A defect in NCC leads to Gitelman’s syndrome. ATP, adenosine triphosphate.

Figure 6–5. Major transport pathways for sodium, chloride, and water in principal cells of the cortical collecting duct (CCD). The principal cells are the major cell type in the CCD. Sodium reabsorption is via apical sodium channels (ENaC). Activity of ENaC is controlled by the hormone aldosterone (see Figure 7–11 and associated text in Chapter 7). Chloride reabsorption is passive via the paracellular pathway. Water reabsorption is via aquaporins, the activity of which is controlled by the antidiuretic hormone (ADH). ATP, adenosine triphosphate.
reabsorption is also associated with another class of collecting duct cells, the intercalated cells (see Figure 9–3).4

The principal cells reabsorb sodium; the luminal entry step is via epithelial sodium channels. Regulation of this entry step is enormously important for whole-body physiology, and we expand on this topic in Chapter 7. Some sodium chloride reabsorption continues in the medullary collecting ducts, probably via some form of epithelial sodium channels.

What about water reabsorption in tubular segments beyond the loop of Henle? The water permeability of the distal convoluted tubule is always very low and unchanging, similar to that of the ascending limbs of Henle’s loop. Accordingly, as fluid flows through the distal convoluted tubule and sodium chloride reabsorption proceeds, virtually no water is reabsorbed. The result is that the already hypo-osmotic fluid entering the distal convoluted tubule from the thick ascending limb of Henle’s loop becomes even more hypo-osmotic. Thus, the distal convoluted tubule, like the ascending limbs of Henle’s loop, functions as a diluting segment and further separates salt from water.

In contrast, the water permeability of the collecting-duct system—both the cortical and medullary portions—is subject to physiological control by circulating antidiuretic hormone (ADH; see Figure 6–5). The inner medullary collecting duct has at least a finite water permeability even in the absence of ADH, but the outer medullary and cortical regions have a vanishingly low water permeability without ADH.

Depending on levels of ADH, therefore, water permeability for most of the collecting-duct system can be very low, very high, or everything in between. When water permeability is very low, the hypo-osmotic fluid entering the collecting-duct system from the distal convoluted tubule remains hypo-osmotic as it flows along the ducts. When this fluid reaches the medullary portion of the collecting ducts, there is now a huge osmotic gradient favoring reabsorption, which occurs to some extent. That is, although there is little cortical water reabsorption without ADH (where most distal absorption normally occurs), there is still a finite medullary absorption because of the enormous osmotic gradient. However, because there is such a high tubular volume (ie, it was not reabsorbed in the cortex), most of the water entering the medullary collecting duct flows on to the ureter. The result is the excretion of a large volume of very hypo-osmotic (dilute) urine, or water diuresis.

In water diuresis, the last tubular segment to reabsorb large amounts of water is the descending limb of Henle’s loop; in all later segments reabsorption of solute (mainly sodium chloride) continues, but water reabsorption is minimal (although not zero in the inner medulla). Note that even when very little water reabsorption occurs beyond the loop of Henle, the reabsorption of sodium is not retarded to any great extent. Therefore, the intraluminal sodium concentration can be lowered almost to zero in these tubular segments, and the osmolality can approach

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4Different types of intercalated cells, besides mediating chloride reabsorption, also play a significant role in maintaining acid-base homeostasis (see Chapter 9).
50 mOsm/kg. (This is possible because these tubular segments are “tight” epithelia, and there is very little back-leak of sodium from interstitium to tubular lumen despite the large electrochemical gradient favoring diffusion.)

What happens when the collecting-duct system’s water permeability is very high instead of very low? As the hypo-osmotic fluid entering the collecting-duct system from the distal convoluted tubule flows through the cortical collecting ducts, water is rapidly reabsorbed. This is because of the large difference in osmolality between the hypo-osmotic luminal fluid and the iso-osmotic (285 mOsm/kg) interstitial fluid of the cortex. In essence, the cortical collecting duct is reabsorbing the large volume of water that did not accompany solute reabsorption in the ascending limbs of Henle’s loop and distal convoluted tubule. In other words, the cortical collecting duct reverses the dilution carried out by the diluting segments. Once the osmolality of the luminal fluid approaches that of the interstitial fluid, the cortical collecting duct then behaves analogously to the proximal tubule, reabsorbing approximately equal proportions of solute (mainly sodium chloride) and water. The result is that the tubular fluid, which leaves the cortical collecting duct to enter the medullary collecting duct, is iso-osmotic compared with cortical plasma, and its volume is greatly reduced compared with the amount entering from the distal tubule.

In the medullary collecting duct, solute reabsorption continues but water reabsorption is proportionally even greater. In other words, the tubular fluid becomes more and more hyperosmotic and reduced in volume in its passage through the medullary collecting ducts because the interstitial fluid of the medulla is very hyperosmotic, for reasons discussed later.

How does ADH convert epithelial water permeability from very low to very high? An alternative name for ADH is vasopressin, because the hormone can constrict arterioles and thus increase arterial blood pressure, but ADH’s major renal effect is antidiuresis (ie, “against a high urine volume”). In the absence of ADH, the water permeability of the cortical and outer medullary collecting duct is very low, and little if any water is reabsorbed from these segments, resulting in water diuresis. On the other hand, in the presence of high plasma concentrations of ADH, the water permeability of all regions of the collecting ducts is great, and only a small volume of maximally hyperosmotic urine is excreted. The tubular response to ADH is not all or none, however, but shows graded increases as the plasma concentration of ADH is increased over a certain range, thus permitting fine adjustments of collecting-duct water permeability and, hence, water reabsorption. (The control of ADH secretion is described in Chapter 7.) ADH acts in the collecting ducts on the principal cells, the same cells that reabsorb sodium (and, as is seen in Chapter 8, secrete potassium). The renal receptors for ADH (vasopressin type 2 receptors) are in the basolateral membrane of the principal cells and are different from the vascular receptors (vasopressin type 1). The binding of ADH by its receptors results in the activation of adenylate cyclase, which catalyzes the intracellular production of cyclic adenosine monophosphate. This second messenger then induces, by a sequence of events, the migration of intracellular vesicles to, and their fusion with, the luminal membrane. Recall from Chapter 4, this is one
of the ways of regulating membrane permeability. The vesicles contain an isoform of a water channel protein, aquaporin 2, through which water can move, so the luminal membrane becomes highly permeable to water. In the absence of ADH, the aquaporins are withdrawn from the luminal membrane by endocytosis. (As stated earlier, the water permeability of the basolateral membranes of renal epithelial cells is always high because of the constitutive presence of other aquaporin isoforms; thus, the permeability of the luminal membrane is rate limiting.)

**URINARY CONCENTRATION: THE MEDULLARY OSMOTIC GRADIENT**

The kidneys can produce urine that is hypo-osmotic, iso-osmotic, or hyperosmotic. The production of hypo-osmotic urine is, we hope by now, an understandable process: The tubules (particularly the thick ascending limb of Henle’s loop) reabsorb relatively more solute than water, and the dilute fluid that remains in the lumen is excreted. The production of hyperosmotic urine is also straightforward in that reabsorption of water from the lumen into a hyperosmotic interstitium concentrates that luminal fluid, leaving a concentrated urine to be excreted. The question is, how do the kidneys generate a hyperosmotic medullary interstitium? Not only is the medullary interstitium hyperosmotic, but there is a gradient of osmolality, increasing from a nearly iso-osmotic value at the corticomedullary border, to a maximum of greater than 1000 mOsm/kg at the papilla. This peak value is not rigidly fixed; it is a variable that changes depending on conditions. It is highest during periods of water deprivation and dehydration, when urinary excretion is lowest, and is “washed out” to only about half of that during excess hydration and urinary excretion is high. Some aspects of how the kidneys generate a medullary osmotic gradient are still uncertain. However, the essential points are clear, and it is these essential points on which we now focus.

We should first differentiate between the development of the medullary osmotic gradient, as opposed to its maintenance once established. In the steady state, there must be mass balance, ie, every substance that enters the medulla via tubule or blood vessel must leave the medulla via tubule or blood vessel. However, during development of the gradient there are transient accumulations of solute, and during washout of the gradient there are losses. In describing the medullary osmotic gradient, it is easiest conceptually to start from a condition in which there is no gradient, and then follow its development over time. The main components of the system that develops the medullary osmotic gradient are (1) active NaCl transport by the thick ascending limb; (2) the unusual arrangement of blood vessels and nephron segments in the medulla, with descending components in close apposition to ascending components; and (3) the recycling of urea between the medullary collecting ducts and the deep portions of the loops of Henle (Figure 5–3).

To develop the osmotic gradient in the medullary interstitium, there must be deposition of solute in excess of water. It is reabsorption of sodium and chloride by the thick ascending limb that accomplishes this task. At the junction between the
inner and outer medulla, the ascending limbs of all loops of Henle, whether long or short, turn into thick regions and remain thick all the way back until they reach the original Bowman’s capsules from which they arose in the cortex. As they re-absorb solute without water and dilute the luminal fluid, they simultaneously add solute without water to the surrounding interstitium. This action of the thick ascending limb is absolutely essential and is the key to everything else that happens. If transport in the thick ascending limb is inhibited (by loop diuretics that block the Na-K-2Cl symporter), then the lumen is not diluted and the interstitium is not concentrated, and the urine becomes iso-osmotic. For those portions of the thick ascending limb in the cortex, the reabsorbed solute simply mixes with material re-absorbed by the nearby proximal convoluted tubules. Because the cortex contains abundant peritubular capillaries and a high blood flow, the reabsorbed material immediately moves into the vasculature and returns to the general circulation. However, in the medulla, the vascular anatomy and blood flow are quite different, and reabsorbed solute that is deposited in the outer medullary interstitium during the establishment of the osmotic gradient is not immediately removed, ie, it accumulates. The degree of accumulation is a function of the arrangement of the vasa recta, their permeability properties and the volume of blood flowing within them.

As described in Chapter 1, blood enters and leaves the outer medulla through parallel bundles of descending and ascending vasa recta. These vessels are permeable to sodium and they take up most of the sodium that is being transported by the thick ascending limbs into the interstitium of the outer medulla. The ascending vasa recta return sodium to the general circulation, but the descending vessels distribute it down into the inner medulla, where it diffuses out across the endothelia of the vasa recta and the interbundle capillaries that they feed, thereby raising the sodium content (and osmolality) throughout the medulla. It is here that the anatomy of the vasculature becomes particularly important. If medullary blood with its somewhat elevated sodium concentration simply flowed into a venous drainage system, very little additional increase in sodium concentration would occur. However, the interbundle capillaries drain into ascending vasa recta that lie right next to descending vasa recta. The walls of the ascending vasa recta are fenestrated, allowing rapid and thorough equilibration of water and small solutes between plasma and interstitium. As the total sodium content of the medulla rises, blood in the ascending vessels takes on an increasingly higher sodium concentration, while blood entering the medulla always has a normal sodium concentration (about 140 mEq/L). Accordingly, some of the medullary sodium begins to re-circulate, diffusing out of ascending vessels and reentering nearby descending vessels. The process of crossing between ascending and descending vessels is called countercurrent exchange. At this point, sodium is entering descending vasa recta from two sources—re-circulated sodium from the ascending vasa recta and new sodium from the thick ascending limbs. Over time, the concentration of sodium in the ascending vessels and medullary interstitium rises until a steady state is reached in which the amount of new sodium entering the interstitium from thick ascending limbs matches the amount of sodium leaving the interstitium.
in ascending vasa recta and returning to the general circulation. At its peak, the concentration of sodium in the inner medulla may reach 300 mEq/L, more than double its value in the general circulation. As sodium is accompanied by an anion, mostly chloride, the contribution of salt to the medullary osmolality is about 600 mOsm/kg.

What happens to water in the medulla during this time? Although solute can accumulate without a major effect on renal volume, the amount of water in the medullary interstitium must remain relatively constant; otherwise, the medulla would undergo significant swelling or shrinking. The endothelial cells of descending vasa recta, although not as leaky as the fenestrated endothelium of ascending vasa recta, contain aquaporins, allowing water to be drawn osmotically into the medullary interstitium by the high salt content in a manner similar to water being drawn out of tubular elements. This loss of water from descending vasa recta decreases the plasma volume of blood penetrating deeper into the medulla and raises its osmolality, thereby reducing the tendency to dilute the inner medullary interstitium. Water leaving descending vessels diffuses across to nearby ascending vasa recta and is removed from the medulla. Just as there is countercurrent exchange of solute between descending and ascending vessels, there is countercurrent exchange of water. In descending vessels water leaves and solute enters, while in ascending vessels water enters and solute leaves (see Figures 1–6 and 6–6). Water also enters the medullary interstitium by reabsorption from thin descending limbs and from medullary collecting ducts. As there is no water secretion by the tubules, all water entering the medullary interstitium from tubules and descending vasa recta must leave the medulla via ascending vasa recta.

The magnitude of blood flow in the vasa recta is a crucial variable. If blood flow is very high, water from the isosmotic plasma entering the medulla in descending vasa recta dilutes the hyperosmotic interstitium. But medullary blood flow is always low, and even lower in conditions where medullary osmolality is highest. Therefore, the diluting effect of water diffusing out of descending vasa recta during periods of minimal blood flow is not great.\(^5\)

There is one additional major player involved in the development of the medullary osmotic gradient—urea. As indicated above, the peak osmolality in the renal papilla reaches over 1000 mOsm/kg. About half of this is accounted for by sodium and chloride, and most of the rest is (500–600 mOsm/kg) accounted for by urea. To develop such a high concentration of urea (remember that the normal plasma concentration is only about 5 mmol/L), there must be a process of recycling. This involves the tubules as well as the vasa recta.

We described this recycling process in Chapter 5 and review it here. Urea is freely filtered and about half is reabsorbed in the proximal tubule. Urea is secreted in the loop of Henle (thin regions), driven by the high urea concentration in the medullary interstitium. This essentially restores the amount of tubular urea back

\(^5\)We have not explicitly included the roles of the thin portions of the loop of Henle in the development of the medullary osmotic gradient, but since the thin descending limb reabsors water and thin ascending limb reabsors salt, they must in principle exert an influence. Although their quantitative contribution is not clear, it is most assuredly much smaller than that of the thick ascending limbs.
to the filtered load. From the end of the thin limbs to the inner medullary collecting ducts, little urea transport occurs, so whatever urea arrives at the thick ascending limb is still there at the start of the inner medullary collecting ducts. Because the vast majority of water has been reabsorbed before the inner medullary collecting ducts (by the cortical and outer medullary collecting ducts), the luminal urea concentration has risen up to 50 times its plasma value (i.e., 500 mmol/L or more). In the inner medullary collecting ducts, some urea is reabsorbed via specialized urea uniporters and the rest (typically about half the filtered load) is excreted. Because blood flow in this region is low, the reabsorbed urea accumulates and raises the interstitial concentration close to that in the lumen, i.e., 500 mmol/L or more depending on conditions. (It is this high interstitial concentration that drives secretion in the thin limbs). The combination of high urea, along with the high sodium and chloride, brings the medullary osmolality to a value exceeding 1000 mOsm/kg. The importance of urea in contributing to the medullary osmotic gradient is emphasized in the case of low protein intake, which results in a greatly

Figure 6–6. Renal water handling in states of maximum antidiuresis and maximum diuresis. Numbers to the right indicate interstitial osmolality; numbers in the tubules indicate luminal osmolality. The dashed line indicates the corticomedullary border. Arrows indicate sites of water movement. In both antidiuresis and diuresis, most (65%) of the filtered water is reabsorbed in the proximal tubule and another 10% in the descending loop of Henle. The greater relative reabsorption of solute versus water by the loop as a whole and distal tubule results in luminal fluid that is quite dilute (110 mOsm) as it enters the collecting ducts. During antidiuresis (A), the actions of antidiuretic hormone (ADH) permit further water reabsorption in the cortical and medullary collecting tubules. The equilibration of tubular fluid with the high medullary osmolality results in final fluid that is very hyperosmotic (1200 mOsm).
reduced metabolic production of urea. In this condition, the ability of the kidneys to produce highly concentrated urine is reduced.

To summarize the generation of the renal osmotic gradient: salt (without water) is deposited in the interstitial by the thick ascending limb. That salt accumulates because of a combination of low blood flow and countercurrent exchange between ascending and descending vasa recta. Adding to the osmolality of the medulla is urea, which recycles from the inner medullary collecting ducts to the thin limbs of the loop of Henle. Urea also participates in countercurrent exchange between ascending and descending vasa recta for the same reasons that salt does.

As already mentioned, the magnitude of the medullary osmotic gradient (actually the maximum osmolality found in the inner medulla) varies according to states of hydration. A key regulator of this varying osmolality is ADH, which in addition to raising water permeability in the cortical and medullary collecting ducts, also raises urea permeability by stimulating a specific ADH-sensitive isoform of the urea uniporters. But it does this only in the inner medullary collecting
ducts. Consider how this affects the medullary osmotic gradient. When a person is dehydrated, glomerular filtration rate (GFR) is somewhat low and levels of ADH are high. The extraction of water in the cortical collecting duct removes most of the water from the lumen (and makes it iso-osmotic with the cortical interstitium ie, about 300 mOsm/kg). Then, as the remaining but greatly reduced, volume flows through the high osmolality medulla, further concentration occurs. The increased urea permeability signaled by ADH greatly assists in generating the medullary osmotic gradient by permitting the recycling of urea.

Contrast this with a state of overhydration as after a beverage drinking contest. Some of the medullary solute is washed out and the magnitude of the osmotic gradient is reduced. How does this occur? In states of overhydration levels of ADH are low. GFR is substantial. Only a small amount of the tubular fluid entering the cortical collecting ducts is reabsorbed. Therefore, tubular urea does not become concentrated very much. A high volume of very dilute fluid with a modest urea concentration is delivered to the inner medullary collecting ducts. In contrast to the cortical and outer medullary collecting ducts, which are nearly water impermeable in the absence of ADH, the inner medullary collecting duct has a finite water permeability in the absence of ADH. Although this water permeability is not large, the osmotic driving force is huge, so substantial amounts of water are reabsorbed. (However, even more is not reabsorbed, and so the urine volume is still very large.) Not much urea is reabsorbed; in fact, it may be secreted initially because the luminal urea concentration is lower than in the medullary interstitium. The result of the water reabsorption and the low (or absent) urea reabsorption is that the inner medulla is partially diluted (ie, the urea concentration and total osmolality of the medullary interstitium decrease over time). The osmolality falls to about half of its value, from well over 1000 mOsm/kg down to 500–600 mOsm/kg (Table 6–5). A major factor contributing to the washout is an increase

**Table 6–5.** Composition of medullary interstitial fluid and urine during the formation of a concentrated urine or a dilute urine

<table>
<thead>
<tr>
<th>Interstitial fluid at tip of medulla (mOsm/L)</th>
<th>Urine (mOsm/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concentrated urine</td>
<td></td>
</tr>
<tr>
<td>Urea = 650</td>
<td>Urea = 700</td>
</tr>
<tr>
<td>Na⁺ + Cl⁻ = 750a</td>
<td>Nonurea solutes = 700 (Na⁺, Cl⁻, K⁺, urate, creatinine, etc.)</td>
</tr>
<tr>
<td>Dilute urine</td>
<td></td>
</tr>
<tr>
<td>Urea = 300</td>
<td>Urea = 30–60</td>
</tr>
<tr>
<td>Na⁺ + Cl⁻ = 350a</td>
<td>Nonurea solutes = 10–40 (Na⁺, Cl⁻, K⁺, urate, creatinine, etc.)b</td>
</tr>
</tbody>
</table>

a Some other ions (eg, K⁺) contribute to a small degree to this osmolarity.
b Depending on the sodium balance state, sodium in the urine can vary between undetectable and the majority of the osmolytes.
in medullary blood flow, due in part to removal of the vasoconstricting action of ADH. Figure 6–6 depicts renal water fluxes in the two extremes of maximum diuresis and antidiuresis.

Let us conclude this chapter by addressing two issues that often confuse students. First, one might expect that under conditions of high ADH, there should be even more water reabsorbed from the medullary collecting ducts than there is with low ADH, and that this water would dilute the interstitium and abolish the osmotic gradient. Water does enter the medullary interstitium from medullary collecting ducts aided by the actions of ADH, but so little water remains in the tubule after passage through the cortex that the amount remaining to be reabsorbed is quite small. Also, as described earlier, water enters from descending thin limbs of the loops of Henle and from descending vasa recta. Although there is a tendency for all of this water to dilute the interstitium, there is also a continuing deposition of new solute by the thick ascending limb. The competing tendencies to dilute the interstitium with water and to concentrate the interstitium with salt reach a balance in which osmolality is high. It is this balance that sets the upper limit on medullary osmolality.

Another issue that appears paradoxical concerns medullary water reabsorption during diuresis when ADH is low and the body is excreting large amounts of water. In this condition, more water is reabsorbed in the medulla than during antidiuresis when ADH is high and the body is conserving water. This seeming paradox is resolved by realizing that during diuresis there is little cortical water

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**Figure 6–7.** Osmolarity of the tubular fluid and the percentage of filtered water remaining at different sites along the tubule. The latter numbers, of course, are derived simply from the numbers given in Table 6–2 for the percentage of water reabsorbed by each tubular segment. ADH, antidiuretic hormone.
reabsorption, so the amount of medullary water absorption is greatly exceeded by the amount not reabsorbed, ie, excreted.

Figure 6–7 summarizes the previously described changes in volume and osmolality of the tubular fluid as it flows along the nephron and emphasizes how, once fluid enters the collecting-duct system, the osmolality depends very much on the levels of ADH.

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**KEY CONCEPTS**

1. The reabsorption of most of the filtered water, anions, and osmotic content is linked to the active reabsorption of sodium.

2. In all conditions, the vast majority of the filtered volume is reabsorbed iso-osmotically in the proximal tubule in a manner that is entirely dependent on active sodium reabsorption.

3. The capacity to generate a variable-osmolality urine depends on “separating salt from water” in the diluting segments.

4. Reabsorption of water remaining in the lumen beyond the loop of Henle is variable, depending on hydration status, allowing the kidneys to excrete either a high-volume dilute urine, a low-volume concentrated urine, or anything in between. Levels of ADH determine whether the hypo-osmotic fluid leaving the diluting segments is excreted largely as is or whether most of this fluid is subsequently reabsorbed.

5. The existence of the medullary osmotic gradient depends on (1) transport of salt without water into the medullary interstitium by the thick ascending limb, (2) recycling of urea, and (3) low-volume countercurrent blood flow in the vasa recta.

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**STUDY QUESTIONS**

6–1. A man ingests 12 g of sodium/day. His nonrenal loss (gastrointestinal tract and sweat) is 0.4 g/day. In the steady state, what amount of sodium chloride is excreted daily in the urine?

6–2. Mannitol is a substance sometimes infused to reduce cerebral edema. It is handled by the kidneys similarly to inulin. What effect would a large mannitol infusion have on sodium excretion?
   A. No effect
   B. Increase sodium excretion
   C. Reduce sodium excretion
6–3. Would complete inhibition of active sodium and chloride transport by the thick ascending limb of Henle's loop eliminate the ability to excrete a concentrated urine or a dilute urine?

6–4. Increasing the passive permeability of the thick ascending limb of Henle's loop to sodium and chloride would reduce the maximal concentrating ability of the kidney. True or false?

6–5. Active reabsorption of sodium and chloride by the descending thin limb of Henle's loop is a component of the countercurrent multiplier system. True or false?

6–6. In conditions of maximum levels of ADH, there is net bulk flow of fluid from medullary interstitium into the vasa recta. True or false?

6–7. In conditions of minimum levels of ADH, there is net bulk flow of fluid from medullary interstitium into the vasa recta. True or false?

6–8. A drug is given that blocks all sodium channels and transporters in the luminal membrane all along the tubule but does not act on the Na-K-ATPase pumps in the basolateral membrane. What happens to sodium reabsorption?

6–9. The osmolality in the renal papilla in a healthy young person with excellent renal function who is not taking any medications is always 1200 mOsm/kg or more. True or false?
Control of Sodium and Water Excretion: Regulation of Plasma Volume and Plasma Osmolality and Renal Control of Systemic Blood Pressure

OBJECTIVES

The student describes the renal regulation of extracellular fluid volume, total-body sodium balance, total-body water balance, and blood osmolality and their relationship to systemic blood pressure:

- Describe the 3 temporal domains of blood pressure control and the major mechanisms associated with them.
- Describe the relationship between renin and angiotensin II.
- Describe the detectors that can alter renin secretion.
- Define pressure natriuresis and diuresis.
- Define tubuloglomerular feedback and describe the mechanism for tubuloglomerular feedback and autoregulation of glomerular filtration rate.

The student describes the renal regulation of total body sodium balance:

- State the formula relating filtration, reabsorption, and excretion of sodium.
- Describe the nature and locations of receptors (“sensors”) in sodium-regulating reflexes.
- List the major factors that regulate sodium excretion.
- State the tissue origin of aldosterone, its renal sites of action, and its effect on sodium reabsorption.
- List the factors controlling aldosterone secretion and state which is normally most important.
- State the origin of atrial natriuretic peptides, the stimulus for their secretion, and their effect on sodium reabsorption and glomerular filtration rate.
- State the effect of antidiuretic hormone on sodium reabsorption.
- State all direct and indirect effects of catecholamines and angiotensin II on sodium reabsorption.
- Describe how intrarenal physical factors influence sodium reabsorption; state how changes in filtration fraction influence sodium reabsorption; predict the changes in physical factors that occur with changes in sodium or fluid balance and how they alter sodium and water reabsorption.
- Define glomerulotubular balance and describe its significance.
The student understands the renal regulation of total-body water balance and control of plasma osmolality:

- Describe the origin of antidiuretic hormone and the 2 major reflex controls of its secretion; define diabetes insipidus; state the effect of antidiuretic hormone on arterioles.
- Distinguish between the reflex changes that occur when an individual has suffered iso-osmotic fluid loss because of diarrhea as opposed to a pure water loss (ie, solute-water loss as opposed to pure-water loss).
- Describe the control of thirst.
- Diagram in flow-sheet form the pathways by which sodium and water excretion are altered in response to sweating, diarrhea, hemorrhage, high-salt diet, and low-salt diet.

The regulation of salt and water excretion by the kidneys involves all of the endothelial and epithelial transport mechanisms we have described in previous chapters, and either directly or indirectly all of those mechanisms are subject to physiological control. Although it might appear to be an overwhelming task to understand their regulation, there is indeed a logic to regulation that we can easily grasp. First, net salt and water excretion is the sum of all the transport processes in the glomerulus and various nephron segments, and second, that sum maintains balance. However, and here is the key point, balance is not regulated on the basis of input, using physiological “water meters” or “salt meters” directing the kidneys to excrete at rates that match input. Rather, renal excretion is regulated in response to the consequences of input and loss. Those consequences are manifested primarily in the cardiovascular system. There are many interactions between the kidneys and the cardiovascular system in which the action of one affects the other. Accordingly, we can paint a “big picture” of renal control in the following way. Control of salt and water excretion serves to (1) maintain a body fluid volume appropriate for filling the vascular tree, (2) maintain an osmolality of that fluid appropriate for the function of cells bathed in it, and (3) allow the heart to generate the arterial pressure necessary to perfuse peripheral tissues. In one way or another, all of the processes discussed in the rest of this chapter fit into this picture.

Regulating the amount of salt and water in the body is conceptually straightforward. In regulating total body salt and water, the kidneys are actually regulating 4 quantities simultaneously: water balance, salt balance, osmolality, and blood pressure. When the body takes on a load of water or salt or both, the kidneys excrete these loads to maintain balance. Regulating osmolality involves a juggling act. Osmolality is not a substance that has an input and output; rather, it is the ratio of substances (solute and water). Therefore, the kidneys have to independently handle water loads and salt loads, all the while keeping their ratio in the body within rather tight limits. Regulating blood pressure is clearly the most complex of the various renal tasks. As mentioned above, it centers on maintaining a fluid volume that appropriately fills the vascular tree, but includes other actions of the kidneys.
REGULATION OF BLOOD PRESSURE

We choose to organize the control of sodium and water excretion around the topic of blood pressure for 2 reasons. First, because pressures in various parts of the vascular system have such a powerful influence on renal function, and second, because renal actions so strongly affect blood pressures. In doing so, we will encounter many important concepts and components. We briefly outline them here and expand them in the ensuing discussion. First is the concept of a set-point, which is the value that blood pressure should be at any moment (similar to the setting for temperature on the thermostat in your house). Second are detectors of blood pressure (“pressure gauges”), which assess the level of blood pressure at any moment. Third are signals generated in response to changes in blood pressure sensed by the detectors that regulate the fourth component: effectors, which change what they do in response to the signals in order to raise or lower blood pressure and return it to the setpoint. The effectors of blood pressure regulation are (1) the heart, which has a variable contractility and beat rate; (2) peripheral arterioles, which determine resistance to flow in the peripheral vasculature; (3) large veins, which change their compliance to vary the capacity of the vascular system to hold blood; and (4) the kidneys, which vary their output of salt and water. We will elaborate on the renal involvement in these effectors as we go along.

The various blood pressure regulatory processes occur over different time spans. There are immediate (within seconds) cardiovascular reflexes that are for the most part nonrenal in nature. Then, there are slower processes spanning time scales of minutes to days centered on renal regulation of salt and water (ie, fluid volume and osmolality). We can arbitrarily divide regulation into short-term, intermediate-term, and long-term processes, recognizing that those in one time domain overlap with those in others and thus each process can interact with the others. Despite this overlap between these systems, it still helps to conceptualize them, as we will do below, as separate (but interacting) processes. Figure 7–1 summarizes these relationships.

Short-Term Regulation of Blood Pressure: Cardiovascular Reflexes

Arterial blood pressure is regulated around a setpoint controlled by a set of brainstem nuclei often called the vasomotor center. There are 2 major sets of detectors for the short-term control of blood pressure. The most important are baroreceptors that mediate the classic baroreceptor reflex. These are afferent nerve cells (mechanoreceptors) with sensory endings located in the carotid arteries and arch of the aorta. They report arterial blood pressure to the vasomotor center via sensory neural pathways. They do this continuously on a heartbeat-by-heartbeat basis. The second set of baroreceptors is the cardiopulmonary baroreceptors. These are also nerve cells, with sensory endings located in the cardiac atria and parts of the pulmonary vasculature. They, like the arterial baroreceptors, send afferent neural information to the brainstem vasomotor center. They are often called low-pressure baroreceptors because they assess pressures in regions of the vascular tree where pressures are much lower than in the arteries. The cardiopulmonary
baroreceptors serve as de facto blood volume detectors in the sense that pressures in the atria and pulmonary vessels rise when blood volume increases and fall when blood volume decreases. Their most important role lies in regulating salt and water excretion, but their actions mix with those of the arterial baroreceptors.

On the basis of the inputs from the arterial and cardiopulmonary baroreceptors, the vasomotor center sends regulatory signals to effector systems: the heart, blood vessels, and kidneys via the autonomic nervous system. Changes in the activity of the brainstem vasomotor center lead to changes in sympathetic signals that directly regulate the actions of our first effector system: cardiac contractility and heart rate. At the same time, these signals are sent in parallel to our second effector system: vasoconstriction or dilation of all systemic arterioles (including those of the kidneys), with consequent changes in peripheral vascular resistance. When we express mean arterial pressure (MAP) as the product of cardiac output (CO) and total peripheral resistance (TPR), ie, MAP = CO × TPR, it becomes clear that adjusting either CO or vascular resistance directly changes MAP.

Sympathetic output from the vasomotor center is also directed to our third effector system: the large peripheral veins. These veins contain about two thirds of the total blood volume. When blood volume changes, almost all the change occurs in the volume of peripheral venous blood. The compliance of the veins (ease of being stretched) allows them to accommodate moderate changes in blood volume.
volume. Furthermore, their compliance is a regulated variable (via contraction or relaxation of smooth muscle in their walls). Stimulatory sympathetic signals reduce venous compliance, i.e., make the veins less stretchy. This has the effect of squeezing on the blood in the veins and raising its pressure, a de facto shrinkage in the capacity of the venous tree to hold blood. In contrast, a reduction in sympathetic signals raises the venous compliance, allowing the system to hold more blood. These adjustments are very important in terms of keeping central venous pressure (pressure at the right atrium) appropriate for filling the cardiac chambers between beats.

A pathological increase in venous compliance, as in certain forms of circulatory shock, has the same effect as a major hemorrhage, because this creates an overcapacity of the vascular system relative to its actual volume, with a resulting drop in central venous pressure and insufficient filling of the cardiac chambers.

All of these fast effector mechanisms of the heart, arterioles, and large veins act very rapidly when pressure begins to change as a result of muscle activity or simple changes in posture. The result is to stabilize arterial pressure at its setpoint, the MAP, which for most people is slightly less than 100 mm Hg. The setpoint is not rigidly fixed; however, it varies during the day, depending on the activity and levels of excitement, and decreases about 20% during sleep.1 As we will expand on shortly, a complication lies in the fact that the value of the setpoint over the long term is highly influenced by renal processes because the kidneys regulate blood volume. That is, the renal processing of salt and water, via its control of blood volume, ultimately determines the average value of the setpoint for blood pressure of the brainstem vasomotor center. As long as the kidneys regulate salt and water excretion appropriately, the average value of blood pressure over the course of a day will be normal. However, if renal excretion is inappropriate and remains so for several days, then the setpoint becomes reset to a new value.

As the name implies, short-term control of blood pressure through the baroreceptor reflex and signals from the cardiopulmonary baroreceptors is a rapidly acting system that can respond to external perturbations in pressure on a time scale of a few seconds (1 or 2 heartbeats). Both of these types of pressure detectors work in concert to produce sympathetic signals that maintain blood pressure nearly constant in the short term through fast vascular and cardiac effector responses. However, besides initiating rapid responses, changes in the sympathetic signals also have effects on the kidney that contribute to initiating the intermediate-term regulation of blood pressure.

**Intermediate-Term Regulation of Blood Pressure: Renal Control of Vascular Resistance**

In the event that the short-term regulation of blood pressure does not completely restore blood pressure to its normal setpoint within a few tens of seconds, then the kidneys are capable of strongly reinforcing the short-term vascular effects of the

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1As an example of this variation, some patients experience “white coat hypertension,” a phenomenon in which their blood pressure is normal while resting calmly at home but rises when a white-coated physician measures it in an office setting.
vasomotor center if a deviation in blood pressure is maintained. This reinforcement involves direct vascular actions. The major detectors involved in the kidney’s ability to regulate vascular resistance are the previously described baroreceptors, and another set of pressure-sensitive cells within the kidney, often referred to as intrarenal baroreceptors. These baroreceptors sense renal afferent arteriolar pressure. Anatomically, these structures are not neural baroreceptors (ie, they are not nerve cells and do not send signals to the brainstem vasomotor center) but rather are specializations of the cells of the afferent arteriole: granular cells (also called juxtaglomerular cells) that form part of the juxtaglomerular apparatus. They act entirely within the kidney.2 Although granular cells acting as intrarenal baroreceptors do not send signals centrally, neural signals originating in the vasomotor center (generated in response to vascular baroreceptors) reach the granular cells via the renal sympathetic nerve. Thus, the activity of the granular cells is affected both by direct sensing of pressure in the renal artery and by pressures sensed by neural baroreceptors elsewhere in the body. Baroreceptors and their key actions are summarized in Figure 7–2.

In response to changes in pressures sensed by baroreceptors, a number of renal events are set in motion that have powerful effects on the vasculature and on

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2Although intrarenal baroreceptors are not neural afferents, there are afferent nerves originating in other regions of the kidneys that impact autonomic function.
sodium excretion The most critical involve signaling pathways known as renin-angiotensin systems (RAS).

Renin-Angiotensin Systems

If it is possible to single out one substance as being the most important in the control of sodium excretion and blood pressure, then that substance would be angiotensin II. It is a potent vasoconstrictor and a mediator of multiple actions in the kidneys that affect sodium excretion. Thus, it affects blood pressure directly as a vasoconstrictor and indirectly via regulation of renal sodium excretion.

There are many local RAS in individual tissues, including the kidneys, brain, and the heart. There is also a global or systemic RAS which we normally consider to be the major renal regulator of blood pressure. All RAS, whether global or local, consist of a large protein substrate called angiotensinogen, several enzymes and several products. The key product is angiotensin II. When angiotensin II binds to cell surface receptors, it initiates actions that affect blood pressure and excretion of sodium. The first key enzyme in all RAS is renin. It acts on angiotensinogen to produce a small (10-amino-acid) product called angiotensin I. Angiotensin I is acted upon by another enzyme, angiotensin-converting enzyme (ACE), to produce the highly active 8-amino-acid peptide angiotensin II. In the global RAS, the source of angiotensinogen circulating in the blood is the liver. The source of circulating renin is the granule cells in the kidney. Renin is secreted both into the interstitium of the kidney and into the lumen of the afferent arterioles, where it acts on circulating angiotensinogen to produce circulating angiotensin I. ACE, which is expressed on the luminal surface of endothelial cells in many parts of the vasculature, particularly in the lungs, then converts angiotensin I to angiotensin II.

We will describe the regulation of RAS in terms of the global system, in part because more is known about it. Investigators agree, however, that locally produced angiotensin II (and some related peptides) is the more important source in terms of regulating the kidneys because the levels of angiotensin II in renal tissues are far higher than can be accounted for by a systemic source. It is presumed that global angiotensin II arriving in the renal blood supply acts synergistically with local angiotensin II as a regulator of function.

Circulating levels of angiotensinogen are usually fairly high and ACE activity usually converts most angiotensin I into angiotensin II. Therefore, the primary determinant of circulating levels of angiotensin II is the amount of renin available to convert angiotensinogen to angiotensin I. Consequently, to understand angiotensin II regulation of blood pressure, we require an understanding of the regulation of renin secretion. What determines how much renin is secreted? Two primary regulators have been described. The first are the neural baroreceptors, which produce signals via the renal sympathetic nerves that stimulate granular cells: Activation of β1-adrenergic receptors on the granular cells stimulates renin secretion via a cyclic adenosine monophosphate and protein kinase A-dependent process. (In addition, the activity of renal sympathetic nerves causes afferent arteriolar constriction and reduction in renal blood flow.) The second regulators of
renin secretion are the intrarenal baroreceptors, ie, granular cells that deform in response to changes in afferent arteriolar pressure; when the pressure falls, renin production increases. Thus, as mentioned previously, granular cells act both as detectors (of renal arteriolar pressure) and as signal generators (releasing renin) in response to changes in pressure and sympathetic activity. The signals from the vasomotor system to the renin-producing granular cells ensures that there is tight coordination between the rapid activity of the baroreceptor reflex and the slower acting RAS; ie, the short-term regulation and the intermediate-term regulation have at least one common set of detectors. However, the intrarenal pressure detector can function in the absence of renal innervation (eg, after a renal transplant).

There is also a third detector mechanism that regulates renin release. It is also intrarenal, but it does not detect blood pressure. Rather, it measures the amount of sodium chloride that leaves the thick ascending limb, directly bathing the macula densa cells of the juxtaglomerular apparatus and delivered to the distal convoluted tubule. This amount of sodium chloride depends on both the rate of filtration and the rate of sodium reabsorption in all the nephron elements preceding the macula densa. When sodium chloride delivery (a combination of concentration and flow rate) to the luminal surface of macula densa cells increases, renin production decreases. This is due to increased uptake of NaCl by the cells with subsequent osmotic swelling. Osmotic swelling (Figure 7–3) causes the release of transmitter agents (see later discussion) that inhibit renin release. This load detector, therefore, does not generate signals that directly regulate blood pressure. However, it does contribute to the regulation of renin secretion.

Figure 7–3. Responses of macula densa cells to changes in delivery of NaCl load. The macula densa cells (arrowheads) are in close apposition to the glomerulus (G). Macula densa cells swell in response to increasing tubular NaCl concentration from 25 (total osmolality = 210 mOsm/kg H2O) on the left to 135 mmol (total osmolality = 300 mOsm/kg H2O). Bar = 10 μm. (From Peti-Peterdi J et al, Am J Physiol Renal Physiol. 2002;283;F197. Used with permission.)
Thus, there are three separate, redundant mechanisms regulating renin secretion (neural signals, afferent arteriolar pressure, and NaCl at the macula densa). This redundancy reflects the importance of the RAS and angiotensin II, in particular, in regulating blood pressure. Among the most significant actions of circulating angiotensin II produced by the global RAS is general arteriolar vasoconstriction. This vasoconstriction acts in parallel with sympathetically mediated neural control. This raises total peripheral resistance, thereby increasing blood pressure. The importance of this system makes the RAS a natural target for pharmacological intervention to reduce high blood pressure. A number of blood pressure–lowering pharmacological agents are aimed at components of the RAS, including ACE inhibitors and blockers of the arteriolar smooth muscle receptors for angiotensin II. Figures 7–4 to 7–7 illustrate the various features of the RAS described in the text and show how the system responds to a major fall in blood pressure resulting from a hemorrhage.

Besides these primary mechanisms, angiotensin II acts in a negative feedback manner to inhibit renin production by acting directly on granular cells (by interacting with AT1 receptors on granular cells to increase intracellular Ca concentration, which inhibits renin production).

Most of the time it is appropriate for the vasoconstrictive and sodium retaining actions of angiotensin II to be exerted in parallel. However, by having both a global and an intrarenal RAS, it is possible to separate these actions, such that changes in sodium excretion can be effected without, at the same time, altering vascular resistance elsewhere in the body.

CONTRIBUTION OF THE KIDNEY TO THE REGULATION OF SODIUM EXCRETION AND BLOOD PRESSURE

Despite the strength and efficacy of the vascular baroreceptor reflex and the potency of renin-induced angiotensin II in regulating vascular smooth muscle tone, these mechanisms are not the ultimate determinants of blood pressure. That is, the average value of blood pressure (or perhaps the average value of the setpoint around which the baroreceptor reflex operates) is fixed not by the brainstem vasomotor center but rather by the kidneys. Guyton and colleagues, in their classic experiments, surgically cut the neural pathways between the baroreceptors and the vasomotor center of anesthetized dogs. After recovery, the dogs’ blood pressure varied widely from moment to moment, far more so than normal, but the mean value eventually returned to baseline. Various investigators ultimately showed that the kidneys are responsible for determining the setpoint for mean blood pressure. It does this, as should be clear by now, by controlling the amount of sodium, and hence volume, in the vascular space on a long-term basis.

It is worth emphasizing the time lag between volume changes and pressure changes. For example, increasing volume by ingesting a large amount of liquid or decreasing volume by sweating during a tennis match on a hot day does not immediately cause changes in blood pressure. This is because tendencies to change pressure are buffered immediately by the classic baroreceptor reflex and by renal
output of salt and water. However, if the kidneys do not match their output to input, and changes in extracellular fluid (ECF) volume are sustained, then pressure gradually creeps toward a new elevated or depressed value. In the face of sustained changes in volume, the baroreceptor reflex cannot forever keep pressure normal. We are normally unaware of the kidneys’ role in the control of blood pressure because the baroreceptor reflex is very effective on a short-term basis in buffering changes and because healthy kidneys do such a good job of adjusting their volume output in the face of changes in input.
Hemorrhage

- Activity of renal sympathetic nerves
- Pressure sensed by intrarenal baroreceptors
- Macula densa flow

Granular cells

- Renin secretion
- Plasma renin concentration
- Plasma angiotensin II concentration

**Figure 7-5.** Schematic diagram showing the increase in renin secretion and the increased production of angiotensin II in response to a major hemorrhage. Three primary mechanisms activate renin secretion: (1) increased renal sympathetic nerve activity; (2) decreased pressure sensed by intrarenal baroreceptors; and (3) decreased sodium chloride delivery to the macula densa. The first two mechanisms directly stimulate renin release, whereas the third mechanism reduces inhibitory feedback, allowing more renin release. Renin promotes the formation of angiotensin II, which produces strong vasoconstriction and helps to correct the decrease in blood pressure that resulted from the hemorrhage.

**The Connection between Sodium, Water, and Blood Pressure**

At this point, we have described signals affecting the first three effector mechanisms for blood pressure control, i.e., cardiac performance, vascular resistance, and venous compliance. All of these three mechanisms can generally be thought of as adjusting the properties of the vascular system to match the available volume of blood. The fourth renal mechanism for controlling blood pressure is to adjust the volume of blood to fit the vascular system. Because control of blood volume is arguably the most complex of these effector systems, it is worth elucidating the logical connection between renal sodium excretion and blood volume before further describing the mechanisms of control per se.

Let us pose the following question: What does sodium have to do with blood pressure? Pressures in the vascular tree require an appropriate volume of blood (to fill both the highly elastic venous system and the chambers of the heart). With insufficient volume, the heart can neither fill nor pump. Blood pressure in the long term depends on blood *volume*. Blood volume, in turn, depends on total ECF
volume (ie, the volume of blood plasma and fluid in the interstitial spaces of the tissues throughout the body). Fluid in the interstitial spaces acts as a buffer for plasma volume, protecting the vascular compartment from immediate changes associated with drinking, sweating, and so on. However, over time, sustained changes in ECF volume lead to parallel changes in blood volume and ultimately arterial pressure. If the vascular system is inappropriately filled on a prolonged basis, the setpoint gradually drifts. To keep arterial pressure normal, the ECF volume must be kept normal. In many ways, regulating the ECF volume to a level appropriate for the vascular system is the most important function of the kidneys.

The relationship between blood volume and total body water may appear obvious, but the relationship between total body sodium content and blood volume may not. However, as discussed in Chapter 4, there is a simple relation between the volume of a compartment (essentially the amount of water) and its osmolarity: osmolarity = total osmoles/volume. In other words, volume = total osmoles/osmolarity. Therefore, the ECF volume is determined by the total osmotic content.

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Figure 7–6. Schematic diagram showing the vascular response to a major hemorrhage. The baroreceptor reflex increases sympathetic activity. Besides the effect of sympathetic neurotransmitters on β1-adrenergic receptors to stimulate renin release, they also stimulate α1-adrenoreceptors (like those present on other vascular smooth muscle cells) to cause afferent arteriolar contraction and a reduction in renal blood flow. In the kidney, most of this reduction in blood flow is blunted by tubuloglomerular feedback (see Figure 7–14). GFR, glomerular filtration rate.
Figure 7–7. The macula densa NaCl load sensor. Macula densa cells in the thick ascending limb sense sodium chloride delivery by changing the uptake of salt with subsequent osmotic swelling (see Figure 7–3). Changes in cell volume lead to the release of chemical transmitters that alter renin secretion from the granular cells. When sodium chloride delivery increases, renin production decreases. GFR, glomerular filtration rate.

and osmolarity. If the body regulates the total osmotic content of the ECF and regulates its osmolarity, it has accomplished the task of regulating its volume. This is precisely what the kidneys do. They regulate ECF osmolarity and total osmotic content. Recall that more than 90% of the ECF osmotic content is accounted for by sodium and the equal number of anions that must accompany it. To a first approximation, total ECF osmotic content = sodium content × 2. The other 10% of the ECF solute is accounted for by substances such as potassium, glucose, urea, and so on. The regulation of solutes other than sodium occurs for purposes unrelated to control of ECF osmolality, so that the regulation of osmotic content amounts to the regulation of sodium content. Figure 7–8 shows how the ECF volume changes when the body takes on sodium loads and Figure 7–9 shows the excretory response to those loads.

In simple terms, long-term regulation of arterial blood pressure involves long-term control of body sodium content. If the body controls sodium content and plasma osmolarity (the water content containing the sodium), it controls volume.
Figure 7–8. Relation between sodium and extracellular fluid (ECF) volume. Each large rectangle represents total-body water divided into an ICF (open areas) and ECF (shaded areas). The ECF is further subdivided into interstitial and plasma volumes. Excess sodium is almost always accompanied by water, so that excess sodium causes an expansion of the ECF volume. If there is no change in osmolality, as shown in the middle example, the expansion is entirely in the ECF and there is no change in ICF volume. If there is excess sodium without excess water, as shown in the bottom example, water is drawn from the ICF to maintain equal osmolalities between compartments. In both cases of excess sodium, the increase in ECF volume causes an increase in both plasma and interstitial volumes. ICF, intracellular fluid; ECF, extracellular fluid.

If it controls volume, then it controls pressure. This raises the following question: How do the kidneys know about sodium content so that they can respond to changes? Surprisingly, in detecting total body sodium, the primary variable that the kidney monitors is not a direct measure of the amount of sodium in the body or plasma sodium concentration but rather pressures in various parts of the vascular tree and in the kidneys that we have already described. Pressure changes at any of these sites are interpreted as a change in total body sodium because, except for pathophysiological circumstances, blood pressure, blood volume, and total body sodium march in lockstep.

Sodium content and blood pressure can be too high or too low. Some of the mechanisms that control sodium excretion mainly serve to correct elevated pressure/high sodium content, while others mainly correct low pressure/low sodium content. Still others come into play with deviations in either direction. This bidirectional responsiveness applies to the first control mechanism we discuss—control of glomerular filtration rate (GFR).

**Control of Glomerular Filtration Rate**

Because sodium excretion represents the difference between filtration and reabsorption, it is not surprising that one of the major controls over sodium excretion is the regulation of GFR. A change in the amount of sodium filtered resulting from a change in GFR is also accompanied by a change in the amount of water filtered. Therefore, any change in GFR represents a mechanism for altering ECF volume.

The reflex control of GFR is mediated mainly by changing the resistance of the afferent and efferent arteriolar resistance. The changes in resistance are produced by changes in renal sympathetic nerve activity and circulating levels of
Cardiopulmonary baroreceptors

**A**
- **↑Volume**
- **↓Sympathetic drive from vasomotor center**
- **↑GFR**
- **↓Proximal tubule Na reabsorption**
- **↑Salt and water delivery to distal nephron**
- **↑Salt excretion**
- **↑Water excretion**

**B**
- **↑Salt in ECF**
- **↑Volume (transfer from ICF)**
- **↑ECF osmolality**
- **↓Sympathetic drive from vasomotor center**
- **↑GFR**
- **↓Proximal tubule Na reabsorption**
- **↑Salt and water delivery to distal nephron**
- **↑Salt excretion**
- **↑ADH**
- **Collecting duct water reabsorption**
- **No change in water excretion**

Salt and water in ECF

Osmoreceptors

Cardiopulmonary baroreceptors

Na reabsorption

GFR

Proximal tubule

Collecting duct
on by external signals. The high pressure reduces levels of intrarenal angiotensin II. The number of Na-H exchangers in the apical membrane is strongly influenced by angiotensin II. When its levels fall, Na-H exchangers are withdrawn, along with a concomitant reduction in the activity of the basolateral Na-K-ATPase. The result of the reduction in angiotensin II in response to high renal artery pressure is less sodium absorption and more presentation of sodium to the loop of Henle, and therefore more excretion (see Figure 7–10). Pressure natriuresis and diuresis

Figure 7–10. Response of the kidneys to an increase in blood pressure (natriuresis/diuresis). Part of the intermediate-term response to increases in blood pressure is to reduce blood volume (in an attempt to match blood volume with the capacity of the vascular tree). There are several mechanisms for this response. By far, the most important is a reduction in proximal tubular sodium reabsorption because of a reduction in the number of functional transporters (Na-H antiporters) in the apical membrane of the proximal tubule epithelial cells. The reduction is probably in response to reduced levels of angiotensin II. There is also an increase (usually small) in glomerular filtration rate (GFR) and an increase in peritubular hydrostatic pressure and renal interstitial pressure that favor reduced absorption of salt and water in the cortex (particularly from the proximal nephron). ECF, extracellular fluid.
serves as a kind of backup system that comes into play if fast-acting reflex systems of regulating blood pressure fail to completely correct large increases.

If peritubular levels of angiotensin II are kept constant by experimental means, pressure natriuresis and diuresis are strongly blunted or even eliminated. The effect of maintaining constant sympathetic transmitters is similar but not so pronounced. Thus, the same agents that directly affect vascular peripheral resistance to correct blood pressure (sympathetic transmitters and angiotensin II) also affect tubular reabsorption to correct ECF volume.

A key feature of pressure natriuresis and diuresis is that the degree of salt and water excretion for a given rise in pressure varies with the volume status of the body. Even though pressure natriuresis is turned on strictly by intrarenal mechanisms, the amount that occurs can be dampened by external factors. If the ECF volume is normal or high and the renal artery pressure rises, pressure natriuresis and diuresis are very effective in increasing excretion of sodium and water and reducing blood volume. On the other hand, if ECF volume is low and the renal artery pressure rises, there is much less salt and water loss. It appears that the volume status of the body acts as a gain control on pressure natriuresis and diuresis.\(^4\) There is potent pressure natriuresis and diuresis when ECF volume is high, and much less pressure natriuresis and diuresis when ECF volume is depleted. Under normal conditions, pressure natriuresis and diuresis is a proximal nephron mechanism that is very important for dumping sodium and water when blood pressure is too high. It does this by reducing isotonic reabsorption of salt and water from the proximal convoluted and straight tubule.

**Peritubular-Capillary Starling Factors and the Role of Renal Interstitial Hydraulic Pressure**

Changes in GFR, besides directly affecting the filtered volume, also affect reabsorption of that volume. A rise in either peritubular capillary pressure or interstitial pressure reduces net reabsorption (and therefore causes more excretion). From the viewpoint of Starling forces acting on the capillary, it should be obvious that high capillary pressure opposes reabsorption. But high interstitial pressure should favor reabsorption, so why does it also oppose it? First, an increased interstitial pressure causes back-leak of reabsorbed fluid from the interstitial space across the tight junctions into the tubule. Thus, this pressure does not alter the cellular transport mechanisms for sodium and water but rather reduces the net reabsorption achieved by these mechanisms, particularly in the “leaky” proximal tubule. In effect, if the interstitium gets “too full,” then it is difficult to transport more fluid into it. Put another way, high interstitial pressure does more to oppose the movement of fluid from tubule to interstitium than it does to promote the movement of fluid from interstitium to capillary.

A decrease in peritubular-capillary oncotic pressure (\(\pi_{PC}\)) also opposes reabsorption. Of course, the new question is: How do changes in GFR cause changes in \(P_{PC}\) and \(\pi_{PC}\)? We already know the answers to this question from Chapter 2:

\(\text{The only known way this can occur is via signals originating in cardiopulmonary baroreceptors and transmitted to the kidneys via renal nerves. However, there are probably other factors.}\)
$P_{PC}$ is set by (1) arterial pressure and (2) the combined vascular resistances of the afferent and efferent arterioles, which determine how much of the arterial pressure is lost by the time the peritubular capillaries are reached. $\pi_{PC}$ is set by (1) arterial oncotic pressure and (2) filtration fraction (GFR/RPF), which determine how much of the oncotic pressure increases from its original arterial value during passage through the glomeruli.

Teleologically, it makes sense that $P_{PC}$ and $\pi_{PC}$ influence interstitial pressure and, hence, sodium reabsorption because these phenomena are simply a logical continuation of the flow diagrams we have used previously for studying the homeostatic control of GFR. Events initiated by fluid loss from the body end with 3 changes that lower GFR: increased constriction of the afferent and efferent arterioles (induced by the renal nerves and angiotensin II), decreased arterial hydraulic pressure, and increased arterial oncotic pressure. Figure 7–10 illustrates how these same 3 factors also decrease renal interstitial hydraulic pressure and, hence, increase sodium reabsorption. Thus, homeostatic responses that tend to lower GFR in response to a reduction in body sodium also usually increase sodium reabsorption, the “desired” homeostatic event of preserving volume in response to bodily fluid depletion.

The same logic applies when the desired homeostatic responses are increased GFR and decreased sodium reabsorption so as to eliminate excess sodium from the body. Thus, when a high-salt diet or expansion of the ECF volume from some other physiological cause is the primary event, the following occurs: (1) decreased plasma oncotic pressure (resulting from dilution of plasma proteins), (2) increased arterial pressure, and (3) renal vasodilation secondary to decreased activity of the renal sympathetic nerves and decreased angiotensin II. Simultaneously, then, the GFR increases a small amount and so does interstitial pressure, which reduces fluid reabsorption. Figure 7–10 illustrates these natriuretic responses to a rise in arterial pressure.

**Glomerulotubular Balance**

As stated earlier, in the regulation of sodium excretion, the control of tubular sodium reabsorption is more important than control of GFR. One reason for this is that a change in GFR automatically induces a proportional change in the reabsorption of sodium by the proximal tubules, so that the fraction reabsorbed (but not the total amount) remains relatively constant (Table 7–1). This phenomenon has the rather ungainly name of *glomerulotubular balance*. In response to a primary change in GFR, the percentage of the filtered sodium reabsorbed proximally remains approximately constant (about 65%). The fraction not reabsorbed also remains approximately constant (about 35%). Therefore, a change in GFR is still reflected as a change in the sodium and water presented to the loop of Henle. Glomerulotubular balance does not mean that proximal reabsorption is always exactly 65% of filtered sodium. It only says that when the fraction reabsorbed is changed, the change is caused by processes other than changes in GFR. Several mechanisms are manifested in the proximal tubule to stimulate sodium reabsorption (raise the percentage reabsorbed above 65%) or inhibit sodium reabsorption (lower the percentage below 65%).
The mechanisms responsible for matching changes in tubular reabsorption to changes in GFR are completely intrarenal (ie, glomerulotubular balance requires no external neural or hormonal input; indeed, the presence of such input usually obscures the existence of glomerulotubular balance, as described previously).

Glomerulotubular balance is actually a second line of defense preventing changes in renal hemodynamics per se from causing large changes in sodium excretion. The first line of defense is autoregulation of GFR, described in Chapter 2 and in the prior discussion of tubuloglomerular feedback. GFR autoregulation prevents GFR from changing too much in direct response to changes in blood pressure, and glomerulotubular balance blunts the sodium-excretion response to whatever GFR change does occur. Thus, tubuloglomerular feedback and glomerulotubular balance mediated by GFR autoregulation are processes that allow a large fraction of the responsibility for homeostatic control of sodium excretion to reside in those primary inputs that act to influence tubular reabsorption of sodium independently of GFR changes.

Before describing the mechanisms of long-term control in the next section we want to point out 2 key features of the renal handling of sodium. First, interactions between the various mechanisms we have described thus far allow the kidneys to be true integrators of signals that are sometimes in conflict. A good example is the case of prolonged aerobic exercise, specifically marathon running. Well-trained, well-hydrated athletes running marathons on cool days (thus eliminating excessive loss of sodium as a confounding factor) exercise intensely for well over 2 h with an elevated blood pressure. Systolic pressure is typically elevated by 50%, while MAP is elevated about 20%. Acting alone this rise in pressure should induce vigorous pressure natriuresis. But it does not. If anything, renal excretion of sodium is decreased in these conditions because other signals override pressure natriuresis.

We also want to point out that all the mechanisms described so far lead to co-regulation of solute and water, ie, they tend by themselves to increase or decrease the excretion of sodium and water in exact parallel. This is very effective as a coarse control over ECF volume. However, sodium ingestion

Table 7–1. Effect of “perfect” glomerulotubular balance on the mass of sodium leaving the proximal tubule

<table>
<thead>
<tr>
<th>GFR (L/min)</th>
<th>$P_{\text{Na}}$ (mmol/L)</th>
<th>Filtered (mmol/min)</th>
<th>Reabsorbed proximally (66.7% of filtered; mmol/min)</th>
<th>Leaving proximal (mmol/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.124</td>
<td>145</td>
<td>18</td>
<td>12</td>
<td>6</td>
</tr>
<tr>
<td>0.165</td>
<td>145</td>
<td>24</td>
<td>16</td>
<td>8</td>
</tr>
<tr>
<td>0.062</td>
<td>145</td>
<td>9</td>
<td>6</td>
<td>3</td>
</tr>
</tbody>
</table>

The net result of fixed fractional reabsorption is to reduce the magnitude of difference in sodium leaving the proximal tubule.
and water ingestion are both highly variable and often unrelated to each other. If one is ingested in excess of the other, the body has to excrete more of whichever one is in excess. Such independent control requires additional mechanisms not operative in the proximal tubule. Most of the processes for independent control of sodium and water balance occur in the distal nephron (not surprising because the distal nephron represents mammalian evolutionary adaptation to a terrestrial environment).

**Long-Term Control: Aldosterone Regulation of Sodium Balance**

In the face of a constant rate of ingestion of salt and water, correction of a sustained decrease in blood pressure requires a decrease in renal excretion of salt and water until the transient positive fluid balance returns blood volume to normal. A major control over the reabsorption of sodium in the distal nephron involves the hormone aldosterone. The primary effect of aldosterone is to increase sodium reabsorption in the connecting tubules and collecting ducts. Aldosterone-stimulated sodium retention is an effector system that is vital in correcting prolonged reductions in blood pressure. The most important physiological factor controlling circulating levels of aldosterone is the circulating level of angiotensin II. Thus, a decrease in blood pressure produces a rapid short-term baroreceptor-mediated vascular response followed by the intermediate-term renal-mediated release of renin and production of angiotensin II, which reinforces the initial short-term vascular response. However, even if the blood pressure returns to near normal, the circulating angiotensin II will stimulate the adrenal cortex to produce aldosterone. This targets the distal nephron to increase sodium reabsorption and thus increase total body sodium and blood volume to produce a long-term correction to total body sodium content and mean blood pressure.

Aldosterone stimulates sodium reabsorption mainly in the cortical connecting tubule and cortical collecting duct, specifically by the principal cells. An action on this late portion of the nephron is what one would expect for fine-tuning the output of sodium, because more than 90% of the filtered sodium has already been reabsorbed by the time the filtrate reaches the collecting-duct system.

The total quantity of sodium reabsorption dependent on the influence of aldosterone is approximately 2% of the total filtered sodium. Thus, all other factors remaining constant, in the complete absence of aldosterone, a person would excrete 2% of the filtered sodium, whereas in the presence of maximal plasma concentrations of aldosterone, virtually no sodium would be excreted. Two percent of the filtered sodium may seem small but is actually large because of the huge volume of glomerular filtrate:

$$\text{Total filtered Na/day} = \text{GFR} \times \text{PNa}$$
$$= 180 \text{ L/day} \times 145 \text{ mmol/L}$$
$$= 26,100 \text{ mmol/day}$$

*Circulating angiotensin II has a very short plasma half-life; thus, continued stimulation of aldosterone secretion requires the continued production of angiotensin II.*
Thus, aldosterone controls the reabsorption of $0.02 \times 26,100 \text{ mmol/day} = 522 \text{ mmol/day}$. In terms of sodium chloride, the form in which most sodium is ingested, this amounts to the control of approximately 30 g NaCl/day, an amount considerably more than the average person consumes. Therefore, by control of the plasma concentration of aldosterone between minimal and maximal, the excretion of sodium can be finely adjusted to the intake so that total-body sodium and ECF volume remain constant. (Interestingly, aldosterone also stimulates sodium transport by other epithelia in the body, namely, sweat and salivary ducts and the intestine. The net effect is the same as that exerted on the kidney: movement of sodium from lumen to blood. Thus, aldosterone is an all-purpose stimulator of sodium retention.) In the kidney, aldosterone acts like many other steroid hormones. As a molecule, it has enough lipid character to freely cross principal cell membranes, after which it combines with mineralocorticoid receptors in the cytoplasm. Aldosterone-bound receptors undergo a change in conformation that reveals a formerly hidden nuclear localization signal. After being transported to the nucleus, the receptor acts as a transcription factor that promotes gene expression and synthesis of messenger RNA (mRNA). The mRNA mediates the translation of specific proteins. The effect of these proteins is to increase the activity or number of luminal membrane sodium channels and basolateral membrane Na-K-ATPase pumps to exactly supply what is needed to promote increased reabsorption of sodium (Figure 7–11).

![Figure 7–11. Mechanism of aldosterone action.](image)

Aldosterone enters principal cells and interacts with cytosolic aldosterone receptors. The aldosterone-bound receptors interact with nuclear DNA to promote gene expression. The aldosterone-induced gene products activate sodium channels and sodium pumps to increase sodium reabsorption. Glucocorticoids such as cortisol are also capable of binding to the aldosterone receptor. However, they are inactivated by 11β-hydroxysteroid dehydrogenase (11β-HSD).
Control of Aldosterone Secretion

Several inputs to the adrenal gland regulate aldosterone secretion and play a role in electrolyte balance. The most important is angiotensin II produced by the global RAS. In addition, elevated plasma potassium concentration, as described in Chapter 8 in the context of the renal handling of potassium, stimulates aldosterone secretion, while the atrial natriuretic factors (discussed later) inhibit aldosterone secretion.

As described earlier, the plasma concentration of angiotensin II is determined mainly by the plasma concentration of renin. Accordingly, control of aldosterone secretion in sodium-regulating reflexes is determined by those factors that regulate renin secretion (ie, intrarenal baroreceptors, macula densa, and renal sympathetic nerves). Thus, when plasma volume is reduced, eg, by a low-sodium diet, hemorrhage, or diarrhea, renin secretion is stimulated, which leads, via angiotensin II, to an increased aldosterone secretion. This hormone then stimulates sodium reabsorption (Figure 7–12). In contrast, when a person ingests a high-sodium diet, renin secretion is reduced, which leads, via a reduced plasma angiotensin II, to decreased aldosterone secretion.

Tubuloglomerular Feedback and Autoregulation Revisited

Responses to cardiopulmonary, arterial, and intrarenal baroreceptors are extremely effective mechanisms for controlling blood volume. Part of this control causes changes in GFR, mediated by changes in afferent arteriolar resistance. Although this change in afferent resistance has the effect of altering GFR in a manner necessary to correct blood volume, it has the additional effect of altering RBF and pressure in the glomerular capillaries that may have the deleterious consequences described in Chapter 2. Substantial reductions in RBF severely compromise already oxygen-poor regions of the kidney like the medulla. Substantial increases in glomerular capillary pressures are likely to damage the glomeruli. In addition, the ability of the kidney to correct total body electrolyte and water imbalances depends on keeping tubular flow (ie, GFR) within a certain limited range. Therefore, the kidneys have specific mechanisms for blunting responses that would otherwise lead to excessively large changes in GFR or RBF. These mechanisms are autoregulation and tubuloglomerular feedback. It is important to emphasize that these mechanisms do not block changes in GFR and renal blood flow; they simply keep the changes from becoming excessive.

Autoregulation of GFR involves local production of prostaglandins in conditions when strong vasoconstriction might by itself reduce GFR and renal blood flow too much (high sympathetic stimulation and high levels of angiotensin II). Intrarenal (autoregulatory) prostaglandin production opposes the actions of angiotensin II on the kidneys, ie, prostaglandins lead to vasodilation of arterioles and relaxation of mesangial cells (Figure 7–8). Increased local (intrarenal) angiotensin II concentrations associated with renin release and increased sympathetic input stimulate the production of prostaglandins. The vasodilatory effect...
of prostaglandins dampens the effect of angiotensin II and sympathetic input on renal arterioles and permits a reasonable, but reduced blood flow and GFR to continue (see Figure 7–13).

Tubuloglomerular feedback, alternatively, is associated with the macula densa sodium chloride load detector and plays a major role in conditions when GFR is very high (eg, volume overload). Recall from the previous discussion that large loads of sodium chloride in the thick ascending limb lead to inhibition of renin release. The macula densa cells at the end of the thick ascending limb have Na-K-2Cl symporters that can avidly take up Na, Cl, and K and cause the cells to swell dramatically when GFR (NaCl delivery) is high (see Figure 7–9). The increased Na and Cl in the lumen of the thick ascending limb stimulate the Na-H antiporter and depolarizes the cells (as in thick ascending limb cells, the K recycles via K channels). This depolarization leads to Ca entry across the basolateral membrane. The rise in Ca leads to the release of ATP from the basolateral surface of
the cells in close proximity to the glomerular mesangial cells. This ATP stimulates purinergic P2 receptors on the mesangial cells and afferent arteriolar smooth muscle cells. P2 receptor stimulation increases Ca in these cells and promotes contraction. In addition, it is the increased Ca in the afferent arteriolar cells that reduces renin secretion. The ATP may also be metabolized to adenosine, which can stimulate adenosine receptors that produce the same result as the P2 receptors (in contrast to the vasodilatory actions of adenosine in most other tissues). Contraction of mesangial cells decreases the effective filtration area, which decreases GFR. Contraction of the afferent arteriolar smooth muscle cells increases afferent resistance and decreases RBF and GFR (see Figures 7–14 and 7–15).

Figure 7–13. Prostaglandins mediate autoregulatory responses. Production of prostaglandins (mostly PGE₂) near the glomerulus relaxes the afferent arteriole and thus counteracts the contractile effects of renal sympathetic nerve activation and angiotensin II.

6The actions of adenosine in a given cell depend on the type of purinergic receptor and the signaling pathway initiated on binding of adenosine, similar to the situation with adrenergic receptors, in which an array of receptor types permits a variety of responses to any given agonist.

7The increase in intracellular Ca of the granular cells inhibits their production of intrarenal renin, thus reducing local production of angiotensin II and of prostaglandins, which would normally counteract the vasoconstrictive effects of the purinergic agonists. Another mediator—nitric oxide (NO)—is not a factor in initiating tubuloglomerular feedback but does appear to play a secondary role to sustain the tubuloglomerular feedback once it has been initiated. The net effect of tubuloglomerular feedback is that the pressure natriuretic and diuretic responses are blunted (but not eliminated).
Figure 7–14. Mechanism of tubuloglomerular feedback. Tubuloglomerular feedback acts to prevent changes in renal artery pressure from causing extreme changes in sodium delivery to the macula densa. This mechanism acts in the opposite direction to the other reflexes and thus partially reduces or blunts their effectiveness. However, the overall effect of an increase in renal artery pressure is still a net increase in sodium excretion (compare with Figure 7–10). GFR, glomerular filtration rate; $P_{GC}$, hydrostatic pressure in glomerular capillaries.

The set of events just described is admittedly confusing, so the bottom line is this: high salt content in the thick ascending limb of a given nephron generates signals that reduce filtration in that nephron, thus blunting (but not eliminating) the tendency to raise sodium excretion initiated by other process in conditions (eg, volume expansion) where the appropriate overall response is increased sodium excretion.
Although there are several other renal mechanisms for controlling sodium balance independent of water balance, under normal physiological circumstances none is as important as aldosterone. Only under certain pathophysiological conditions do these other mechanisms contribute significantly to the regulation of sodium balance.

**Natriuretic Peptides**

Several tissues in the body synthesize members of a hormone family called *natriuretic peptides*, so named because they promote excretion of sodium in the urine. Key among these are atrial natriuretic peptide (ANP) and brain natriuretic peptide (BNP; named as such because it was first discovered in the brain). The main source of both natriuretic peptides is the heart. The natriuretic peptides have both vascular and tubular actions. They relax the afferent arteriole, thereby promoting increased filtration, and act at several sites in the tubule. They inhibit the release of renin, inhibit the actions of angiotensin II that normally promote reabsorption of sodium, and act in the medullary collecting duct to inhibit sodium absorption. The major stimulus for increased secretion of the natriuretic peptides is distention of the atria, which occurs during plasma volume expansion. This is probably the stimulus for the increased natriuretic peptides that occurs in persons on a high-salt diet. Although most experts assume that these peptides play some physiological role in the regulation of sodium excretion in this and other situations in which
plasma volume is expanded, it is not currently possible to quantitate precisely their contribution, although it is surely less than aldosterone. As described later, these peptides are greatly elevated in patients with heart failure and serve as diagnostic indicators.

**Antidiuretic Hormone**

As described in Chapter 6, the major function of ADH is to increase the permeability of the cortical and medullary collecting ducts to water, thereby decreasing the excretion of water. In addition to this effect, ADH also increases sodium reabsorption by the cortical collecting duct, one of the same segments influenced by aldosterone. This effect is particularly evident when plasma aldosterone is elevated, and ADH’s action seems to synergize with the action of this steroid hormone. This makes teleological sense because, as discussed later, the secretion of ADH, like that of aldosterone, is stimulated when plasma volume is reduced.

**Other Hormones**

Many well-known hormones not normally associated with renal function can exert an influence on sodium reabsorption. Cortisol, estrogen, growth hormone, thyroid hormone, and insulin enhance sodium reabsorption, whereas glucagon, progesterone, and parathyroid hormone decrease it. When the level of any of these hormones is elevated (e.g., estrogen during pregnancy), it will exert a significant influence on sodium reabsorption and thus excretion. However, the secretion of these hormones, unlike the hormones described earlier, is not reflexively controlled specifically for the homeostatic regulation of sodium balance.

**Summary of the Control of Sodium Excretion**

The control of sodium excretion depends on the control of 2 variables of renal function: GFR and rate of sodium reabsorption (Tables 7–2 and 7–3). The latter is controlled by the renin-angiotensin-aldosterone hormonal system, renal sympathetic nerves, direct effects of arterial blood pressure on the kidneys (pressure natriuresis), and atrial natriuretic factors. The renal interstitial hydraulic pressure and several renal paracrine agents play important roles in regulating sodium reabsorption.

When considering mechanisms of sodium excretion, it is useful to consider 2 conceptually different categories of mechanisms: (1) proximal nephron mechanisms (control of GFR, pressure natriuresis, and, to a lesser extent, changes in Starling forces) that lead to coupled changes in sodium and water excretion and (2) distal nephron effects in which sodium can be reabsorbed independently of water. The proximal mechanisms are primarily involved in excreting excess ECF volume, whereas the distal mechanisms alter sodium excretion when ingestion of sodium is not balanced by ingestion of water. Both types of mechanisms can alter blood pressure because of the intimate relationship among total body sodium and water, blood volume, and blood pressure.

There is great flexibility in such a multifactor system. Thus, e.g., although the renal sympathetic nerves influence GFR, renin secretion, renal interstitial hydraulic pressure, and the tubular cells themselves, a transplanted and, therefore, denervated kidney maintains sodium homeostasis quite well because of the other
known nonneural factors involved. Overall, the one input whose absence causes the greatest difficulty in sodium regulation is aldosterone.

In normal persons, the mechanisms for regulating sodium excretion are so precise that sodium balance does not vary by more than a small percentage despite marked changes in dietary intake or losses caused by sweating, vomiting, diarrhea, hemorrhage, or burns.
CONTROL OF WATER EXCRETION

Over time, water excretion must meet the constraint of balance: matching output to input. However, there is no physiological “water meter” to measure input. So output is not controlled by input. Instead, output is regulated by factors relating to the major “big picture” goals described in the introduction to this chapter. That is, maintain a volume sufficient to fill the vascular space, and set an osmolality appropriate for a healthy environment of tissue cells. Then, it is not surprising that the major signals regulating water excretion originate from baroreceptors that assess vascular fullness and osmoreceptors that assess plasma osmolality.

Water excretion conceptually consists of 2 major components: a proximal nephron component, in which water is absorbed along with sodium as an isotonic fluid, and a distal nephron component, in which water can be reabsorbed independent of sodium. The proximal nephron component is primarily a mechanism to regulate ECF volume in response to changes in blood pressure, while the distal nephron rate of water reabsorption is independent of sodium reabsorption. It is determined mainly by ADH, which increases the water permeability of the collecting ducts, thereby increasing water reabsorption and, hence, decreasing water excretion. Accordingly, total-body water is regulated mainly by reflexes that alter the secretion of ADH.

ADH is a peptide produced by a discrete group of hypothalamic neurons whose cell bodies are located in the supraoptic and paraventricular nuclei and whose axons terminate in the posterior pituitary gland, from which ADH is released into the blood. The most important of the inputs to these neurons are from cardiovascular baroreceptors and osmoreceptors.

Baroreceptor Control of ADH Secretion

A decreased extracellular volume (eg, resulting from diarrhea or hemorrhage) reflexively produces an increased aldosterone secretion. It also induces increased ADH secretion. The reflex is mediated by neural input to the ADH-secreting neurons from both cardiopulmonary and arterial baroreceptors.

Decreased cardiovascular pressures cause less firing by the baroreceptors. Via afferent neurons from the baroreceptors and ascending pathways to the hypothalamus, this decreased baroreceptor firing causes stimulation of ADH secretion. Conversely, the baroreceptors are stimulated by increased cardiovascular pressures, and this results in the inhibition of ADH secretion. The adaptive value of these baroreceptor reflexes is to help restore ECF volume and, hence, blood pressure (Figure 7–16).

There is a second adaptive value to this reflex: Large decreases in plasma volume elicit, by way of the cardiovascular baroreceptors, such high concentrations of ADH—much higher than those needed to produce maximal antidiuresis—that the hormone is able to exert direct vasoconstrictor effects on arteriolar smooth muscle. The result is increased total peripheral resistance, which helps raise arterial blood pressure independently of the slower restoration of body fluid volumes.
Renal arterioles and mesangial cells also participate in this constrictor response, and so a high plasma concentration of ADH, quite apart from its effect on tubular water permeability, may promote the retention of both sodium and water by lowering GFR.

**Osmoreceptor Control of ADH Secretion**

We have seen how changes in ECF volume simultaneously elicit reflex changes in the excretion of both sodium and water. This is adaptive because the situations causing ECF volume alterations are often associated with loss or gain of both sodium and water in approximately proportional amounts. In contrast, we see now that changes in total-body water in which no change in total-body sodium occurs are compensated by alterations in water excretion but not in sodium excretion.

The major effect of gaining or losing water without corresponding changes in sodium is a change in the osmolality of the body fluids. This is a key point because, under conditions of gain or loss of water without solute, the receptors that initiate the reflexes controlling ADH secretion are osmoreceptors: receptors responsive to...
changes in osmolality. Most osmoreceptors are located in tissues surrounding the 3rd cerebral ventricle. These tissues contain fenestrated capillaries, which allow rapid adjustment of interstitial composition when plasma composition changes. The hypothalamic cells that secrete ADH receive neural input from the osmoreceptors. Via these connections, an increase in osmolality stimulates them and increases their rate of ADH secretion. Conversely, decreased osmolality inhibits ADH secretion (see Figure 7–16). For example, when a person drinks 1 L of water, the excess water lowers the body fluid osmolality, which reflexively inhibits ADH secretion via the hypothalamic osmoreceptors. As a result, water permeability of the collecting ducts becomes very low, little or no water is reabsorbed from these segments, and a large volume of extremely dilute (hypo-osmotic) urine is excreted. In this manner, the excess water is eliminated.

Conversely, when a pure-water deficit occurs (eg, because of water deprivation), the osmolality of the body fluids is increased, ADH secretion is reflexively stimulated, water permeability of the collecting ducts is increased, water reabsorption is maximal, and a very small volume of highly concentrated (hyperosmotic) urine is excreted. By this means, relatively less of the filtered water than solute is excreted, which lowers body fluid osmolality toward normal.

We have described 2 different major afferent pathways controlling the ADH-secreting hypothalamic cells: one from baroreceptors and one from osmoreceptors. These hypothalamic cells are, therefore, true integrators, whose rate of activity is determined by the total synaptic input to them. Thus, a simultaneous increase in plasma volume and decrease in body fluid osmolality cause strong inhibition of ADH secretion. Conversely, a simultaneous decrease in plasma volume and increase in osmolality produce very marked stimulation of ADH secretion. However, what happens when baroreceptor and osmoreceptor inputs oppose each other (eg, if plasma volume and osmolality are both decreased)? In general, because of the high sensitivity of the osmoreceptors, the osmoreceptor influence predominates over that of the baroreceptor when changes in osmolality and plasma volume are small to moderate. However, a very large change in plasma volume will take precedence over decreased body fluid osmolality in influencing ADH secretion; under such conditions, water is retained in excess of solute, and the body fluids become hypo-osmotic (for the same reason, plasma sodium concentration decreases). In essence, it is more important for the body to preserve vascular volume and thus ensure an adequate CO than it is to preserve normal osmolality.

The ADH-secreting cells also receive synaptic input from many other brain areas. Thus, ADH secretion and, hence, urine flow can be altered by pain, fear, and a variety of other factors, including drugs such as alcohol, which inhibits ADH release. However, this complexity should not obscure the generalization that ADH secretion is determined over the long term primarily by the states of body fluid osmolality and plasma volume.

Recent evidence suggests that there are both osmolality-sensing and direct sodium concentration-sensing cells in the brain, the latter via a special class of sodium channels located primarily in glial cells. In addition, the ADH-secreting cells in the hypothalamus may also respond to osmolality in the cerebrospinal fluid, which responds on a slower time scale to changes in plasma composition.
Several diseases (e.g., diabetes insipidus, which is different from diabetes mellitus, also known as sugar diabetes) illustrate what happens when the ADH system is disrupted. Diabetes insipidus is characterized by a constant water diuresis, as much as 25 L/day. In most cases, people with diabetes insipidus have lost the ability to produce ADH because of damage to the hypothalamus or have lost the ability to respond to ADH because of defects in principal cell ADH receptors. Thus, collecting-duct permeability to water is low and unchanging regardless of extracellular osmolality or volume. In contrast, other diseases (e.g., head trauma or brain tumors) are associated with inappropriately large secretion of ADH. As is predictable, patients with these diseases manifest decreased plasma osmolality (and sodium concentration) because of the excessive reabsorption of pure water.

Figure 7–17 summarizes the major factors known to control renal sodium and water excretion in response to severe sweating. Sweat is a hypo-osmotic solution containing mainly water, sodium, and chloride. Therefore, sweating causes both a decrease in ECF volume and an increase in body fluid osmolality. The renal retention of water and sodium helps to compensate for the water and salt lost in the sweat.

**Thirst and Salt Appetite**

Large deficits of salt and water can be only partly compensated by renal conservation of these substances, and ingestion is the ultimate compensatory mechanism. The centers that mediate thirst are located in the hypothalamus (very close to those areas that produce ADH). The subjective feeling of thirst, which drives one to obtain and ingest water, is stimulated both by reduced plasma volume and by increased body fluid osmolality. The adaptive significance of both are self-evident. Note that these are precisely the same changes that stimulate ADH production, and the receptors—osmoreceptors and the nerve cells that respond to the cardiovascular baroreceptors—that initiate the ADH-controlling reflexes are near those that initiate thirst. The thirst response, however, is significantly less sensitive than the ADH response.

There are also other pathways controlling thirst. For example, dryness of the mouth and throat causes profound thirst, which is relieved by merely moistening them. Also, when animals such as the camel (and humans, to a lesser extent) become markedly dehydrated, they will rapidly drink just enough water to replace their previous losses and then stop. What is amazing is that when they stop, the water has not yet had time to be absorbed from the gastrointestinal tract into the blood. Some kind of metering of the water intake by the gastrointestinal tract has occurred, but its nature remains a mystery.

Angiotensin II is yet another factor that stimulates thirst: by its direct effect on the brain. This hormone constitutes one of the pathways by which thirst is stimulated when ECF volume is decreased.

Salt appetite, which is the analogue of thirst, is also an extremely important component of sodium homeostasis in most mammals. It is clear that salt appetite in these species is innate and consists of 2 components: (1) hedonistic appetite and (2) regulatory appetite. In other words, (1) animals like salt and eat it whenever
they can regardless of whether they are salt deficient, and (2) their drive to obtain salt is markedly increased in the presence of deficiency.

The significance of these animal studies for humans, however, is unclear. Salt craving does seem to occur in humans who are severely salt depleted, but the contribution of such regulatory salt appetite to everyday sodium homeostasis in
normal persons is probably slight. On the other hand, humans do seem to have a strong hedonistic appetite for salt, as manifested by almost universally large intakes of sodium whenever it is cheap and readily available. Thus, the average American intake of salt is 10–15 g/day even though humans can survive quite normally on less than 0.5 g/day. As pointed out previously, a large salt intake may be a contributor to the pathogenesis of hypertension in susceptible individuals.

Congestive Heart Failure and Hypertension: Cardiovascular Pathologies That Involve Altered Sodium Excretion

Congestive heart failure and hypertension involve perturbed renal handling of sodium. In congestive heart failure and in most cases of hypertension, the perturbed renal function seems to lie in inappropriate signaling to the kidneys rather than pathology of renal transport mechanisms per se.

Congestive heart failure is characterized by weak cardiac muscle that cannot increase CO to meet the demands of exercise and, more importantly, can only provide an adequate resting CO in the presence of excessive neurohumoral drive (something like a car operating on 2 cylinders that can only keep up speed when the accelerator is pushed to the floor). The neurohumoral drive is characterized by high levels of renin, angiotensin II, aldosterone, catecholamines, and other mediators. Fluid volume is increased, leading to edema in the lungs, peripheral tissues, or both, which is why this is called congestive heart failure. Because of the high fluid volume, atrial pressures sensed by the cardiopulmonary baroreceptors are high. The high atrial pressures should lead to decreased ADH secretion and decreased sympathetic drive to the kidneys. Instead, these signals are increased, and the kidneys operate at a new setpoint in which normal sodium excretion only occurs at the expense of an excessive body fluid volume. If fluid volume is somehow restored to normal levels, renal excretion of sodium drops to very low levels. Another characteristic of congestive heart failure is high levels of natriuretic peptides. This is an appropriate response to the high atrial pressures and partially counteracts the sodium-retaining signals to the kidneys but does not restore sodium output to a level that would occur in a healthy person who transiently developed the high fluid volume that exists chronically in the heart failure patient. The high fluid volumes of congestive heart failure are deleterious to pulmonary function and over time often lead to structural changes in the heart (dilation) that only exacerbate the defective pumping. Therapy for congestive heart failure includes the use of diuretics to reduce the high fluid volume and drugs that inhibit the generation of angiotensin II (ACE inhibitors) or block the actions of angiotensin II (angiotensin receptor antagonists). In addition, synthetic natriuretic peptides are becoming tools to promote diuresis.

Hypertension is also a disease of abnormal sodium balance. Hypertension must always be associated with a blood volume and total body sodium content that is

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9In contrast to atrial pressures that are high, arterial pressure is usually within the normal range, and heart failure cannot be diagnosed based on arterial blood pressure. It may be that arterial pressure is actually a little below the setpoint of the brainstem vasomotor center, leading to the sympathetic drive associated with heart failure. However, there is no way to measure the setpoint and compare it with the pressures that actually exist in the arterial tree.
too high for the volume of the vascular tree. In some cases, the reason for the excess blood volume is clear. For example, renal glomerular disease often leads to inappropriate release of renin with subsequent increases in angiotensin II, aldosterone, collecting-tubule sodium reabsorption, and finally an increase in blood pressure; a tumor of the adrenal cortex can lead to excess production of aldosterone and increases in blood pressure; or a specific gain of function mutation in the sodium reabsorptive mechanism in the collecting duct also leads to excess sodium reabsorption and profound hypertension. These 3 examples illustrate 3 types of defects in the complicated control mechanism for maintaining total body sodium and mean blood pressure. The first, excess renin production, is a problem with the sensing mechanism for the renal component of the blood pressure control system. The second, excess production of aldosterone, is a defect in the signaling mechanism that lies between the sensor (pressure sensors in the large vessels and the kidney) and the effector (distal nephron sodium reabsorption). The third example is a defect in the effector system (distal nephron sodium reabsorption). In these cases, the defect is more or less obvious, and correcting the underlying pathology usually corrects the hypertension (ie, ACE inhibitors reduce the effect of excess renin, spironolactone will inhibit aldosterone receptors, and administration of amiloride will reduce sodium reabsorption by epithelial sodium channels). Renin levels, angiotensin levels, and aldosterone levels are normal or even reduced, and yet blood pressure is elevated as if the setpoint of the control loop from sensing blood pressure to sodium reabsorption is inexplicably high. The relatively normal levels of circulating renin, angiotensin II, and aldosterone imply that the defect in the regulation of sodium reabsorption lies subsequent to aldosterone interaction with the cells of the collecting tubule. One holy grail of hypertension research has been to define the components responsible for controlling sodium reabsorption that are induced by aldosterone in an effort to intervene in the process that leads to increased total body sodium and increased blood pressure.

### KEY CONCEPTS

1. **Multiple overlapping mechanisms regulate sodium and water excretion; most are related to blood pressure.**

2. **The medullary vasomotor center regulates blood pressure on a moment-to-moment basis via the baroreceptor reflex and also regulates renal excretion of sodium and water.**

3. **Angiotensin II is a crucial regulator of sodium excretion and blood pressure via its actions in the kidneys, peripheral vasculature, and adrenal glands.**

4. **The regulation of sodium content is the ultimate determinant of blood pressure in the long term via control of extracellular fluid (ECF) volume.**
7–1. In a canine experiment, a dog’s filtered load of sodium in an isolated pump-perfused kidney is found to be 15 mmol/min. (1) How much sodium do you predict remains in the tubule at the end of the proximal tubule? (2) If its GFR is suddenly increased by 33%, how much sodium now is left at the end of the proximal tubule?

7–2. Normally aldosterone stimulates the reabsorption of approximately 33 g of sodium chloride/day. If a patient loses 100% of adrenal function, will 33 g of sodium chloride be excreted per day indefinitely?

7–3. A patient has suffered a severe hemorrhage and the plasma protein concentration is normal. (Not enough time has elapsed for interstitial fluid to move into the plasma.) Does this mean that the peritubular-capillary oncotic pressure is also normal?

7–4. If the right renal artery becomes abnormally constricted, what will happen to renin secretion by the right kidney and the left kidney?

7–5. A patient is suffering from primary hyperaldosteronism (ie, increased secretion of aldosterone caused by an aldosterone-producing adrenal tumor). Is plasma renin concentration higher or lower than normal?

7–6. An agent that increases sodium and water excretion is called a diuretic (even though natriuretic is probably a better term). Block of sodium reabsorption in the proximal tubule, loop of Henle, distal tubule, or collecting duct all exert a diuretic effect. True or false?

7–7. A person is given a drug that dilates both the afferent and efferent arterioles. Assuming no other action of the drug, what will happen to the percentage of filtered sodium that this person’s proximal tubule reabsorbs?

7–8. A new drug is found to have dual actions: It blocks sodium entry pathways in the proximal tubule epithelium, and it binds to ADH receptors in the collecting ducts and mimics the actions of ADH. Will the final urine contain excess or low amounts of sodium and excess or low amounts of water, and will it be hyperosmotic, isosmotic, or hypo-osmotic?

7–9. Another new drug also has dual actions, this time blocking sodium entry pathways in the thick ascending limb and exerting ADH-like actions as in Question 7–8.
Now will the final urine contain excess or low amounts of sodium and excess or low amounts of water, and will it be hyperosmotic, iso-osmotic, or hypo-osmotic?

7–10 Angiotensin II that is capable of regulating the kidneys is formed by enzymatic action in which locations (choose all that apply)?

A. The adrenal glands
B. The kidneys
C. The hypothalamus
D. The lungs

SUGGESTED READING


Renal Regulation of Potassium Balance

OBJECTIVES

The student understands the internal exchanges of potassium:

- States the normal balance and distribution of potassium between cells and extracellular fluid.
- Describes how potassium moves between cells and the extracellular fluid, and how, on a short-term basis, the movement protects the extracellular fluid from large changes in potassium concentration.
- Describes how plasma levels of potassium do not always reflect the status of total-body potassium.
- States how insulin and epinephrine influence the cellular uptake of potassium and identifies the situations in which these hormonal influences are most important.

The student understands the renal regulation of potassium excretion:

- States generalizations about renal potassium handling for persons on high- or low-potassium diets.
- States the relative amounts of potassium reabsorbed by the proximal tubule and thick ascending limb of Henle’s loop regardless of the state of potassium intake.
- Describes how nephron segments beyond the thick ascending limb can manifest net secretion or reabsorption; describes the role of principal cells and intercalated cells in these processes.
- Lists inputs that control the rate of potassium secretion by the distal nephron.
- Describes the actions of ROMK and BK potassium channels in conditions of low, normal, and high potassium excretion.
- Describes the mechanism by which changes in potassium balance influence aldosterone secretion.
- States the effects of most diuretic drugs and osmotic diuretics on potassium excretion.
- Describes the association between perturbations in acid-base status and the plasma potassium level.

Potassium, like all other important ions, is distributed between the intracellular fluid and extracellular fluid (ECF) of the body. But unlike sodium, the vast majority of potassium is intracellular, and only about 2% of total-body potassium is extracellular. This small fraction, however, is absolutely
crucial for body function, and the concentration of potassium in the ECF is a closely regulated quantity. Major elevations or depressions (called hyperkalemia and hypokalemia) away from the normal value of 4 mEq/L are cause for medical intervention. The importance of maintaining this concentration relatively constant stems primarily from the role of potassium in the excitability of nerve and muscle. The resting membrane potentials of these tissues are strongly influenced by the ratio of intracellular to extracellular potassium concentration. Raising the extracellular potassium concentration depolarizes the resting membrane potential, thus perturbing cell excitability. Conversely, lowering the extracellular potassium concentration usually hyperpolarizes cell membranes.1

REGULATION OF POTASSIUM BETWEEN THE INTRACELLULAR AND EXTRACELLULAR COMPARTMENTS

Given that the vast majority of body potassium is within cells, the extracellular potassium concentration is crucially dependent on (1) the total amount of potassium in the body and (2) the distribution of this potassium between the extracellular and intracellular fluid compartments. Total-body potassium is determined by the balance between potassium intake and excretion. Normal individuals remain in potassium balance, as they do in sodium balance, by excreting an amount of urinary potassium equal to the amount of potassium ingested minus the small amounts eliminated in the feces and sweat. Normally, potassium losses via sweat and the gastrointestinal tract are small, but very large quantities can be lost from the digestive tract during vomiting or diarrhea. Again, the control of renal function is the major mechanism by which total-body potassium is maintained in balance.

The fact that most body potassium is intracellular follows strictly from the size and properties of the intracellular and extracellular compartments. About two thirds of the body fluids are intracellular (the collective cytosolic volume of all the cells in the body); typical cytosolic potassium concentrations are about 140–150 mEq/L. One third of the body fluids are extracellular, with a potassium concentration of about 4 mEq/L. In a clinical setting, only the extracellular concentration can be measured (the intracellular potassium is, in a sense, hidden behind the wall of the cell membrane). Furthermore, the extracellular value does not necessarily reflect total-body potassium. A patient may, for example, be hyperkalemic (high plasma potassium concentration) and yet at the same time be depleted of total-body potassium.

The high level of potassium within cells is maintained by the collective operation of the sodium-potassium-adenosine triphosphatase (Na-K-ATPase) plasma membrane pumps, which actively transport potassium

1Whether a given change in potassium concentration makes an excitable cell more or less excitable is not always obvious. Depolarization resulting from elevated extracellular potassium may make a cell more excitable, but very strong depolarization actually decreases excitability (because of inactivation of voltage-gated sodium channels), a situation called a depolarizing block. In cases of very high extracellular potassium, a depolarizing block in the heart prevents propagation of the electrical signal that causes synchronized contraction.
into cells. Because the amount of potassium in the extracellular compartment is so small (40–60 mEq total), even very slight shifts of potassium into or out of cells can produce large changes in extracellular potassium concentration. Similarly, a meal rich in potassium (e.g., steak, potato, and spinach) could easily double the extracellular concentration of potassium if most of that potassium were not transferred from the blood to the intracellular compartment. It is crucial, therefore, that dietary loads be taken up into the intracellular compartment rapidly to prevent major changes in plasma potassium concentration. The tissue contributing most to the sequestration of potassium is skeletal muscle, simply because it contains the largest collective intracellular volume. Muscle effectively buffers extracellular potassium by taking up or releasing it and keeping the plasma potassium concentration close to normal. On a moment-to-moment basis, this is what protects the ECF from large swings in potassium concentration. Major factors involved in these homeostatic processes include insulin and epinephrine, both of which cause increased potassium uptake by muscle (and certain other cells) through stimulation of plasma membrane Na-K-ATPase. Another influence is the GI tract, which contains an elaborate neural network (the “gut brain”) that sends signals to the central nervous system. It also contains a complement of endocrine cells that release an array of peptide hormones. Together these neural and hormonal signals affect many target organs, including the kidneys (see later discussion) in response to dietary input. While details of GI signaling systems are still under investigation, it is well known that orally administered loads of substances like potassium and glucose are handled differently from identical loads that are administered intravenously.

The rise in plasma insulin concentration after a meal is a crucial factor in moving ingested and absorbed potassium into cells rather than allowing it to accumulate in the ECF. This new potassium then slowly comes out of cells between meals to be excreted. Moreover, a large increase in plasma potassium concentration facilitates insulin secretion at any time, and the additional insulin induces greater potassium uptake by the cells, a negative feedback system for opposing acute elevations in plasma potassium concentration. In the natural order of things, insulin also stimulates glucose uptake and metabolism by cells: a necessary source of energy to drive the insulin-activated Na-K-ATPase responsible for moving potassium into cells.

The effect of epinephrine on cellular potassium uptake is probably of greatest physiological importance during exercise and trauma. In exercise, potassium moves out of muscle cells that are rapidly firing action potentials, and damaged cells leak potassium. In both cases, this raises extracellular potassium concentration. However, at the same time, exercise or trauma increases adrenal secretion of epinephrine, and this hormone’s stimulation of potassium uptake by other cells partially offsets the outflow from the exercising or damaged muscle cells.

\[\text{During very intense intermittent exercise (e.g., wind sprints), extracellular potassium can easily double as a result of release from exercising muscle. As soon as exercise stops, the muscle takes it back up rapidly, within 1 minute or so. Without the influence of catecholamine hormones on sensitive tissues such as the heart, the suddenly high levels of potassium reached during intense exercise would be quite dangerous.}\]
Still another influence on the distribution of potassium between the intracellular fluid and ECF is the ECF hydrogen ion concentration: An increase in ECF hydrogen ion concentration (acidosis; see Chapter 9) is often associated with net potassium movement out of cells, whereas a decrease in ECF hydrogen ion concentration (alkalosis) causes net potassium movement into them. It is as though potassium and hydrogen ions were exchanging across plasma membranes (ie, hydrogen ions moving into the cell during acidosis and out during alkalosis and potassium doing just the opposite), but the precise mechanism underlying these “exchanges” has not yet been clarified. However, like the effect of insulin, it probably involves an inhibition (acidosis) or activation (alkalosis) of the Na-K-ATPase.

RENAL POTASSIUM HANDLING

Overview

Although other tissues play an important role in the moment-to-moment control of plasma potassium concentration, in the final analysis, the kidney determines total-body potassium content. Therefore, understanding potassium handling by the kidneys is the key to understanding potassium balance. Potassium is freely filtered into Bowman’s space. Under all conditions almost all the filtered load (~90%) is reabsorbed by the proximal tubule and thick ascending limb of the loop of Henle. Then, if the body is trying to conserve potassium, most of the rest is reabsorbed in the distal nephron and medullary collecting duct, leaving almost none in the urine. In contrast, if the body is ridding itself of potassium, a large amount is secreted in the distal nephron, resulting in a large excretion. When secretion occurs at high rates, the amount excreted may exceed the filtered load. The chief means of regulation lies in control over secretion in parts of the nephron beyond the loop of Henle. Let us look at potassium handling by various nephron segments and then address the issue of control.

Since potassium is freely filtered, a normal plasma level of 4 mEq/L and GFR of 150 L/day or more results in a daily filtered load of about 600 mEq/day. The subsequent events in various tubule segments are summarized in Table 8–1. In the proximal tubule, about 65% of the filtered load is reabsorbed, mostly via the paracellular route. Some of the flux is driven by the concentration gradient set up when water is reabsorbed (thus concentrating all solutes remaining in the tubular lumen). Some may also move by entrainment with the rapidly reabsorbed water (solvent drag). Either way this accounts for major potassium absorption in an essentially unregulated manner.3 In the loop of Henle there is additional reabsorption. The major events take place in the thick ascending limb, where the Na-K-2Cl multiporter in the luminal membrane reabsorbs potassium (see Figure 6–3). Some of this potassium is returned to the lumen across the apical membrane via potassium channels, and the rest exits the cells across the basolateral membrane

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3Virtually every process in the kidney is regulated. However, for potassium in the proximal tubule, as is the case for a number of other solutes, reabsorption is tied to sodium reabsorption, and regulation is directed more toward sodium rather than other solutes.
by a combination of passive flux through channels and through symporters with chloride, resulting in net transcellular reabsorption. Some potassium is also reabsorbed by the paracellular route in this segment, driven by a lumen-positive voltage. Usually about 25% of the filtered load is reabsorbed in the thick ascending limb, so that only about 10% is passed on to the distal nephron.4

In the distal nephron there is continuous reabsorption if dietary loads are very small, but a major superimposed secretion that greatly exceeds the reabsorption when dietary loads are high. It is in these distal segments where most regulation of potassium excretion is exerted. The distal nephron is composed of a number of segments, including the distal convoluted tubule, connecting tubule, initial collecting tubule, and cortical collecting duct, ie, all of the tubule segments between the end of the thick ascending limb and the medullary collecting duct. It is not possible to finely differentiate between these segments in terms of function, although the connecting tubule stands out as being particularly important in potassium handling because of its rich complement of transport elements. It appears that most of the potassium secretion occurs before segments where most of the water is absorbed (cortical collecting duct). Finally, the medullary collecting ducts reabsorb potassium modestly under all conditions. When the sum of upstream processes has already reabsorbed almost all the potassium, the medullary collecting ducts bring the final urine excretion down to a few percent

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4Although the loops of Henle as a whole exhibit net reabsorption, long loops of Henle (from juxtamedullary nephrons) secrete potassium in the thin descending limb; this potassium comes from the medullary interstitium. In turn, interstitial potassium comes from potassium reabsorption in the medullary collecting ducts. Thus, there is some potassium recycling analogous to the urea recycling described earlier.
of the filtered load, for an excretion of about 10–15 mEq/day. On the other hand, if upstream segments are secreting avidly, the modest reabsorption in the medullary collecting ducts does little to prevent an excretion that can reach 1000 mEq/day. Figure 8–1A and B depict the overall renal handling of potassium in different tubule regions in conditions of high and low potassium excretion.

A complication in the renal handling of potassium in all regions, specifically including the proximal tubule and thick ascending limb, is that its active transport is always coupled with the active transport of another solute. Active influx of potassium across the basolateral membrane via the ubiquitous Na-K-ATPase is coupled with efflux of sodium, while influx of potassium across luminal membranes via H-K antiporters is accompanied by efflux of protons. Thus in describing the renal handling of potassium in various segments, we always have to keep in mind the fate of these other solutes. In the proximal tubule, the Na-K-ATPase in the basolateral membrane is very active in moving sodium from the cell to the interstitium, necessitating that potassium be simultaneously taken up from the interstitium. As we know that potassium is being put into the interstitium surrounding the proximal tubule, this pumped potassium must therefore recycle right back to the interstitium by passive flux through channels in the basolateral membrane.

**Figure 8–1.** A, Potassium transport in different regions of the tubule under conditions in which potassium excretion is low. In the proximal tubule (1) the majority of filtered potassium is reabsorbed, mainly by the paracellular route. In the thick ascending limb (2) most of the rest is reabsorbed, mostly by the transcellular route. In the cortical (3) and medullary collecting duct (4), there is some additional reabsorption via intercalated cells. Some of the potassium reabsorbed into the medullary interstitium recycles back into the thin limbs of the loop of Henle (5). B, Potassium reabsorption under conditions calling for high potassium excretion. The events in most regions of the tubule are the same as when there is little potassium excretion, but in the distal nephron, particularly in the connecting tubule, there is major secretion (6) that in some cases is greater than the sum of the reabsorptive processes.
In the thick ascending limb the interaction with sodium is even more complicated. As mentioned above, potassium is actively transported into the cells across both membranes and exits the cells passively across both membranes. It is pumped into the epithelial cells from the tubular lumen with sodium via Na-K-2Cl antiporters and from the interstitium via the Na-K-ATPase. As there is far less potassium than sodium in the lumen, potassium must recycle back to the lumen by passive channel flux to keep a supply of potassium available to run the multiporter with sodium. Otherwise, sodium reabsorption would be limited only to the amount of potassium present in the tubular fluid. Quantitatively, the sum of all these transcellular and paracellular processes is net reabsorption of about 20% of the filtered load.

**Secretion in the Distal Nephron and its Regulation**

As described in Chapter 6, there are two cell types in the epithelium of the distal nephron: those cell types being principal cells (about 70% of the cells) and intercalated cells. The intercalated cells are further subdivided into type A (most numerous) and type B (sparse) intercalated cells. The principal cells secrete potassium at highly variable rates, whereas the type A intercalated cells reabsorb potassium. The principles governing both secretion and reabsorption are straightforward. Secretion of potassium by principal cells involves the uptake of potassium from the interstitium via the Na-K-ATPase and secretion into the tubular lumen through channels (Figure 8–2). Type A intercalated cells reabsorb potassium via the H-K-ATPase in the luminal membrane, which actively takes up potassium

![Figure 8–2.](image)

**Figure 8–2.** Generalized pathway for potassium secretion by principal cells. Potassium secretion is tied to sodium reabsorption via the Na-K-ATPase. The drug amiloride inhibits sodium entry, and therefore inhibits potassium secretion. Aldosterone stimulates both sodium and reabsorption and potassium secretion at multiple points.
from the lumen and then allows potassium to enter the interstitium across the basolateral membrane via potassium channels.

Regulation of potassium excretion involves multiple controls over the secretory processes in the distal nephron, something like a passenger van where everyone in it has an accelerator and a brake pedal. As is the case with regulation of sodium excretion, we cannot predict just how these controls operate in every situation. Fortunately, with potassium as well as sodium, the healthy kidneys do a remarkable job of “doing the right thing,” ie, increasing potassium excretion in response to high dietary loads and reducing excretion in the face of restricted diets. Even with experimental manipulations, it is hard to get healthy kidneys to do otherwise (but as described later, certain pathologies interfere with normal potassium regulation). Much of the regulation involves controlling the activity of potassium channels. The kidneys and other body organs express numerous potassium channel species; for simplicity, we do not usually differentiate between types. (Otherwise, we would have a whole book devoted entirely to potassium channels!) However in principal cells of the distal nephron two types of channels stand out as being those that secrete potassium in a regulated manner: ROMK (standing for renal outer medulla, because that is where they were first indentified) and BK (as each channel has a “big” capacity to secrete potassium). Owing to different combinations of channel subunits and alternative splicing of transcripts, several isoforms of each type of channel exist. Although ROMK and BK channels both conduct potassium, they play different roles and are regulated by quite different mechanisms. At very low dietary loads of potassium, there is virtually no secretion by either kind of channel. ROMK channels are sequestered in intracellular vesicles and BK channels are closed. At normal potassium loads, ROMK channels are moved to the luminal membrane and secrete potassium. BK channels are still closed, held in reserve and ready to respond to appropriate signals when needed. At high excretion rates, both types of channel are present in the luminal membrane and avidly secreting potassium (Figure 8–3).

Figure 8–4 shows factors known to influence the secretion, and thus the ultimate excretion of potassium. The following text provides a brief description of how specific factors affect potassium excretion. (1) Plasma potassium. The role of plasma potassium is the most understandable influence. First, the filtered load is directly proportional to plasma concentration. Second, the environment of the principal cells, ie, the cortical interstitium, has a potassium concentration that is nearly the same as in plasma. The Na-K-ATPase that takes up potassium is highly sensitive to the potassium concentration in this space, and varies its pump rate up and down when potassium levels in the plasma vary up and down. Thus, plasma potassium concentration does exert an influence on potas-

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5BK channels require either an elevation in intracellular calcium or a large membrane depolarization in order to open them. ROMK channels are regulated by a multitude of intracellular signaling cascades. Some affect expression of the channels, some move channels back and forth between the luminal membrane and storage vesicles, while others alter the probability that the channel is in an open configuration. Although progress is being made in sorting out the details of these intracellular cascades, it is still a challenge to relate them to extrarenal signals that turn them on and off.
Aldosterone. We discussed the role of aldosterone in regulating sodium excretion in Chapter 7. Here, we describe its role in potassium excretion. One stimulator of aldosterone secretion is an elevated level of plasma potassium (see Figure 7–11). This is a direct action of potassium on the adrenal cortex and does not involve the renin-angiotensin system. Aldosterone, as well as increasing expression of the Na-K-ATPase, also stimulates the expression of ROMK channels in the distal nephron. Both actions have the effect of increasing potassium secretion. Greater pumping by the Na-K-ATPase supplies more potassium from the interstitium to the cytosol of the principal cells, and more ROMK channels provide more pathways for secretion. 

Figure 8–3. Activity of ROMK and BK potassium channels in principal cells under different conditions. When the body is conserving potassium and little is being excreted, ROMK channels are mostly sequestered in intracellular vesicles and BK channels are closed; thus, there is virtually no secretion. Under modest potassium loads (normal conditions), ROMK channels secrete potassium, while BK channels remain closed. With high potassium excretion, ROMK channel activity is maximized and BK channels are open, allowing substantial secretion.

Figure 8–4. Factors that increase the secretion of potassium by principal cells.
before the distal nephron determines how much is sent on from the thick ascending limb, i.e., delivered to the distal nephron. Changes in upstream handling of sodium include changes in filtered load and reabsorption in prior segments. Sodium delivery influences potassium secretion because more sodium delivered means more sodium taken up by principal cells; therefore, more sodium pumped out by the Na-K-ATPase, in turn causing more potassium to be pumped in. The increased potassium can just recycle back to the interstitium, but the usual result is more potassium secretion. (4) **Distal nephron flow rate.** The role of flow rate in regulating potassium secretion is a story by itself. Increased flow is detected by mechanosensitive elements of the principal cells. This includes bending of the central cilium that protrudes from the apical surface into the tubule lumen. Bending of the central cilium initiates intracellular release of calcium and activation of BK channels. Under most conditions, increased delivery of sodium is the chief cause of increased flow, because sodium is accompanied by water. Thus increased delivery of sodium implies increased flow. Increased flow has another effect. By sweeping away potassium that reaches the tubule by secretion, luminal potassium concentration is kept low enough to preserve a favorable concentration gradient for secretion. (5) **Concentration of nonchloride anions.** In order for principal cells to secrete potassium there must be a route (channels) and a driving force (electrochemical gradient). Under conditions where the reabsorption of sodium is restricted because some of the luminal chloride has been replaced with anions that are not usually in high concentration and cannot accompany the sodium (because their permeability is less than that of chloride), one effect is a depolarization of the luminal membrane (usually described as increasing luminal negativity). This increases the driving force for potassium secretion. We will discuss this further in the section on perturbations of potassium excretion. (6) **Dietary potassium.** The influence of dietary potassium on renal function is the most obvious regulator of potassium excretion, yet the least understood. A major task of the kidneys is to maintain potassium balance by increasing and decreasing potassium excretion in parallel with dietary load. The healthy kidneys do this very well. The problem is in understanding the signaling—how do the kidneys know how much potassium a person has consumed? Although large potassium loads can raise plasma potassium somewhat, the changes in excretion associated with diet do not seem to be accounted for on the basis of either changes in plasma potassium or the other identified factors. However, the previously mentioned gastrointestinal signals influence not only the cellular uptake of potassium absorbed from the GI tract, but also renal handling of potassium, and seem to be one of the links between dietary load and excretion. One of the manifestations of changing dietary loads is to regulate the distribution of ROMK channels between the luminal membrane and intracellular storage, i.e., high-potassium diets lead to insertion of luminal channels and therefore higher potassium secretion. In contrast, during periods of prolonged

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6This potential problem may be less than previously thought if most of the potassium secretion occurs early in the distal nephron (connecting tubule), rather than in the more distal regions (collecting ducts). This is because the major reabsorption of water that concentrates luminal potassium and other solutes in the collecting ducts has not yet occurred in the connecting tubule.
low potassium ingestion there are few ROMK channels in the apical membrane. Another adaptation to prolonged periods of low potassium ingestion is an increase in H-K-ATPase activity in intercalated cells, resulting in even more efficient reabsorption of filtered potassium.

**Perturbations in Renal Handling of Potassium**

We begin this section by mentioning a potential problem in the renal handling of potassium that in healthy kidneys is *not* a problem. The potential problem is the simultaneous balance of sodium and potassium. Normal human diets contain variable amounts of sodium and potassium, and the kidneys have to handle any combination of loads: both high, both low, one high, and the other low. Given that so much of the transport of these ions is by coupled mechanisms it is remarkable that the kidneys can deal with every combination. This is all the more remarkable when we consider that aldosterone is a regulator of both. If a person is consuming little salt of any kind (low loads of both sodium and potassium), then we expect, in order to preserve body stores of sodium, to see aldosterone levels high enough to stimulate avid reabsorption of sodium. But this should also lead to avid secretion of potassium, which is an unwanted action as the body is also trying to conserve potassium. The answer is simply that potassium cannot be secreted unless there are open channels. If the actions of intracellular signaling cascades has caused most of the ROMK channels to be sequestered in intracellular vesicles, then potassium that is taken up from the interstitium via the Na-K-ATPase recycles back to the interstitium and is not secreted.

**Effects of Diuretics**

Diuretics are agents that increase urine volume and reduce ECF volume. Most diuretics, although effective at their designated task of increasing water and sodium excretion, have the unwanted side effect of increasing the renal excretion of potassium. Potassium excretion is almost always increased in individuals undergoing osmotic diuresis (high filtration of solute that is not reabsorbed) or treatment with diuretics that block sodium reabsorption in the proximal tubule, loop of Henle, or distal convoluted tubule (ie, block of sodium reabsorption upstream from the principal cells). The potassium loss may cause severe potassium depletion.

The increased potassium excretion is due partly to the fact that, as noted earlier, potassium reabsorption in the proximal tubule and Henle’s loop is linked to sodium reabsorption. Accordingly, diuretics that act on these sites inhibit not only sodium reabsorption but also potassium reabsorption. However, *most* of the increased potassium excretion is due not to this decreased reabsorption but rather to increased potassium secretion by the distal nephron. In all these diuretic states, the delivery of sodium and the volume of fluid flowing to the distal nephron per unit time are increased by the upstream inhibition of sodium and water reabsorption. It is this increased flow and increased delivery of sodium that drives increased potassium secretion and, hence, excretion (Figure 8–5).

To reinforce this point further, let us integrate this information with our understanding of the action of aldosterone. Elevated aldosterone in individuals with
heart failure or other diseases of secondary hyperaldosteronism generally does not cause potassium hypersecretion because these patients simultaneously have low fluid delivery to the distal nephron. However, what happens when such persons are treated with diuretics to eliminate their retained sodium and water? The diuretics increase fluid delivery to the distal nephron, and now the patients have both increased aldosterone and increased flow. This combination tends to cause marked increases in potassium secretion and excretion. To prevent this combination, drugs that block the renal actions of aldosterone may be given; such drugs are weak diuretics because they block aldosterone’s stimulation of sodium reabsorption (with its small amount of associated water reabsorption). However, unlike other diuretics, they are “potassium sparing” because they simultaneously block aldosterone’s stimulation of potassium channels that promote potassium secretion. Another class of “potassium-sparing” diuretics blocks sodium channels in
the principal cells of the cortical collecting duct; this prevents sodium entry from lumen to cell and effectively prevents the basolateral membrane Na-K-ATPase pumps from transporting either sodium or potassium and blocks the apical exchange of sodium for potassium ions. Blocking sodium absorption upstream from the distal nephron increases potassium secretion; however, blocking sodium reabsorption in the distal nephron does not.

**Effects of Acid-Base Changes on Potassium Excretion**

Primary acid-base disturbances are a major cause of secondary potassium imbalances (and, as discussed in Chapter 9, imbalances in body potassium can perturb acid-base status). These topics are fraught with difficulty because the effects are not consistently seen. Nevertheless, the existence of an elevated plasma pH (alkalosis) is often (i.e., frequently but not always) associated with hypokalemia (low plasma potassium concentration). Similarly, low plasma pH (acidosis) is usually associated with hyperkalemia. Whether these relations between acid-base and potassium are actually seen in a particular patient depends on many factors, including the cause of the acid-base disturbance.

There are 2 known reasons for the effects of acid-base status on potassium. First, elevations and depressions in the extracellular concentration of hydrogen ions lead to a de facto exchange of these ions with cellular cations, the most important of which is potassium. During an alkalosis, e.g., the low extracellular hydrogen ion concentration induces the efflux of hydrogen ions that are normally bound to intracellular buffers. The loss of the positively charged hydrogen ions is balanced by the uptake of other cations, in this case potassium. Thus, an alkalosis (with hydrogen ions leaving tissue cells to replenish the loss from the ECF) induces cells to take up potassium, causing a hypokalemia. Conversely, a low pH (with a concomitant cellular uptake of hydrogen ions) often leads cells to dump potassium, causing a hyperkalemia.

In addition to these exchanges of potassium for hydrogen ions, there is an effect of intracellular pH on cellular Na-K-ATPase and potassium channel activity. Low intracellular pH inhibits pumps everywhere, allowing potassium to escape from cells (particularly muscle cells) to increase plasma potassium. Ordinarily, the increase in plasma potassium would stimulate potassium uptake by the Na-K-ATPase in principal cells, but low intracellular pH also inhibits the pumps here as well as luminal membrane potassium channels. Therefore, the principal cells respond inappropriately and do not effectively secrete the excess plasma potassium (paradoxical potassium retention). A high intracellular pH reverses these effects and relieves this inhibition (effectively stimulating the pump and the potassium channels). Alkalosis promotes potassium loss and contributes to the production of a hypokalemia. Thus, a patient suffering from alkalosis (induced, e.g., by vomiting) will manifest increased urinary excretion of potassium solely as a result of the alkalosis and will, therefore, become potassium deficient.

Finally, it should be emphasized that, although alkalosis is often associated with hypokalemia and acidosis with hyperkalemia, this is not always the case.
8–1. Control of potassium excretion is achieved mainly by regulating the rate of which of the following?

A. Potassium filtration
B. Potassium reabsorption
C. Potassium secretion

8–2. When on a high-potassium or high-sodium diet, is it possible to excrete more potassium or more sodium in the urine than is filtered?

8–3. Indicate whether each statement is true or false.

A. In the proximal tubule, the major pathway of reabsorption for both sodium and potassium is paracellular.

B. In the thick ascending limb, the major pathway of reabsorption for both sodium and potassium is via the Na-K-2Cl multiporter.

C. In the thick ascending limb, equal amounts of sodium and potassium are absorbed.

8–4. The presence of high amounts of nonreabsorbed solute (eg, glucose) in the proximal tubule inhibits proximal tubule potassium reabsorption. True or false?

8–5. The presence of high amounts of nonreabsorbed solute (eg, glucose) in the collecting tubule inhibits potassium secretion. True or false?
8–6. A patient has a tumor in the adrenal gland that continuously secretes large quantities of aldosterone (primary hyperaldosteronism). Is the rate of potassium excretion normal, high, or low?

8–7. A patient with severe congestive heart failure is secreting large quantities of aldosterone. Is the rate of potassium excretion normal, high, or low?

8–8. A person on a high-potassium diet is excreting large amounts of potassium. This is accomplished mainly by what mechanism?
   A. Reduced reabsorption in the proximal tubule
   B. Reduced reabsorption in the thick ascending limb
   C. Reduced reabsorption in the connecting tubule and collecting ducts
   D. Increased secretion in the connecting tubule and collecting ducts

8–9. In the face of a potassium-rich meal, the key action of insulin to prevent a large rise in plasma potassium concentration is what?
   A. Decreased absorption of potassium from the GI tract
   B. Increased uptake of potassium by tissue cells
   C. Increased renal excretion of potassium
OBJECTIVES

The student understands the interrelation among (1) input and output of acids and bases, (2) regulation of plasma buffers, and (3) value of plasma pH:

- States the major sources for the input of fixed acids and bases into the body, including metabolic processes and activities of the gastrointestinal tract.
- Describes why carbon dioxide levels affect the concentration of hydrogen ions, and differentiates carbon dioxide input (volatile acid) from the input of fixed acids.
- States the Henderson-Hasselbalch equation for the carbon dioxide—bicarbonate buffer system.
- Describes how the input of fixed acids and bases affects the body levels of bicarbonate.
- Explains why body levels of carbon dioxide are usually not altered by the input of fixed acids and bases.
- Explains why some low pH fluids alkalinize the blood after they are metabolized.

The student understands the general renal handling of acids and bases:

- Describes the reabsorption of filtered bicarbonate by the proximal tubule, including the role of carbonic anhydrase and apical membrane Na-H antiporters.
- Describes how bicarbonate is excreted in response to an alkaline load:
- Describes how the kidneys respond to an acid load:
  - Defines the concept of urinary titratable acidity.
  - Defines how the titration of filtered buffers is one means of excreting acid.
  - Describes how the conversion of glutamine to ammonium and subsequent excretion of ammonium accomplishes the goal of excreting acid.
  - States how total acid excretion is related to titratable acidity and ammonium excretion.

The student understands the nature of acid-base disturbances and the meaning of compensation:

- Defines the 4 categories of primary acid-base disturbance.
- Defines the meaning of compensation.
- Describes the renal response to respiratory acid-base disorders.
- Describes the respiratory response to changes in arterial pH.
- Identifies nonrenal problems that may cause the kidneys to generate a metabolic alkalosis.
The topic of acid-base physiology has vexed students for generations. Understanding acid-base physiology is greatly aided by keeping in mind several fundamental principles outlined below and always viewing any complexities of acid-base regulation through the lens of these fundamentals.

It is essential for the body to regulate the concentration of free protons in the extracellular fluid (ECF) to a value close to 40 nM (pH 7.4) in order for proteins exposed to the ECF to function properly. This regulation is called acid-base balance. The essence of acid-base balance comes down to 2 related processes: (1) Matching the excretion of acid/base equivalents to their input, and (2) regulating the ratio of weak acids to their conjugate bases in buffer systems. Matching excretion to input keeps the body content of these substances constant, ie, keeps the body in balance. Regulating the ratio of weak acids to their conjugate bases clamps the pH to a constant value that is buffered; ie, is protected against rapid changes in pH. Excretion of acid-base equivalents is the job of the kidneys. Balancing total-body input and output of acid or base and regulating physiological buffer concentrations are intimately related, but it is easy to lose sight of one perspective while seeing things from the other.

**Guideline 1: Acids and Bases Obey the Balance Principle**

Acids and bases are subject to the same constraints of input and output balance as other substances (eg, sodium, urea, and water). Each day, physiological processes add acids and bases to our body fluids, tending to raise or lower the concentration of hydrogen ions (change the pH). On the other hand, normal kidneys excrete acids and bases to exactly match the daily input, thereby keeping the body in acid-base balance. Despite the intricacies of acid-base balance, the basic principle of balance always holds. One reason to emphasize the balance concept is that, unlike substances such as sodium, there are multiple routes for the entry of acids or bases, including (1) processing of ingested food, (2) secretions of the gastrointestinal (GI) tract, and (3) de novo generation of acids and bases from metabolism of stored fat and glycogen. In addition, of course, the kidneys are factors in the overall maintenance of hydrogen ion balance.

Another reason to emphasize the balance concept is to avert a common misconception about acid-base disturbances, situations in which either there is an unusually high input or output of acid or the plasma pH is abnormal (eg, diabetic ketoacidosis). Although the body is sometimes transiently out of balance for acids and bases (just as it is sometimes transiently out of balance for many other substances), acid-base disturbances do not mean there is a persistent imbalance. In a prolonged metabolic acidosis, eg, there may be a high input of acid and an equally high output. There is never a situation in which acid or base pours into the body for an extended period of time without being balanced by an equivalent output. However, being in acid-base balance (ie, the same input of acid as output) does not necessarily mean that there are no changes in body chemistry. As in the case
for sodium balance, persistent excess sodium reabsorption does not continuously and indefinitely increase total-body sodium. Rather, other factors are stimulated to bring sodium back into balance, but the new balance state only comes at the price of elevated blood pressure. Input and output of hydrogen ion may be equal (in balance) during metabolic disorders that produce excess acid, but the balance comes about only after there has been a significant change in blood pH or bicarbonate concentration.

**Guideline 2: Body Fluids Are Buffered**

Acids and bases that enter the body must be excreted at the same rate to maintain balance. There is often a lag between input and output, allowing a transient accumulation of acid or base. Buffer systems prevent large changes in pH when these transient accumulations occur. A buffer system consists of three substances that have defined relations to each other according to the equilibrium constant: a weak acid, its conjugate base, and free protons. In such a system, the free aqueous concentration of protons is only a trivial fraction of the concentration of the acid and is determined by the ratio of acid to its conjugate base. Simple mass action chemistry (Equation 9–1) describes how a weak acid dissociates into its conjugate base and a free hydrogen ion. At equilibrium, the concentration of free hydrogen ions is determined by the ratio of the concentrations of conjugate base to the weak acid (Equation 9–2) or in the more familiar pH form (the Henderson-Hasselbalch equation) in Equation 9–3.

\[
\text{Acid} \leftrightarrow \text{conjugate base} + \text{H}^+ \quad (9-1)
\]

\[
[H] = K \frac{[\text{acid}]}{[\text{base}]} \quad (9-2)
\]

\[
\text{pH} = pK + \log \frac{[\text{base}]}{[\text{acid}]} \quad (9-3)
\]

Consider what would happen if we did not have buffers. If we added strong acid (eg, hydrochloric acid) to water that contained no buffers, the concentration of free protons would equal the concentration of the acid. If we had 10 mmol/L acid, we would have 10 mmol/L protons (ie, pH 2 blood). However, if the same amount of acid is added to a buffer system, most of the protons combine with the conjugate base, resulting in only a small rise in concentration of free protons. Buffer systems by themselves can only exert a kind of delaying action and reduce the magnitude of pH changes upon addition of acid/base equivalents. They do not eliminate added acid or base equivalents, but only ameliorate the effect of the equivalents on blood pH. In the face of persistent imbalance between input and output, the acid form of the buffer or its conjugate base is gradually consumed. Eventually acid or base equivalents added to the body, even if associated with blood buffers, have to be excreted by the kidneys to maintain balance.

What are the buffer systems in the body, and where are they? Buffers exist in the extracellular fluid, the intracellular fluid (the cytosol of the various cells in the body), and in the matrix of bone. Although these buffers are in different
compartments, the buffers in all of these compartments communicate with each other. Phosphate and albumin in the plasma are important ones in the ECF. Hemoglobin in red blood cells is also important, because changes in plasma pH lead to uptake or release of protons from red blood cells. For several reasons, the most important buffer system in the body turns out to be the CO$_2$–bicarbonate buffer system. Fortunately, we can understand acid-base balance by looking at this single buffer system alone and ignore the others, because all buffer systems must have ratios of weak acid to conjugate base that result in the same pH. In other words, if we control one buffer system, the others all fall into line.

One thing that sets the CO$_2$–bicarbonate buffer system apart from other buffer systems is that the concentrations of CO$_2$ and bicarbonate are both physiologically regulated, and they are regulated independently. And because their concentrations are regulated, the ratio of their concentrations is regulated. As it is the ratio of weak acid to conjugate base that sets pH, this system therefore regulates pH, and that is one of the goals of the regulation.

In the CO$_2$–bicarbonate buffer system, CO$_2$ is not a weak acid per se, but it acts like a weak acid because when it combines with water it releases protons. (CO$_2$ is often called a volatile acid because it can evaporate. All other acids, eg, sulfuric, lactic, are called fixed acids.) The combination of CO$_2$ with water forms carbonic acid, which dissociates like any other weak acid into a proton and its conjugate base, which is bicarbonate. Considered this way, and given the ubiquitous presence of water in our body, it is clear that carbon dioxide is effectively an acid. The concentration of carbonic acid in our blood is miniscule (about 3 µmol/L), and at first glance it appears that this system has little effective buffering capacity. However, the supply of CO$_2$ is effectively infinite, so that any carbonic acid consumed in a reaction is replaced by new generation from existing CO$_2$.

$$\text{CO}_2 + \text{H}_2\text{O} \rightleftharpoons \text{H}_2\text{CO}_3 \rightleftharpoons \text{HCO}_3^- + \text{H}^+ \quad (9-4a)$$

$$\text{CO}_2 + \text{H}_2\text{O} \rightleftharpoons \text{HCO}_3^- + \text{H}^+ \quad (9-4b)$$

(carbonic anhydrase)

The reaction on the left-hand side of Equation 9–4a to form carbonic acid is rather slow, but most tissues express one or several isoforms of the enzyme, carbonic anhydrase, intracellularly, extracellularly, or both. This is an enzyme that greatly speeds the reaction between CO$_2$ and water to form bicarbonate and a hydrogen ion. In doing, so it actually skips the step of forming carbonic acid, as

---

1ECF buffers react essentially instantaneously to an acid or base load. The intracellular buffers, greater in buffering capacity than those in the ECF, consist of intracellular proteins and various phosphates that buffer acid and base loads on a time scale of hours. The enormous amount of protein in skeletal muscle is a key component, along with hemoglobin in red blood cells. Buffering by bone is also significant (and rather complicated) on an even slower time scale (see Chapter 10).

2The actual reaction involves combining CO$_2$ with a hydroxyl ion already bound to the enzyme, resulting in the immediate formation of bicarbonate (HCO$_3^-$. As the bicarbonate dissociates from the enzyme, a water molecule binds to it. The water is then split into a hydrogen ion that dissociates from the enzyme and a hydroxyl ion that stays behind on the enzyme. The end result is that a CO$_2$ molecule and a water molecule have been converted to a hydrogen ion and a bicarbonate, the same as if they had gone through the slower uncatalyzed reaction of first forming a carbonic acid molecule.
shown in Equation 9–4b. However, as with all enzyme catalyzed reactions, the enzyme increases the velocity of the reaction but not the equilibrium concentrations of reactants and products. Thus, with or without carbonic anhydrase, the equilibrium concentrations of all components are the same.

**Guideline 3: Input and Output of Acids Alter Bicarbonate But Not the Partial Pressure of CO₂**

Unlike the other buffer systems in the body, where addition or loss of hydrogen ions changes the concentration of the weak acid, in the CO₂–bicarbonate system, the concentration of the weak acid (CO₂) is essentially constant. This is because the partial pressure of arterial CO₂ (Paco₂) is regulated by our respiratory system to be about 40 mm Hg. This partial pressure corresponds to a CO₂ concentration in blood of 1.2 mmol/L. Any rise or fall in Paco₂ resulting from the addition or loss of hydrogen ions as depicted in Equation 9–4 is sensed by the respiratory centers in the brainstem that alter the rate of ventilation to restore the concentration. There are times when the Paco₂ differs from 40 mm Hg, but this reflects activity of the respiratory system, not a change in Paco₂ in response to addition or loss of hydrogen ions.

Although adding or removing hydrogen ions from a source other than CO₂ does not change Paco₂, such changes do change the concentration of bicarbonate. Adding hydrogen ions drives the reaction in Equation 9–4 to the left and reduces bicarbonate on a nearly mole-for-mole basis. Removing hydrogen ions drives the reaction to the right and raises bicarbonate in the same way. There are many ways of adding or removing hydrogen ions, but, regardless of the process, the result is to change the concentration of bicarbonate. Let us consider the implications of such changes in bicarbonate.

When any process puts hydrogen ions into the blood, most of the hydrogen ions, as we have emphasized, combine with bicarbonate (and other buffers to some extent). Now a protonated bicarbonate is simply a molecule of carbonic acid. When the concentration of carbonic acid rises, the carbonic acid dissociates into CO₂ and water. The CO₂ formed in this manner mixes with metabolic CO₂ and is exhaled, thus restoring the concentration of CO₂ and carbonic acid to their former values, but some bicarbonate has been lost. Therefore, when we add hydrogen ions by diet or some physiological process, we lose some bicarbonate but we do not change the Paco₂ or concentration of carbonic acid. Suppose we remove hydrogen ions (eg, by adding strong base). CO₂ and water combine to generate a hydrogen ion (replacing the one lost) and a bicarbonate. The CO₂ is supplied from the enormous store of metabolic CO₂. The Paco₂ remains constant (if it starts to change, then ventilation adjusts to restore it). We end up with a gain in bicarbonate and no change in Paco₂. Thus, addition or removal of hydrogen ion alters total-body bicarbonate. The problem of maintaining hydrogen ion balance becomes one of maintaining bicarbonate balance. For every hydrogen ion added to the body, one bicarbonate disappears; therefore, to maintain balance it is necessary to generate a new bicarbonate to replace the one that was lost. Generation of new bicarbonate is the responsibility of the kidneys.
We have established that \( \text{CO}_2 \) is effectively an acid. Let us be sure we understand why normal metabolic production of \( \text{CO}_2 \) does not keep acidifying the body. An enormous amount of \( \text{CO}_2 \) is generated from metabolism each day. It is produced in our body at a rate of about 9 mmol/min. However, it is eliminated at the same rate, so there is no net addition. As arterial blood flows into tissue capillaries, the majority of the \( \text{CO}_2 \) entering the blood immediately combines with water to form hydrogen ions and bicarbonate, catalyzed by carbonic acid in red blood cells. Most of the hydrogen ions then combine with non-bicarbonate buffers (eg, hemoglobin), so the change in pH is not great, although there is a small decrease. The concentration of bicarbonate rises by about 1 mmol/L (from 24 to 25 mmol/L). When this blood carrying the newly loaded \( \text{CO}_2 \) (now venous blood) reaches the capillaries of the lungs, the processes that occurred in the tissue capillaries are reversed. Bicarbonate and hydrogen ions combine to generate \( \text{CO}_2 \) and water, and the \( \text{CO}_2 \) diffuses into the air spaces of the lungs. The pH rises a little, and the concentration of bicarbonate falls by about 1 mmol/L (back to 24 mmol/L).

**Guideline 4: Excretion of \( \text{CO}_2 \) and Bicarbonate Are Independent of Each Other**

Another situation that confuses students, and one that we also want to clarify immediately, is that the input and output of \( \text{CO}_2 \) and bicarbonate are handled independently: One cannot be excreted as the other. If there is an excess generation of \( \text{CO}_2 \) (eg, if there is a rise in metabolism not matched by an increase in ventilation), the \( \text{CO}_2 \) cannot be converted to fixed acid and excreted by the kidneys. Increased \( \text{CO}_2 \) input must be balanced by increased \( \text{CO}_2 \) exhalation from the lungs. Similarly, if there is excess input of fixed acid, the body cannot convert this acid to \( \text{CO}_2 \) and excrete it through the lungs. The reason is that every proton derived from a fixed acid that combines with bicarbonate to form \( \text{CO}_2 \) removes that bicarbonate and lowers its concentration. Although the \( \text{CO}_2 \) is simply exhaled, the deficit in bicarbonate remains. A continuous input of fixed acid would soon reduce the bicarbonate concentration to zero. Thus, an input of fixed acid must be balanced by renal output.

**Sources of Acids and Bases**

**Metabolism of Dietary Protein**

Although the oxidative metabolism of most foodstuff is acid-base neutral, protein contains some amino acids that contribute acid or base. When sulfur-(or phosphorus-) containing amino acids and those with cationic side chains are metabolized to \( \text{CO}_2 \), water, and urea, the end result is addition of fixed acid. Similarly, the oxidative metabolism of amino acids with anionic side chains adds base (consumes hydrogen ions). Depending on whether a person’s diet is high in either meat or fruit and vegetables, the net input can be acid or base. For typical American diets the input is usually acidic.

**Metabolism of Dietary Weak Acids**

Fruits and vegetables, particularly citrus fruit, contain a lot of weak acids and the salts of those acids (ie, the conjugate base plus a cation, usually potassium). We
all know that citrus fruit is acidic, with some fruit juices having a pH below 4.0. Interestingly, metabolism of these acidic substances alkalinizes the blood, sometimes called the fruit juice paradox. The complete oxidation of the protonated form of an organic acid (eg, citric acid) to CO2 and water is acid-base neutral, no different in principle from the oxidation of glucose. However, the complete oxidation of the base form adds bicarbonate to the body. One can think of this as taking a hydrogen ion from the body fluids to protonate the base, converting it to the acid, and then oxidizing the acid. Acidic fruits and vegetables contain a mixture of organic acids in the protonated form and base form. Before oxidation, the mixture is acidic, but on complete oxidation to CO2 and water, the result is addition of base.

**GI Secretions**

The GI tract, from the salivary glands to the colon, is lined with an epithelium that can secrete hydrogen ions, bicarbonate, or a combination. In addition, the major exocrine secretions of the pancreas and liver that flow into the duodenum contain large amounts of bicarbonate. To accomplish these tasks the GI tract (and the kidneys as we discuss later) use the CO2–bicarbonate system in an ingenious way. When we generate bicarbonate and protons from CO2 and water in a given medium, say in the blood or in a cell, the result is always acidification, because the concentration of protons rises. However, cells of the GI tract separate the protons from the bicarbonate. They transport protons out of the cell into one medium (eg, the lumen of the GI tract), and bicarbonate into another (the interstitium bathing the basolateral surface). Therefore, the lumen becomes acidified and the surroundings (and therefore the blood leaving the tissue) becomes alkalinized (see Figure 9–1). In other regions of the GI tract the cells reverse the direction of these processes, ie, they transport bicarbonate into the lumen (alkalinizing it) and protons into the surroundings. Thus, different regions of the GI acidify and alkalinize the blood. Normally, the sum of GI tract secretions is nearly acid-base neutral (ie, the secretion of acid in one site, eg, the stomach) is balanced by the secretion of bicarbonate elsewhere (eg, the pancreas). Typically, there is a small net secretion of bicarbonate into the lumen of the GI tract, resulting in the addition of protons to the blood. However, in conditions of vomiting or diarrhea, one kind of secretion may vastly exceed the other, resulting in a major loss of acid or base to the outside world complete with a major retention of base or acid in the blood.

**Anaerobic Metabolism of Carbohydrate and Fat**

The normal oxidative metabolism of carbohydrate and fat is acid-base neutral. Both carbohydrate (glucose) and triglycerides are oxidized to CO2 and water. Although there are intermediates in the metabolism (eg, pyruvate) that are acids or bases, the sum of all the reactions is neutral. However, some conditions lead to the production of fixed acids. The anaerobic metabolism of carbohydrate produces a fixed acid (lactic acid). In conditions of poor tissue perfusion, this can be a major acidifying factor, and the metabolism of triglyceride to β-hydroxybutyrate and acetoacetate also adds fixed acid (ketone bodies). These processes normally do not add much of an acid load but can add a huge acid load in unusual metabolic conditions (eg, diabetes).
There are other ways of adding acids or bases (e.g., by ingestion of certain drugs or other foreign materials and by intravenous infusions). Typically, there is a small net load of acid or base resulting from normal metabolism of foodstuff and from GI processes. This load can be greatly increased in unusual circumstances. If the kidneys are working properly, they excrete the load, small or large, and keep the body in balance.

**RENAL HANDLING OF ACIDS AND BASES**

A simplified overview of the renal processing of acids and bases is as follows: In the early part of the nephron (mostly proximal tubule), the kidneys reabsorb the enormous filtered load of bicarbonate (thereby resulting in no addition or loss) from the plasma and, under appropriate conditions, can secrete organic bases or weak organic acids and acid equivalents. Then, in the distal nephron (mostly the collecting tubules), the kidneys secrete either protons or bicarbonate to balance the net input into the body (summarized in Table 9–1).

The first task is to reabsorb filtered bicarbonate. Bicarbonate is freely filtered at the renal corpuscles. How much is normally filtered per day? Given a typical plasma concentration of 24 mmol/L and a glomerular filtration rate (GFR) of 180 L/day, this amounts to 4320 mmol/day. Excretion of this bicarbonate would be equivalent to adding more than 4 L of 1 N acid to the

![Diagram](image-url)
body! It is essential, therefore, that virtually all the filtered bicarbonate be reabsorbed or the body fluids would become profoundly acidic. Thus, the reabsorption of bicarbonate is an essential conservation process.

Bicarbonate reabsorption is an active process, but it is not accomplished in the conventional manner of importing bicarbonate across the luminal membrane and exporting it across the basolateral membrane. Rather, the mechanism by which bicarbonate is reabsorbed involves the tubular secretion of hydrogen ions.

An enormous amount of hydrogen ion secretion occurs in the proximal tubule, with additional secretion in the thick ascending limb of Henle’s loop and collecting-duct system. In contrast to the situation for handling sodium, water, and potassium, the collecting-duct cells that secrete hydrogen ion are the Type A intercalated cells, not the principal cells.

The basic pattern followed in all these tubular segments is the same (although the precise transporters involved differ to some extent) and is illustrated in Figure 9–1 without indicating any specific transporters. Within the cells, a hydrogen ion and a bicarbonate are generated from CO₂ and water, catalyzed by carbonic anhydrase. The hydrogen ion is actively secreted into the tubular lumen. For every hydrogen ion secreted, one bicarbonate ion remains within the cell. The cellular bicarbonate is transported across the basolateral membrane into the interstitial fluid and then into the peritubular capillary blood. The net result is that, for every hydrogen ion secreted into the lumen, a bicarbonate ion enters the blood in the peritubular capillaries. There must be a 1-for-1 match between hydrogen ions secreted and bicarbonate ions transported into the interstitium.

Figure 9–2 illustrates how the process of hydrogen ion secretion achieves bicarbonate reabsorption (especially in the proximal tubule). Once in the tubular lumen, the secreted hydrogen ion combines with a filtered bicarbonate to form water and carbon dioxide, which diffuse into the cell. As mentioned above, the CO₂ and water that is now within the cell combines to form bicarbonate and hydrogen ion.

### Table 9–1. Normal contributions of tubular segments to renal hydrogen ion balance

<table>
<thead>
<tr>
<th>Tubular Segment</th>
<th>Contributions</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Proximal tubule</strong></td>
<td>Reabsorbs most filtered bicarbonate (normally about 80%)*</td>
</tr>
<tr>
<td></td>
<td>Produces and secretes ammonium</td>
</tr>
<tr>
<td><strong>Thick ascending limb of Henle’s loop</strong></td>
<td>Reabsorbs second largest fraction of filtered bicarbonate (normally about 10–15%)*</td>
</tr>
<tr>
<td><strong>Distal convoluted tubule and collecting-duct system</strong></td>
<td>Reabsorbs virtually all remaining filtered bicarbonate as well as any secreted bicarbonate (Type A intercalated cells)*</td>
</tr>
<tr>
<td></td>
<td>Produces titratable acid (Type A intercalated cells)*</td>
</tr>
<tr>
<td></td>
<td>Secretes bicarbonate (Type B intercalated cells)</td>
</tr>
</tbody>
</table>

* Processes achieved by hydrogen ion secretion.
The hydrogen ion is secreted across the apical membrane to combine with another luminal bicarbonate and the cellular bicarbonate leaves the cell across the basolateral membrane to enter the plasma. The overall result is that the bicarbonate filtered from the blood at the renal corpuscle has disappeared, replaced by the bicarbonate that reenters the plasma from inside the cell. Thus, no net change in plasma bicarbonate concentration has occurred as all bicarbonate filtered has combined with secreted hydrogen ion and subsequently ended up first inside the cell and then in plasma. It may seem inaccurate to refer to this process as bicarbonate reabsorption because the bicarbonate that appears in the peritubular capillary is not the same bicarbonate that was filtered. Yet the overall result is the same as it would be if the filtered bicarbonate had been more conventionally reabsorbed, like a sodium or potassium ion.

It is also important to note that the hydrogen ion that was secreted into the lumen is not excreted in the urine. It has been incorporated into water. Any secreted hydrogen ion that combines with bicarbonate in the lumen to cause bicarbonate reabsorption does not contribute to the urinary excretion of hydrogen ions but only to the conservation of bicarbonate.

Specific transporters are required for the transmembrane movements of both the hydrogen ion and the bicarbonate. Active transport of hydrogen ion across the luminal membrane from cell to lumen is achieved by several distinct luminal membrane transporters. First, particularly prominent in the proximal tubule is

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**Figure 9–2.** Predominant proximal tubule mechanism for reabsorption of bicarbonate. Hydrogen ions and bicarbonate are produced intracellularly. The hydrogen ions are secreted via an Na-H antiporter (member of the NHE family), while the bicarbonate is transported into the interstitium via an Na-3HCO₃ symporter (member of the NBC family). As more sodium enters via the Na-H antiporter than leaves via the Na-3HCO₃ symporter, additional sodium is removed via the Na-K-ATPase.
a specific isoform of the Na-H antiporter NHE3 as described in Chapter 6 and shown in Figure 9–2. This transporter is the major means not only of hydrogen ion secretion but also of sodium uptake from the proximal tubule lumen. Second, a primary active H-ATPase exists in all the hydrogen ion–secreting distal tubular segments. The Type A intercalated cells of the collecting-duct system, in addition to their primary active H-ATPase, possess a primary active H-K-ATPase, which simultaneously moves hydrogen ions into the lumen and potassium into the cell, both actively (Figure 9–3). Note that, as described in Chapter 8, the luminal membrane H-K-ATPase also mediates active potassium reabsorption by these cells and contributes to potassium homeostasis.

The basolateral membrane exit step for bicarbonate is via Cl-HCO3 antiporters or Na-HCO3 symporters (Figures 9–2 and 9–3), depending on the tubular segment. In both cases, the movement of bicarbonate is down its electrochemical gradient (ie, the transport is passive). Symport with sodium is the dominant means of extruding bicarbonate in the proximal tubule and is particularly interesting because efflux of sodium is up its electrochemical gradient. This is a rare case of a secondary active transporter that does not use the sodium gradient as the energy source, but actually moves sodium up its electrochemical gradient.

Through its secretion of hydrogen ions, the proximal tubule reabsorbs 80–90% of the filtered bicarbonate. The thick ascending limb of Henle’s loop reabsorbs another 10%, and almost all the remaining bicarbonate is normally reabsorbed by the distal convoluted tubule and collecting-duct system (except for alkalotic individuals, who will excrete some of the bicarbonate; see later discussion).

Throughout the tubule, intracellular carbonic anhydrase is involved in the reactions generating hydrogen ion and bicarbonate. In the proximal tubule, carbonic anhydrase is also located in the lumen-facing surface of apical cell membranes, and this carbonic anhydrase catalyzes the intraluminal generation of CO2 and water from the large quantities of secreted hydrogen ions combining with filtered bicarbonate.

**RENNAL EXCRETION OF ACID AND BASE**

If all the filtered bicarbonate is reabsorbed, there is no acid-base consequence for the body of filtering a large amount of bicarbonate; it is as though none had been filtered in the first place. When base has been added to the body fluids, the effect is to increase the plasma concentration of bicarbonate. Whether this increased bicarbonate comes from metabolism of protein that contains many anionic amino acids or from ingestion of baking soda (sodium bicarbonate), the result is the same: The body fluids contain more bicarbonate. The renal handling of such base loads is relatively straightforward: We excrete enough bicarbonate in the urine to match the input. If the addition of base to the body is 30 mEq and the kidneys put out 30 mmol of bicarbonate, the kidneys have achieved their goal: balance. The kidneys do this in 2 ways: (1) allow some filtered bicarbonate to pass through to the urine and (2) secrete bicarbonate via Type B intercalated cells. The Type B intercalated cells, which are found only in the cortical collecting duct, do indeed secrete bicarbonate. In essence, the Type B intercalated cell is a “flipped-around” Type A
Figure 9–3. Type A and Type B intercalated cells. A, Predominant collecting tubule mechanisms in Type A intercalated cells for the secretion of hydrogen ions that result in the formation of titratable acidity. The apical membrane contains H-ATPases, which transport hydrogen ions alone or in exchange for potassium. B, The Type B intercalated cell secretes bicarbonate and simultaneously transports hydrogen ions into the interstitium. The difference between this cell type and the Type A cell and those in the proximal tubule is that the location of the transporters for hydrogen ions and bicarbonate are switched between apical and basolateral membranes. ATP, adenosine triphosphate.
intercalated cell (Figure 9–5). Within the cytosol, hydrogen ions and bicarbonate are generated via carbonic anhydrase. However, the H-ATPase pump is located in the basolateral membrane, and the Cl-HCO3 antiporter is in the luminal membrane. Accordingly, bicarbonate moves into the tubular lumen, whereas hydrogen ion is actively transported out of the cell across the basolateral membrane and enters the blood, where it can combine with a bicarbonate ion. Thus, the overall process achieves the disappearance of excess plasma bicarbonate and the appearance of bicarbonate in the urine, with resulting acidification of the plasma and alkalinization of the urine and maintenance of bicarbonate balance.

How do the kidneys excrete an acid load? For all individuals who ingest animal protein of any kind, excretion of excess acid is more typical than the production and removal of excess base. This is a more complex process than excretion of base, but it obeys the principles we have developed earlier. Recall that the net result of addition of acid to the body reduces the amount of bicarbonate on an almost mole-for-mole basis. Therefore, the task for the kidney is to replace the lost bicarbonate by generating new bicarbonate from CO2 and water (being careful to excrete the hydrogen ion that is created at the same time). In essence, the process is as follows: Hydrogen ions and bicarbonate are produced from carbon dioxide and water within cells. Hydrogen ions are secreted and combine with the conjugate base of buffers in the tubular lumen other than bicarbonate, thereby generating the acid form of the buffer. The acid form of that buffer is excreted in the urine. The process of producing and secreting hydrogen ions generated new bicarbonate that goes into the blood and replaces the bicarbonate lost when the acid load entered the body. The key is generation of new bicarbonate to replace the bicarbonate that was lost. If we just reabsorb filtered bicarbonate, nothing is changed. We must generate new bicarbonate.

HYDROGEN ION EXCRETION ON URINARY BUFFERS

We emphasized earlier how hydrogen ion secretion achieves bicarbonate reabsorption and how this process prevents loss of filtered bicarbonate. Now we see that the identical transport process of hydrogen ion secretion can also achieve acid excretion and addition of new bicarbonate to the blood. At first glance, this seems like a contradiction: How can the same process produce 2 different end results? The answers lies in the fate of the hydrogen ion once it is in the lumen. If the secreted hydrogen ion combines with bicarbonate, then we are simply replacing bicarbonate that would have left the body. In contrast, if the secreted hydrogen ion combines with a non-bicarbonate buffer in the lumen (or, to an extremely small degree, remains free in solution), the hydrogen ion is excreted. The bicarbonate produced in the cell and transported across the basolateral membrane is new bicarbonate, not a replacement for existing bicarbonate.

There are two sources of tubular non-bicarbonate buffers: filtration and synthesis. Normally, the most important of the filtered buffers is phosphate, while ammonia is the most important synthesized buffer. Ammoniagenesis is crucial to renal acid excretion because its rate can be greatly increased in the face of large acid loads, whereas the availability of filtered buffers, while somewhat variable, is
not regulated for purposes of acid excretion. Figure 9–4 illustrates the sequence of events that achieves hydrogen ion excretion on filtered phosphate and the addition of new bicarbonate to the blood. The process of hydrogen ion secretion in this sequence is exactly the same as described previously, but the secreted hydrogen ion combines with filtered phosphate rather than with filtered bicarbonate. Therefore, the bicarbonate generated within the tubular cell (that always occurs when hydrogen ions are secreted) enters the plasma and constitutes a net gain of bicarbonate by the blood, not merely a replacement for filtered bicarbonate. Thus, when a secreted hydrogen ion combines in the lumen with a filtered buffer other than bicarbonate, the overall effect is not merely bicarbonate conservation but rather addition to the blood of new bicarbonate, which raises the bicarbonate concentration of the blood and the pH to a value similar to what it was before the addition of fixed acids.

Figure 9–4 also illustrates a point we want to emphasize: namely, that the renal contribution of new bicarbonate to the blood is accompanied by the excretion of an equivalent amount of buffered hydrogen ion in the urine. In this case, in contrast to the reabsorption of bicarbonate, the secreted hydrogen ion remains in the tubular fluid, trapped there by the buffer, and is excreted in the urine. This should reinforce the concept that bicarbonate can always be generated from CO$_2$ and water, but to add this new bicarbonate to the blood (and alkalinize the blood), the kidneys must separate the hydrogen ion from the bicarbonate and excrete the hydrogen ion that is created at the same time.

Figure 9–4. Excretion of hydrogen ions on filtered phosphate. Divalent phosphate (base form) that has been filtered and not reabsorbed reaches the collecting tubule, where it combines with secreted hydrogen ions to form monovalent phosphate (acid form) and is then excreted in the urine. The bicarbonate entering the blood is new bicarbonate, not merely a replacement for filtered bicarbonate. ATP, adenosine triphosphate.
It must be emphasized also that neither filtration per se nor excretion of free hydrogen ions make a significant contribution to hydrogen ion excretion. First, the filtered load of free hydrogen ions, when the plasma pH is 7.4 (40 nM/H⁺), is less than 0.1 mmol/day. Second, there is a minimum urinary pH—approximately 4.4—that can be achieved. This corresponds to a free hydrogen ion concentration of 0.04 mmol/L. With a typical daily urine output of 1.5 L, the excretion of free hydrogen ions is only 0.06 mmol/day, a tiny fraction of the normal 50–100 mmol of hydrogen ion ingested or produced every day. To excrete these additional amounts of protons, they must associate with tubular buffers.

**PHOSPHATE AND ORGANIC ACIDS AS BUFFERS**

Filtered phosphate is normally the most important non-bicarbonate urinary buffer. Most free plasma phosphate exists in a mixture of monovalent and divalent forms. In Equation 9–5, monovalent dihydrogen phosphate (on the left) is a weak acid, and divalent monohydrogen phosphate (on the right) is its conjugate base.

\[
H_2PO_4^- + H^+ \rightarrow HPO_4^{2-} \quad (9–5)
\]

We can write this in the form of the Henderson-Hasselbalch equation:

\[
\text{pH} = 6.8 + \log \frac{[\text{HPO}_4^{2-}]}{[H_2\text{PO}_4^-]} \quad (9–6)
\]

At the normal pH of plasma (7.4) and, therefore, of the glomerular filtrate, we find that about 80% of the phosphate is in the base (divalent) form and 20% is in the acid (monovalent) form. As the tubular fluid is acidified in the collecting ducts, most of the base form combines with secreted hydrogen ions. By the time the minimum intratubular pH of 4.4 is reached, virtually all the base (HPO_4^{2-}) has been converted to acid (H_2PO_4^-). Therefore, secreted hydrogen ions that combined with the base form are excreted, and the bicarbonate that was generated intracellularly in the process enters the blood. How much phosphate is available for this process? The amount is somewhat variable, depending on a number of factors (see Chapter 10), but a typical plasma concentration is about 1 mmol/L, of which about 90% is free (the rest being loosely bound to plasma proteins). At a GFR of 180 L/day, the total filtered load of phosphate is about 160 mmol/day. The fraction reabsorbed is also variable: from 75% to 90%. Thus, unreabsorbed divalent phosphate available for buffering is roughly 40 mmol/day. In other words, the kidneys can excrete hydrogen ions, using the phosphate buffer system, at a rate of about 40 mmol/day. However, the availability of phosphate cannot be easily upregulated to increase acid excretion.

There are other organic buffers in the urine, and under certain conditions these may appear in the tubular fluid in sufficient quantities to allow them also to act as important buffers. A particularly important example is a patient with uncontrolled diabetes mellitus. Because of metabolic processes that result from insulin deficiency, such a patient may become extremely acidic due to the excess production of acetoacetic acid and β-hydroxybutyric acid. At normal plasma pH,
these species completely dissociate to yield the anions β-hydroxybutyrate and acetoacetate (and hydrogen ions). These anions are filtered at the renal corpuscle but are only partly reabsorbed because they are present in great enough quantities to exceed the renal reabsorptive Tm.s. Accordingly, they are available in the tubular fluid to buffer a portion of the hydrogen ions being secreted by the tubules. However, their usefulness in this role is limited by the fact that their pKs are low: approximately 4.5, meaning that only at very low pH will the secreted hydrogen ions combine with them. At the limiting urine pH of 4.4 only half of them have actually combined with a hydrogen ion. The rest remain in the base form.

**HYDROGEN ION EXCRETION ON AMMONIUM**

Ordinarily, hydrogen ion excretion associated with phosphate and other filtered buffers is no greater than about 40 mmol/day. This amount is not sufficient to balance the normal hydrogen ion production of 50–100 mmol/day or take care of any unusually high (usually pathological) production of acid loads. To excrete the rest of the hydrogen ion and achieve balance, there is a second means of excreting hydrogen ions that involves ammoniagenesis and excretion of hydrogen ions as ammonium. Quantitatively, far more hydrogen ions can be excreted by means of ammonium than via organic buffers. There are many nuances to hydrogen ion excretion via ammonium, but the basic concepts are straightforward.

As described in Chapter 5, the catabolism of protein and oxidation of the constituent amino acids by the liver generates CO₂, water, urea, and some glutamine. Protein catabolism, which occurs constantly, even in starvation, requires the continuous excretion of urea by the kidneys to prevent uremia. Although, as described earlier, metabolism of the side chains of amino acids can lead to the addition of acid or base, the processing of the core of an amino acid—the carboxyl group and amino group—is acid-base neutral. After many intermediate steps, processing of the carboxyl group of the amino acid produces a bicarbonate, and processing of the amino group produces an ammonium ion. Processing does not stop there, however, because ammonium in more than miniscule levels is quite toxic. Ammonium is further processed by the liver to either urea or glutamine. In both cases, each ammonium consumed also consumes a bicarbonate. Thus, the bicarbonate produced from the carboxyl group is just an intermediate, consumed as fast as it is made, and the process as a whole is acid-base neutral. We can write this process as follows:

\[
2 \text{ amino acids (+oxygen)} \rightarrow 2\text{NH}_4^+ + 2\text{HCO}_3^- \rightarrow \text{urea or glutamine (+CO}_2 \text{ and water)}
\]  

(9–7)

When the urea (or glutamine) is excreted, the body has completed the catabolism of protein in a manner that promotes total body protein nitrogen balance, but is acid-base neutral.
Renal handling of urea is somewhat complicated from the osmotic point of view, as described in earlier chapters, but the handling is acid-base neutral. Glutamine, however, is different. Although the production of glutamine by the liver is acid-base neutral, it is important to recognize that glutamine can be thought to contain 2 components: a base component (bicarbonate) and an acid component (ammonium). Ammonium is the protonated form of ammonia and is an acid because it contains a dissociable proton, as shown in Equation 9–8, though an extremely weak acid. The pK of ammonium is near 9.2. At physiological pH over 98% of the total exists as ammonium, and less than 2% exists as ammonia. For renal acid-base purposes, this is a good thing because virtually all excreted ammonia is in the protonated form and takes a hydrogen ion with it.

\[
\text{NH}_4^+ \leftrightarrow \text{H}^+ + \text{NH}_3 \quad (9–8)
\]

Glutamine released from the liver is taken up by proximal tubule cells, both from the lumen (filtered glutamine) and from the renal interstitium. The cells of the proximal tubule then convert the glutamine back to bicarbonate and NH$_4^+$. In essence, the proximal tubule reverses what the liver has done. The NH$_4^+$ is secreted into the lumen of the proximal tubule, and the bicarbonate exits into the interstitium and then into the blood (Figure 9–5A). This is new bicarbonate, just

*Figure 9–5. Ammoniagenesis and excretion. A, Ammonium production from glutamine. Glutamine is originally synthesized in the liver from NH$_4^+$ and bicarbonate. When it reaches the proximal tubule cells, it is converted back to NH$_4^+$ and bicarbonate. There are more biochemical steps in the conversion of glutamine to ammonium and bicarbonate than indicated here; only the end result is shown.*
Figure 9–5. (Cont.) B, Ammonium reabsorption in the thick ascending limb. Ammonium reaches the thick ascending limb from two sources. Most comes from secretion in the proximal tubule. Some also enters the thin limbs from the medullary interstitium in the form of neutral ammonia and is subsequently reprotonated in the lumen (ammonium recycling). Ammonium is reabsorbed in the thick ascending limb by several mechanisms, the predominant one being entrance via the NKCC multiporter (where ammonium substitutes for potassium). C, Ammonium secretion in the inner medulla. Several mechanisms are involved. A prominent one involves uptake and secretion of neutral ammonia via specific transporters in parallel with hydrogen ion secretion, resulting in reformation of ammonium in the lumen. In the innermost medulla, the high interstitial ammonium concentration allows ammonium to substitute for potassium on the Na-K-ATPase.
like the new bicarbonate generated by titrating non-bicarbonate buffers. Further processing of the \( \text{NH}_4^+ \) is complicated, but eventually the ammonium is excreted (Figure 9–5C).

Ammonium ion is interesting in that it can masquerade as other ions, in some cases as a hydrogen ion and in other cases as a potassium ion. This is because some transporters and some channels are not completely selective for the species they usually move compared to ammonium. As the concentration of ammonium rises, there is an increasing tendency for ammonium to substitute for these other ions and “piggy back” its way across membranes.

Also, whenever ammonium is present in body fluids, a small fraction always dissociates into a proton and ammonia because the dissociation, although limited, is nearly instantaneous. Ammonium, being a small hydrated ion, is essentially impermeant in lipid bilayers and must be handled by channels or transporters if it is to move across membranes, but the neutral ammonia has a finite permeability. In terms of cellular handling, cells sometimes transport ammonium as such and at other times transport ammonia and a proton in parallel; the end result being the same in both cases.

It would “make sense” if the ammonium secreted into the proximal tubule simply stayed in the lumen and was excreted, but the kidneys have evolved a more complicated way of doing things for several reasons. An array of channels and transporters participate in moving ammonium or ammonia into or out of the tubule in various segments. As long as all the ammonium that is produced from glutamine and secreted in the proximal tubule ends up being excreted, the process accomplishes the goal of excreting acid, even if ammonium is transported as such in some places and moved as \( \text{H}^+ \) and \( \text{NH}_3 \) separately in other places. But if ammonium is returned to the circulation, it is metabolized by the liver back to urea, consuming a bicarbonate in the process, thereby nullifying the renal generation of bicarbonate.

Most of the ammonium synthesized from glutamine in the proximal tubule is secreted via the NHE-3 antiporter in exchange for sodium (with ammonium substituting for a hydrogen ion); but some may also diffuse into the lumen as ammonia and then combine with a secreted hydrogen ion. The next major transport event occurs in the thick ascending limb. In this segment about 80% of the tubular ammonium is reabsorbed, mostly by the Na-K-2Cl multiporter (with ammonium now substituting for potassium). In the medullary portions of the thick ascending limb, this reabsorption results in the accumulation of ammonium (and therefore some ammonia) in the interstitium, with the concentration progressively increasing toward the papilla, analogous to the osmotic gradient. The high interstitial concentration surrounding the loops of Henle leads to some secretion of ammonia into the thin descending limbs, which become protonated in the lumen. Therefore there is a certain amount of recycling, with the consequence that a considerable amount of ammonium is trapped in the medullary interstitium (similar to the situation with urea). Finally, in the medullary collecting ducts there is secretion again, mainly by parallel transport of hydrogen ions and ammonia. Thus, the ammonium that was reabsorbed in the thick ascending limb
and accumulated in the medullary interstitium is now put back into the tubule and excreted.  

A comparison of Figures 9–4 and 9–5 demonstrates that the overall result of renal generation of a new bicarbonate is the same regardless of whether it is achieved by hydrogen ion secretion and excretion on filtered buffers (Figure 9–4) or by glutamine metabolism with ammonium excretion (Figure 9–5). It is convenient, therefore, to view ammonium excretion as representing H⁺ excretion in the form of an H⁺ bound to NH₃, just as the former case constitutes hydrogen ions bound to phosphate or other non-bicarbonate buffers. In this manner, we can, in both cases, quantitatively equate the terms H⁺ excretion and renal contribution of new bicarbonate. It is interesting to note that while protein metabolism is a major source of excess total body hydrogen ions, protein metabolism also produces excess nitrogen as ammonia. By excreting ammonium, the kidney is removing both the excess ammonia nitrogen and excess hydrogen ion.

**QUANTIFICATION OF RENAL ACID-BASE EXCRETION**

We can now quantify the kidneys’ contribution to hydrogen ion balance. In other words, we can calculate the kidneys’ net bicarbonate addition to the body or elimination from it. This value is, again, identical to the kidneys’ net excretion of hydrogen ions (“acid”). Such a calculation is made by answering 3 questions:

1. How much bicarbonate is excreted in the urine? This represents bicarbonate loss from the body. It is measured simply by multiplying the urine flow rate by the urinary bicarbonate concentration.

2. How much new bicarbonate is contributed to the plasma by secretion of hydrogen ions that combine in the tubular lumen with non-bicarbonate urinary buffers? This can be measured by titrating the urine with NaOH to a pH of 7.4, the pH of the plasma from which the glomerular filtrate originated. This simply reverses the events that occurred within the tubular lumen when the tubular fluid was titrated by secreted hydrogen ions. Thus, the number of milliequivalents of sodium hydroxide required to raise the pH back to 7.4 must equal the number of milliequivalents of hydrogen ion added to the tubular fluid that combined with phosphate and organic buffers. This value is known as the *titratable acid*.

3. How much new bicarbonate is returned to the plasma by secretion of hydrogen ions that are excreted as ammonium? The titratable acid measurement will not titrate hydrogen ions in NH₄⁺ because ammonium is such a weak acid with a pK of the ammonia-ammonium reaction so high (9.2) that titration with alkali to pH 7.4 will not remove hydrogen ions from NH₄⁺.

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3The secretory processes in the medullary collecting ducts are highly complex. Ammonia taken up from the medullary interstitium and secreted across the apical membrane involves both diffusive flux and transporter-mediated flux. In both cases, the secreted ammonia is matched by a secreted proton to reform ammonium in the tubular lumen. In the deepest portions of the medulla, there is probably a parallel route of ammonium transport in which ammonium is taken by the Na-K-ATPase, with ammonium substituting for potassium. By reabsorbing ammonium in the medullary thick ascending limb, luminal ammonium in the cortical collecting duct does not become concentrated to the high level it would otherwise as a result of the major reabsorption of water. In the inner medulla, however, the tubular concentration can be rather high, with secretion still favored because of the high interstitial ammonium concentration.
Therefore, to know how much new bicarbonate is contributed by glutamine metabolism with ammonium excretion, urinary ammonium excretion (urine flow rate times urinary ammonium concentration) must be measured separately, remembering that for every ammonium excreted a new bicarbonate was added to the blood.

Thus, the data required for a quantitative assessment of the renal contribution to acid-base regulation in any person are as follows:

1. Titratable acid excreted
2. Plus NH₄⁺ excreted (ie, HCO₃⁻ lost from the body because of incomplete reabsorption or HCO₃⁻ secretion)
3. Minus HCO₃⁻ excreted (mmol/day) (lost from body)

The total equals net excretion of hydrogen ion or the net HCO₃⁻ gain or loss to the body (negative values equal loss, positive values equal gain).

Note that there is no term for free hydrogen ion in the urine because, even at a minimum urine pH of 4.4, the number of free hydrogen ions is trivial.

Typical urine data for the amounts of bicarbonate contributed to the blood by the kidneys in three potential acid-base states are given in Table 9–2. Note that in response to acidosis, as emphasized previously, increased production and excretion of NH₄⁺ is quantitatively much more important than increased formation of titratable acid.

It should also be emphasized that the data shown for alkalosis are typical of “pure” alkalosis (ie, alkalosis uncomplicated by other electrolyte abnormalities). As we will see, electrolyte imbalances frequently complicate alkalosis so that the expected values are different from those actually measured.

Table 9–2. Renal contribution of new bicarbonate to the blood in different states

<table>
<thead>
<tr>
<th></th>
<th>Alkalosis</th>
<th>Normal state</th>
<th>Acidosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Titratable acid (mmol/day)</td>
<td>0</td>
<td>20</td>
<td>40</td>
</tr>
<tr>
<td>plus NH₄⁺ excreted (mmol/day)</td>
<td>0</td>
<td>40</td>
<td>160</td>
</tr>
<tr>
<td>minus HCO₃⁻ excreted (mmol/day)</td>
<td>80</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Total (mmol/day)</td>
<td>–80</td>
<td>59</td>
<td>200</td>
</tr>
<tr>
<td></td>
<td>(lost from body)</td>
<td>(added to body)</td>
<td>(added to body)</td>
</tr>
<tr>
<td>Urine pH</td>
<td>8.0</td>
<td>6.0</td>
<td>4.6</td>
</tr>
</tbody>
</table>

The suffixes “-osis” and “-emia” (meaning “in the blood”), as in acidosis or acidemia, are technically different but are often used interchangeably or inappropriately. Acidemia means high hydrogen ion concentration in the blood, essentially a pH below the normal range. Acidosis means a process adding acid to the blood. We can have an acidosis but not an acidemia if the pH is within the normal range and the kidneys are excreting the high acid load as fast as it is added. The analogous argument applies to alkalemia and alkalosis. As will be seen, the proper distinction between “-osis” and “-emia” is usually violated in the common description of acid-base disorders.
It is clear that renal acid-base processing is regulated in response to different body conditions. Most importantly, the regulatory signal that determines the magnitude of hydrogen ion excretion (= the production of new bicarbonate) is the concentration of free hydrogen ion in the fluids to which the various transport elements are exposed, i.e., the pH of the ECF and cytosol within renal cells. In effect, the kidneys act as “pH meters” and adjust their transport of hydrogen ion and ammonium excretion accordingly. To some extent, transport is also affected by aldosterone. A comprehensive description of how this occurs is beyond the scope of this text, so we instead cover a few of the basic concepts.

When acid-base status is normal (Table 9–2), and there is no net input of acid or base, the tubules should secrete exactly enough hydrogen ions to achieve complete reabsorption of all filtered bicarbonate. When we have the more common case of a small acid load, additional hydrogen ions will be secreted that titrate buffers in the tubular lumen (titratable acid) and produce some excreted ammonium, thereby returning new bicarbonate to the blood. (Recall that our diet usually produces net hydrogen ions that combine with bicarbonate to reduce total body bicarbonate. The amount of bicarbonate lost titrating hydrogen ion added to the blood must be generated by the kidneys to maintain balance.) During alkalosis, tubular secretion of hydrogen ion should be too low to completely reabsorb the filtered bicarbonate. Then bicarbonate can be lost in the urine; no titratable acid is formed because no extra secreted hydrogen ions are available to combine with non-bicarbonate buffers, and so no new bicarbonate is contributed to the blood. During acidosis, tubular hydrogen ion secretion should increase to reabsorb all filtered bicarbonate and have enough hydrogen ions left to convert most of the base form of titratable buffers to the acid form. Furthermore, glutamine production by the liver and its subsequent metabolism by the proximal nephron to produce ammonium should increase in order to excrete as ammonium the hydrogen ion that is not excreted as titratable acid. As should be clear by now, both titration of filtered buffer and production of ammonium contribute new bicarbonate to the blood.

An increase in PaCO₂, as occurs during respiratory acidosis (eg, caused by shallow breathing after chest trauma), will produce a decrease in plasma pH and, thereby, signal an increased tubular hydrogen ion secretion. A decrease in PaCO₂, as occurs during respiratory alkalosis (eg, high altitude hyperventilation), causes a decrease in secretion. The effects are not due to the CO₂ molecule itself but to the effects of an altered PaCO₂ on renal intracellular pH. Thus, because the tubular membranes are quite permeable to CO₂, an increased arterial Pco₂ causes an equivalent increase in PCO₂ within the tubular cells. This, in turn, causes elevated intracellular hydrogen ion concentration by driving the reactions shown in Equation 9–4 to the right. It is this change that, via a sequence of intracellular

---

5Here again, misuse of the “-osis” versus “-emia” suffixes can confuse the reader. A respiratory acidosis, as described later, simply means a high PaCO₂ (>40 mm Hg). It does not mean that CO₂ is entering the body at an elevated rate. If the PaCO₂ is high but bicarbonate is normal, then, by the Henderson-Hasselbalch equation, the pH must be low.
events, increases the rate of hydrogen ion secretion. This probably occurs in most, if not all, the tubular segments that secrete hydrogen ions.

The second signal that influences hydrogen ion secretion in a homeostatic manner is a change in extracellular pH unrelated to PaCO$_2$. The generalization is that a decreased extracellular pH acts directly on the tubular cells, at least in part by changing intracellular pH, to stimulate hydrogen ion secretion. An increased extracellular pH does the opposite. As with PaCO$_2$, these effects are probably exerted on most, if not all, the tubular segments that secrete hydrogen ions.

We can see that these renal responses are appropriate. If the PaCO$_2$ is high (causing a drop in plasma pH), the increased hydrogen ion secretion raises plasma bicarbonate, thereby restoring plasma pH to normal (despite the continued high PaCO$_2$). Similarly, if the pH is low because of low bicarbonate, the new bicarbonate restores the bicarbonate (and, therefore, the pH) to normal.

**CONTROL OF RENAL GLUTAMINE METABOLISM AND NH$_4^+$ EXCRETION**

In addition to regulating hydrogen ion secretion per se, there are several homeostatic controls over the production and tubular handling of NH$_4^+$. First, the generation of glutamine by the liver is increased by low extracellular pH. In this case, the liver shifts some of the disposal of ammonium ion from urea to glutamine. Second, the renal metabolism of glutamine is also subject to physiological control by extracellular pH. A decrease in extracellular pH stimulates renal glutamine oxidation by the proximal tubule, whereas an increase does just the opposite. Thus, an acidosis, by stimulating renal glutamine oxidation, causes the kidneys to contribute more new bicarbonate to the blood, thereby counteracting the acidosis. This pH responsiveness increases over the first few days of an acidosis and allows the glutamine–NH$_4^+$ mechanism for new bicarbonate generation to become the predominant renal process for opposing the acidosis. Conversely, an alkalosis inhibits glutamine metabolism, resulting in little or no renal contribution of new bicarbonate via this route.

In conclusion, acidosis increases renal NH$_4^+$ synthesis and excretion, whereas alkalosis does the opposite. This explains the spectrum of changes in NH$_4^+$ excretion summarized previously. These effects are summarized in Table 9–3.

**Table 9–3.** Homeostatic control of the processes that determine renal compensations for acid base disturbances

<table>
<thead>
<tr>
<th></th>
<th>Glutamine metabolism and NH$_4^+$ excretion are increased during acidosis and decreased during alkalosis. The signal is unknown.</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>Tubular hydrogen ion secretion is</td>
</tr>
<tr>
<td></td>
<td>a Increased by the increased blood P$<em>{CO_2}$ of respiratory acidosis and decreased by the decreased P$</em>{CO_2}$ of respiratory alkalosis.</td>
</tr>
<tr>
<td></td>
<td>b Increased, independently of changes in P$_{CO_2}$, by the local effects of decreased extracellular pH on the tubules; the opposite is true for increased extracellular pH.</td>
</tr>
</tbody>
</table>
Chapter 9

INTRAVENOUS SOLUTIONS: LACTATED RINGER’S

One more way in which acid-base loads can enter the body is via intravenous solutions. Hospitalized patients receive a variety of intravenous solutions, the most common being physiological saline (0.9% NaCl) and 5% dextrose monohydrate (D5W). Physiological saline is iso-osmotic with normal body fluids (osmolality, 287 mOsm/kg), whereas D5W is slightly hypotonic (osmolality, 263 mOsm/kg). Neither has any acid-base content. Another common solution is lactated Ringer’s solution, a mixture of salts that contains lactate at a concentration of 28 mEq/L. The pH is about 6.5. However, this is an alkalinizing solution for the same reason described as the fruit juice paradox earlier. Lactate is the conjugate base of lactic acid. When lactate is oxidized to CO₂ and water, it takes a hydrogen ion from the body fluids (and leaves a bicarbonate).

Table 9–4 provides a summary of the processes, other than CO₂ production, of adding acids and bases to the body fluids. We omit CO₂ production because, except for transient states, CO₂ production is always matched by CO₂ excretion via the lungs. The unifying and, therefore, simplifying principle is that all processes of acid or base addition boil down to addition or loss of bicarbonate. All processes

Table 9–4. Summary of processes that acidify or alkalinize the blood

<table>
<thead>
<tr>
<th>Nonrenal mechanisms of acidifying the blood</th>
</tr>
</thead>
<tbody>
<tr>
<td>Consumption and metabolism of protein (meat) containing acidic or sulfur-containing amino acids</td>
</tr>
<tr>
<td>Consumption of acidic drugs</td>
</tr>
<tr>
<td>Metabolism of substrate without complete oxidation (fat to ketones and carbohydrate to lactic acid)</td>
</tr>
<tr>
<td>GI tract secretion of bicarbonate (puts acid in blood)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
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GI, gastrointestinal.

The effect of lactated Ringer’s (alkalinizes the body) should not be confused with a lactic acidosis (acidifies the body) associated with exercise. A lactic acidosis results from the anaerobic conversion of glucose to lactic acid (equal amount of lactate and hydrogen ion), whereas lactated Ringer’s solution contains miniscule amounts of hydrogen ion compared with lactate.
that acidify the blood end up removing bicarbonate, and all processes that alka-
linize the blood end up adding bicarbonate.

**SPECIFIC CATEGORIES OF ACID-BASE DISORDERS**

To help sort out the many acid-base disorders, clinicians assign them to 4 cat-
egories: (1) respiratory acidosis, (2) respiratory alkalosis, (3) metabolic acidosis,
and (4) metabolic alkalosis. For reference, we again write this in the form of the
Henderson-Hasselbalch equation for the CO₂–bicarbonate buffer system.

\[
pH = 6.1 + \log \frac{[\text{bicarbonate}]}{0.03 P_{\text{CO}_2}}
\]

The definition of acid-base disorders is simple conceptually. If there are respi-
ratory disorders, the \( P_{\text{CO}_2} \) is high or low; if there are metabolic disorders, the
bicarbonate is high or low. As we see in the Henderson-Hasselbalch equation,
changing either the \( P_{\text{CO}_2} \) or changing the bicarbonate concentration raises or
lowers the pH.

In a respiratory acidosis (eg, resulting from pulmonary insufficiency), the
low ventilation causes an increase in \( P_{\text{CO}_2} \), in turn causing a decrease in
pH. It should be clear from the Henderson-Hasselbalch equation that
the pH could be restored to normal if the bicarbonate could be elevated to the
same degree as the elevation in \( P_{\text{CO}_2} \). It is the job of the kidneys to cause this
bicarbonate increase by contributing new bicarbonate to the blood. An elevation
in bicarbonate in response to an altered \( P_{\text{CO}_2} \) is called compensation. Compensation
occurs because (1) \( \text{NH}_4^+ \) production and excretion are increased and (2) the rise in
\( P_{\text{CO}_2} \) and drop in extracellular pH both stimulate renal tubular hydrogen ion
secretion so that all filtered bicarbonate is reabsorbed, and increased amounts of
secreted hydrogen ion are left over for the formation of titratable acid.

Renal compensation varies in effectiveness. If the elevation in bicarbonate is
great enough to bring the pH back to within the normal range, the situation is
well compensated. Generally, compensation is not complete (ie, when a new steady
state is reached, the plasma bicarbonate is usually not elevated to quite the same
degree as the \( P_{\text{CO}_2} \)). Consequently, blood pH is not returned completely to nor-
mal. If the pH remains quite low, this is an uncompensated or partially compen-
sated case. Note that with a well-compensated case, even though the pH does not
indicate that something is wrong, the elevated \( P_{\text{CO}_2} \) and elevated bicarbonate
indeed indicate that things are not normal.

The renal compensation in response to respiratory alkalosis is just the opposite.
Respiratory alkalosis is the result of hyperventilation, as occurs at high altitude, in
which the person transiently eliminates carbon dioxide faster than it is produced,
thereby lowering \( P_{\text{CO}_2} \) and raising pH. Thereafter, even though ventilation

---

The reader is reminded that this does not mean the body is being flooded with \( \text{CO}_2 \) that is somehow
retained. Although there was a transient period of \( \text{CO}_2 \) accumulation to raise the \( P_{\text{CO}_2} \), the patient with
a respiratory \( P_{\text{CO}_2} \) disorder is not producing\( \text{CO}_2 \) faster than normal or eliminating it more slowly than
normal.
remains high, CO₂ production and its excretion are normal. The decreased PaCO₂ and increase in extracellular pH reduce tubular hydrogen ion secretion, so that bicarbonate reabsorption is not complete. In addition, bicarbonate secretion is stimulated. Bicarbonate is, therefore, lost from the body, and the loss results in decreased plasma bicarbonate and a return of plasma pH toward normal. There is no titratable acid in the urine (the urine is alkaline in these conditions), and there is little or no NH₄⁺ in the urine because the alkalosis inhibits NH₄⁺ production and excretion.

**RENAL RESPONSE TO METABOLIC ACIDOSIS AND ALKALOSIS**

There are many possible causes of metabolic disturbances, including the kidneys themselves. These include (1) increased input of acid by ingestion, infusion, or production; (2) decreased renal production of bicarbonate, as in renal failure, so that normal acid input is not excreted; or (3) direct loss of bicarbonate from the body (as in diarrhea). The result is the same regardless of whether there is loss of bicarbonate or addition of hydrogen ions, i.e., a lower concentration of bicarbonate and a lower plasma pH. The kidneys’ response is an attempt to raise the plasma bicarbonate concentration back toward normal, thereby returning pH toward normal. To do this, the kidneys must reabsorb all the filtered bicarbonate and contribute new bicarbonate through increased formation and excretion of NH₄⁺ and titratable acid. This is precisely what normal kidneys do, but if the acid load is too great or the problem is in the kidneys themselves, the bicarbonate concentration will remain low.

Just as there is renal compensation for a respiratory acid-base disturbance, there is respiratory compensation for a metabolic disturbance. Specifically, a decrease in arterial pH stimulates ventilation, thereby lowering PaCO₂, whereas a rise in arterial pH retards ventilation, allowing Paco₂ to rise.

By now the astute reader has recognized a potential problem in the interpretation of acid-base disorders. When any acid-base disorder is well compensated, both the PaCO₂ and bicarbonate are elevated or depressed in the same direction (e.g., in a well-compensated respiratory acidosis, both the PaCO₂ and bicarbonate are high). Thus, is this actually a respiratory acidosis with renal compensation, or is it a metabolic alkalosis with respiratory compensation? Similarly, in a well-compensated metabolic acidosis, both the PaCO₂ and bicarbonate are low. However, in a real-life situation, renal compensation for respiratory acid-base disturbances can be nearly complete, whereas respiratory compensation is usually only partial (because an alteration in breathing to correct blood pH also alters the uptake of oxygen, leading to an independent modulation of ventilation via changes in PaO₂). In addition, it would be rare in a clinical setting not to have additional information. For example, the high PaCO₂ of an emphysema patient is, in all likelihood, a respiratory acidosis resulting from impaired ventilation, not a compensation for a metabolic alkalosis. Nevertheless, real-life mixed acid-base disorders often present a challenge in the clinic.
FACTORS CAUSING THE KIDNEYS TO GENERATE OR MAINTAIN A METABOLICALKALOSIS

We conclude this chapter with several examples of how otherwise normally functioning kidneys transport hydrogen ions inappropriately and thereby either generate or maintain an acid-base disorder; in these cases, metabolic alkalosis. In any metabolic alkalosis, by definition the plasma bicarbonate concentration is elevated. This problem is not a defect in the ability of the kidneys to excrete bicarbonate; if a person is fed a large load of bicarbonate, the kidneys can excrete the load without a major rise in bicarbonate levels. The problem seems to be in regulation of bicarbonate excretion. The most important situations in which this occurs are (1) volume contraction, (2) chloride depletion, and (3) the combination of aldosterone excess and potassium depletion. The key event in all these situations is oversecretion of hydrogen ion (and sometimes of NH₄⁺ as well), either producing a metabolic alkalosis or failing to respond as usual to an existing metabolic alkalosis. These cases represent exceptions to the normal situation where the kidneys handle any combination of requirements to increase or decrease excretion of salts and acid-base components.

Influence of Extracellular Volume Contraction

The presence of total-body volume contraction because of salt loss interferes with the ability of the kidneys to handle bicarbonate appropriately. Because the bicarbonate concentration is high in any metabolic alkalosis, the normal renal response should be to turn down hydrogen ion secretion to a level that falls short of complete bicarbonate reabsorption, thereby allowing the excess bicarbonate to be excreted. However, the presence of the extracellular volume contraction stimulates not only sodium reabsorption but also hydrogen ion secretion because the transport of these ions is linked via the Na/H antiporters in the proximal tubule. In addition, the renin-angiotensin system is usually activated, resulting in the stimulation of aldosterone secretion. Besides stimulating sodium reabsorption, aldosterone stimulates hydrogen ion secretion by Type A intercalated cells. The net result is that all the filtered bicarbonate is reabsorbed so that the already elevated plasma bicarbonate associated with the preexisting metabolic alkalosis is locked in, and the plasma pH remains high. The urine, instead of being alkaline, as it should be when the kidneys are normally responding to a metabolic alkalosis, is somewhat acid. The generation or maintenance of a metabolic alkalosis in volume contraction may also occur when the volume is normal or high but the body “thinks” volume is low, specifically in congestive heart failure and advanced liver cirrhosis.

Influence of Chloride Depletion

We referred to extracellular volume contraction without distinguishing between sodium and chloride losses as the cause because loss of either of these ions will lead to extracellular volume contraction. However, we emphasize that specific chloride depletion, in a manner independent of and in addition to extracellular volume contraction, helps maintain metabolic alkalosis by stimulating hydrogen
Figure 9–6. Pathway by which overuse of diuretics leads to a metabolic alkalosis. $\text{NH}_4^+$ production and excretion are also increased by the presence of a high aldosterone and potassium depletion. The extracellular volume contraction, via both aldosterone and as yet unidentified nonaldosterone mechanisms, helps to maintain the alkalosis once it has been generated. If the diuretics have also caused chloride depletion, this too will contribute to the maintenance of the metabolic alkalosis (not shown).
ion secretion. The most common reasons for chloride depletion are chronic vomiting and heavy use of diuretics. The result is that bicarbonate excretion remains essentially zero, and the metabolic alkalosis is not corrected.

**Influence of Aldosterone Excess and Simultaneous Potassium Depletion**

As noted, aldosterone stimulates hydrogen ion secretion. Potassium depletion, by itself, also weakly stimulates tubular hydrogen ion secretion and NH$_4^+$ production. However, the combination of potassium depletion of even moderate degree and high levels of aldosterone stimulates tubular hydrogen ion secretion markedly (NH$_4^+$ production also goes up significantly). As a result, the renal tubules not only reabsorb all filtered bicarbonate but also contribute inappropriately large amounts of new bicarbonate to the body, thereby causing metabolic alkalosis. Note that there may have been nothing wrong with the acid-base balance to start with: The alkalosis is actually generated by the kidneys themselves. Of course, if alkalosis were already present, this high aldosterone-potassium depletion combination would not only prevent the kidneys from responding appropriately but also would make the alkalosis worse. This phenomenon is important because the combination of a markedly elevated aldosterone and potassium depletion occurs in a variety of clinical situations, the most common of which is the extensive use of diuretic drugs (eg, from extensive, inappropriate diuretic use for weight loss) that can generate a metabolic alkalosis (Figure 9–6).

**KEY CONCEPTS**

1. To maintain acid-base balance, the kidneys must excrete acid or base at a rate that matches net input.
2. The regulation of body pH consists of regulating the concentrations of CO$_2$ (Pco$_2$) and bicarbonate.
3. The addition or loss of fixed acids and bases is equivalent to removing or adding bicarbonate.
4. Fixed acids and bases can enter the body via GI processes, metabolism, intravenous infusions, and renal processes.
5. Under all conditions, the kidneys must recover virtually all filtered bicarbonate proximally and then add acid or base distally depending on input.
6. The kidneys excrete acid by titrating (acidifying) filtered base.
The kidneys also excrete acid by converting glutamine to bicarbonate and ammonium, excreting the ammonium, and returning the bicarbonate to the blood.

Primary acid-base disorders that change either PCO₂ or bicarbonate can be compensated by changing the other variable in the same direction, thereby preserving the ratio of bicarbonate to PCO₂.

Some situations, including volume contraction and aldosterone excess, can cause the kidneys to excrete too much acid, generating a metabolic alkalosis.

**STUDY QUESTIONS**

9–1. Even if the urine pH is neutral (7.4), the kidneys can still excrete acid in the form of ammonium. True or false?

9–2. A patient is observed to excrete 2 L of alkaline (pH, 7.6) urine having a bicarbonate concentration of 28 mmol/L. What is the rate of titratable acid excretion?
   A. 56 mmol
   B. Negative
   C. Cannot determine without data for ammonium

9–3. Which of the following is an acid load per se or generates an acid load that must be excreted by the kidneys?
   A. Prolonged vomiting of stomach secretions
   B. Eating unsweetened grapefruit juice
   C. Eating sweetened grapefruit juice
   D. Intravenous infusion of sodium lactate

9–4. Proximal tubular reabsorption of filtered bicarbonate involves a pair of transporters: one that imports bicarbonate across the apical membrane and another that exports bicarbonate across the basolateral membrane. True or false?

9–5. During a metabolic acidosis, such as a diabetic ketoacidosis, the renal excretion of acid decreases well below normal levels. True or false?

9–6. Two patients have plasma pH values of 7.39 and 7.41, respectively. What is their acid-base status?
   A. One is acidotic; the other is alkalotic.
   B. They are both normal.
   C. There is not enough information.

9–7. An emphysema patient has had serious difficulty breathing for a long time. Which of the situations below are likely?
   A. His PCO₂ is elevated.
   B. His bicarbonate is low.
   C. His urine has an elevated amount of titratable acidity.

9–8. From an acid-base perspective, 1 mEq of titratable acid in the urine is the same as 1 mmol of ammonium. True or false?
Calcium is the fifth most abundant element in the human body (behind oxygen, carbon, hydrogen, and nitrogen), with phosphorous right behind in sixth place. Calcium is distributed in 3 body compartments. The vast majority exists as a structural component of bone. A much smaller, but critical fraction is dissolved in the extracellular fluid (ECF), where its concentration is vital for the conformation, and hence function, of membrane proteins exposed to the external medium. A third compartment of calcium is within cells, where it functions as a key component of intracellular signaling cascades. Intracellular calcium is further compartmentalized, with most calcium being sequestered in organelles such as mitochondria and the endoplasmic reticulum.
Chapter 10

Calcium obeys the principles of input–output balance (as do all the other substances we have discussed), but its regulation is fundamentally different from other substances we have considered because its balance is regulated predominantly by the gastrointestinal (GI) tract, although the kidneys play an important role. In addition, like potassium, plasma calcium is strongly “buffered” by the large amounts of calcium in bone that is readily exchangeable with ECF calcium.

There are 2 time scales of calcium balance to consider: rapid transfer of calcium between the ECF and other tissues of the body and the slow rate of calcium ingestion into and excretion from the body. Calcium in the ECF represents only a tiny fraction of total-body calcium, but its level is crucial for the function of cells in the body. Thus, it is essential to maintain the concentration of free calcium in our ECF within a narrow range. Too low a concentration leads to the life-threatening condition of low-calcium tetany (discussed later).

Moment-to-moment regulation of extracellular calcium is achieved by shifting calcium in and out of bone. Bone stores of calcium serve as an enormous buffer system capable of keeping plasma calcium nearly constant. In essence, balance for calcium in the ECF is balance to and from bone, not the outside world. Besides the rapid exchange of calcium into or out of bone, there is also a long-term regulation of total calcium in bone that is important for bone growth during childhood and bone integrity in adult life. Here, the kidneys play an important but indirect role because they excrete calcium in the urine and are involved in forming the active form of vitamin D.

The dominant regulation of total-body calcium balance is less focused on output and more focused on input from the GI tract. Interestingly, and in contrast to most substances considered thus far, absorption of dietary calcium is only partial. In fact, most calcium that we eat simply passes through the GI tract to the feces. Calcium is both reabsorbed and secreted in the gut. The secretory rate is more or less constant, but the fraction absorbed is regulated to produce anywhere from small to moderate net absorption. Some net absorption is essential for the long-term maintenance of adequate bone calcium. Lack of calcium during childhood growth leads to a weak bone condition called rickets, and loss of calcium in an adult leads to osteomalacia or osteoporosis.¹

Normal levels of plasma calcium are about 10 mg/dL (2.5 mmol/L or 5 mEq/L). This calcium exists in 3 general forms. First, almost half is in the free ionized (Ca²⁺) form. This is the only form that is biologically active in target organs. Second, about 15% is complexed to anions with relatively low molecular weights, such as citrate and phosphate. Third, the remaining 40% is reversibly bound to plasma proteins.

¹Rickets, osteomalacia, and osteoporosis are characterized by low calcium content in bone. Rickets and osteomalacia are commonly associated with a low supply of calcium, typically because of low vitamin D. Osteoporosis seems to represent improper regulation, so that the ongoing bone-forming and bone-dissolving processes are dominated by bone dissolution.
The Chemistry of Calcium

The role of calcium in the physiology of the body is critically dependent on its chemistry. Calcium is a divalent cation, and like all ions in solution it is surrounded by a shell of water molecules. Because of the relatively large atomic size of calcium, the water molecules in the shell are not tightly held, so that it is relatively easy for calcium to shed them, thereby increasing the reactivity of calcium with other ions. Accordingly, calcium avidly forms complexes with small anions such as phosphate and oxalate, and associates reversibly with anionic groups on proteins. Calcium behaves as a “sticky” ion. This accounts for the fact that only about half the plasma calcium is actually in the free, dissolved form. The complexes with small anions easily precipitate out of solution, limiting their combined concentrations (low solubility product). High concentrations of such anions tend to remove calcium from solution. Similarly, proteins with many groups available to bind calcium also remove calcium from solution. How does all this affect the physiology of the body? (1) The operation of functional proteins like transporters, enzymes, and channels is governed by their shape and charge. The relative amount of calcium bound to such proteins strongly affects their charge, and hence their function. The body takes advantage of this property by using reversible binding of calcium to components of intracellular signaling pathways to turn cellular processes on and off. (2) All body cells regulate their cytosolic calcium concentrations to very low levels in order to prevent formation of calcium complexes with the many forms of phosphate within cells, and to prevent inappropriate activation of signaling pathways. They do this with a variety of active transport systems (ATPases and antiporters) that remove calcium from the cytosol, either to the external medium or to intracellular organelles. (3) In bone, the body takes advantage of the tendency of calcium to form complexes. Much of the hardness and strength of bone comes from its high content of hydroxyapatite, a complex of calcium, phosphate, and hydroxyl groups.² (4) Because calcium forms complexes with anions, the plasma concentration of anions like phosphate cannot be allowed to get too high; otherwise calcium phosphate complexes precipitate out. This tendency to precipitate becomes a problem in the kidneys under conditions of very high urinary calcium content when complexes of calcium phosphate and calcium oxalate can form renal calculi (kidney stones) that can grow large enough to block the ureter.

Calcium in the ECF is the source for calcium entering cells through calcium channels and is the calcium that triggers rapid exocytosis of hormones and neurotransmitters, and signals contraction in smooth and cardiac muscle cells. A second role of extracellular calcium is to regulate the threshold for excitation in nerve and muscle cells. This role is completely separate from its role as a cation that enters through membrane channels. The effect of reversible calcium binding to

²Apatites are a class of compounds of variable structural formula that usually include calcium, phosphate, and an anion such as fluoride, chloride, or hydroxide. The apatites in bone are a mixture of these, but the dominant one is hydroxyapatite
membrane proteins is strongly exerted on voltage-gated sodium channels. Low levels of calcium fool sodium channels into sensing more depolarization than actually exists, leading to spontaneous firing of motor neurons. In turn, this firing triggers inappropriate muscle contraction, called low-calcium tetany. If severe enough, then it leads to respiratory arrest because of spasms in the ventilatory muscles.3

One of the most important influences on the degree of calcium binding to nerve membranes is the plasma pH. Serum albumin has many anionic sites that reversibly bind both protons and calcium. These ions compete for occupancy of the binding sites. As pH rises, protons dissociate and calcium ions take their place, thereby lowering the concentration of free calcium ions. In turn, this tends to cause reduced binding of calcium to cell membranes. Thus, a patient with an acute alkalosis is more susceptible to tetany, whereas someone with acidosis will not manifest tetany at levels of total plasma calcium low enough to cause symptoms in normal people.

Elevated plasma calcium (hypercalcemia) is also a serious medical problem, particularly if the rise occurs rapidly. Hypercalcemia has multiple effects, generally resulting from the deterioration of excitable cell function, including CNS depression, muscle weakness, and GI tract immotility.

**EFFECTOR SITES FOR CALCIUM BALANCE**

**GI Tract**

As mentioned, and in contrast to sodium, chloride, and potassium, most ingested calcium is not normally absorbed from the intestine and simply leaves the body along with the feces. Accordingly, changes in the active transport system that moves calcium from intestinal lumen to blood can result in large increases or decreases in calcium absorption. Hormonal control of this absorptive process is the major means for homeostatically regulating total-body calcium balance. Calcium enters from the intestinal lumen passively through calcium-selective channels, binds reversibly to mobile cytosolic calcium-binding proteins (called calbindins), and is then actively transported out the basolateral side via a Ca-ATPase and Na-Ca antiporter. Calbindins contain multiple binding sites for calcium and are free to diffuse throughout the cytosol. They act as ferry boats for calcium, permitting large amounts of calcium to move from place to place within a cell, in this case from apical to basolateral membrane, all the while keeping the concentration of free calcium at a low level.

**Kidneys**

The kidneys handle calcium by filtration and reabsorption. About 60% of the plasma calcium is filterable; the remainder is bound to plasma proteins. Most calcium reabsorption occurs in the proximal tubule (about 60% of the filtered load).

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3Even though calcium in the extracellular medium is required to trigger the release of neurotransmitter from motor neurons (absence of calcium blocks this process completely), low calcium causes excessive muscle stimulation because hyperexcitability is manifested at calcium levels well above the extremely low levels needed to block exocytosis.
and the remainder in the thick ascending limb of Henle’s loop, distal convoluted tubule, and collecting-duct system. Overall, reabsorption is normally 97–99%.

Calcium reabsorption in the proximal tubule and thick ascending limb of Henle’s loop is largely passive and paracellular, and the electrochemical forces driving it are dependent directly or indirectly on sodium reabsorption as they are for so many other substances. In contrast, calcium reabsorption in the more distal segments is active and transcellular. It uses the same mechanism as in the GI tract, ie, entrance via calcium-specific channels, diffusion bound to calbindins and exit across the basolateral membrane actively by a combination of Ca-ATPase and Na-Ca antiport activity (Figure 10–1). The distal tubule is the site of endocrine control of renal calcium handling.

The amount of calcium excreted in the urine is, on average, equal to the net addition of new calcium to the body via the GI tract; thus, the kidneys help maintain a stable balance of total-body calcium. However, the change in renal excretion in response to changes in dietary input is much less than the equivalent responses to dietary sodium, water, or potassium. For example, only about 5% of an increment in dietary calcium appears in the urine, whereas virtually all of an increased ingestion of water or sodium soon appears in the urine. The reason is that most of the dietary increment never gains entry to the blood because it fails

**Figure 10–1.** Generic method of transcellular calcium transport in the GI tract and kidney. In all body cells, the free intracellular calcium concentration must be kept to minuscule levels to prevent the formation of complexes and activation of deleterious processes, even though the concentrations of calcium at the two external surfaces of the cells are thousands of times higher. The epithelial cells solve this by using diffusible calbindins. As calcium enters the cells through channels in the luminal surface, this raises the calcium concentration in the microenvironment near the channels, thus promoting binding to calbindins. At the basolateral surface this process is reversed. Active extrusion of calcium via ATPases and sodium-calcium antiporters lowers the calcium concentration in the local microenvironment, promoting dissociation of calcium from the calbindins.
to be absorbed from the GI tract. In contrast, when dietary intake of calcium is reduced to extremely low levels, there is a slow reduction of urinary calcium, but some continues to appear in the urine for weeks.

How do the renal homeostatic mechanisms operate? Because calcium is filtered and reabsorbed, but not secreted,

\[ \text{Ca excretion} = \text{Ca filtered} - \text{Ca reabsorbed} \]

Accordingly, excretion can be altered homeostatically by changing either the filtered load or the rate of reabsorption. Both occur. For example, what happens when a person increases calcium intake? Transiently, intake exceeds output, positive calcium balance ensues, and plasma calcium concentration may increase. (However, recall that bone acts as a huge calcium buffer, so the rise in plasma calcium is slight.) A rise in plasma calcium increases both the filtered mass of calcium and excretion. Simultaneously, as we will see, the increased plasma calcium triggers hormonal changes that cause a diminished reabsorption. The net result of these responses is increased calcium excretion.

A wide variety of factors not designed to maintain calcium homeostasis can also influence urinary calcium excretion, mainly by stimulating or inhibiting tubular reabsorption. These include a large number of hormones, ions, acid-base disturbances, and drugs.

One of the most important of these influences on calcium reabsorption is sodium. An increase or decrease in urinary calcium excretion can be induced simply by administering or withholding salt, respectively. This fact used clinically as an emergency procedure when calcium levels in the blood get alarmingly high, the treatment consisting of administering large amounts of saline, with the consequence that large amounts of calcium-containing fluid pass through the kidneys to the urine.

A second very important factor that influences tubular calcium reabsorption, but is not designed to maintain calcium homeostasis is the presence of an acidosis. The mechanism is not clear, but acidosis markedly inhibits calcium reabsorption and, hence, causes increased calcium excretion. Alkalosis tends to do just the opposite: enhance calcium reabsorption and reduce excretion.

**Bone**

Bone is a complex tissue structurally and physiologically. It is the least understood, but in some ways the most important of the major effector systems for calcium management. The literature about bone physiology and its regulation is filled with uncertainty and disagreement. However, 2 things are abundantly clear. (1) Bone serves as a powerful short-term calcium-buffering system that prevents large swings in plasma calcium. About 0.5 g of calcium passes back and forth between bone and the blood plasma each day. (2) Bone is also the repository of calcium that keeps blood supplied with calcium during times of negative whole-body calcium balance. What remains controversial are the mechanisms by which calcium moves in and out of bone and how they are regulated by hormones.
The majority of the bone mass is made up of a tough proteinaceous framework, mostly collagen, on which is deposited hard mineral crystals of hydroxyapatite. Although very hard, bone is not uniformly solid, like brick. Rather, it is penetrated by a labyrinth of tiny passageways containing fluid (bone fluid), cells (mostly osteocytes), and, in the larger passageways, blood vessels. It is believed that the osteocytes deep within bone communicate with each other and the surface cells via long cellular extensions containing gap junctions. Calcium can move back and forth between blood and the inner recesses of bone via this cellular network.

Synthesis and degradation of hydroxyapatite is fundamentally different from that of the other substances that form the bulk of the nonwater mass of the body (ie, protein and triglyceride). First, unlike those substances, hydroxyapatite is synthesized extracellularly. Second, and also unlike those substances, it is not synthesized by a series of enzyme-catalyzed steps; rather it occurs by chemical mass action. The equilibrium between crystalline hydroxyapatite and its dissolved components is highly labile, dependent on the concentrations of calcium, phosphate, hydrogen ions, and specific noncollagenous proteins. Thus, the structure and physiology of bone is determined by the chemistry of calcium and the modulation of the calcium-containing bone fluid. Bone fluid is separated from the ECF by a layer of cells called the “bone membrane.” These cells are mostly flattened versions of the active osteoblasts involved in forming bone. The actions of cells of the bone membrane are crucial in regulating the balance between synthesis and degradation of hydroxyapatite via control over the fluid environment of the bone matrix.

Movement of calcium across the bone membrane constitutes the rapid buffering system that protects the blood plasma from short-term swings in calcium concentration. This process does not require hormonal signals. However, the set point for plasma calcium maintained by the rapid buffering system is critically regulated by hormonal control, as discussed later. The second flux process involving calcium is called bone remodeling and affects calcium stores on a slower time scale. Remodeling involves the paired actions of giant, multinucleated cells called osteoclasts that erode little pits in the bone matrix and their partners, nearby osteoblasts, which follow behind and fill in the pits with new bone matrix. The osteoclasts pump hydrogen ions and create an acidic microspace directly underneath them that solubilizes hydroxyapatite. The calcium and phosphate freed up by this process is then transported transcellularly to the ECF. The daily flux of calcium via remodeling is much less than that associated with rapid flux across the bone membrane. Normally, the fluxes associated with remodeling result in no net gain or loss of calcium, but imbalance in resorption of bone matrix relative to replacement causes gradual loss of bone density and pathology such as osteoporosis.

A long-standing question has been whether the bone membrane acts as a seal that maintains the ionic composition of bone fluid quite different from the ECF surrounding bone, or is a more porous barrier that permits paracellular flux of calcium and other small ions. Recent evidence favors the latter view. A related question is whether hormonal stimulation is exerted by increasing transcellular calcium transport across cells of the bone membrane or by raising the solubility of hydroxyapatite, thereby releasing calcium that moves passively across the bone membrane via the paracellular route. Again, recent evidence favors the latter view.
HORMONAL CONTROL OF EFFECTOR SITES

The regulation of calcium is achieved mostly through the actions of 2 hormones: the active form of vitamin D—1,25-(OH)₂D—and parathyroid hormone (PTH), a peptide hormone produced by the parathyroid glands. The active form of vitamin D acts mainly to stimulate intestinal absorption of calcium and phosphate. In growing children, this ensures a supply of substrate for bone formation, and in adults it ensures a supply to replace the ongoing dissolution of bone. PTH has several actions, a key one being to dissolve bone and move calcium into the blood. PTH stimulates the bone membrane to release calcium on a short-term basis, and via paracellular signals from osteoblasts, also stimulates osteoclasts to resorb bone (Figure 10–2). These processes protect the body from low-calcium tetany. Simply stated, the active form of vitamin D regulates what comes into the body and PTH regulates what is in the ECF.

Vitamin D

The term vitamin D denotes a family of closely related molecules that are derived from cholesterol. One member of this family, called vitamin D₃ (cholecalciferol), is synthesized by the action of ultraviolet radiation on 7-dehydrocholesterol in the skin. The 7-dehydrocholesterol precursor is normally present in adequate amounts to avoid limiting the production of vitamin D₃. Thus, the synthesis of vitamin D₃ itself is strongly dependent on exposure to sunlight, which is, in turn, dependent on sunlight exposure.

Figure 10–2. Mechanism of calcium reabsorption. The distal convoluted tubule is the major site for regulated reabsorption. Ca enters via apical Ca channels (under the control of parathyroid hormone [PTH]) and is actively transported across the basolateral membrane via Na-Ca antiport and via a Ca-ATPase. The apical membrane also contains the Na-Cl symporter (NCC), which is the target for inhibition by thiazide diuretics. Interestingly, the inhibition of NCC with thiazide diuretics promotes calcium reabsorption (probably by enhancing the basolateral sodium gradient and increasing Na-Ca exchange). Thus, thiazides may reduce the calcium loss associated with osteoporosis.
on climate, latitude, clothing, and so on. Vitamin D₃, like cholesterol, is a 27-carbon molecule containing 1 hydroxyl group. Another member of the family is vitamin D₂, which is ingested in food, specifically food derived from plants. Vitamin D₂ (ergocalciferol) differs chemically from vitamin D₃ by an additional methyl group and a double bond between 2 of the carbons. It is a form of vitamin D often added as a supplement to foods. These 2 members of the vitamin D family act by identical mechanisms, although vitamin D₃ is more potent. Vitamin D means either vitamin D₂ or vitamin D₃.

Vitamin D as such is inactive (ie, neither ingested vitamin D₂ nor the vitamin D₃ formed in the skin has any significant biological activity). It must undergo metabolic changes within the body before it can influence its target cells. Circulating vitamin D is hydroxylated at the 25 position by the liver and then hydroxylated again at the 1 position by proximal tubular cells within the kidneys to yield a cholesterol derivative containing 3 hydroxyl groups. (When formed from vitamin D₃, it is called calcitriol.) This dihydroxy form of vitamin D—1,25-dihydroxyvitamin D, or 1,25-(OH)₂D—exerts actions on target cells. From this description, it should be evident that the molecular species that is actually exerting actions on target tissues is a hormone, not a vitamin, because it is made in the body. Therefore, we usually call this hormone the active form of vitamin D. Because the active form is generated in the renal epithelium, the kidneys are major regulators of calcium homeostasis in the GI tract and bone as well as regulators via urinary excretion.

The major action of vitamin D is to stimulate active absorption of calcium and phosphate by the intestine. A role of vitamin D is to stimulate synthesis of the proteins involved in the steps described earlier. In addition, vitamin D has some independent actions on bone that are not entirely clear. As well, it stimulates the renal-tubular reabsorption of calcium and phosphate, again by increasing the synthesis of the protein components in the transport pathway. The influences of vitamin D on bone and the kidney are far less important than its actions on the GI tract to stimulate absorption of calcium and phosphate.

The major event in vitamin D deficiency is decreased gut calcium absorption, resulting in decreased availability of calcium for bone formation or reformation. In children, the newly formed bone protein matrix fails to be calcified normally because of the low availability of calcium, leading to the disease rickets.

**PTH**

PTH (parathyroid hormone) is an 84-amino peptide hormone secreted by the parathyroid glands. The GI tract, kidneys, and bone are all subject to direct or indirect control by PTH. It is a substance essential for life, for without PTH, plasma calcium falls to lethal levels within a few days. All of its normal activity is contained in the first 34 amino acids, and synthetic PTH can be made containing only this component. The PTH half-life in the plasma is very short (<10 min), mostly due to rapid degradation in the liver, with renal filtration and uptake.

In breaking down PTH, the liver releases peptide fragments that are active hormones in their own right, but with actions different from PTH. These fragments act to oppose the normal actions of PTH.
playing a secondary role. PTH secretion is controlled on a moment-to-moment basis by the calcium concentration of the ECF bathing the cells of the parathyroid glands. Decreased plasma calcium concentration stimulates PTH secretion, and increased plasma concentration inhibits secretion. Extracellular calcium concentration acts directly on the parathyroid glands by binding to a novel class of G protein–linked receptors whose ligands are divalent cations. The calcium receptor couples via an intracellular G protein to a signaling cascade that inhibits the secretion of PTH. Thus, low extracellular calcium stimulates PTH secretion by removing a tonic inhibition. This is a sensitive control system designed to keep free plasma calcium at about 5 mg/dL.

Phosphate also affects PTH secretion: Elevated phosphate stimulates PTH secretion by stimulating the capacity of the parathyroid gland to synthesize PTH, so that chronically high levels of phosphate lead to elevated PTH. Vitamin D has slower inhibitory effects (discussed later), but calcium is the primary acute regulator.

PTH exerts at least 4 distinct effects on calcium homeostasis (summarized in Figure 10–3):

1. PTH actions on bone normally increase the movement of calcium from bone into the ECF. They do this by stimulating the release of calcium across the bone membrane. Without PTH, the equilibrium between hydroxyapatite and free calcium is shifted toward the formation of hydroxyapatite, causing the bone buffering system to take up calcium from the ECF and reduce plasma calcium concentration to a level incompatible with life. On a slower basis, PTH also stimulates resorption carried out by osteoclasts. It does this via a complex interaction between osteoblasts, which contain receptors for PTH, and osteoclasts. In this manner, the immense store of calcium contained in bone is made available for the regulation of extracellular calcium concentration.

2. PTH stimulates the activation of vitamin D as described earlier (and this hormone then increases intestinal absorption of calcium). The blood concentration of vitamin D is subject to physiological control by PTH. The major control point is the second hydroxylation step, the one that occurs in the kidneys. This step is stimulated by PTH. This action is highly adaptive. If plasma calcium falls acutely, the subsequent increase in PTH immediately stimulates calcium transport from bone, thus restoring plasma calcium levels, and also stimulates calcium uptake (via vitamin D) from the GI tract. This ensures that enough new calcium will enter the body to replace that “borrowed” from bone.

3. PTH increases renal-tubular calcium reabsorption, mainly by an action on the distal convoluted tubule and connecting tubule. At these locations, it acts rapidly through activation of kinases that phosphorylate regulatory

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6As alluded to in Footnote 4, PTH either directly stimulates transport or causes release of calcium from hydroxyapatite by increasing its solubility. Regardless of the mechanism, the result is an increased efflux of calcium from bone.
proteins on a short-term basis. It also acts, on a slower time scale, to increase the synthesis of all the components of the transport pathway. The increased uptake of calcium from the tubular lumen stimulates basolateral extrusion (by a combination of Ca-ATPase activity and Na-Ca antiporter activity) and thus decreases urinary calcium excretion.\(^7\)

\(^7\)The strongest and most commonly prescribed diuretics (called “loop” diuretics because they inhibit sodium reabsorption in the thick ascending limb of the loop of Henle) have the secondary effect of also decreasing calcium absorption. Another class of diuretics, called the thiazides, are somewhat weaker diuretics. They inhibit sodium reabsorption not in the loop but in the distal tubule by blocking the Na-Cl symporter. Interestingly, they increase Ca reabsorption by a mechanism that hyperpolarizes the cells and stimulates influx of Ca across the apical membrane.
4. It reduces the proximal tubular reabsorption of phosphate, thereby raising urinary phosphate excretion and lowering extracellular phosphate concentration.

The adaptive value of the first 3 effects should be obvious: They all result in a higher extracellular calcium concentration and thus compensate for the lower calcium concentration that originally stimulated PTH secretion. Regarding the fourth effect, when PTH acts on bone, both calcium and phosphate are released into the blood. Similarly, the active form of vitamin D enhances the intestinal absorption of both calcium and phosphate, so that the processes that are restoring calcium to its normal level are simultaneously tending to raise the plasma phosphate above normal. But this is an unwanted action because an increase in dissolved phosphate causes a decrease in dissolved calcium. Under the influence of PTH plasma phosphate does not actually increase, because of PTH’s inhibition of tubular phosphate reabsorption. Indeed, this effect is so potent that plasma phosphate may actually decrease when PTH levels are elevated. The simultaneous presence of high levels of both calcium and phosphate can create pathologies in a number of tissues, including the heart and blood vessels, because this increases the formation of insoluble calcium phosphate complexes.

All of the above describes the effects of a rise in PTH induced by a fall in plasma calcium. An increase in extracellular calcium concentration reduces PTH secretion, thereby producing increased urinary and fecal calcium loss and net movement of calcium from the ECF into bone.

PTH has other functions in the body, but the 4 effects discussed previously constitute the major mechanisms by which it integrates various organs and tissues in the regulation of extracellular calcium concentration.

The actions of PTH on bone are dependent on the pattern of its plasma concentration over time. It can either promote resorption of hydroxyapatite (its usual action) or promote deposition. Primary hyperparathyroidism, resulting from a primary defect in the parathyroid glands (eg, a hormone-secreting tumor), generates a continuous excess hormone level and causes enhanced bone resorption. This leads to bone thinning and the formation of completely calcium-free areas or cysts. In this condition, plasma calcium often increases and plasma phosphate decreases; the latter is caused by increased urinary phosphate excretion. The increased plasma calcium modulates the activity of various body tissues. A seeming paradox is that urinary calcium excretion is increased despite the fact that tubular calcium reabsorption is enhanced by PTH. The reason is that the elevated plasma calcium concentration induced by the effects of PTH on bone (and via vitamin D on the load of calcium entering the body from the GI tract) causes the filtered load of calcium to increase even more than the reabsorptive rate. Because the filtered load is so great, there is also an increased amount not reabsorbed (ie, excreted). This result nicely illustrates the necessity of taking both filtration and

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8 The chemistry of the Ca phosphates is highly complex because Ca and phosphate can combine in different stochiometric ratios, and the tendency of any of these to precipitate depends on the concentrations not only of Ca and phosphate but also of other substances as well.
reabsorption (and secretion, if relevant) into account when analyzing excretory changes of any substance. And as mentioned earlier, the high urinary calcium content promotes the formation of stones.

In contrast to what happens with the continuous presence of elevated PTH that accelerates bone resorption and release of calcium, intermittent rises (produced by infusions once per day) actually increases deposition of calcium in bone. Intermittent infusion of PTH is used therapeutically to increase bone density in osteoporosis patients.

**OVERVIEW OF RENAL PHOSPHATE HANDLING**

The renal handling of phosphate is almost always mentioned in the context of other topics, such as sodium reabsorption or urine acidification. We now review certain key aspects of renal phosphate handling because control of urinary phosphate excretion is a major pathway for the homeostatic regulation of total-body phosphate balance.

Approximately 5–10% of plasma phosphate is protein bound, so that 90–95% is filterable at the renal corpuscle. Normally, approximately 75% of this filtered phosphate is actively reabsorbed, almost entirely in the proximal tubule (in support with sodium).

As with other substances handled by filtration and tubular reabsorption, the rate of phosphate excretion can be changed by altering the mass filtered per unit time or the mass reabsorbed per unit time. Indeed, even relatively small increases in plasma phosphate concentration (and, hence, filtered load) can produce relatively large increases in phosphate excretion. This occurs when plasma phosphate concentration increases as a result of increased dietary phosphate intake or release of phosphate from bone. Phosphate is reabsorbed by a tubular maximum-limited ($T_m$) system, and the normal filtered load is just a little higher than the $T_m$. Thus, most filtered phosphate is reabsorbed, but some spills into the urine. (Recall that this phosphate is responsible for accepting hydrogen ion in the collecting duct and is the primary ion responsible for titratable acidity.) As the reabsorptive capacity is saturated at normal filtered loads, any increase in filtered load simply adds to the amount excreted. As mentioned, systemic acidosis promotes the release of calcium and phosphate from bone. The increase in plasma phosphate and the consequent increase in filtered load of phosphate means that there is more titratable buffer in the collecting tubule to help remove the excess hydrogen ion that promoted the phosphate release.

Changes in PTH and vitamin D are not “designed” to mediate the homeostatic association between dietary phosphate and tubular phosphate reabsorption. Nevertheless, as we have seen, whenever PTH is increased or decreased, tubular phosphate reabsorption is powerfully inhibited or stimulated, respectively. Other hormones, too, are known to alter phosphate reabsorption, for example, insulin increases it and glucagon decreases it.

Much of the physiology we have described is illustrated by the case of chronic renal failure, in which a low glomerular filtration rate limits the ability of the
The kidneys excrete a number of substances, specifically phosphate. An almost universal complication of chronic renal failure is elevated plasma phosphate (hyperphosphatemia). Another common finding is elevated levels of PTH. The high PTH stimulates excessive bone resorption, leading to osteoporosis. This is an example of secondary hyperparathyroidism (not primary) because the pathology is not in the gland itself, but in the signals that drive it. One goal in the treatment of hyperphosphatemia associated with chronic renal failure is the reduction of phosphate absorption from the GI tract. This is accomplished by feeding the patient high doses of calcium. The calcium forms complexes with phosphate in the GI tract, reducing the availability of absorbable phosphate. The high levels of PTH in a normal patient should signal the kidneys to form vitamin D, but in chronic renal failure a further complication is a reduced ability to synthesize it. Another clinical intervention in this case is to provide exogenous vitamin D. This hormone suppresses the expression of the PTH gene in the parathyroid gland. The vitamin D should increase GI absorption of phosphate, the very thing we are trying to inhibit, but its ability to lower the synthesis of PTH is more important because this reduces the excessive resorption of bone stimulated by PTH. Thus, administering vitamin D is a useful clinical tool.

KEY CONCEPTS

1. Moment-to-moment regulation of plasma calcium primarily involves calcium flux between bone and plasma rather than input and output from the body.

2. The most important action of vitamin D is to ensure adequate absorption of calcium from the GI tract.

3. PTH is essential both to maintain proper calcium flux between bone and plasma and to maintain adequate levels of vitamin D.

4. Keeping phosphate levels in the normal range allows normal calcium retrieval from bone.
10–1. The most important action of vitamin D is to stimulate:
   A. Calcium deposition in bone
   B. Calcium resorption from bone
   C. Calcium absorption from the GI tract
   D. Calcium reabsorption in the renal tubule

10–2. PTH stimulates phosphate secretion in the proximal tubule. True or false?

10–3. In response to a sudden decrease in plasma calcium, where does most of the calcium come from to restore plasma levels?
   A. The renal tubules
   B. Bone
   C. The GI tract

10–4. When a reduction in PTH signals the kidneys to increase excretion of calcium, they do so by what mechanism?
   A. Increased glomerular filtration
   B. Reduced reabsorption in the proximal tubule
   C. Increased secretion in the distal tubule
   D. Reduced reabsorption in the distal tubule

10–5. Chronic failure of the kidneys leads to what problem?
   A. High plasma PTH
   B. Low plasma phosphate
   C. Inability to transfer calcium from bone to the blood
   D. High plasma vitamin D
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CHAPTER 1

1–1. The correct answer is false. All glomeruli are in the cortex.

1–2. The correct answer is none. Most medullary blood is supplied by efferent arterioles near the corticomedullary border. A small fraction may be supplied from cortical radial arteries. All of these vessels are in the cortex.

1–3. The correct answer is no. It is possible that substance $T$ was filtered, but substance $T$ might also enter tubules by secretion into the tubules.

1–4. All substances in the general circulation enter the kidneys via the renal arterial system and flow through the renal microvasculature. It is possible that it was neither secreted nor filtered, accounting for its not being in the urine, but it is more likely that whatever substance $V$ reaches the lumen via filtration or secretion is completely reabsorbed. Many substances fall into this category.

1–5. The answer is none. In glomerular filtration, the filtered substances pass around the endothelial cells (through fenestrations) and around the podocytes. Topologically, Bowman’s space is continuous with the external environment; thus, a filtered substance can move from blood to the bladder and be excreted without crossing through any plasma membrane.

1–6. The answer is no. Freely filtered means the substance is filtered in the same proportion that volume is filtered. If 20% of the volume is filtered, then 20% of a freely filtered substance is filtered, meaning that 80% is not filtered and passes on to the efferent arterioles and peritubular capillaries.

1–7. You would find label X in the cortex. Cells of the macula densa are part of the juxtaglomerular apparatus, just next to the glomeruli that are all in the cortex. You would find label Y in the cortex and outer medulla. Thick ascending limbs of all nephrons begin in the outer medulla and continue back to the glomerulus from which that nephron arose.

1–8. The answer is yes. Under all circumstances, there is net volume reabsorption in the medulla. By mass balance, this requires that the reabsorbed volume leave the medulla. The only way out for reabsorbed substances is in the blood. Thus blood flow out is slightly greater than blood flow in.
CHAPTER 2

2–1. The answer is false. Freely filtered means no restriction or sieving by the filtration barrier. Because normally about 20% of the plasma volume is filtered, then about 20% of substance \( X \) would be filtered.

2–2. The answer is 125 mg/min. The amount of any substance filtered per unit time is given by the product of the GFR and the filterable plasma concentration of the substance, in this case, 125 mL/min \( \times \) 100 mg/100 mL (1 dL = 100 mL).

2–3. Approximately 40% of the calcium in plasma is bound to proteins and so is not filterable.

2–4. If the protein were freely filtered, there would be 100 mg/L \( \times \) 100 L/day = 10 g/day. However, no exact value can be calculated from these data because the molecular weight is high enough so that some “sieving” would occur. Some would be filtered, but less than 10 g/day.

2–5. (1) Constrict glomerular mesangial cells and, hence, reduce \( K_f \). (2) Lower arterial pressure and, hence, \( P_{GC} \). (3) Constrict the afferent arteriole and, hence, reduce \( P_{GC} \). (4) Dilate the efferent arteriole and, hence, reduce \( P_{GC} \).

2–6. It might be increasing \( K_f \) (ie, changing the hydraulic permeability of the glomerular membranes or the surface area available for filtration).

2–7. RBF will show no change because the drug has no effect on total renal vascular resistance. GFR will increase because of a large increase in \( P_{GC} \). Filtration fraction will, therefore, increase. (Now back up and think a bit more about the GFR: Because filtration fraction increases, there will be a larger than average increase in \( \pi_{GC} \) along the glomeruli, and this will offset some of the GFR-increasing effect of the increased \( P_{GC} \); therefore, GFR will not increase as much proportionately as the \( P_{GC} \).)

2–8. The answer is C. Arterial pressure decreases by 33%, but autoregulation prevents the RBF from decreasing in direct proportion. Autoregulation is not perfect, so some decrease, but less than 33%, will occur.

CHAPTER 3

3–1. The answer is C. Clearance units are volume per time, not mass per time.

3–2. The answer is lower. Metabolic clearance rate includes all routes of elimination; its value is the renal clearance plus any others.

3–3. The answer is C. \( C_{in} = U_{in} V/P_{in} \). When \( P_{in} \) increases, there is no change in \( C_{in} \) because \( U_{in} \) rises by an identical amount. In other words, the mass of inulin filtered and excreted increases, but the volume of plasma supplying this inulin (ie, completely cleared of inulin) is unaltered.

3–4. Substance \( A \) could be (1) poorly filtered at the glomerulus, (2) taken up and degraded by the tubular epithelium, or (3) reabsorbed and returned to the blood.
3–5. Substance B is secreted by the tubules.
3–6. PAH; creatinine; inulin; urea; sodium; glucose
3–7. The answer is 0.01. The excreted amount of sodium is as follows: 75 mmol/L × 2 mL/min × 0.001 L/mL = 0.15 mmol/min. Because sodium is freely filtered, the filtered amount is the GFR × P_{Na}, and GFR is the same as C_{in}, GFR = C_{in} = 50 mg/L ÷ 1 mg/L × 2 mL/min = 100 ml/min, or 0.1 L/min. Filtered sodium = 150 mmol/L × 0.1 L/min = 15 mmol/min. Therefore, the fraction of the filtered load that is excreted, fractional excretion, is 0.15 mmol/min ÷ 15 mmol/min = 0.01.
3–8. The answer is A. With only one-fifth of the excretory capacity remaining, the plasma concentration will rise by a factor of 5 to achieve the same excretion.

CHAPTER 4

4–1. The answer is false. Flux by a uniporter is always passive, down the electrochemical gradient. Flux by a symporter may be active depending on direction. Flux via an ATPase is always active. (Theoretically, it could pump downhill, but this does not normally occur.)
4–2. Most regions of the nephron have tight junctions that are far less leaky than those found in the proximal tubule, and most apical membranes are far less permeable to water. As a result, it is possible to sustain much larger osmotic gradients across the epithelium in tubular regions beyond the proximal tubule. Reabsorption beyond the proximal tubule is generally not iso-osmotic.
4–3. The answer is true. The tubular epithelium is quite permeable to many (but not all small solutes), but the peritubular endothelium is even more permeable.
4–4. Failure to move fluid from the interstitium to peritubular capillary as a result of low plasma oncotic pressure quickly leads to a backup of fluid in the interstitial space. Since the interstitial space contains a small fraction of total volume of the kidneys, only a small increase in fluid volume is required to generate a rise in hydrostatic pressure. Once interstitial pressure rises significantly, this drives an increasing back-leak.
4–5. The answer is false. Not all solutes are alike osmotically. Proteins, eg, exert far more osmotic effect mole for mole than do saccharides. In addition, saccharides exert a somewhat higher osmotic effect than simple salts. In this text, we sometimes simplify things by using osmolarity, when technically we should use osmolality. In most cases, this does not introduce a large error.
4–6. Despite the volume flow, most solutes are close to diffusional equilibrium between plasma and interstitium. If the interstitium starts to become depleted of a substance as a result of secretion, net diffusion from the plasma will soon replenish it.
The answer is B. Normally, all the filtered glucose is reabsorbed, meaning that the filtered load does not saturate the transporter capacity. If either A or C was true, there would be at least some glucose in the urine.

Gating is a process that changes the probability that a channel is open. Changes in interstitial osmolality are not known to gate channels directly. However, if a change in interstitial osmolality causes a cell to swell or shrink, the resulting change in mechanical stretch of mechano-sensitive channels could gate them (this is thought to be how hypothalamic cells detect changes in plasma osmolality).

The answers are A, B, and C. These waste products are all normally excreted in large amounts; a decreased GFR would cause their plasma concentrations to increase until the filtered load was increased enough to reestablish normal excretion. In contrast, the reabsorption $T_m$'s for glucose, amino acids, and many other organic compounds that are not waste products are usually so high relative to normal filtered loads that, even with a 50% loss of nephrons, virtually all the filtered loads are reabsorbed. Accordingly, their plasma concentrations are virtually independent of renal function (i.e., the kidneys do not participate in the setting of their plasma concentrations).

The answer is no. The overall tubular handling of urea is reabsorption. Urinary urea concentration is higher than that of plasma because relatively more water has been reabsorbed than urea, thereby concentrating the urea in the tubular fluid leaving the kidney.

This is simply the GFR (180 L/day) times the concentration in the filtrate: $180 \text{ L/day} \times 0.005 \text{ g/dL} \times 10 \text{ dL/L} = 9 \text{ g/day}$.

The answer is A. The luminal membranes of nephron segments beyond the proximal tubule do not express glucose transporters, so no further reabsorption occurs regardless of conditions.

Decreased pH (acidify the urine). Weak bases like quinine become protonated, and thus less permeable, at low pH. This would prevent its passive reabsorption and increase its excretion.

(a) About 50%; (b) about 100% (due to secretion in the thin limbs); (c) about 50%
6–3. The answer is both. The luminal fluid and the medullary interstitium would both become isosmotic, and so would the final urine.

6–4. The answer is true. Allowing back-leak of chloride would reduce the paracellular reabsorption of sodium, and thus reduce the overall reabsorption of both sodium and chloride and prevent concentrating the medullary interstitium.

6–5. The answer is false. There is no reabsorption of sodium or chloride by the descending thin limb of Henle’s loop.

6–6. The answer is true. There is always net reabsorption of both sodium and water in the medulla from long loops of Henle and medullary collecting ducts. Under the influence of ADH, the reabsorption of water in the inner medullary collecting ducts is further stimulated. This solute and water must be removed from the medullary interstitium.

6–7. The answer is true. As with maximum levels of ADH, there is always net reabsorption of sodium and water in the medulla. With minimum levels of ADH, there is still net reabsorption of water in the inner medullary (not cortical) collecting ducts. As described in the text, with low levels of ADH (and hence little reabsorption in the cortex), the gradient for water reabsorption in the inner medulla is quite large, and there is always some finite water permeability in this region.

6–8. It ceases completely. Even though the active step is not altered by the drug, there will be no sodium entering the cell to be acted on by the pumps.

6–9. The answer is false. The medullary osmolality reaches its highest value only during extreme dehydration. The normal value is somewhat below the maximum, and fluctuates throughout the day in response to dietary input.

CHAPTER 7

7–1. (1) 5 mmol/min. Approximately two-thirds of filtered sodium is reabsorbed by the proximal tubule. (2) 6.6 mmol/min. Filtered sodium rises from 15 to 20 mmol/min. Glomerulotubular balance maintains fractional sodium reabsorption at approximately two-thirds of the filtered load.

7–2. The answer is no. As soon as the person starts to become sodium deficient as a result of the increased sodium excretion, the usual sodium-retaining reflexes will be set into motion. They will, of course, be unable to raise aldosterone secretion, but they will lower GFR and alter the other factors that influence tubular sodium reabsorption to compensate at least partially for the decreased aldosterone-dependent sodium reabsorption.

7–3. The answer is no. It will probably be slightly above normal because of increased filtration fraction (ie, reduction in GFR and an even greater reduction in renal blood flow secondary to renal arteriolar constriction mediated by the renal sympathetic nerve and angiotensin II).
7–4. The right kidney will have increased secretion because of decreased renal perfusion pressure acting via the intrarenal baroreceptor and decreased flow to the macula densa. This increased secretion will result in elevated systemic arterial angiotensin II and elevated arterial blood pressure, both of which will inhibit renin secretion from the left kidney.

7–5. Plasma renin concentration is lower. The increased aldosterone causes the body to retain sodium, which reflexively inhibits renin secretion. Thus, one observes high plasma aldosterone and low plasma renin, a strong tip-off to the presence of the disease because, in almost all other situations, renin and aldosterone change in the same direction (because the renin-angiotensin system is the major control of aldosterone secretion).

7–6. The answer is true. Although some diuretic drugs are more potent than others, blockage at any site results in at least mild diuresis. Because less than 2% of the filtered load is normally excreted, it does not require a huge reduction in the percentage reabsorbed to result in a large increase in the amount of sodium that is excreted.

7–7. It will decrease. This question focuses on the effect of peritubular factors on proximal sodium reabsorption. Although the GFR will remain about the same, the renal blood flow increases. The peritubular capillary pressure will, therefore, rise. At the same time, the peritubular oncotic pressure will decrease because of the decreased filtration fraction. Both of these effects tend to reduce fluid reabsorption from the interstitium, which reduces proximal sodium and water reabsorption.

7–8. There will be excess sodium. We cannot be sure of the net effect on water because there are opposing influences: the excess sodium leading to increased water excretion and the ADH-like effect leading to decreased water excretion. The osmolality, for sure, will be hyperosmotic.

7–9. There will be excess sodium, excess water, and an iso-osmotic urine. Blocking sodium reabsorption in the thick ascending limb is what “loop diuretics” do. Not only do they lead to excess sodium and water in the urine, but they prevent the kidneys from generating a medullary osmotic gradient. Even with the ADH-like actions of the drug, the urine cannot become more concentrated than the now iso-osmotic medullary interstitium.

7–10. Both B and D are correct.

CHAPTER 8

8–1. The answer is C: potassium secretion.

8–2. For potassium, yes. High rates of secretion may exceed reabsorption. For sodium, no.

8–3. A. False: Most potassium reabsorption is paracellular, but all sodium reabsorption is transcellular. B. True. C. False: Even though the multiporter moves equal amounts of sodium and potassium, most of the potassium leaks back and is recycled.
8–4. The answer is true. The excess solute retains water, thus diluting tubular potassium and reducing the driving force for reabsorption.

8–5. The answer is false. The high amounts of nonreabsorbed solute increase the sodium content of the luminal fluid, with water accompanying it. This dilutes potassium. This stimulates potassium secretion both by the dilution effect and by the high rate of sodium reabsorption.

8–6. The answer is high. The increased aldosterone stimulates potassium secretion and, thereby, excretion. Moreover, once enough sodium has been retained to increase GFR and to cause partial inhibition of proximal reabsorption, the increased delivery of fluid to the cortical collecting duct further enhances potassium secretion. There is no potassium escape similar to the sodium escape from aldosterone.

8–7. The answer is relatively normal. One may have answered “high,” assuming that the increased aldosterone would stimulate potassium secretion, as in Question 8–6. However, this effect is more than balanced by the fact that the patient has a decrease in flow of fluid into the cortical collecting duct (because of decreased GFR and increased proximal and loop reabsorption). Recall that potassium secretion is impaired when the amount of fluid flowing through the cortical collecting duct is reduced. This explains why patients with the diseases of secondary hyperaldosteronism with edema do not lose large quantities of potassium, whereas those with primary hyperaldosteronism do.

8–8. The answer is D. There is little control over potassium transport before the connecting tubule. When excretion is high, the major process leading to high excretion is high secretion by the connecting tubule and cortical collecting duct.

8–9. The answer is B. While insulin exerts many actions, its chief action in limiting the rise in plasma potassium due to a high-potassium meal is to stimulate tissue uptake, particularly by skeletal muscle.

CHAPTER 9

9–1. The answer is true.

9–2. The answer is B. If the urine has a pH greater than 7.4, clearly there is no titratable acid excreted. Indeed, there is negative titratable acid excretion. Ammonium does not contribute to titratable acid and may be ignored in the calculation of titratable acid.

9–3. None of them are acid loads. Vomiting of stomach acid adds bicarbonate to the blood. Fruit juice, when oxidized to CO₂ and water, adds bicarbonate. Sweetening it makes no difference because the metabolism of saccharides is acid-base neutral. Lactate, when metabolized, adds bicarbonate.

9–4. The answer is false. Filtered bicarbonate is not transported into the epithelial cells; rather, it is converted in the lumen to CO₂ and water when it combines with secreted protons. Bicarbonate is generated within
the epithelium on a one-for-one basis as the secreted protons, and this bicarbonate is transported across the basolateral membrane.

9–5. The answer is false. Renal acid excretion rises. The acidosis is caused by increased generation of metabolic acids, not failure of the kidneys to excrete acid. The kidneys respond by increasing their excretion of acid. A steady state is reached when input and output are both elevated and plasma bicarbonate is low.

9–6. The answer is C. The pH values are both within the normal range, so both patients could be perfectly normal. However, pH is set by the ratio of bicarbonate to Pco₂. Both values could be elevated or depressed, yielding a normal ratio but an acidosis or alkalosis that is well compensated.

9–7. Answer A is certainly true. This a respiratory acidosis. Answer B is false. In response to the prolonged acidosis, the patient’s bicarbonate would be elevated as compensation, not depressed. Answer C is false. Although the development of compensation would require increased acid excretion, the maintenance would only require a urinary excretion to match the input of fixed acid. CO₂, no matter how much it is elevated, cannot be excreted as urinary acid.

9–8. The answer is true. The excretion of titratable acid and ammonium both involve secreting a hydrogen ion and generating a bicarbonate that goes into the blood.

CHAPTER 10

10–1. The answer is C. Vitamin D probably acts in a permissive manner for all of these processes, but stimulation of calcium absorption from the GI tract is the most important direct action.

10–2. The answer is false. PTH inhibits phosphate reabsorption, resulting in increased excretion. The increased excretion is the same result that would occur if secretion were increased, but phosphate is not secreted.

10–3. The answer is B: bone.

10–4. The answer is D. PTH stimulates calcium reabsorption in the distal tubule; thus, a reduction in PTH reduces reabsorption and permits more excretion.

10–5. The answer is A. Chronic renal failure reduces phosphate excretion. As a consequence, levels of PTH increase, causing excessive resorption of calcium from bone.
### Table A–1. Summary of reabsorption and secretion by major tubular segments

<table>
<thead>
<tr>
<th></th>
<th>Proximal tubule</th>
<th>Henle's loop</th>
<th>Distal convoluted tubule</th>
<th>Collecting duct system</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>R</td>
<td>S</td>
<td>R</td>
<td>S</td>
</tr>
<tr>
<td>Organic nutrients</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urea</td>
<td>X</td>
<td>(X)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Proteins, peptides</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phosphate</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sulfate</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Organic anions</td>
<td>X</td>
<td>(can also be reabsorbed and/or secreted passively along tubule)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Organic cations</td>
<td>X</td>
<td>(can also be reabsorbed and/or secreted passively along tubule)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urate</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sodium</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Chloride</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Water</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Potassium</td>
<td>X</td>
<td>X</td>
<td>(X)</td>
<td>X</td>
</tr>
<tr>
<td>Hydrogen ions</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Bicarbonate</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Ammonium</td>
<td>X</td>
<td>(X)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcium</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
</tbody>
</table>

R, reabsorption; S, secretion.
### Table A–2. Major functions of the various collecting-duct cells

<table>
<thead>
<tr>
<th>Principal cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Reabsorb sodium (stimulated by aldosterone)</td>
</tr>
<tr>
<td>2 Secret potassium (stimulated by several signals, including aldosterone)</td>
</tr>
<tr>
<td>3 Reabsorb water (stimulated by antidiuretic hormone)</td>
</tr>
<tr>
<td>Comment: Processes 1 and 2 are linked by a basolateral membrane Na-K-ATPase.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Type A intercalated cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Secrete hydrogen ions, which effect reabsorption of bicarbonate and excretion</td>
</tr>
<tr>
<td>2 Reabsorb potassium</td>
</tr>
<tr>
<td>Comment: These 2 processes are linked by a luminal membrane H-K-ATPase.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Type B intercalated cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Reabsorb chloride (? stimulated by chloride depletion)</td>
</tr>
<tr>
<td>2 Secrete bicarbonate (stimulated by increased extracellular pH)</td>
</tr>
<tr>
<td>Comment: These 2 processes are linked by a luminal membrane Cl-bicarbonate</td>
</tr>
<tr>
<td>countertransporter.</td>
</tr>
</tbody>
</table>

* Functions of the inner medullary collecting-duct cells are not presented. Only the most important physiological regulators are given.
### Table B–1. Classes of diuretics

<table>
<thead>
<tr>
<th>Class</th>
<th>Mechanism</th>
<th>Major site affected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbonic anhydrase</td>
<td>Inhibit secretion of hydrogen ions, which causes less reabsorption of bicarbonate and sodium</td>
<td>Proximal tubule</td>
</tr>
<tr>
<td>Loop diuretics</td>
<td>Inhibit Na, K, 2Cl cotransporter in luminal membrane</td>
<td>Thick ascending limb of Henle's loop</td>
</tr>
<tr>
<td>Thiazides</td>
<td>Inhibit Na,Cl cotransporter in luminal membrane</td>
<td>Distal convoluted tubule</td>
</tr>
<tr>
<td>Potassium-sparing diuretics*</td>
<td>Inhibit action of aldosterone</td>
<td>Cortical collecting duct</td>
</tr>
<tr>
<td></td>
<td>Block sodium channels in Collecting-duct system luminal membrane</td>
<td></td>
</tr>
</tbody>
</table>

*Except for this category, diuretics increase potassium excretion as well as sodium excretion (see text for discussion of the reasons for this increase). Aldosterone antagonists do not increase potassium excretion because they inhibit aldosterone's stimulation of potassium secretion. The sodium channel blockers also inhibit potassium secretion, in this case by reducing the amount of sodium entering the collecting duct cell for transport across the basolateral membrane by the Na-K-ATPase pumps; this reduces the activity of the pumps and, hence, the active transport of potassium into the cell.
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