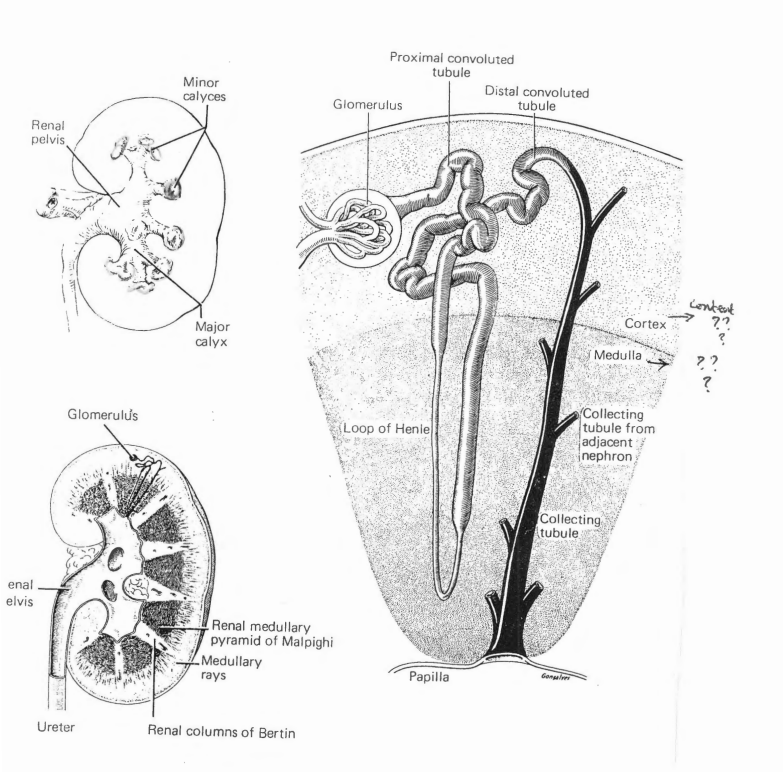
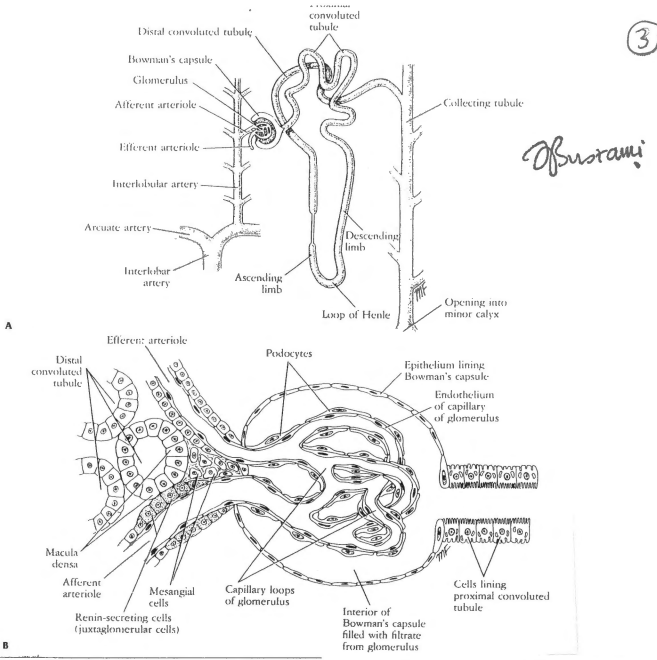


* The fresh Kidney can easily be divided into a dark reddish-brown outer cortex and a lighter-coloured inner medulla. * The medulla -> is composed of about a dozen renal Pyramids, each with its base oriented toward the cortex and its apex (the renal papilla) projecting into a minor * The cortex -> Extends into the medulla between adjacent Pyramids as the renal columns. * Extending from the bases of the renal Pyramids into the correx are Striations Known as medullary rays (400-500 in number; Each consists of a straight collecting tubule into which the distal convoluted pubules of many neighbouring nephrons empty their contents through arched collecting tubules. * A RENAL LOBE ?? may be defined as a renal Pyramid together with the cortical tissue overlying its base and lying along its sides * A Renal lobule ?? is a medullary ray and the

* If renal counter, a sleeve of nephrons draining into these associated tubules (a sleeve of nephrons draining into these tubules) and is reparated from its neighbour by the interlobular arteries.





Udniferous Tubules

The kidney is composed of large numbers of microscopic units called *uriniferous tubules*. Each tubule is composed of two functional regions, the *nephron*, which produces an excretion known as urine, and

the collecting tubule, which concentrates the urine and conveys it to the calyces (Fig. 13-3).

Nephron

There are over a million nephrons in one kidney. Each consists of four distinct parts: (1) the renal corpuscle, which contains the glomerulus, (2) the proximal convoluted tubule, (3) the loop of Henle, and (4) the distal convoluted tubule (see Fig. 13-3). The parts of the nephron form a continuous tubule that measures about 50 mm in length and runs from the cortex to the medulla and then returns to the

RENAL CORPUSCLE. The *renal corpuscle* is situated in the cortex. It is formed by the upper end of the uriniferous tubule, which is expanded into a structure called a *Bowman's capsule* (Figs. 13-4-13-7; see Fig. 13-3). The renal corpuscle contains the glomerulus, which is a network of capillaries into which blood enters by an *afferent arteriole* and leaves through a smaller *efferent arteriole*.

The glomerulus indents the wall of the Bowman's capsule as a fist might press into the side of a balloon (Fig. 13-8). The epithelial cells that form the wall of the Bowman's capsule also serve as a covering for the glomerulus. The renal corpuscle thus consists of the Bowman's capsule and the glomerulus (see Figs. 13-4-13-7).

The outer wall of the Bowman's capsule is lined with simple squamous epithelium that abruptly

changes into cuboidal epithelium at the start of the proximal convoluted tubule. Where the capsular wall is reflected onto the glomerulus, the squamous cells change into star-shaped cells with multiple processes. These cells, called *podocytes*

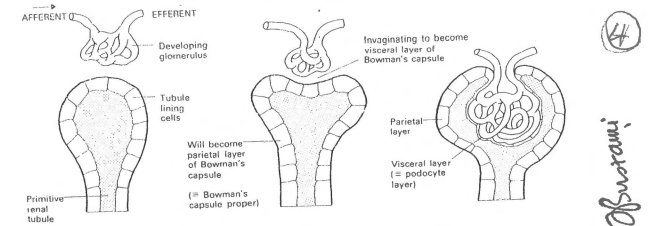
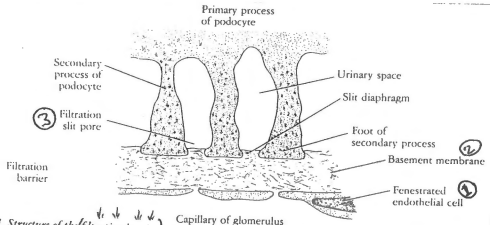
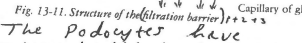


Fig. 16.8 Development of the renal corpuscle

This diagram illustrates in a highly schematic manner the mode of development of the renal corpuscle. The nephrons develop from the embryological metanephros as blind-ended tubules consisting of a single layer of cuboidal epithelium. The ends of the tubules dilate and become invaginated by a tiny mass of tissue which differentiates to form the glomerulus. The layer of invaginated epithelium flattens and differentiates into podocytes which become closely applied to the surface of the knot of glomerular capillaries. The intervening connective tissue disappears so that the basement membrane of glomerular endothelial cells and podocytes effectively fuse forming the glomerular basement membrane. A small amount of connective tissue nevertheless remains to support the capillary loops and differentiates to form the mesangium. Where the mesangium stretches between the capillary loops, its surface is directly invested by podocyte cytoplasm with podocyte basement membrane lying between the two. When examining ultra-thin light microscope specimens as in Figure 16.11 and electron micrographs as in Figure 16.14, the podocytes, endothelial cells and mesangium are identified most easily by tracing out the podocyte and endothelial cell basement membranes.





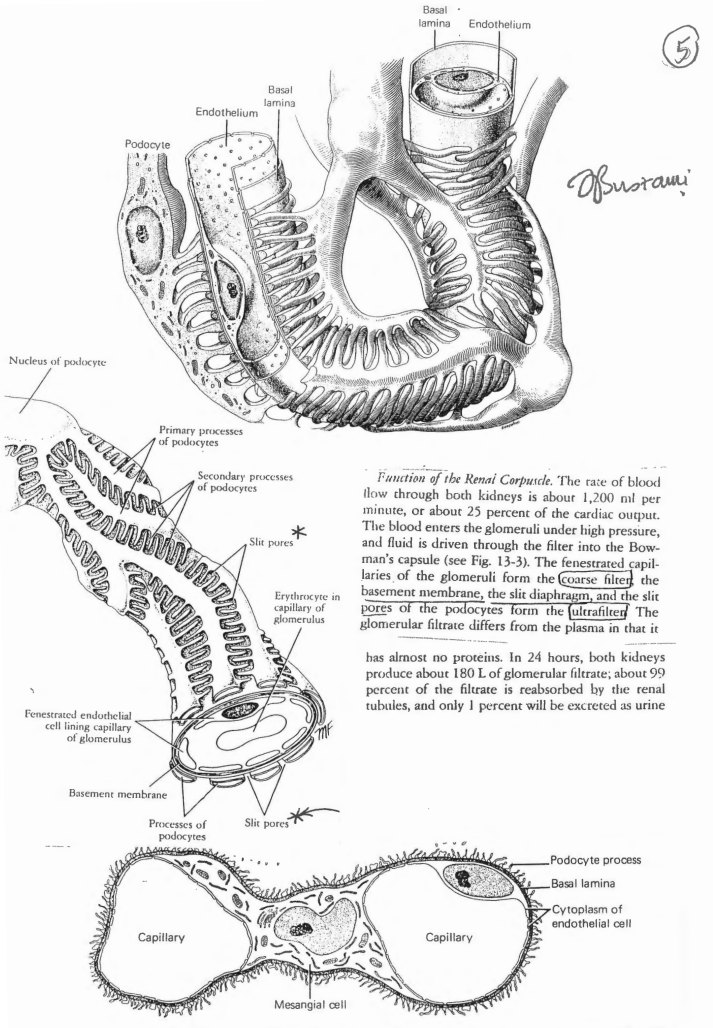
primary processes that tightly clasp the glomerular capillaries (Figs. 13-9 and 13-10). From the primary processes, smaller secondary processes arise that interdigitate with the secondary processes of other podocytes. This arrangement leaves small slitlike gaps between the processes that measure about 25 nm

across and are called *slit pores* (Fig. 13-11). The secondary processes end in *feet* that are applied firmly to the basement membrane of the capillary wall of the glomerulus. Extending across the slit pores between adjacent feet is a thin *slit diaphragm* about 6 nm thick (Fig. 13-12).

The blood in the glomerular capillaries is separated from the cavity of the Bowman's capsule by: (1) the fenestrated endothelial cells lining the capillaries (Fig. 13-13), (2) a thick basement membrane (Fig. 13-14), and (3) the slit pores of the podocytes. Together these structures are known as the *filtration barrier* (see Fig. 13-11). The holes, or fenestrae, in the endothelial cells permit the passage of plasma but hold back the cells of the blood. The smaller molecules of the plasma readily pass through the basement membrane and the slit diaphragm of the podocytes to enter the cavity of the Bowman's capsule. Particles with a molecular weight greater than 160,000 are held back by the slit diaphragm. The plasma protein albumin, which has a molecular weight of 69,000, would be expected to pass through without difficulty. We know, however, that in a normal individual, it does not. The probable explanation is that the filtration mechanism is blocked by proteins with larger molecules and that the electric charge on the filter repels the albumin molecules. The fluid that finally crosses the filtration barrier and enters the capsular space is called the glomerular filtrate.

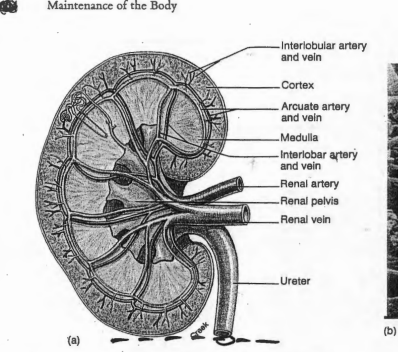
Lying between the glomerular capillaries are small groups of star-shaped cells that are contractile and capable of phagocytosis. These cells are called *mesangial cells* (see Fig. 13-3) and <u>support the capillary</u> walls by producing intercellular substance. They are also thought to remove by phagocytosis any mac-

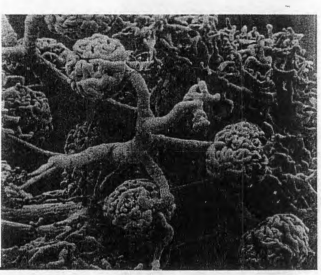
romolecules that escape from the capillaries into the tissue space.



Mesangial cells of glomerular capillaries. They are located between 2 capillary lumens, enveloped by the basal lamina

Renal artery -> anterior & Posterior divisions, arteries Each (segmental) -> 5 segmental artery divides into (LOBAR) arteries (Usually One for each pyramid) = Each lubar artery divides into 2-3 (INTERLOBAR) arteries (which run on each Pyramid) => At the corticomedullary the Side d2 interlobar arteries divide dichotomously Junction the which Arch over the bases ARCUATE) arteries into Pyramids The arcuste arteries give off R the Which IN TERLOBULAR run Fidinely Arteries of (AFFERENT GLOMERULAR ARTERioLES cortex giving the ARTERIOLES divides -SOON > the EFFERENT GLOMULLAR Peritubular capillary plexus around tubules distal convoluted the Proximal of Efferent arteriale Glomerulus Capsule the after of justice medullary Proximal convoluted Afferent arterio tubule Distal convoluted glomerule, Efferent tubule enters a pyramid of Loop of Henle. Collecting duct-Cortex divides into 12-24 Vasa lecta Glomerulus Interlobular arteries breaks up to form Distal convolutedcapillary plexus around Arcuate arterv tubule loops of Hende of Interlobar artery Collecting ducts and vein Vasa recta Medulla At the venous end Loop of Henle Renal column the c. plexus gives rise to ascending vasa Collecting tubule (duct) Peritubular capillarias recta Descending vasa recta fibrous Papilla (arterioles) (7) ascending aprule Vasa Hetta (Venules) - afferent arteriale (4) artery interlabular form the basis 06 arcuste artery Counter current exchange multiplier system Do NOT anastomose with each other . (interlobar arteries)





Kessel and Kardon

Figure 19.7 The vascular structure of the kidneys. (a) An illustration of the major arterial supply and (b) a scanning electron micrograph of the glomeruli.

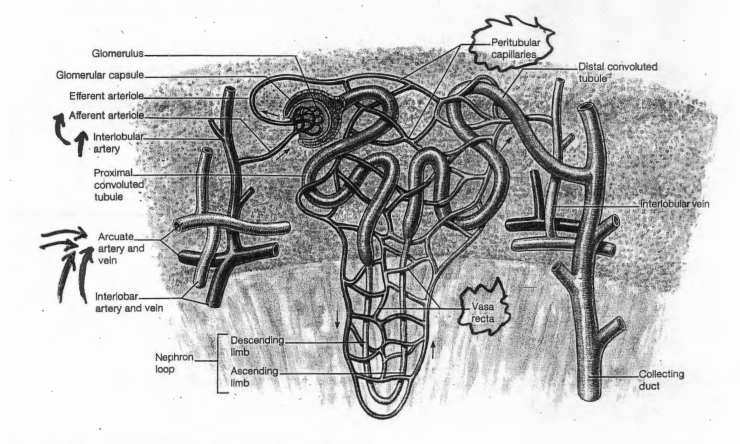
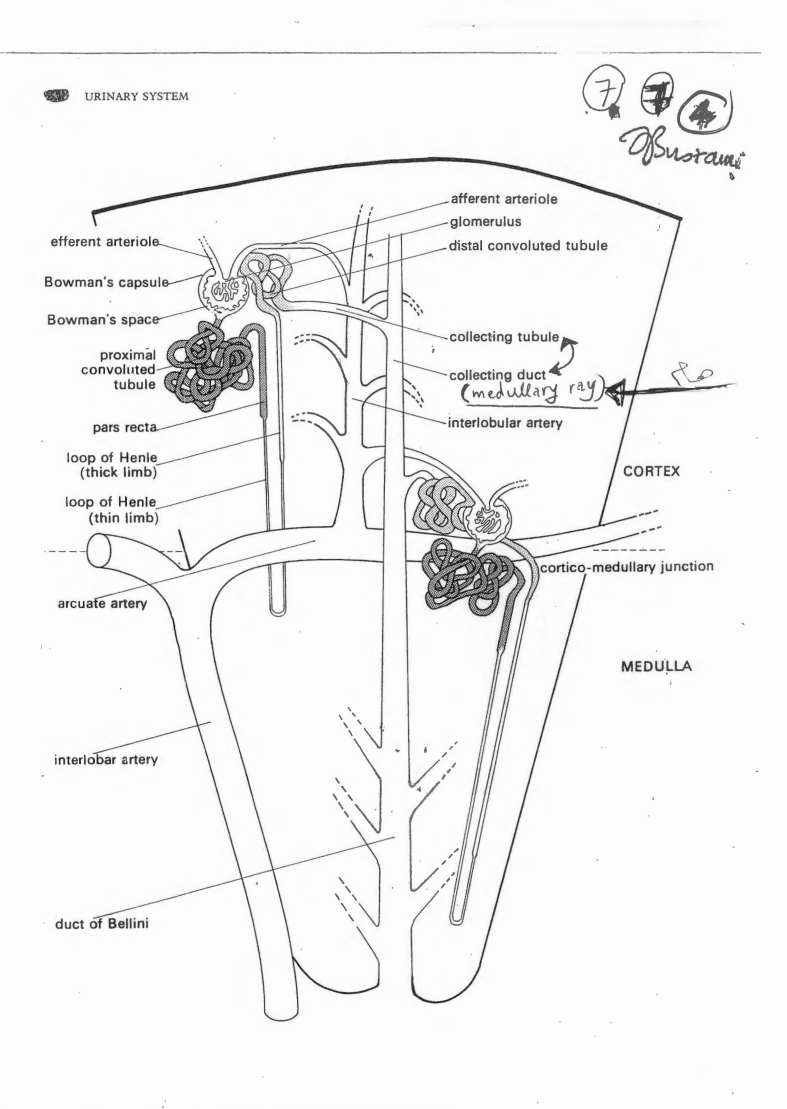


Figure 19.8 A simplified illustration of blood flow from a glomerulus to an efferent arteriole, to the peritubular capillaries, to the venous drainage of the kidneys.

Maintenance of the Body



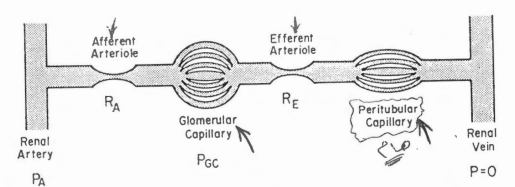


FIG. 4-11. Locations of the two arteriolar resistances and the glomerulus.

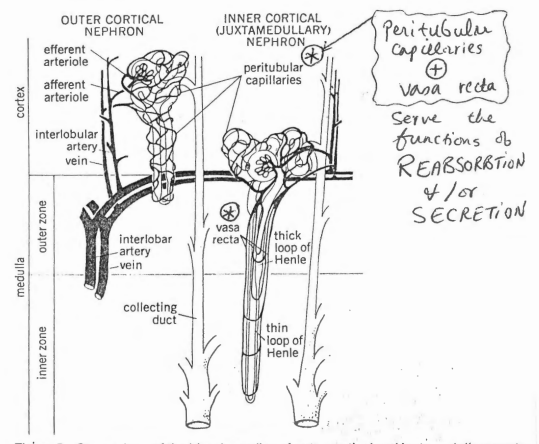
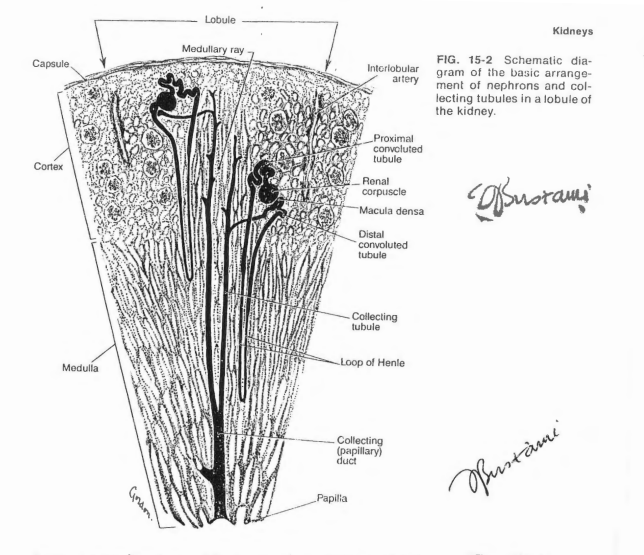


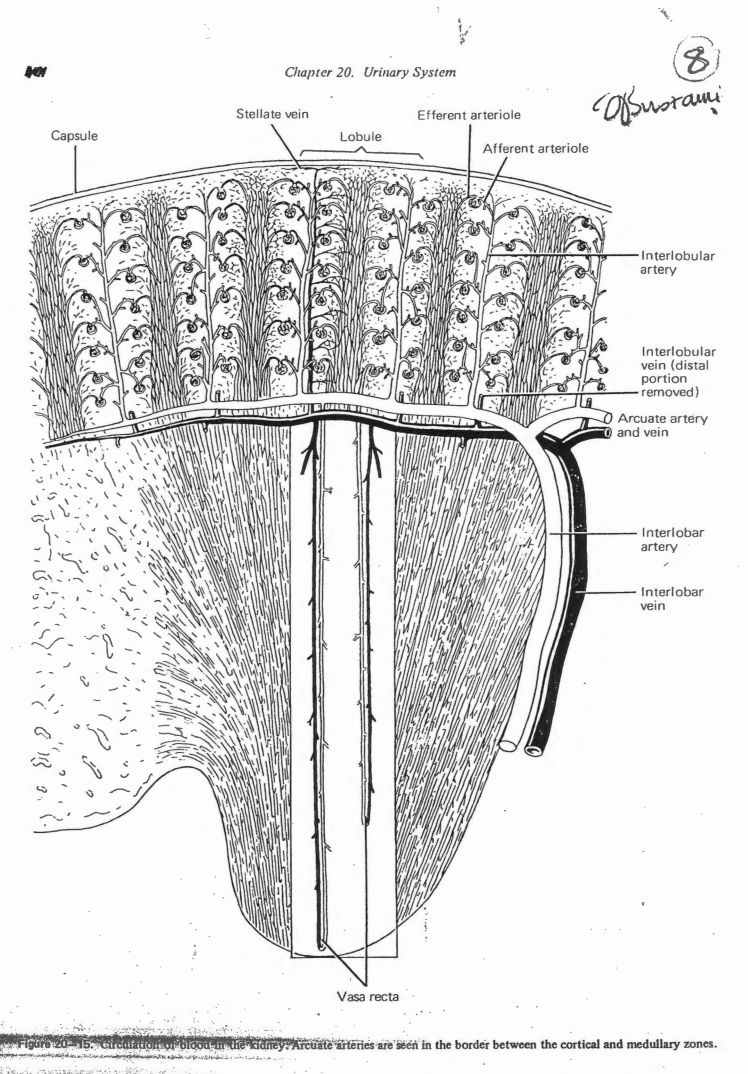
Figure 5 Comparison of the blood supplies of outer cortical and juxtamedullary nephrons. This figure is misleading in one regard in that it suggests that the peritubular capillaries of a tubule are derived only from the efferent arteriole leaving that tubule's glomerulus of origin; In fact, they are derived from efferent arterioles coming from* many glomeruly. (Redrawn from R. F. Pitts, Physiology of the Kidney and Body Fluids,

Notice the presence of 2 arteriales Afferent (larger) efferent ", ", ", ", ", ", Sets of Capillaries glomerular capillaries "Peritubular" & Vasa recta play very important role in makin to in relation to - Important role Secretion eabsurphin

Renal blood flow Bustam Blood entering the Kidneys Passes through f (2) Capillary beds in series > The lower capilling Because of the high glomerular capillary Pressure in the Peritubular capillaries results in only Fressure > only Reabsorption occurring at the plasma filtration Peritubular capillaries occurs out the glomerular capillaries The Vasa recta arise from the Juxtamedullary glomeruli allowing a small amount 5% of renal blood flow to perfuse the renal medulla Total renal blood flow Glomerular capillaries about (100)ml/min averages Afferent arteriole Efferent arteriole is Plasma of it Peritubular Renal plasma flow is Bowman's capsule capillary So approximately 625 ml/min the <u>plasma</u> entering the Kidney is About 20%. 07 at the renal glomerulus a glomerular filtered) Filtration rate & (GFR) of (125ml/min) - Between 80% and (99%) of the glomerular filtration is reabsurped so the final Urinary flow rate varies between 0.4 milimin to 20 ml/min. and usually averages about (I ml/min

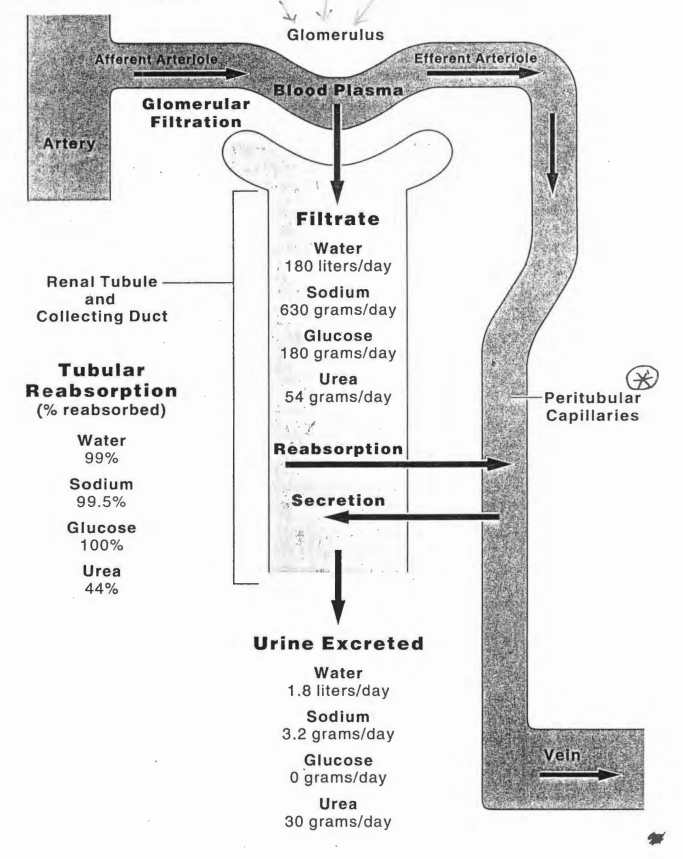


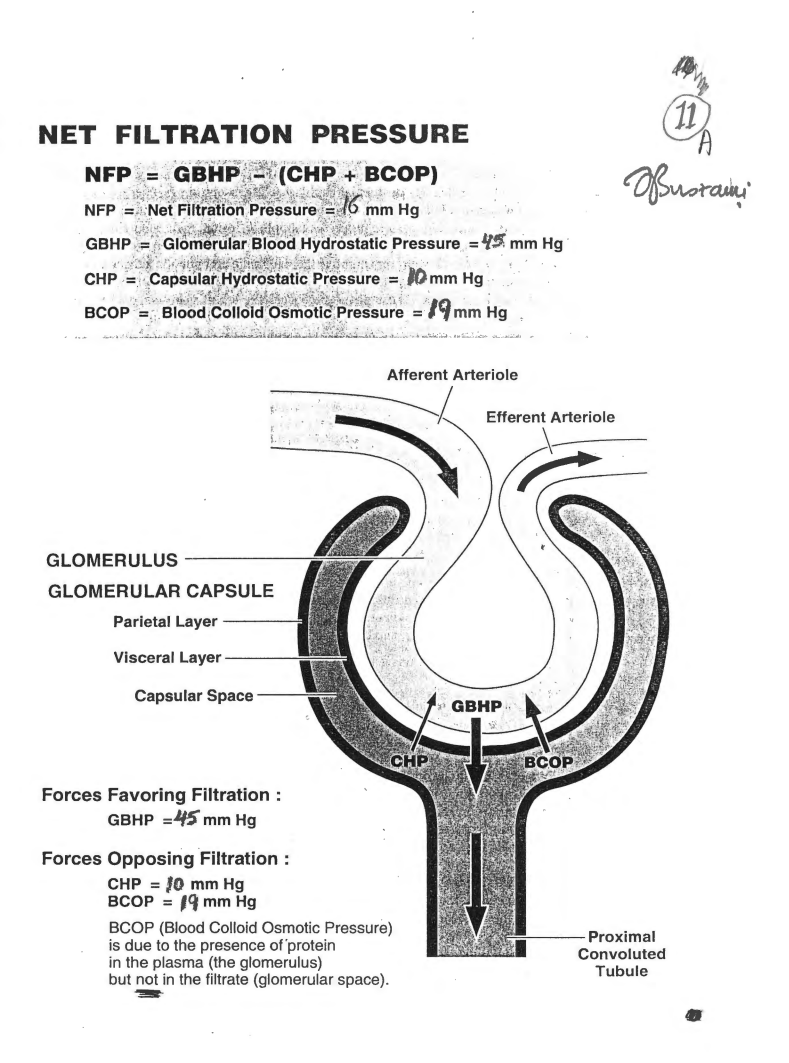
Renal terminology is imprecise and confusing. The structural unit of a kidney is a lobule. This has a central core of collecting tubules - the medullary ray of the cortex, surrounded by a sleeve of nephrons draining into these There is no line of demarcation between lobules. tubules. As the medullary rays approach the renal sinus, space between them gets Less, there is no further space for the sleeve of nephrons and cortex changes to medulla. The merging of the medullary rays form the pyramids and the pyramids in turn merge to form the prominent papillae. The nephrons near the surface have short loops of Henle and are referred to as Cortical nephrons. Those nephrons lying deeply. at the bottom of the nephron sleeve are near the medulia, have long loops of Henle and are referred to as juxta-medullary The the short loops of the Cortical nephrons do not reach nephrons. into the medulla. The long loops of the juxta-medullary nephrons run into the medulla parallel to the collecting ducts and in association with the vasa recta (100)



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URINE FORMATION Diagrammatic





Bustam

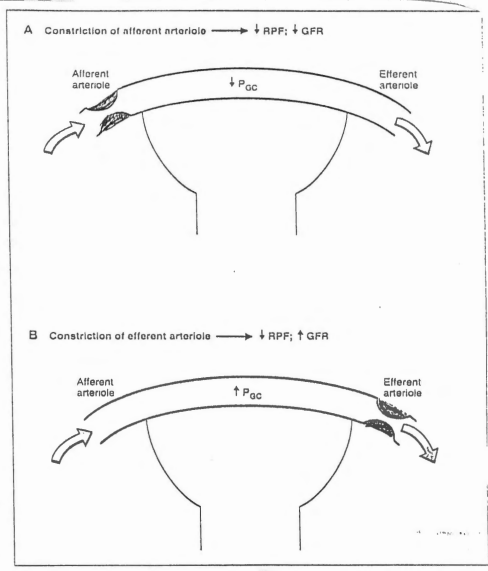


FIGURE 6-11. Effects of constricting afferent (A) and efferent (B) arterioles on renal plasma flow (RPF) and glomerular filtration rate (GFR). $P_{r,C}$, hydrostatic pressure in the glomerular capillary.

TABLE 6-5. Effect of Changes in Starling Forces on RPF, GFR, and the Filtration Fraction

Effect	RPF	GFR	Filtration Fraction (GFR/RPF)
Constriction of afferent arteriole	Ţ	Ţ	N.C.
Constriction of ellerent arteriole	Ţ	Î	Î
Increased plasma protein concentration	N.C.	Ļ	Ţ
Decreased plasma protein concentration	N.C.	Ŷ	î
Constriction of the ureter	N.C.	Ļ	1

GFR, glomerular filtration rate; N.C., no change; RPF, renal plasma flow.

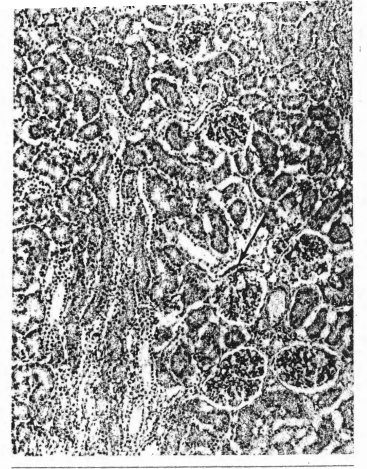


Fig. 13-6. Photomicrograph of the cortex of the kidney, showing several glomeruli and proximal and distal convoluted tubules. Note a macula densa (arrow). (H&E; ×100.)



Fig. 13-17. Photomicrograph showing many proximal convoluted tubules cut in oblique and cross sections. Note that each tubule is lined with cuboidal epithelium and the cytoplasm stains strongly with eosin because of the many mitochondria (not shown). The nuclei are centrally placed, and the luminal cell surfaces have indistinct brush borders formed of microvilli. Three distal convoluted tubules are also present (D). Note that the cytoplasm of the cuboidal cells lining the distal convoluted tubules stains lighter with eosin. (H&E; $\times 400.$)

A Proximal convoluted tubule

Distal convoluted tubule

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Proximal convoluted tubules 1. Most common tubules found in the cortex 2. Have Stellate-shaped lumen bounded by a distinct brush-border 3. The cells are mainly cuboidal or low columnar in shape and have indistinct lateral cell boundaries 4. Not all cells of a given tubule show & nuclear Profile due to the large size of the cells 5. The cytoplasm Stains intensely with cosin (due to the large number of mitochondria within the cell). 6. PAS-positive basal lamine is seen around the proximal

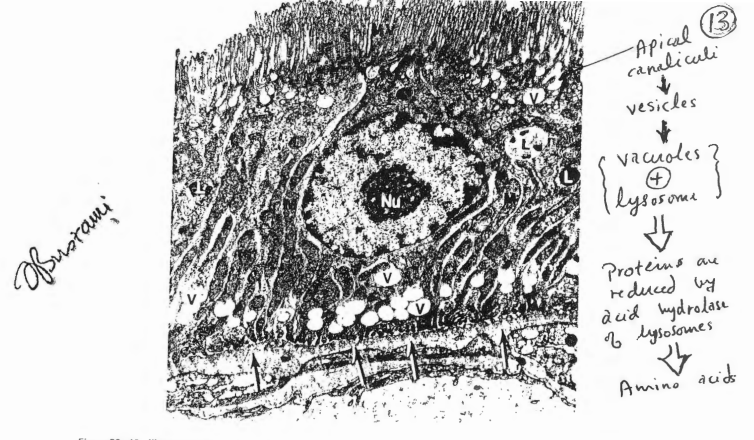
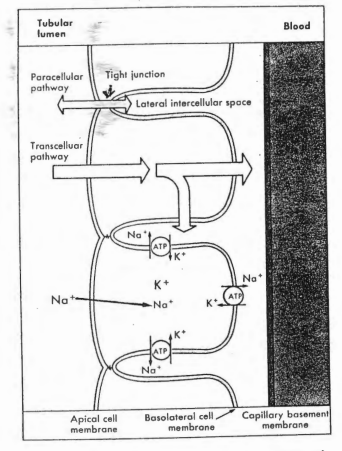
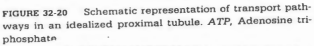


Figure 20-10. Electron micrograph of a proximal convoluted tubule wall. Observe the microvilli (MV), the hysosomes (L), the vacuole (V), the nucleolus (Nu), and the mitochondria (M). The arrows point to the basal lamina, \times 10,500.

Electron microscopic appearance of PCT a) a Golgi apparatus on the apical side of the nucleus b) numerous rod-like mitochondria in the basal cytoplasm c) The plasma membrane, especially on the base of the cell, show much INFOLDING and INTERDIGITATING with neighbouring cells d) The microvilli are long and densely pecked at the apex of the cell e) there are small clefts between the bases of the microvilli => APICAL CANALICULI -> give rise to a series of small vericles -> Coalesce to form larger vacuoles (D Endocytic Complex) Apical canaliculi? involved in Vacuales Strotein absorption (D) Vacuales condense and firse with hysosomes, the acid hydralaser of which reduce the absorped protein to its constituents amino acids which are then released into the blood Stream. + Other functions of proximal convoluted tubule - Absorption = H20 (65% of glomerular filtrate), Na, cl, glucose, amino acids, vit

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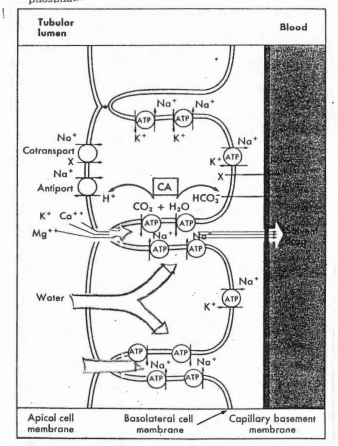
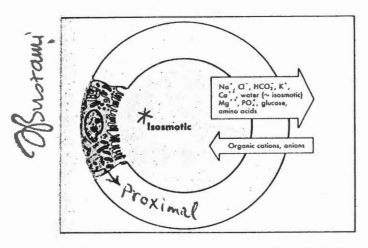


FIGURE 32-22 Schematic representation of the proximal tubule. For the Na⁺-X co-transport protein, X represents either glucose, amino acids, phosphate, chloride, or lactate. CO_2 and H_2O combine inside the cells to form H⁺ and HCO₃ in a reaction facilitated by the enzyme carbonic anhydrase (CA). ATP, Adenosine triphosphate.



NGURE 32-21 Schematic representation of a cell in the munal tubule and the primary transport characteristics. Abular fluid is isosmotic.

 \mathfrak{H}^*) secretion, via the Na⁺-H⁺ antiporter, results in bicarbonate reabsorption (Figure 32-22; see Chapter 34). The Na⁺ that enters the cell across the apical membrane leaves the cell across the basolateral membiane via the Na⁺, K⁺-ATPase. The other solutes that enter the cell with Na⁺ exit across the basolateral membrane down their electrochemical gradients.

The reabsorption of Na⁺ and the other solutes just described increases the osmolality of the lateral intertellular space. Because the lateral intercellular space is slightly hyperosmotic (\sim 3 mOsm/kg H₂O) with respect to tubular fluid, and because the proximal tubule is highly permeable to water, water will flow by ismosis across both the tight junctions and the prox-

imal tubular cells into this hyperosmotic compartment (Figure 32-22). Accumulation of fluid within the lateral intercellular space increases the hydrostatic pressure in this compartment and thereby drives fluid into the capillaries. Thus **water reabsorption follows solute transport.** The reabsorbed fluid is essentially *isosmotic* to plasma. '

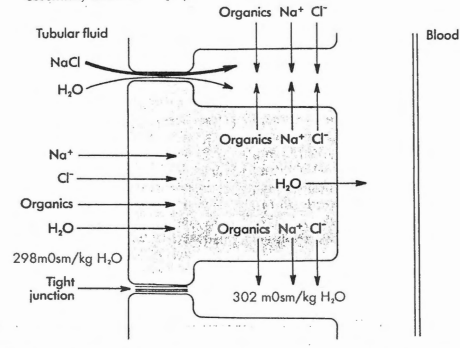


FIGURE 36-5 Routes of water reabsorption across the proximal tubule. Transport of Na⁺, Cl⁻, and organic solutes into the lateral intercellular space increases the osmolality of this compartment, which establishes the driving force for osmotic water reabsorption across the proximal tubule. An important consequence of osmotic water flow across the proximal tubule is that some solutes, especially K⁺, Ca⁺⁺, and Mg⁺⁺, are entrained in the reabsorbed fluid and are thereby reabsorbed by the process of solvent drag.

The second phase of proximal tubular reabsorption involves the reabsorption of Na⁺ with Cl⁻ in the second half of the proximal tubule. This occurs because in the first half of the proximal tubule, Na⁺ is reabsorbed with bicarbonate as the primary accompanying anion, leaving behind a solution that becomes enriched in Cl⁻. The rise in Cl⁻ concentration in the tubular fluid creates a gradient that favors the diffusion of Cl⁻ from the tubular lumen across the tight junctions and into the lateral intercellular space. Movement of the negatively charged chloride ions attracts the positively charged sodium ions. Thus, in the second half of the proximal tubule, some Na⁺ and Cl⁻ are reabsorbed across the tight junctions by passive diffusion.

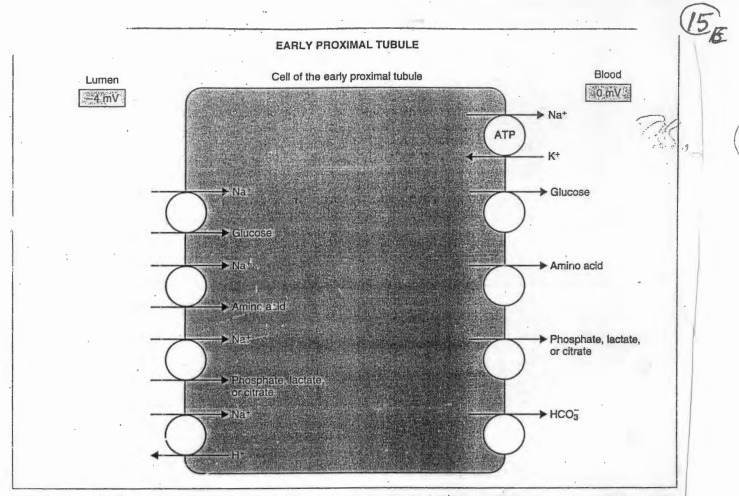
Sodium and chloride reabsorption by the second half of the proximal tubule also occurs by a transcellular route. The pathway for Na^+ and Cl^- transport across the apical membrane is unknown

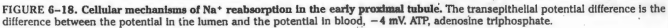
15k Renal tubular epithelial cells Can pransport solutes & water from one side of the tutule to the other Reabsorphion Secretion held together by fight junctions of separated by Intercellular spaces Secretion , across cells - Transcellular Reabsorption between cells -> Paracellular Nethway Nat Reabsorption by transcellar raturnay depends on the operation of Nat-Kt-ATPase 2-Step Process -> Movement àcross àpial membrane (down) an electrochemical gradient established by the Nat-Kt_ATPASE movement across the basalatural membrane against an electrochamical gradient vie +L Nat-K' ATPase writer Proximal tubule Reabsorbs (67%. Nat Kt ALL glucose anino acids (Key clement in reabsorption) > Nat-Kt-ATPase In 1st half > Na + + glucose + aminoacids + Bicarbonne In 2nd half > Na + + CL

During the 1st phase Nat entry into the cell across apical membrane mediated by - Specific transport Proteins (Not by simple diffusion) + Couple movement of Nat with movement of other solutes * Each transp. Protein -> USES the Potential energy released by downhill movement B NAT to POWER the uphill movement of other solutes Antipaters Co-transport protein Nat-Ht antipater (Symporteri) Nat aminorid Nat Aminorid Nat Phosphare Nat CLits searchion results in HCO3T reabsorption leave down their elecnochemical basolat- memb. gradient by Nat-Kt-ATPase * Reabsorption of Nat & other Solutes. Tosmolality of the loten space when will flow by Osmosis across both the tight junctions & apical membrane - tubulan - Accumulation of fluid 2. A hydrostetic pressure within the lateral intercellular in this compartment - Absorped Aluid Strosmotic Driver fluid into the apillances

2nd phase of proximal tubular Reabsonption Reabsonphion of Nat with CLT in 2nd half of prox. tubuk In 1st half of 3-> Nat Reabsorped & HCOS leaves behind a Solution Rich in CL-- Kise of Cl - concentration in tubular fluid CREATES A GRADIENT that favours the diffusion of cl- from tubular lumen ACROSS TIGHT junctions into the lateral intercellular sona Movement & regatively charged Cl-attracts the postitively a Nat -> Mat & Cl- reabsaption by 2nd half of proximal tubuke also occurs by transcellular route -> Mathway is unknown

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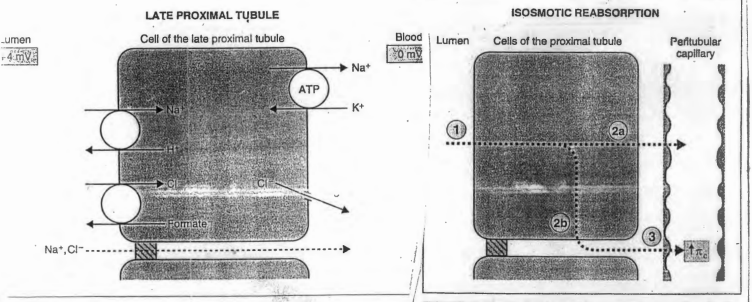
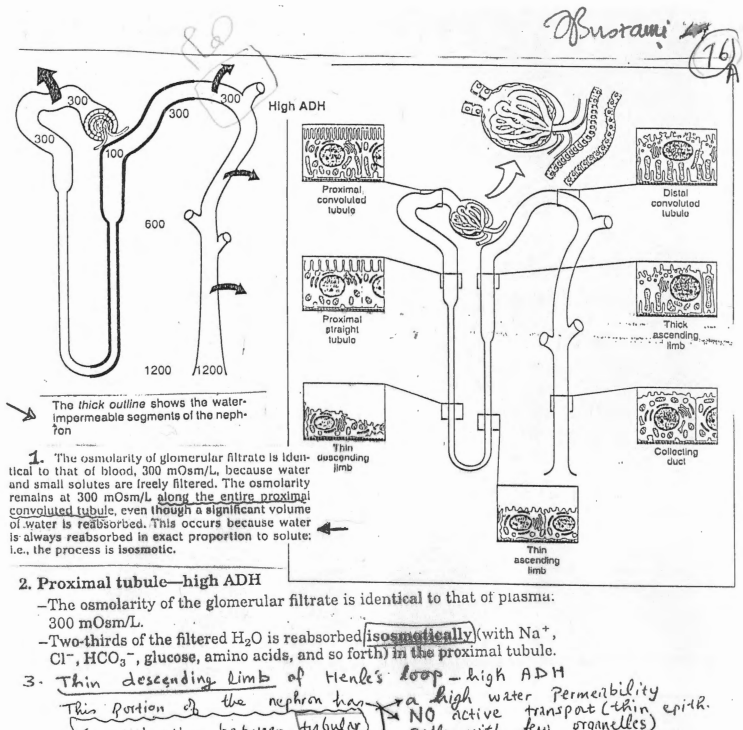


FIGURE 6-20. Mechanism of isosmotic reabsorption in the proximal tubule. Dashed arrows show the pathways for reabsorption; circled numbers correspond to the text. π_{e} , peritubular capillary colloid osmotic pressure.

Bustani



Hubular organelles between few cells with Equilibration rend interstitium Permeabilit very low Solute flirid of the extraction by H20 takes place Nacl increases to , mM 1600 The concentration of the nephron at the base of the loop (the fluid soutside contains (300 mM NAC) of 600 mM ured) in the tensel, interstitium Progressively pertonic + -> tubular fluid becomes Henles loopf+Impermeable to Hzok ascending limb oh 4. Thin Shighly permeable to Nacl that reaches this tubular fluid contains > 600 mm Nace while the surrounding The rephron do the 300 mM Nach - Nach PASSively pation contains renil interstitium interstitium -> 1H20 Cannot follow renal into the Ases. the the rubule becomes HYPOTONICK relative TU inside fluid interstitium

12

Proximal convoluted tubule Distal.convoluted Glomerulus tubule 60810000 00000 0/070 000 Thin segment of loop of Henle Proximal convoluted tubule Distal convoluted tubule 100 Cortex 0 0 0 0 3 Medulla 000 Ø 3 Operstance D Collecting Loop of Henle tubule from O 3 adjacent nephron Collecting Collecting tubule The thin limb of the Loop of Henle (Segment) The terminal straight portion of the PCT suddenly changes to the descending thin limb. With the light microscope, it is usually difficult to distinguish the difference between thin limbs and blood capillaries, even when they are side by side, unless the capillaries contain red cells. When empty, the cytoplasm of the capillaries is slightly thinner Papilla than that of the cells lining the thin limbs, while the nuclei of the thin limb cells are slightly more prominent in that they bulge into the lumen. The difference is quite marked on examination with the EM, since the cytoplasm of the cells of the thin limb not only have microvilli on their surfaces, but are at least twice as thick as those of the capillaries. The (nuclei) appear almost uniformly round, while those of the capillaries are usually oval or irregular in shape. interposed between the proximal and distal convoluted tubules of has descending and The loop 02 Henle ascending limbs which lie together inside the Renal medulla (close to the Vasa recta & the collecting tubules) + thin descending limb (structure)) -> quite long in juxtamedullary rephrons the cells show thigh water permenbility -> Reabsorp water from the rubular fluid (equilibration between the tubular fluid at the surrounding renal interstitium takes place by water extraction) > (hypertonic tubular fluid) (10) > thin ascending limb (structure as above) Impermeable to Highly permeable to Nacl ~ Nacl partively diffuses into the renal interstitium but the cannot follow ~ (hypotonic tubular fluid) for thick ascending limb - impermeable to H20) Concentrating Segment Cotransport & Nat- K+ - 2 clsegment diluting (more hypotonic tubular fluid) (10 Similar in structure to early distal tubule (Lined by cosinophilic cuboidal

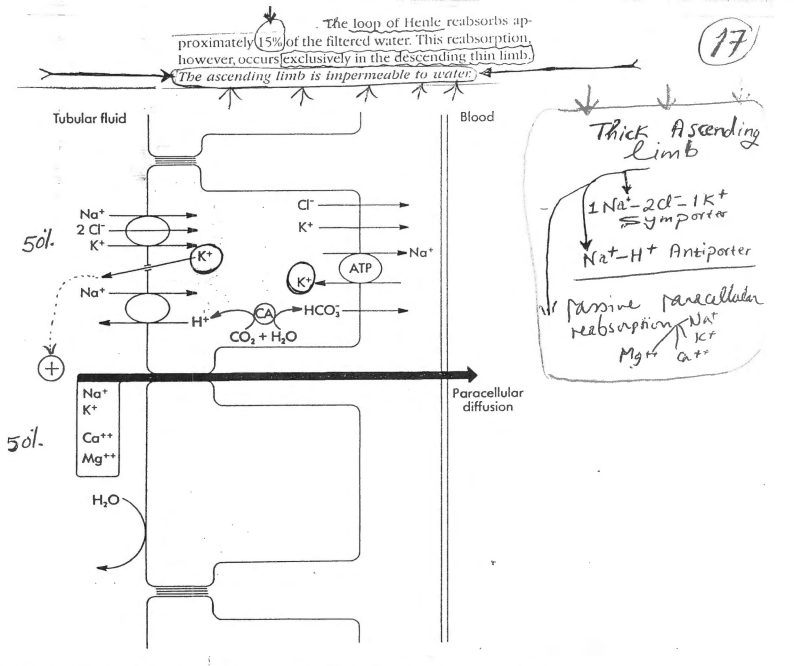


FIGURE 36-7 Transport mechanisms for NaCl reabsorption in the thick ascending limb of Henle's loop. The lumen positive transepithelial voltage results from the diffusion of K^+ from the cell into the tubular fluid, and plays a major role in driving passive paracellular reabsorption of cations.

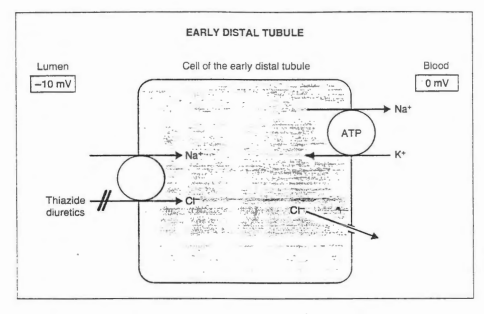
The key element in solute reabsorption by the thick scending limb is the Na⁺-K⁺-ATPase pump in the basoateral membrane (Figure 36-7). As with reabsorption in he proximal tubule, the reabsorption of every solute by he thick ascending limb is linked to the Na⁺-K⁺-ATPase pump. The operation of the Na⁺-K⁺-ATPase pump mainains a low cell [Na⁺]. This low [Na⁺] provides a favorble chemical gradient for the movement of Na⁺ from he tubular fluid into the cell. The movement of Na⁺ cross the apical membrane into the cell is mediated by he 1Na⁺-2Cl⁻-1K⁺ symporter, which couples the movenent of 1Na⁺ with 2Cl⁻ and 1K⁺. This symport protein ses the potential energy released by the downhill movenent of Na⁺ and Cl⁻ to drive the uphill movement of . ⁺ into the cell. An Na⁺-H⁺ antiporter in the apical cell nembrane also mediates Na⁺ reabsorption as well as H⁺ ecretion (HCO, reabsorption) in the thick ascending mb (Figure 36-7). Na⁺ leaves the cell across the basoiteral membrane via the Na⁺-K⁺-ATPase pump, and K⁺, 1, and HCO, leave the cell across the basolateral iembrane by separate pathways

The voltage across the thick ascending limb is positive in the tubular fluid relative to the blood because of the unique location of transport proteins in the apical and basolateral membranes. The important points to recognize are that increased salt transport by the thick ascending limb increases the magnitude of the positive voltage in the lumen, and that this voltage is an important driving force for the reabsorption of several cations, including Na⁺, K⁺, Ca⁺⁺, and Mg⁺⁺, across the



Because the thick ascending limb is very impermeable to water, reabsorption of NaCl and other solutes reduces the osmolality of tubular fluid to less than 150 mOsm/kg H_2O . \rightarrow H γPO

Distal convoluted tubule (DCT) consists of 3 Parts: O Early DCT -> the continuation of the thick segment of Henles loop and has the same histological Structure (lined by Essinophilic Cuboidal epithelium) > Reabsorbs 5%. of the filtered Nat (Nat-cl-cotransporter at the luminal membrane) > impermeable to the clike the thick segment) > called the cortical DILUTING segment (2) The macula densa: Columnu closely Packed cells - may function to Distal convoluted tubule and ascending thick limb of Henle's loop Sense < Nat ?? Concentration in DET - part of J-9 apparatus Oppustani (3) The late or convoluted Pornion: Can be distinguished from PCT by the following Criteria -> 4 The lumen of the DCT is generally WiDER 2) The cells are Shorter and lighter staining 3) Nuclear Profiles are usually Seen in each cell (in part because) many are binucleate) (4) a brush border is lacking Anatomically & functionally the late distal tubule & collecting ducts (tubules) one similar -> 2 major cell types interspersed along these segments: - Principle cells (Light cells) involved in (3% of filtered Nat) - principle cells (dark cells) have VERY DISTINCT - intercalated cells (dark cells) CELL BOUNDARIES K+ secretion involved in the readsorphion (in low dietary k+ content) No. & mitochondria H+ secretion



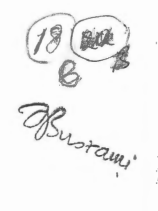
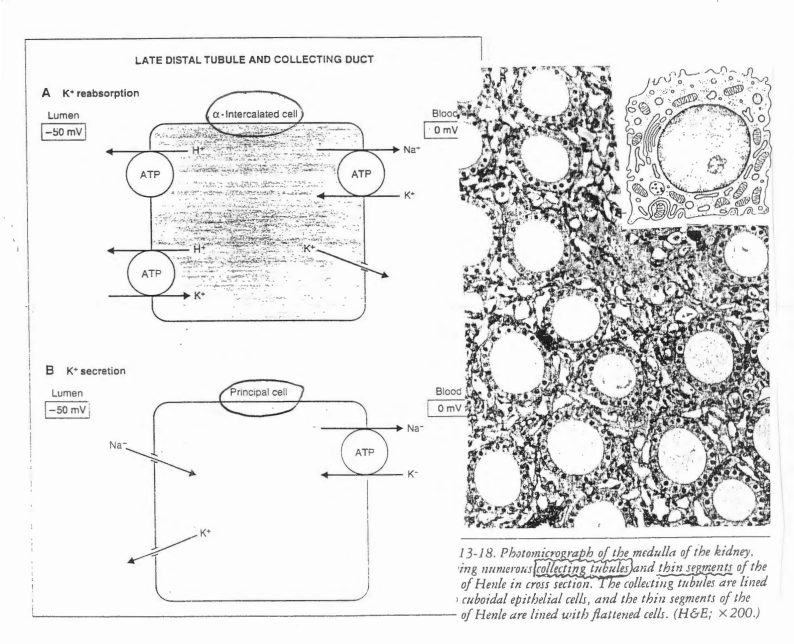


FIGURE 6-23. Cellular mechanism of Na⁺ reabsorption in the early distal tubule. The transepithelial potential difference is -10 mV. ATP, adenosine triphosphate.



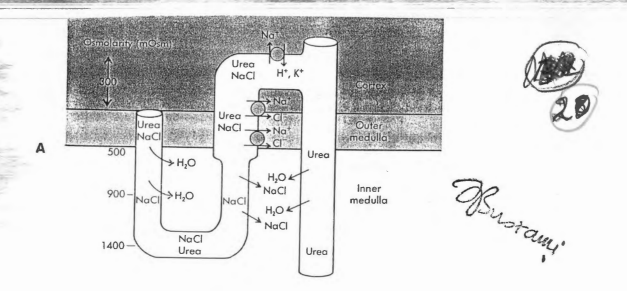


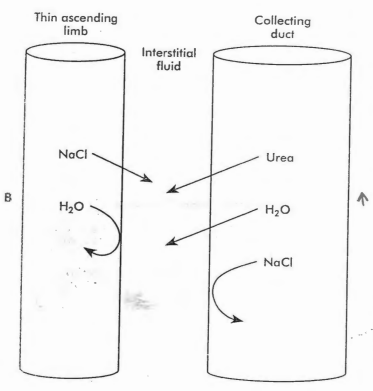
The mechanism for Nat reabsorption in the Principle cells of the late distal tubule + collecting duct lumen Blood 0 mu 50 mu. The Ulminal membrane of the >Nn+ Nat F Principle cells contains Nat channels Principle (ATP > Nat diffuses through these channels -Kt Cell down its electrochemical gradient from the lumen into the cell -> Late distal Nat then is extruded from the tubule of cell viz the Nat-K+ ATPase in the Colle ching basolateral membrane duct ALDOSTERONE acts directly on the Principle cells to >1 Nat reabsorphion A Kt secretion * Aldosterone increases Nat reabsorption in the principle cells by inducing synthesis of the luminal membrane Nat channels & the baschkeral membrane Nat- Kt ATPase > 1 Nat entry into the cell + provides more Nat to Nat- Kt ATPase more Nat is pumped out of the cell + 1 more Kt primped into the cell - A intracellular Kt concentration -> 1 the driving force for K+ secretion from the cell into the lumen Collecting Tubules (a) is the most distal part of the Uriniferous tubule and is NOT part of the nephron (b) Each DCT of a nephron becomes continuous with a collecting tubule that runs a short arched course and. ENTERS & MEDULCARY RAX Here a number of short collecting fubules joing a main collecting tubule as side We presence or absence of ADM) tributaries. collecting tubule then The main passes down in the medullary ray to enter the medullary pyramid When the collecting tubules reach the (Inner Zone) of the pyramid, group of them join at scute angles to form Straigh Papillary ducts that open on the apex of the renal papella into a minor calyx * Chigh unca permeability in the presence The cells lining the collecting tubules are at first CUBOIDAL, later in the straight papillary duces they are TALL COLUMNAR

S(d) The cell borders are regular with few interdigitations S(d) The nuclei are dark staining but the cytoplasm is pake Staining because there are relatively few cytoplasmic organelles (P) on the apex of the renal papilla, the columnar epithelium changes to the transitional papilla, the columnar epithelium Ehanges to the transitional epith. Lining the minor caby x. Functions The collecting tubules (ducts) function in the Conservation B Water and the Production of hypertonic Urine. As the ducts pass through the medulla to the tips of the papillare, they pass through the INCREASINGLY HYPERTONIC ENVIRONMENT ESTABLISHED AND MAINTAINED BY THE LOOPS OF HENLE. The Permedbility of collecting ducts to water is controlled by antidiurchic hormone (ADH). In the Presence of this hormone, the collecting ducts become Permeable to water which is drawn from the tubules (ducts) by OSMOSIS as the result of the hypertonic environment maintained in the medullary interstitium. The Loss of worker from the tribules(ducti) results in a concentrated hypertonic usine. In the absence of ADH -> the Kidney cannot concentrate or form hypertonic usine. This condition is known as Didbetes insipidus (production of large amounts of dilute arise -> severe dehydration of the individual.

How does the kidney produce Urine that is more concentrated than blood & what determines how high the Urine Osmolarity will be ?? Remember the [4 Pareners] within the RENAL MEDULLA Collecting tubules of ducts (Loop of Henle) (Vasa I - Urine becomes hyperosmotic, in the Presence of AD+1 - As the tubular fluid flows down the collecting tubules + ducts -> it is exposed to interstinal fluid with increasingly hyperosmolarity (i.e the corticopanillary osmotic gradient 300 mosm/L. > 600 > 900 > 1200) > water will be reassurped until the tubular fluid equilibrates Osmotically with surrounding interstitial fluid > The final Urine osmularity, in the presence of ADH will be equal to the osmolarity at the bend of the loop of Henle (1200 mOSm/L.)

Nat reabsorption H20 reabsorption Proximal C. 651.~67/= 65% tubule (2/3)(2/3) (PCT) Thin Idescending 15% Х Segment (impermet ble to Na+) of Henles loop Ascending limb of 25% Henle's thin (impermeable to FI20) distal convoluted 10%. 5% tubule (early & ADH needed in } late) Collecting hubbles 4% 91. ADH needed (of duct is a major solute of tubular fluid in the thin ascending limb, Nacl Remember a is a major solute of tubular fluid of the medullary collecting duct the collecting duct is more the thin ascending limb is more permeable to permeable to thea than to Nacl than to med Nacl thea gradient across the Nacl gradient across the collecting duct thin ascending limb both gradients THICK ASCENDING LIMB





Currently the most plausible hypothesis is the two-solute hypothesis (see Figure 19-18). This hypothesis builds on the finding that, along with NaCl, urea makes up a large fraction of the total solute of the medullary interstitial fluid. The high concentrations of NaCl and urea in the medullary interstitial fluid result because (1) NaCl is the major solute of tubular fluid in the thin ascending limb, and urea is a major solute in the tubular fluid of the medullary collecting duct; and (2) the thin ascending limb is more permeable to NaCl than to urea, and the collecting duct is more permeable to urea than to NaCl.

In summary, two driving forces are at work in the two-solute hypothesis (Figure 19-18, B): the MaCl gradient across the thin ascending limb and the urea gradient across the collecting duct. Both of these gradients are created by the active reabsorption of NaCl by the thick ascending limb. Both gradients drive solute into the medullary interstitial

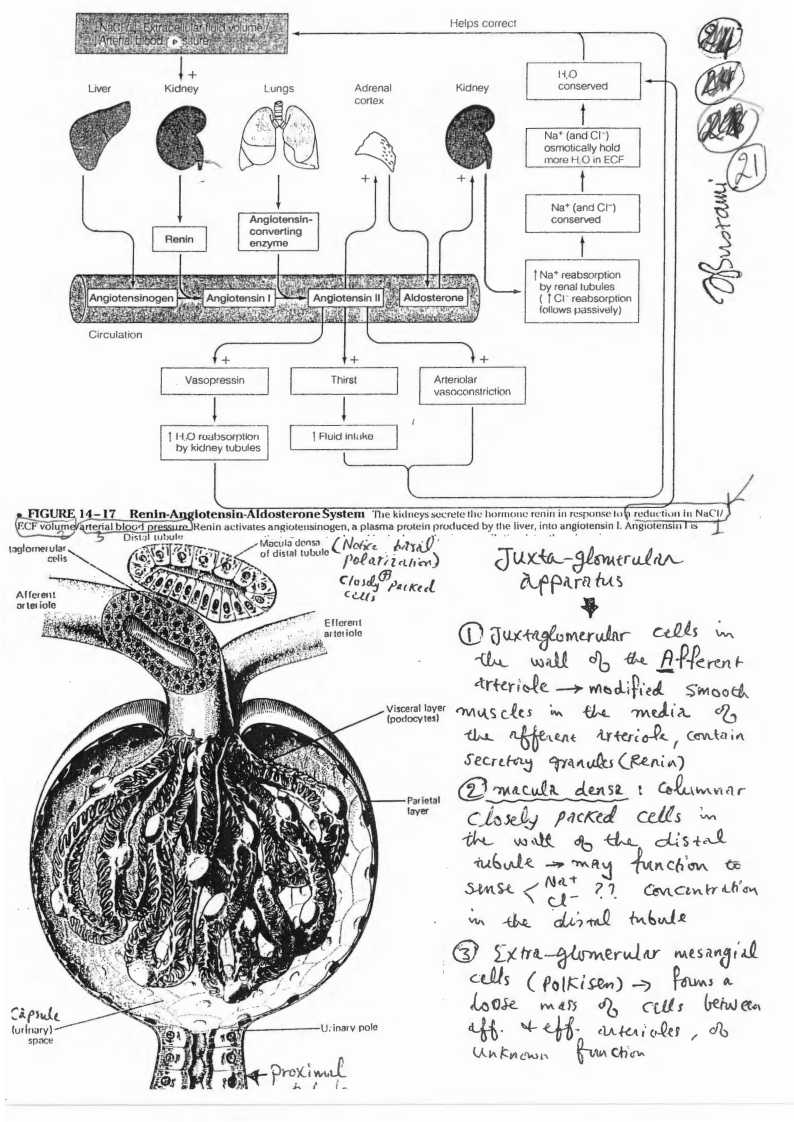
FIGURE 19-18

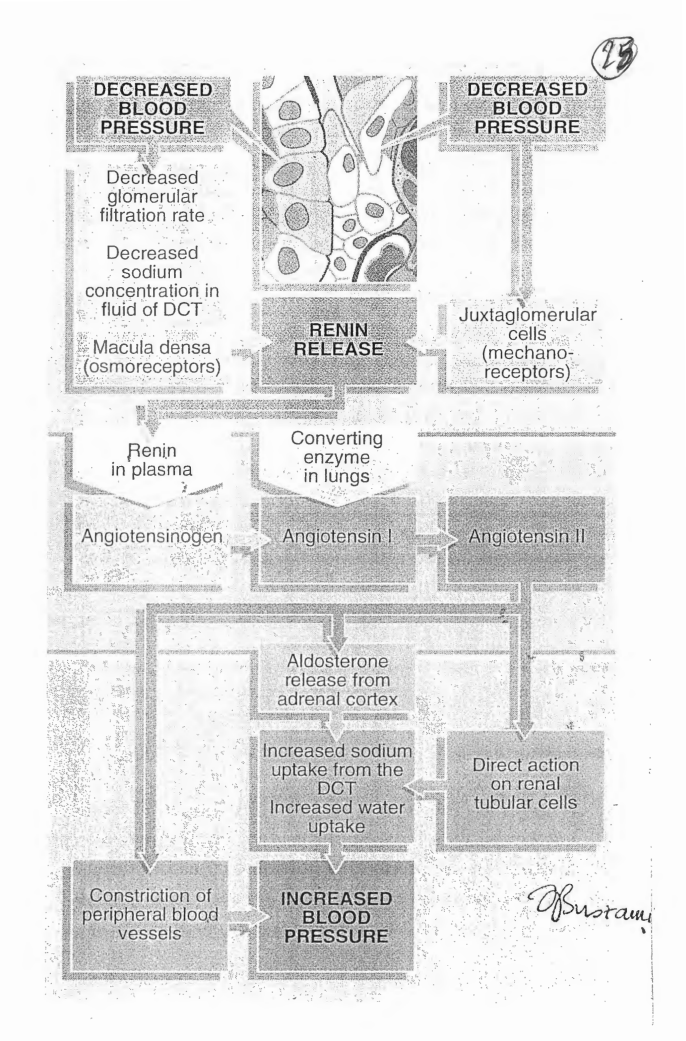
Mechanism of formation of concentrated urine according to the two-solute hypothesis.

Overall view of the loop of Henle, distal tubule and collecting duct: The osmolarity of the interstitial fluid at different levels of the medulla is shown on the scale at the left. The tubular fluid leaving the proximal tubule is isotonic. As the tubular fluid travels through the descending limb of the loop of Henle, water leaves the descending limb, drawn by the increasing osmotic pressure of interstitial fluid in the medulla. As a result, the tubular fluid in the descending limb becomes progressively more concentrated. As the tubular fluid passes through the thin ascending limb, NaCl, but not water, diffuses out, so that the osmotic pressure of interstitial fluid de s. In the thick ascending limb, more salt is removed by active reabsorption. The tubular fluid entering the distal tubule is more dilute than plasma with respect to NaCl, while urea has been concentrated by the reabsorption of water. Urea and water diffuse down their concentration gradients as tubular fluid passes through the collecting duct. The remaining solutes in the tubular fluid are concentrated further by the water reabsorption, and a urine as concentrated as the interstitial fluid at the innermost part of the medulla may be formed if ADH levels are high. If ADH levels are low, a final urine similar to the dilute urine in the distal tubule is excreted.

B The two driving forces that generate a high solute concentration in the medullary interstitial fluid are the NaCl gradient between ISF and thin ascending limb, and the urea gradient between collecting duct and ISF. Water cannot leave the thin ascending limb in response to the osmotic gradient, but can be reabsorbed from the collecting duct in the presence of antidiuretic hormone.

fluid, concentrating both NaCl and urea in the interstitial fluid. The high osmotic concentration of solute in the medulla provides the driving force for water recovery from the medullary collecting duct.





2. Thin Ascending limb impermeable to H20 of Henles loop (by facilianted diffusion of cl-) moderntely permeable to uner No achive MANSPOLI Nad-Mustain' tho. Nacl gradient There is Havourable Hypotonic between the tubulm lumen (600 mM Nacl) renal (interstitium) of renal interstitium (300 mm Nacl) Nacl PASSIVELY diffuses into tenal interstitium 300 mM 600 mm H20 CANNOT Follow (this segment Nacl NACK is nlwrys impermentle to the) H120 fluid inside tubule becomes hypotonik reliative to renal interstitium Thick Ascending limber impermeable to the Alarge amount of Solute transport Nat-Kt-2CF Cotransport inhibited by loop divretics & ethacrynic aud Aurosemide initate medullary osmolar Wadient convoluted tubule) > low furea permenbility in both Distal presence à absence de ADH - corneal collecting duce low (unea) permeability in the Outer medullary collecting duct presence a absence of ADH Ho remeability depends on presence or absence of ADH inner mechellary collecting durch The permerbility (Regulated by ADH) Variable Unea permenbility Thigh in presence of ADH low absence

TABLE 19-2 Effects of Angiotensin II



FUNCTION	RESULT	
Acts as a potent vasoconstrictor	Increased blood pressure	
Facilitates synthesis and release of aldosterone	Resorption of sodium and chloride from lu- men of distal convoluted tubule	
Facilitates release of ADH Resorption of water from lumen of colle tubule		
Increases thirst	Increased tissue fluid volume	
Inhibits renin release	Feedback inhibition	
Facilitates release of prostaglandins	Vasodilation of afferent glomerular arteriole, thus maintaining glomerular filtration rate	

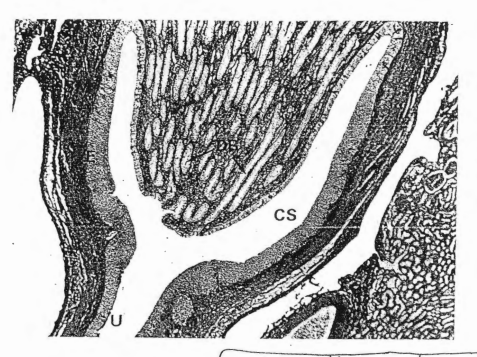


Fig. 16.26 Renal papilla

(Monkey: Azan × 30)

The renal papilla forms the apex of the medullary pyramid where it projects into the calyceal space. The ducts of Bellini DB, the largest of the collecting ducts, converge in the renal papilla to discharge urine into the pelvicalyceal space CS. The renal pelvis is lined by urinary epithelium E, and the wall of the pelvis contains smooth muscle SM which contracts to force urine into the ureter U.



Very small central cavity with Short cleft radiating from it

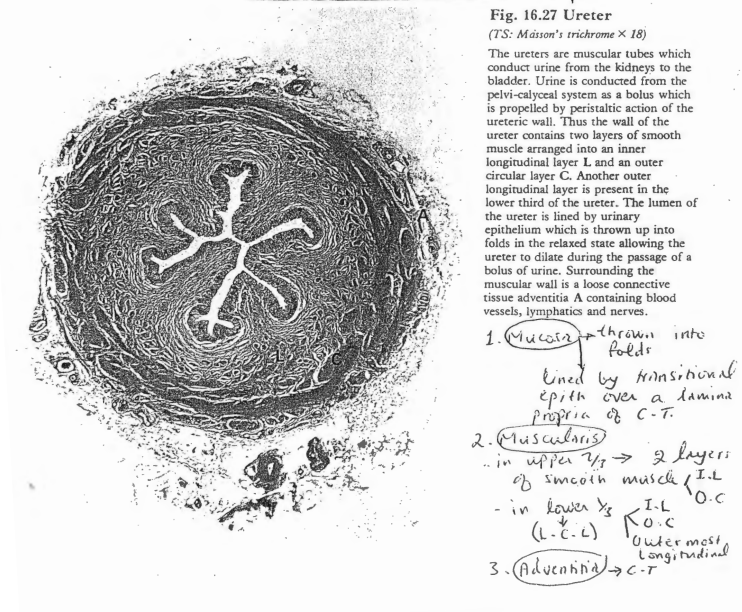
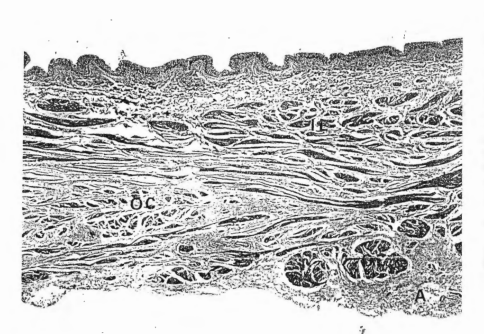


Fig. 16.27 Ureter

(TS: Masson's trichrome × 18)

The ureters are muscular tubes which conduct urine from the kidneys to the bladder. Urine is conducted from the pelvi-calyceal system as a bolus which is propelled by peristaltic action of the ureteric wall. Thus the wall of the ureter contains two layers of smooth muscle arranged into an inner longitudinal layer L and an outer circular layer C. Another outer longitudinal layer is present in the lower third of the ureter. The lumen of the ureter is lined by urinary epithelium which is thrown up into folds in the relaxed state allowing the ureter to dilate during the passage of a bolus of urine. Surrounding the muscular wall is a loose connective tissue adventitia A containing blood vessels, lymphatics and nerves.

1. Mucasa thrown into



URINARY SYSTEM

Fig. 16.28 Bladder

(TS: Masson's trichrome \times 12)

The general structure of the bladder wall resembles that of the lower third of the ureters. The wall of the bladder consists of three loosely arranged layers of smooth muscle and elastic fibres which contract during micturition. Note the inner longitudinal IL, outer circular OC and outermost longitudinal OL layers of smooth muscle. The urinary epithelium lining the bladder is thrown into many folds in the relaxed state. The outer adventitial coat A contains arteries, veins and lymphatics.

The urethra, the final conducting portion of the urinary tract, is discussed as part of the male reproductive tract in Chapter 18.

Bustami

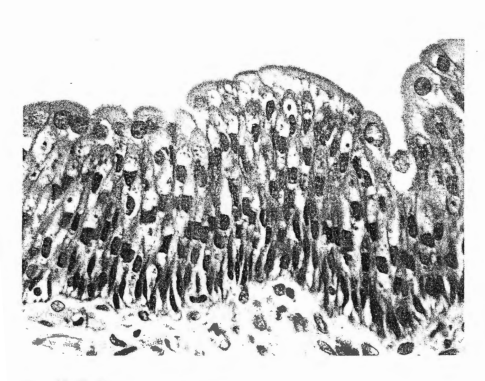


Fig. 16.29 Urinary epithelium

 $(H \& E \times 480)$

Urinary epithelium, also called transitional epithelium or rothelium, is found only within the conducting passages of the urinary system for which it is especially adapted. The plasma membranes of the superficial cells are much thicker han most cell membranes and have a highly ordered ubstructure, thus rendering urinary epithelium npermeable to urine which is potentially toxic. This ermeability barrier also prevents water from being drawn rough the epithelium into hypertonic urine. The cells of rinary epithelium have highly interdigitating cell junctions hich permit great distension of the epithelium without damage to the surface integrity (see also Figs. 5.16 and 5.17).

Urinary epithelium rests on a basement membrane which is often too thin to be resolved by light microscopy and was formerly thought to be absent. The basal layer is irregular and may be deeply indented by strands of underlying connective tissue containing capillaries. This unusual feature led early histologists to believe, mistakenly, that urinary epithelium contradicted the principle that epithelium never contains blood vessels.