



**Damanhour University** 

# **Faculty of Veterianry Medicine**

### **Project Title**

#### **Urine formation**

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#### **Abstract**

Urine is a liquid by-product of metabolism in humans and in many other animals. Urine flows from the kidneys through the ureters to the urinary bladder. Urination results in urine being excreted from the body through the urethra.

Cellular metabolism generates many by-products that are rich in nitrogen and must be cleared from the bloodstream, such as urea, uric acid, and creatinine. These by-products are expelled from the body during urination, which is the primary method for excreting water-soluble chemicals from the body. A urinalysis can detect nitrogenous wastes of the mammalian body.

Urine has a role in the earth's nitrogen cycle. In balanced ecosystems, urine fertilizes the soil and thus helps plants to grow. Therefore, urine can be used as a fertilizer. Some animals use it to mark their territories. Historically, aged or fermented urine (known as lant) was also used for gunpowder production, household cleaning, tanning of leath-er and dyeing of textiles.





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Human urine and feces are collectively referred to as human waste or human excreta, and are managed via sanitation systems. Livestock urine and feces also require proper management if the livestock population density is high.

#### Introduction

Urine is a waste byproduct formed from excess water and metabolic waste molecules during the process of renal system filtration. The primary function of the renal system is to regulate blood volume and plasma osmolarity, and waste removal via urine is essentially a convenient way that the body per-forms many functions using one process.

Urine formation occurs during three processes:

Filtration During filtration, blood enters the afferent arteriole and flows into the glomerulus where filterable blood components, such as water and nitroge-nous waste, will move towards the inside of the glomerulus, and nonfilterable components, such as cells and serum albumins, will exit via the efferent arte-riole. These filterable components accumulate in the glomerulus to form the glomerular filtrate.

Reabsorption

The next step is reabsorption, during which molecules and ions will be reab-sorbed into the circulatory system. The fluid passes through the components of the nephron (the proximal/distal convoluted tubules, loop of Henle, the collecting duct) as water and ions are removed as the fluid osmolarity (ion concentration) changes. In the collecting duct, secretion will occur before the fluid leaves the ure-ter in the form of urine. During secretion, some substances such as hydrogen ions, creatinine, and drugs—will be removed from the blood through the peritubular capillary network into the collecting duct. The end product of all these processes is urine, which is essentially a collection of substances that has not been reab-sorbed during glomerular filtration or tubular reabsorption.





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### **Project Aim and Outline**

- 1. Urine composition
- 2. Kidney and its functionality
  - 2.1. Glomerular filtration
  - 2.2. Tubular reabsorption
  - 2.3. Tubular secretion
- 3. Blood filtiration

#### **Results**

The formation of urine is a process important for the whole organism. Not only acid-base balance is modulated by it, but also blood osmolarity, plasma composition, and fluid volume, and thus it influences all cells in our body.

A healthy adult person produces 1.5-2 liters of urine per day and this process involves three basic mechanisms:

- 1. Glomerular filtration
- 2. Tubular reabsorption
- 3. Tubular secretion

Functional anatomy

The basic functional unit for urine formation is called a nephron. Very important is the arrangement of the nephron: it begins with renal corpuscle (Malpighi) that consists of a glomerulus, which is supplied by afferent glomerular arteriole and drained by efferent glomerular arteriole, and Bowman's capsule (capsula glomeruli, glomerular capsule). Renal tubules have three segments. The proximal tubule, in which we distinguish pars convoluta (initial section) and pars recta, loop of Henle (intermediate tubule), where can be recognized the descending limb and ascending limb (its proximal part is formed by a thick segment of the ascend-ing limb), and distal convoluted tubule (which has conversely first pars recta and then the pars convoluta) that subsequently joins the collecting ducts





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### Functional histology

Glomerulus consists of fenestrated capillaries without diaphragm that forms an important part of a renal filtration barrier. Blood flow and blood pressure in afferent and efferent arteriole are strictly regulated, which allows glomerular filtration into Bowman's capsule. The visceral layer of Bowman's capsule con-sists of podocytes and their pedicels that tightly fit the basement membrane of capillaries. The parietal layer is formed by a single layer of simple squamous epithelium. The renal filtration barrier is composed of the fenestrated capillary endothelium, the basement membrane, and the pedicles of podocytes. Pedicels interdigitate with one another forming filtration slits that are spanned by slit diaphragms (formed by protein nephrin). Due to its negative charge it prevents the filtration of plasma proteins In the glomerulus we can find mesangium that provides mechanical support, has phagocytic activity and secrets prostaglandins. Mesangial cells outside the glomerulus together with the macula densa cells (the distal segment of the ascending limb of the loop of Henle) and the granular cells (modified smooth muscle cells of the afferent arteriole) from juxtaglomerular apparatus. This is the place where the renal corpuscle gets into contact with the renal tubular system.

The proximal tubule is lined by simple cuboidal epithelium with a well-developed brush border on the luminal side, the thin portion of Henle's loop is lined by a simple squamous epithelium (poor in organelles). The distal tubule cells are smaller than those of proximal tubule and lack the brush border. Collecting ducts consist of principal intercalated cells.

#### Glomerular filtration

The volume of liquid filtered per unit time in all glomeruli can be expressed as the glomerular filtration rate (GFR). Its physiological value is 120 ml/min/1,73m2 body surface area, thus 180 l/day. About 99 % of the filtrate gets reabsorbed by the tubular resorption to the extracellular fluid (back into the body), leaving only 1.5-2 l of urine per day. Movement of the fluid through the filtration membrane is controlled and determined by the ratio of the hydrostatic pressure in the capillaries and oncotic pressure of plasma proteins (less by the hydrostatic pressure of the interstitial fluid and oncotic pressure in the filtrate). These forces are called Starling's forces and there are a few differences from the general principles:

1) Fluid is not exchanged between the capillary and the interstitium, but between the capillary and the fluid of Bowman's capsule





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- 2) Hydrostatic pressure in the capillaries is different, the movement is thus only one-sided (in the direction of filtration)
- 3) Filtration barrier (see above) has a unique structure and properties which do not allow passage of proteins into the filtrate (primary urine)

GFR is therefore dependent on the renal blood flow, the filtration pressure, the plasma oncotic pressure, and the size of the filtration area.

Control of glomerular filtration

Its main determinant is the renal blood flow that is directly proportional to the pressure difference between renal artery and renal vein and inversely proportional to the peripheral resistance of the afferent and efferent arteriole and the interlobular artery. We distinguish local and central regulatory mechanisms.

Iocal regulatory mechanisms

Iocal regulatory mechanisms consist mainly of myogenic autoregulation and tubuloglomerular feedback.

Myogenic autoregulation

Elevated blood pressure leads to the contraction of renal blood vessels, thereby increasing peripheral resistance. The reverse process occurs when the blood pressure decreases. Thanks to this regulatory mechanism remains the renal blood flow (and thus the GFR) relatively unchanged during normal fluctuations of the mean arterial blood pressure (80-180 mmHg).

Tubuloglomerular feedback

A decrease in GFR is registers by macula densa (part of the juxtaglomerular apparatus). As an answer to the detection of a low flow of tubular fluid or a reduced amount of sodium ions it sends paracrine chemical signal that causes vasodilation of the afferent arteriole, leading to an increase in a hydrostatic pres-sure and to a restoration of normal GFR.

Central regulatory mechanisms

The central regulatory mechanisms are less important. They are represented by the sympathetic nervous system, epinephrine, angiotensin II, prostaglandins and adenosine.

Postganglionic neurotransmitter of the sympathetic nervous system norepinephrine causes particularly in the afferent arteriole vasoconstriction, thereby reducing the renal blood flow (and thus the GFR) It is important especially in stressful situations, including pain and bleeding. Epinephrine has a similar effect.

Angiotensin II (via angiotensin receptor AT1) acts on both the afferent arteriole and the efferent arteriole in similar way as sympathetic nervous system and epinephrine.





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Iocally produced prostaglandins (especially E2 and I2) reduce the effects of sympathetic nervous system and angiotensin II on both the afferent arteriole and the efferent arteriole.

Adenosine is generally effective vasodilator, in afferent arteriole but acting as vaso-constrictor.

Furthermore, the renal blood flow is increased by atrial natriuretic peptide (ANP), glucocorticoids, nitric oxide or kinins, whereas antidiuretic hormone (ADH), ATP and endothelin cause a reduction in the renal blood flow.

Assessment of the glomerular filtration rate

If we want to determine GFR, which is one of the basic function of our kidneys, we have to use a substance that is excreted from the body only by glomerular filtration (inulin, creatinine) and is not affected by tubular processes. As an example we can mention the calculation of the clearance (plasma volume that is per unit time completely cleaned of marker substances) of endogenous creatinine, whose formula has the following form:

U – urine creatinine concentration in mmol/l

V – volume of urine (diuresis) in ml/s

P – plasma creatinine concentration in mmol/l

In clinical practice, we use more complex calculations, corrected for body surface area (and other physical parameters) – e.g. equation by Cockroft and Gault, equation MDRD etc.

Tubular reabsorption and secretion

As we mentioned above, about 99 % of the filtrate gets reabsorbed by the tubular resorption to the extracellular fluid (back into the body), leaving only 1.5-2 l of urine per day. The main task for renal tubules is therefore an isosmotic tubular reabsorption of primary urine. They absorb water, ions (sodium, chlorides, potassium, calcium, magnesium, bicarbonate or phosphate), urea, glucose and amino acids. All of this is independent on the extracellular fluid volume in the body – we speak about the obligatory resorption. Its primary role is to maintain fluid volume in the body under normal conditions.

Transport can be carried by passive diffusion (in the direction of the concentration or electrical gradient), primary active transport against gradient (needs energy – ATP) or secondary active transport (transport protein uses the concentration gradient created by a primary active transport realized by other transport protein). Substances can be





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transported by paracellular or transcellular routes. Transport of water is always passive. Na+/K+-ATPase located on the basolateral membrane plays important role in the secondary active transport. It creates a concentration gradient for Na+. Transport proteins act as symporters (transport of compound is coupled to the transport of Na+ in the same direction) or antiporters (transport of compound is coupled to the transport of Na+ in the opposite direction). To understand the processes in the tubular system, we must imagine tubular epithelial cells, their apical membrane facing the tubular fluid (primary urine), basolateral membrane, on the other hand, is in contact with the peritubular fluid (here is located the Na+/K+-ATPase).

### The proximal tubule

Reabsorption of sodium ions is in the first half of the proximal tubule coupled with the reabsorption of bicarbonate, glucose, amino acids, lactate, urea and phosphate. Absorbed compounds are osmotically active, thereby draining water from tubules. This leads to an increased concentration of chloride ions in the tubular fluid that is very important for a resorption in other parts of the proximal tubule.

Reabsorption of bicarbonate ions in the proximal tubule

Movement of bicarbonate and hydrogen ions depends on the transport sodium ions. This process is catalyzed by enzyme carbonic anhydrase (located in the apical membrane and in the intracellular part of the epithelial cells). The first step is the secretion of H+ into the tubular fluid through the Na+/H+ antiport, located at the luminal (apical) membrane of proximal tubule cells. Transferred H+ may in the tubular fluid react with filtered bicarbonate ions to form carbonic acid. Carbonic anhydrase facilitates the decomposition of carbonic acid in the tubular fluid to water and carbon dioxide. Both compounds can freely diffuse into the tubule epithelial cells, where carbonic acid is restored by the carbonic anhydrase. Molecules of carbonic acid dissociates into hydrogen and bicarbonate ions. Bicarbonate ions then pass through the basolateral membrane into the interstitial fluid through Na+/3HCO3—cotransporter or anion exchanger (Cl-/HCO3-). H+ returns via antiport with Na+ into the tubular fluid. For each secreted H+, Na+ and HCO3- is absorbed (Na+ is returned to the blood by active transport in exchange for K+ – Na+/K+-ATPase).

Renal (tubular) threshold





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Glucose, amino acid and many other organic compounds are in this part of the tubule completely resorbed under physiological conditions. This transport has some maximum value – so-called renal/tubular threshold. As an example we can mention the renal threshold for glucose. When this renal threshold is exceeded (due to too high plasma concentration – such as 10 mmol/l for glucose), glucose reab-sorption in the proximal tubule is incomplete and some amount of glucose remains in the final urine. Unabsorbed osmotically active molecules drain water molecules to renal tubules, thereby increasing diuresis (osmotic polyuria).

Reabsorption of sodium ions is in the second half of the proximal tubule coupled with the transport of chloride ions, used are both transcellular (on basolateral membrane helps K+/Cl—symport) and paracellular routes. Relatively abundant positively charged ions (sodium, potassium, calcium, magnesium) in the tubular fluid accompany chloride ions in paracellular transport. Transport of ions is followed by passive reabsorption of water.

# Ioop of Henle

Henle's loop absorbs about 25 % of the solutes (thick segment of the ascending limb), but only about 15 % water (descending limb). Its proper function (thick part of the ascending limb is impermeable to water and has active transport of Na+ and Cl-) is essential for the formation of a high osmotic pressure (hyperosmolarity) in the renal medulla that ensures a production of highly concentrated urine. Some mechanisms of reabsorption of ions are similar to those in the proximal tubule. Very important is the specific symport of Na+, K+ and 2 Cl- across the apical membrane. This symport uses energy derived from the transport of sodium and chloride ions in the direction of their concentration gradient for the transport of potassium ions into the cell (against their concentration gradient). Some of these ions leave cells on the basolateral membrane (together with Cl-), some return back into the tubular fluid, thereby creating an electrical imbalance. Due to this, posi-tively charged ions (Na+, K+, Ca2+, Mg2+) are resorbed by paracellular route (very important mechanism for resorption of solutes). This is especially significant for formation of a hypertonic renal medulla. Hypotonic fluid leaves the loop of Henle and enters the distal tubule.

#### Clinical correlation:

Substances that block the symport (e.g. furosemide) are used as very effective diuretic drugs – loop diuretics.

Distal convoluted tubule and collecting duct





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Distal convoluted tubule and collecting duct resorbe about 7 % of solutes (mainly Na+ and Cl–) and approximately 17 % water. Their resorption is affected by hormones (e.g. ADH) – facultative resorption. Hydrogen and potassium ions are secreted here. The distal convoluted tubule and the collecting duct thus play an important role in the formation of the final urine and in the regulation of osmolari-ty and pH. Sodium and chloride ions are absorbed in the first part of the distal convoluted tubule. The distal part of the distal convoluted tubule and the collecting duct consist of two cell types:

- 1) Principal cells responsible for the resorption of sodium ions and water (dependent on ADH) and secretion of K+ ions
- 2) Intercalated cells containing carbonic anhydrase. They are involved in acid-base balance, because they can secrete both hydrogen and bicarbonate ions

About the intercalated cells – see subchapet about acid-base balance.

Calcium and phosphate reabsorption and secretion

Plasma concentration of total calcium is 2.25-2.75 mmol/l and for ionized calcium 1.1-1.4 mmol/l. Only ionized calcium (about 48 % of total) is filterable by kidneys. Resorption takes place by both active (15-20 %) and passive paracellular (80 %) mechanisms. It is localized in the proximal tubule, the ascending part of Henle's loop and partially in the distal convoluted tubule. Parathyroid hormone stimulates the reabsorption by transcellular route in this segment. Calcitriol acts the same way, just mostly in the distal convoluted tubule. In contrast, calcitonin increases the excretion of calcium ions by inhibition of tubular reabsorption.

Serum phosphate concentration is 0.7-1.5 mmol/l, urine concentration is 15-90 mmol/l. Phosphates are also influenced by the parathyroid hormone (inhibits the resorption of phosphates) and by the calcitonin (also reduces the resorption of phosphates). Control of tubular processes

We can distinguish local and central regulatory mechanisms.

Iocal mechanisms

Iocal mechanisms are represented mainly by Starling's forces (increased plasma on-cotic pressure leads to an increased reabsorption of water and solutes from the interstitium into the capillaries, thereby supporting the tubular resorption) and glomerulo-tubular balance (increased GFR leads to an increase in glucose, amino acids, and so-dium ions resorption, these are followed by water – the volume of resorbed fluid increases proportionally with increased GFR).

Central mechanisms





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Central mechanisms are represented by many hormones – such as ADH, aldosterone, angiotensin II, epinephrine, natriuretic peptides (ANP and BNP), or parathyroid hormone. The sympathetic nervous system has a role also.

ADH (antidiuretic hormone, vasopressin) is produced in the hypothalamus and secreted by the posterior pituitary gland in response to increased osmolarity of extracellular fluid (to a lesser extent as an answer to a decrease of extracellular fluid volume). ADH binds to the V2-receptor located on collecting duct cells (partly on distal tubule cells). Its effect increases the number of aquaporins in cell membranes and water molecules can pass along the osmotic gradient into the peritubular fluid (ECF). ADH acts also on transport of urea in the collecting duct and on transport of Na+ and Cl– in the thick segment of the ascending limb of the loop of Henle.

Aldosterone is secreted by the zona glomerulosa of the adrenal cortex in response to increasing plasma concentrations of angiotensin II and potassium ions. It plays therefore an important role in maintaining of a constant level of potassium ions (accelerates secretion of potassium ions in the thick segment of the loop of Henle and in the distal tubule) and in regulation of volume of ECF. As the part of the renin-angiotensin-aldosterone system, it stimulates reabsorption of sodium ions, accompanied by passive water resorption (distal tubule and collecting ducts). This system is activated by decrease in the plasma volume.

Angiotensin II stimulates aldosterone secretion and resorption of sodium ions (and consequently resorption of water molecules) in the proximal tubule.

The sympathetic nervous system and epinephrine stimulate reabsorption of sodium ions and water molecules in the proximal tubule and the thick segment of the loop of Henle.

As the name suggests, natriuretic peptides (ANP – atrial natriuretic peptide and BNP – brain natriuretic peptide) increase natriuresis. They inhibit Na+ reabsorption in the distal tubule, thereby increasing its loss in urine. Sodium ions drain water molecules, the result is increased diuresis. Both peptides are secreted by our hearts. ANP is secreted by atrial cardiomyocytes, the stimulus for its secretion is increased wall stress (increased venous return causes dilation of the heart). BNP is secreted by ventricular cardiomyocytes, the signal is increased tension in the ventricular wall. Natriuretic peptides thus mediate the response of our organism to an excess of Na+ and increased blood volume. Only natriuretic peptides (together with dopamine) increase diuresis. Parathyroid hormone reduces Ca2+ excretion (stimulates reabsorption of Ca2+ from the primary urine) and increases the excretion of phosphates in our kidneys. In a result, it increases calcaemia and decreases phosphatemia.

Control of urine osmolarity





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There are several processes controlling urine osmolarity. Excretion of excess water leads to a formation of hypotonic urine, excretion of excess solutes results in a formation of hypertonic urine.

- 1) Dilution of urine
- a) The loop of Henle creates an osmotic gradient from the cortex to the hypertonic medulla (due to impermeability of the thick segment to water molecules and high reabsorption of solutes)
- b) Production of ADH is reduced
- c) Urea passes from the medulla into the tubular system, thereby reducing hypertonicity of the medulla
- 2) Production of hypertonic urine
- a) The loop of Henle creates an osmotic gradient (hypertonic medulla); Na+, Cl– (see above) and urea plays an important role hypertonicity of the renal medulla reaches its maximum
- b) Production of ADH is increased
- c) Urea circulates in the renal medulla increased hypertonicity of the medulla Acid-base balance and kidneys

The role of our kidneys in acid-base balance is discussed in the subchapter about acid-base balance.

#### Final urine

Final urine is a characteristically malodorous, clear, golden yellow liquid. Its specific gravity varies between 1 003-1 038 kg/m3 and its pH between 4.4-8.0. It contains Na+ (100-250 mmol/l), K+ (25-100 mmol/l), Cl– (about 135 mmol/l), Ca2+, creatinine, vanillylmandelic acid (degradation product of catecholamines), uric acid, urea, etc. Healthy kidneys do not allow a significant amount of proteins and glucose to reach the final urine (they are almost completely reabsorbed). The presence of a high amount of proteins and glucose in the final urine is a pathological finding. Normal diuresis is 1.5-2 l/day. Polyuria is diuresis higher than 2 l/day, oliguria lower than 0.5 l/day, and anuria lower than 0.1 l/day.





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#### **Conclusion**

#### Mechanism of urine Formation

The mechanism of urine formation involves the following steps:

#### Glomerular Filtration

Glomerular filtration occurs in the glomerulus where blood is filtered. This process oc-curs across the three layers- epithelium of Bowman's capsule, endothelium of glomeru-lar blood vessels, and a membrane between these two layers.

Blood is filtered in such a way that all the constituents of the plasma reach the Bow-man's capsule, except proteins. Therefore, this process is known as ultrafiltration.

### Reabsorption

Around 99 percent of the filtrate obtained is reabsorbed by the renal tubules. This is known as reabsorption. This is achieved by active and passive transport.

#### Secretion

The next step in urine formation is the tubular secretion. Here, tubular cells secrete substances like hydrogen ion, potassium ion, etc into the filtrate. By this process, the ionic, acid-base and the balance of other body fluids are maintained. The secreted ions combine with the filtrate and form urine. The urine passes out of the nephron tubule into a collecting duct.

#### Urine

The urine produced is 95% water and 5% nitrogenous wastes. Wastes such as urea, ammonia, creatinine are excreted in urine. Apart from these, the potassium, sodium and calcium ions are also excreted.





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