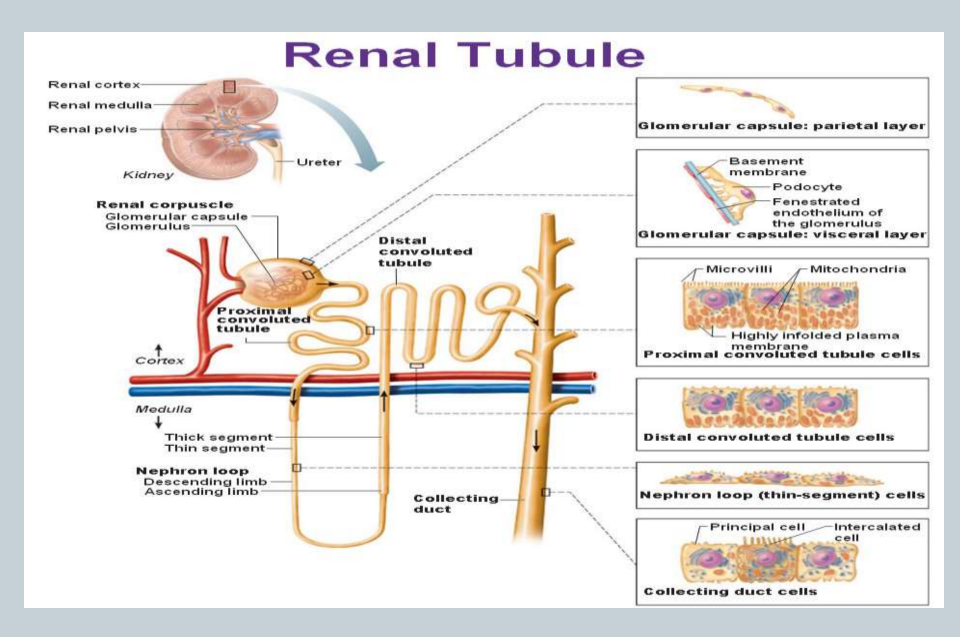
DIURETICS

DEFINITION:

These are drugs which cause a net loss of Na+ and water in urine

There are several categories of diuretics. All diuretics increases the excretion of water from body.



CLASSIFICATION

Diuretics are Classified as:

- 1. High ceiling /Loop diuretics...
- 2. Thiazides.
- 3.Carbonic anhydrase inhibitors.
- 4.Potassium –sparing diuretics.
- 5.Osmatic diuretics.
- 6.Low ceiling diuretics.

DIURETICS CLASSIFICATION

1.HIGH EFFICACY DIURETICS: (Inhibitors of Na+,K+,2Cl- cotransport)

(a) Sulphamoyl deravatives: Furosemide. Bumetanide. Torasemide.

(b) Phenoxyacetic acid derivative: Ethacrynic acid.

MEDIUM EFFICACY DIURETICS

2.Medium efficacy diuretics : (Inhibitors of Na+,Cl- symport)

 (a) Benzothiadiazines(THIAZIDES): Hydrochloro thiazide. Benzthiazide. Hydroflumethe thiazide. Ciopamide.
 (b) Thiazide: Chlorthalidone. Metolazone. Xipamide. Indapamide.

WEAK OR ADJUNCTIVE DIURETICS

3. Weak or adjunctive diuretics: (a) Carbonic anhydrase inhibitors: Acetazolamide. (b) Potassium – sparing diuretics: (i)Aldosterone antagonist: Spironolacton Eplerenone. (ii)Inhibitors of renal epithial Na+ channel: Trimterene. Amiloride.

(c) Osmotic diuretics: Mannitol. Isosorbide. Glycerol.

(d) Xanthines: Theophlline.

ANTI- DIURETICS

1.Anti diuretic homone(ADH) and its analogues: Vasopressin. Desmopressin. Lypressin. Terlipressin. 2. Diuretics: Thiazides. Amiloride. 3. Miscellineous: Chlorpropamide. Carbamazepine.

MECHANISM OF DIURETICS

Mechanism of action of commonly used diuretics

Site of Action	Channel Inhibited	Percent Excreted
Loop of Henle Furosemide, bumetanide, ethacrynic acid	Na/K/2CI	Up to 25
Distal Tubule Thiazides	Na/CI	Up to 3-5
Cortical Collecting Tubule Spironolactone, amiloride, and triamterine	Na channel	Up to 1-2

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MECHANISM OF ACTION

Diuretics: Mechanism of Action

Cortex

Medulla

Thiazides Inhibit active exchange of CI-Na the cortical diluting segment of the ascending loop of Henle

K-sparing

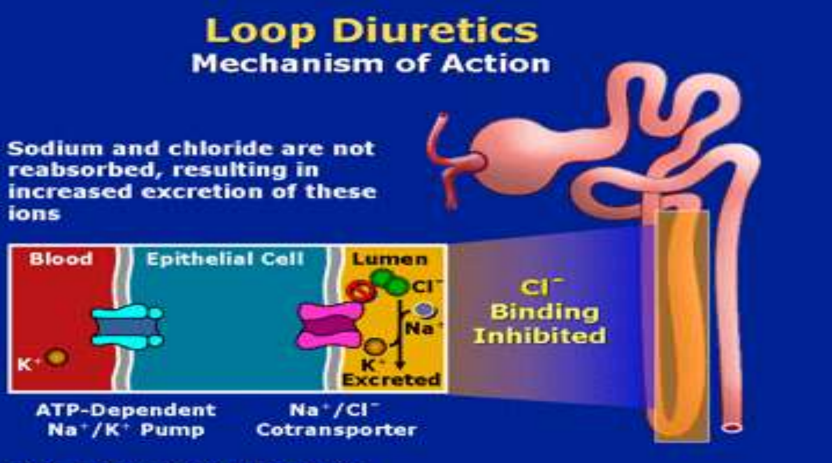
Inhibit reabsorption of Na in the stal convoluted and collecting tubule

Loop diuretics

Inhibit exchange of CI-Na-K in the thick segment of the ascending loop of Henle

Loop of Henle Collecting tubule

MECHANISM OF ACTION OF LOOP DIURETICS



ATP = adenosine triphosphate

Morrison RT. Med Clin North Am, 1997;81:689-704; Brater DC, Am 7 Med Sci. 2000;319:38-50. Elide Siderce Hipertension Online Hipertension problems



Loop diuretics

Site of action – enter via filtration and secretion by the OATs. Act on TAHL

Mechanism - inhibition of Na+/K+/2Clsymporter. Positive luminal potential 1 Mg+, Ca2+ reabsorption 1 Hypochloremia due to NKCC block Large doses abolish osmotic gradient

Renal vascular resistance 1, RBF
via effect on prostaglandins. Kidney is not able to produce dilute urine. After initial strong diuresis - diuretic braking.

Urine - increased excretion of all ions: Na+, Cl-, K+, H+, Mg2+, Ca2+, as well as HCO₃ in case of furosemide (Furosemide is a weak CA inhibitor).

Plasma - hypochloremic alkalosis and hypokalemia (mechanisms are similar to thiazide duiretics and will be considered shortly)

TAHL – thick ascending loop of Henle

impermeable to water!

Loop diuretics

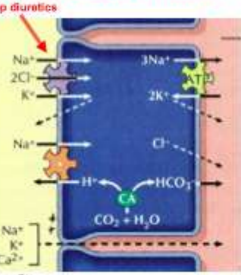
Mg²

Transcellular: Via specialized luminal Na+K+ CI- co-transporters.

Na/H antiporter continues to reabsorb Na+ and excrete H+

Paracellular: Backleak of K+ creates lumen positive 6mV transepithelial gradient which drives paracellular movement of cations out of the lumen.





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INDICATIONS AND SIDE EFFECTS

Indications & Side Effects

- Loop diurctics
 - large volume diuresis
 - isotonic urine (as compared to plasma)
- Indications
 - edema
 - congestive heart failure
 - acute pulmonary edema
 - cirrhosis
 - nephrotic syndrome
 - hypertension
 - hypercalc<u>emia</u>
 - forced diuresis



- excess volume depletion
 - circulatory collapse
 - azotemia & hyperuricemia
- hypokalemia
 - cardiac amhythmias
- hypocalcemia
- hypomagnesemia
- ototoxicity

Thiazides - Mechanism Of Action Act In The Distal Tubule Inhibit Reabsorption Of Sodium And Potassium Stimulate The Reabsorption Of Calcium Loss Of Water As Urine

MECHANISM OF THIAZIDE DIURETICS

Thiazide Diuretics Mechanism of Action

CI Binding

Sodium and chloride are not reabsorbed, resulting in increased excretion of these ions



ATP-Dependent Na⁺/K⁺ Pump Na⁺/Cl⁻ Cotransporter

ATP = adenosine triphosphate

Morrison RT. Med Clin North Am. 1997;81:689-704.

fillder fulselige Hypertexister Ordere Hypertexister



Thiazide Diuretics (HYDROCHLOROTHIAZIDE)

Therapeutic uses:

Absorbed orally. Bound to plasma proteins, Secreted by the OAT in PT Hypertension -reduces blood pressure by reducing volume and producing mild vasodilation

Congestive heart failure

Hypercalciuria- to help prevent renal stones.

Nephrogenic diabetes insipidus (renal insensitivity to ADH): thiazides ↓ plasma volume => lowers GFR -> reabsorption of Na in PT ↑. Less Na+ and water reach CD so overall fluid conservation is obtained.

Adverse effects -

Electrolyte imbalance – hypokalemic metabolic alkalosis, hyponatremia, hypercalcemia, hyperuricemia, hypochloremia, cardiac arrhythmias. Hypokalemia increases risk of torsade de pointes caused by guanidine. Hypotension -due to volume depletion

Hyperglycemia- in patients with diabetes or abnormal glucose tolerance tests. Mechanisms poorly understood

Hyperlipidemia- an increase in the levels of LDL, total cholesterol and total triglycerides

Hypersensitivity

POTASSIUM- SPARING DIURETICS

Spironolactone - Mechanism Of Action

 Competitively Binds The Aldosterone Receptor Preventing The Hormone From Binding To Its Receptor

Aldosterone's Normal Steroid-Nuclear DNA Transcription Is Halted

OSMOTIC DIURETICS Osmotic Diuretics

mannitol

- Raises osmotic pressure of the plasma thus draws H₂0 out of body tissues & produces osmotic diuresis
- Does not effect Na* excretion

Osmotic Diuretics:

Therapeutic Uses

- Used in the treatment of patients in the early, oliguric phase of ARF
- To promote the excretion of toxic substances
- Reduction of intracranial pressure
- Treatment of cerebral edema

PHARMACOLOGICAL ACTIONS OF DIURETICS

HIGH CEILING/LOOP DIURETICS

- High ceiling diuretics may cause a substantial decrease upto 20% of the filtered load of Nacl and water.
- Loop diuretics such as FUROSEMIDE inhibits the body's ability to reabsorb sodium at the ascending loop in NEPHRON.

THIAZIDES

- Thiazide diuretics such as Hydrochlorothiazide act on the distal convoluted tubule and inhibits the sodiumchloride symporter leading to retention of water in the urine.
- Frequent urination is due to the increased loss of water.
- The long term anti –hypertensive action is based on the thiazides which decrease preload and blood pressure.

CARBONIC ANHYDRASE INHIBITORS

- Carbonic anhydrase inhibitors inhibits the enzyme carbonic anhydrase which is found in proximal convoluted tubule.
- This results in several effects including biocarbonate retention in the urine.
- Potassium retention in urine.
- Decreased sodium absorption.
 Eg: Acetazolamide.
 Methazolamine.

POTASSIUM-SPARING DIURETICS

- These are diuretics which do not promote the secretion of potassium into the urine.
- Potassium is retained and not lost as much as with other diuretics.
- The term potassium sparing refers to an effects rather than a mechanism or location.

Eg: Aldosterone antagonists Spironolactone

- Which is a competitive antagonist of aldosterone.
- Aldosterone adds sodium channels in the cells of collecting duct and late distal tubule of the Nephron.
- Spirnolactone prevents aldosterone from entering the cells, and preventing sodium reabsorption.
 Eg: Eplerenone.

Potassium canrenonate.

• Epithelial sodium channel blockers

Eg: Amiloride.

Triamterence.

OSMOTIC DIURETICS

- The compounds as Mannitol are filtered in the glomerulus, but cannot be reabsorbed.
- Their presence lead to an increases in the osmolarity of the filtrate.
- To maintain osmotic balance ,water is retained in the urine.
- Glucose like mannitol behave as an osmotic diuretic.
- Glucosuria causes a loss of hypotonic water & Na+, leading to a hypertonic state with signs of volume depletion.
- Such as Hypotention, Tachycardia.

LOW CEILING DIURETICS

- The low celing diuretics are used to indicate an diuretic has a rapid flatting dose effect curve.
- It refers to a pharmacological profile ,not a chemical structure.

MEDICINAL USES

Diuretics are used to treat

 Heart failures.
 Liver cirrhosis.
 Hypertension.
 Certain kidney diseases

USES AND SIDE EFFECTS OF DIURETICS

