

الميلاد

الأحرار

٨٨

بسم الله الرحمن الرحيم

Drugs and the kidney

Dr Mohammad Salem Hareedy
2025

Kidney diseases and Drugs

- Kidney disease can alter the pharmacokinetics of renally excreted drugs and can increase the possible toxicity of these drugs (especially if the **active drug or metabolites** are renally cleared).
- Drug doses usually should be reduced in renal disease in proportion to the predicted reduction in clearance of the active drug moiety; both patient and drug factors should be considered:
 - 1- Patient factor: the degree of renal impairment and patient size.
 - 2- Drug factors: the **fraction of the drug excreted unchanged in urine** and the **drug's therapeutic index**.

Dose adjustment & therapeutic index

- ❑ For drugs with **narrow therapeutic indices** (Aminoglycosides, warfarin, lithium, digoxin, vancomycin, cyclosporin & phenytoin), even **small changes in drug concentration** can cause **toxicity**.
→ in relation to renal functions
- ❑ These drugs should be dosed using either robust parameters (e.g. clinical response, INR for warfarin, therapeutic drug monitoring (TDM), etc.) OR by empirical calculations of doses (most calculations are not reliable enough to be safe).
↳ renal function alone
- ✓ Conversely, for drugs with a wide therapeutic index (e.g., **beta lactams**), even **large changes in drug clearance** may have only a **modest impact**, and therefore **dose adjustments are less important**.
↳ β -lactams, modest drug

➤ For drugs with **intermediate therapeutic index**: an estimate of **renal function** ~~as~~ an estimate of drug clearance provides **useful guidance to dosing** and can be used together with clinical and biochemical measures of effects (e.g. **serum uric acid** for the anti-gout drug **allopurinol**).

direct relation if complete renal exc.

Dose adjustment according to renal functions

- ❑ For chronic kidney disease, estimates of renal function are used to predict disease outcome.
- ❑ For drug dosing **estimation of renal function** are used to **estimate the renal clearance of the drug** which is used for **further calculation of doses**.

For dose adjustment:

1- Calculate the **drug clearance** based on renal functions.

2- Consider oral **bioavailability** for oral drugs

Both CL and F determine **steady state conc.**

$$\text{Dose} = \text{Desired plasma conc.} \times \frac{\text{Clearance}}{\text{Bioavailability}}$$

❑ Thus, if a drug is **100% renally cleared** and renal function is half-normal, the drug dose should be halved, all other things being equal.

❑ However, many drugs are **inactivated by metabolism** (in the **liver** predominantly), and hence doses of metabolized drugs do not usually require changing in renal disease.

liver
function
test is
needed

Drug-induced nephrotoxicity

❑ Drug-induced nephrotoxicity is the presence of any kidney injury (acute or chronic) caused **directly** or **indirectly** by medication.

❑ Drugs can cause ^{easy to treat} **acute renal injury**, ^{↳ anti-neoplastic drugs w/ its metabolites + statin [same way]} **intrarenal obstruction**, **interstitial nephritis**, **nephrotic syndrome**, **acid-base** and fluid **electrolytes** disorders.

Drug induced acute kidney injury (AKI)

1- Pre-renal AKI

2-Intra-renal or renal AKI:

❑ Drugs causing **Acute Tubular Necrosis** or injury.

❑ Drugs causing **Acute Interstitial Nephritis**

❑ Drugs causing **Glomerulonephritis**

3- Post-renal AKI

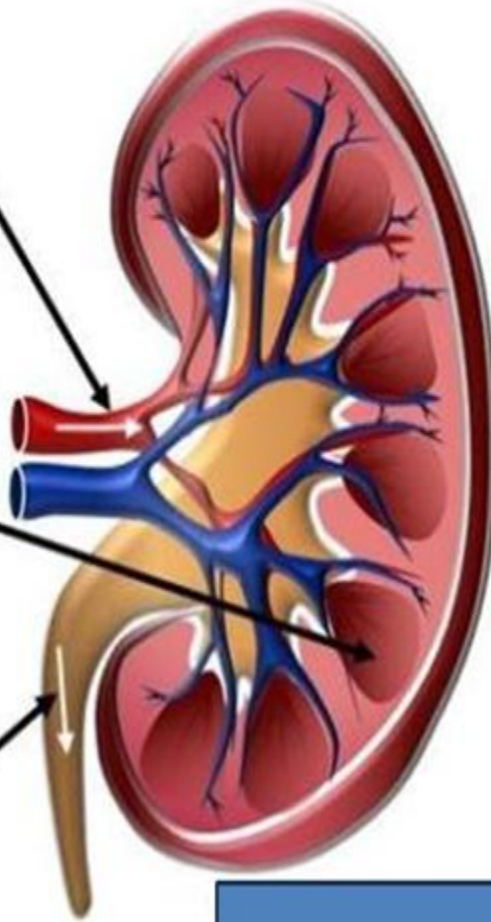
Acute kidney injury (AKI) induced by drugs

Acute Kidney Injury (AKI)

Prerenal: marked decrease in renal blood flow

Renal: damage within the kidney structures

Postrenal: obstruction of urine outflow

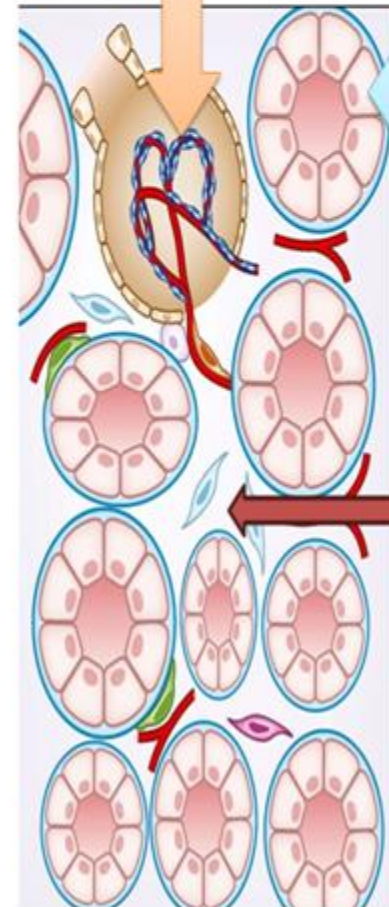


Glomerulus

Same in classification of renal failure

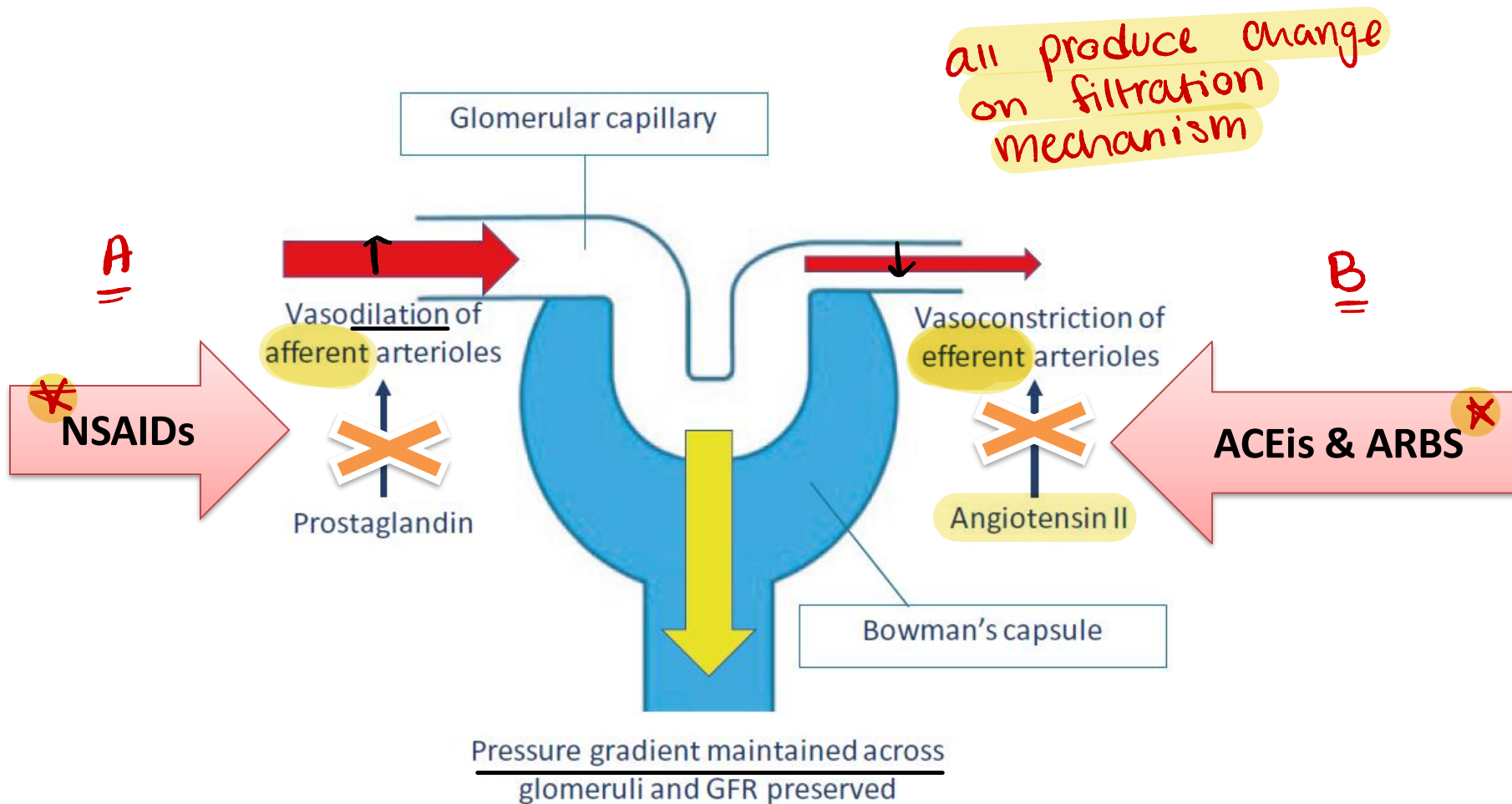
Tubules

Interstitial tissue



1-Drug induced pre-renal AKI

- 1- Reduced circulating volume (e.g., Diuretics).
- 2- Selective reduction in renal perfusion (Drugs that affect glomerular blood flow) like:
 - NSAIDs/COX2 inhibitors – inhibit synthesis of vasodilatory prostaglandins. (A)
 - ACE inhibitors/ARB – block vasoconstrictor effects of angiotensin II. (B)
 - Calcineurin inhibitors (cyclosporin and tacrolimus) which increase vasoconstriction. (B)
- Treatment include maintain vascular volume, Using Vasopressors if necessary.



2- Intra-renal AKI induced by drugs

— tubule
— interstitium
— glomeruli

Intra-renal refer to intrinsic damage to the structure of the kidney (**apoptosis or necrosis**) by ischemia or other cellular mechanisms (e.g. impairing mitochondrial function, interfering with tubular transport or increasing oxidative stress).

A- Acute tubular injury (ATI) or tubular necrosis

Examples: **Aminoglycosides**, **Amphotericin B**, **rifampicin**, **radiocontrast agents**, **cisplatin**, **NSAIDs**, **Loop diuretics**, **Acyclovir**, **Cephalosporins**, **calcineurin inhibitors**, **Paracetamol**, **vancomycin**, ~~mannitol~~.

very common w/ all mechanisms


↳ suppressor after transplant [can damage new kidney]

❑ The important risk factors for acute tubular injury:

1. Exposure to multiple nephrotoxic drugs.
2. A disease that increase the tubular injury (e.g., diabetes, hypertension).
3. Very young or very old age.
4. Pre-existing chronic kidney disease (CKD).
5. Intravascular volume depletion.

B- Acute interstitial nephritis (AIN)

Acute interstitial nephritis (AIN) is an immune-mediated form of kidney injury (infiltration of immune cells in the tubulo-interstitium).

- ^{allergy} Medications are the most common cause of AIN.
- AIN can cause permanent kidney damage from fibrosis formation.
- In drug-induced AIN, drug discontinuation is critical.  NOT dose dependent.
- Management of AIN: corticosteroids are usually prescribed to prevent permanent kidney damage
- Examples of drug induced AIN: Antimicrobials (β -lactams, sulfonamides, quinolones, vancomycin, Aminoglycosides), NSAIDs, Proton Pump Inhibitors, phenytoin, carbamazepine, allopurinol, thiazides, Calcium channel blockers & lithium.

C- Drug induced glomerulonephritis

- ❑ Glomerular damage that occurs after exposure to medications can be caused by direct cellular injury involving the mesangial, endothelial, or visceral epithelial cells (**podocytes**) ^{= ↓ proteins ...}

Drug-induced **podocytopathy** can occur in several situations:

- 1- Interferon (IFN) causes podocyte injury and **nephrotic syndrome** may occur.
- 2- Pamidronate in high doses can cause direct podocyte injury.
- 3- Chronic lithium exposure.
- 4- **Minimal change disease** (MCD) is the most common glomerular lesion observed with **NSAIDs**, which may be because of **shunting of arachidonic acid metabolites** into pathways that alter immune function and promote podocyte injury.

3- Post renal injury by drugs

- Drug induced **Crystalline** nephropathies are characterized primarily by **intra-tubular** crystal deposition (**crystalluria**).
- Urine sediment examination showing **crystal-containing casts** is a helpful non-invasive diagnostic test instead of renal biopsy.

Intra-renal crystal deposition occurs when:

1- The kidney is the major route of a drug/metabolite excretion.

2- Increased excretion of the drug (e.g., excessive drug dosing).

3- Supersaturation of the drug & precipitation within urine due to :

☐ Circulatory volume depletion/dehydration.

☐ the pKa of the drug & the Urine pH that favor drug precipitation:

✱ Examples **acidic pH** for **methotrexate** or **sulfadiazine** and **alkaline pH** for **ciprofloxacin**.
↳ to avoid precipitation?
alkaline urine

4- The presence of underlying kidney disease may further enhance risk for drug-induced crystalline nephropathy.

Culprit Medication	Disease induced	Prevention & Treatment
Methotrexate	<u>Crystalluria</u> , <u>AKI</u> , & chronic kidney disease (<u>CKD</u>)	IV fluids before/during drug, <u>alkalinize urine</u> , <u>adjust drug dose</u> , TDM , <u>folinic acid</u> , or glucarpidase .
Sulfadiazine, sulfamethoxazole	<u>Crystalluria</u> , <u>AKI</u> & <u>nephrolithiasis</u> ! if	<u>Alkalinize</u> urine, <u>adjust dose</u> for kidney function, assure <u>euvolemia</u>
Acyclovir	<u>Crystalluria</u> , <u>AKI</u> , and <u>CKD</u>	Avoid <u>rapid iv bolus</u> , <u>adjust drug dose</u> , assure <u>euvolemia</u>

- **Glucarpidase** is used for treatment of **elevated levels of methotrexate** in **cancer patients** with **impaired kidney functions**.
- Glucarpidase is an **enzyme** that **inactivates methotrexate** rapidly.
- Glucarpidase also **degrades folinic acid** (e.g. Leucovorin) so the two should not be used together (within two hours of one another).

Ciprofloxacin, levofloxacin	<u>Crystalluria</u> and <u>AKI</u>	Assure euvolemia during drug therapy and <u>avoid alkaline urine</u> (if possible)
Triamterene	<u>Crystalluria</u> , AKI, CKD, and nephrolithiasis	Alkalinize urine , assure euvolemia during drug therapy
Cyclophosphamide	Hemorrhagic cystitis	Hydration , continuous bladder irrigation , & prophylactic use of mesna .

❑ **Mesna** has **antioxidant properties**.

❑ Mesna concentrates in the bladder and **conjugated** with **acrolein** and other toxic metabolites

❑ This conjugation reaction **inactivates the toxic compounds** to harmless metabolites.

Indirect drug induced postrenal AKI

Here, AKI is NOT caused by precipitation of the drug itself or its metabolites in urine. Instead, different mechanisms are involved.

Examples:

1- Crystal nephropathy may also result from the use of anticancer chemotherapy due to uric acid and calcium phosphate crystal deposition (due to death of many malignant cells).

2- Drug induced rhabdomyolysis and myoglobinuria (postrenal AKI). Statins, alcohol, Benzodiazepines, methadone and Methamphetamine can cause rhabdomyolysis and AKI.

Drug induced nephrotic syndrome

NSAIDs, gold therapy, probenecid, penicillamine, Tolbutamide, interferon-alfa, **lithium**, and pamidronate

↑ stopped using it

Drug induced renal Acid base disturbances

❑ **Phenformin** and metformin may cause **lactic acidosis**

❑ Proximal renal tubular acidosis by acetazolamide

Drug induced renal water imbalance

❑ **Hyponatremia**, syndrome inappropriate ADH secretion by Chlorpropamide

❑ Nephrogenic diabetes insipidus by lithium.

chronic interstitial nephritis

w/out acute injury

Chronic use of **acetaminophen**, **aspirin**, **diuretics** and **lithium** is associated with chronic interstitial nephritis leading to fibrosis and renal scarring