

## PHARMACOLOGY

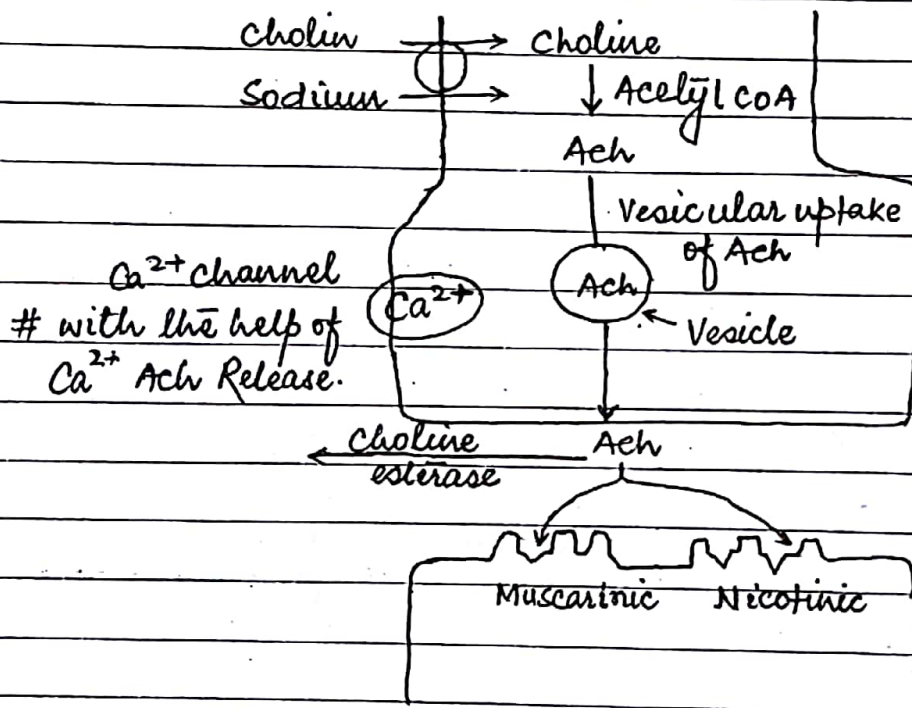
- Sympathetic System Neurotransmitter - Nor-Epinephrine
  - ↳ Thoraco lumbar outflow ( $T_1$  to  $L_3$ )
- Parasympathetic System Neurotransmitter - Acetyl choline
  - ↳ Cranio-sacral outflow ( $III, IV, IX, X, S_2, S_3, S_4$ )

Cholinergic drug:

Choline uptake -  $Na^+$ -choline symport

↳ 1st step → Rate limiting step in synthesis of Ach.

# Source of choline → Serine.



# True cholinesterase → +nt at synapse.

Pseudocholinesterase → +nt in plasma.

Cholinergic drug metabolised by → Pseudocholinesterase.

Choline uptake inhibited by → Hemicholinium.

Vesicular uptake up of Ach blocked by → Vesamicol.

Release of Ach modulated by  $\left\{ \begin{array}{l} \text{Blocked by - Botulinum toxin} \\ \text{Stimulated by - Spider Venom.} \end{array} \right.$

Defect in  $Ca^{2+}$  channel - Lambert Eaton Syndrome.

## Lambert Eaton Syndrome:

Defect is  $Ca^{2+}$  channel Presynaptically.

For t/t we need  $Ca^{2+}$  channel activator  $\rightarrow$  3,4-diamino pyridine  
(Dalf Ampridine)

Also useful for t/t of  
- Multiple Sclerosis

to improve walking capacity.

- It is  $K^+$  channel blocker &  $Ca^{2+}$  channel activator.

## Sites of Release of Ach Neurotransmitter:

at the ① Ganglion

- Preganglionic fibre of sympathetic & parasympathetic Release Ach at ganglion.

② Adrenal Medulla.

③ Neuromuscular junction.

④ Postganglionic Parasympathetic fibre.

# Postganglionic sympathetic fibre normally releases  
- Nor-epinephrine (NE)

Exception:

a) Sweat gland - Release Ach (Sympathetic cholinergic)

# Hyperhydrosis (Excessive Sweating)

t/t  $\leftarrow$  Sympathetic lamy

Botulinum toxin injection.

b) Renal blood flow - Release Dopamine by Sympathetic postganglionic fibre.

Extra point:

① Conversion of NA into Adrenaline by Methylation  
- Eg. of Phase II reaction.

② Conversion of Histamine into methyl histamine by Methylation.

Mast cell secrete histamine.

~~Mast~~ Mastocytosis (Histamine releasing tumour)

↓  
urinary estimation of Methyl histamine - Useful for diagnosis of Mastocytosis.

# Urinary estimation of VMA (Vanillyl Mandelic Acid) -  
Useful for diagnosis of Pheochromocytoma.

Toxins in ANS:

BOTULINIUM TOXIN - A to G Subtype.

Clinical uses of Botulinum A toxin:

- ① Blepharospasm
- ② Strabismus
- ③ Wrinkle (in forehead corrected)
- ④ Cosmetics.

Clinical uses of Botulinum B toxin:

- Used as Muscle relaxant.

↳ Cervical dystonia (Painful muscle spasm)

ONABOTULINUM TOXIN

- Derivative of Botulinum A toxin.

Useful for - ① Prophylaxis of Chronic Migraine.

② Relaxation of Detrusor muscle - Given intravesically.

↓  
Causing Retention of urine  
So useful for t/t of overactive bladder.



Adenyl cyclase Pathway:

2nd Messenger - CAMP.

M<sub>1</sub>, M<sub>3</sub> & M<sub>5</sub> follow G<sub>q</sub> pathway  
 M<sub>2</sub> & M<sub>4</sub> follow G<sub>i</sub> pathway.

Muscarinic Receptors:

M<sub>1</sub>: Location - Stomach

Action - Releasing HCl

Overstimulation of M<sub>1</sub> - Gastritis

Selective M<sub>1</sub> agonist - Oxotremorine.

↳ SE - Gastritis

For Gastric ulcer - Block M<sub>1</sub>.

Selective M<sub>1</sub> antagonist  $\left\{ \begin{array}{l} \text{PIRENZEPINE} \\ \text{TELENZEPINE} \end{array} \right\}$  For t/t of gastric ulcer.

M<sub>2</sub>: Located on Myocardium

↳ Maximally in AV node.

Action: Stimulation of M<sub>2</sub> causes reduction in conduction velocity.

↓  
 Causing Bradycardia

as Vagus (X) fibre is Parasympathetic fibre

↳ act on M<sub>2</sub> receptor → Causes Bradycardia.

# Athletic person → High Vagal tone

# Vagomimetic drug → Causing Bradycardia

Use of M<sub>2</sub> agonist → SVT (Supraventricular Tachycardia).

Selective M<sub>2</sub> agonist - METHACHOLINE  $\left\{ \begin{array}{l} \text{Action} \\ 98-99\% - M_2 \\ 1-2\% - M_1, M_3 \end{array} \right.$

Selective M<sub>2</sub> antagonist - METHOCTRAMINE  
 TRIPITRAMINE

# Methacholine challenge test → Δ of Asthma.  
 ↳ Cause bronchoconstriction.

Digoxin = Vagomimetic property

- Anti-arrhythmic
- Atrial Fibrillation
- Atrial Flutter.

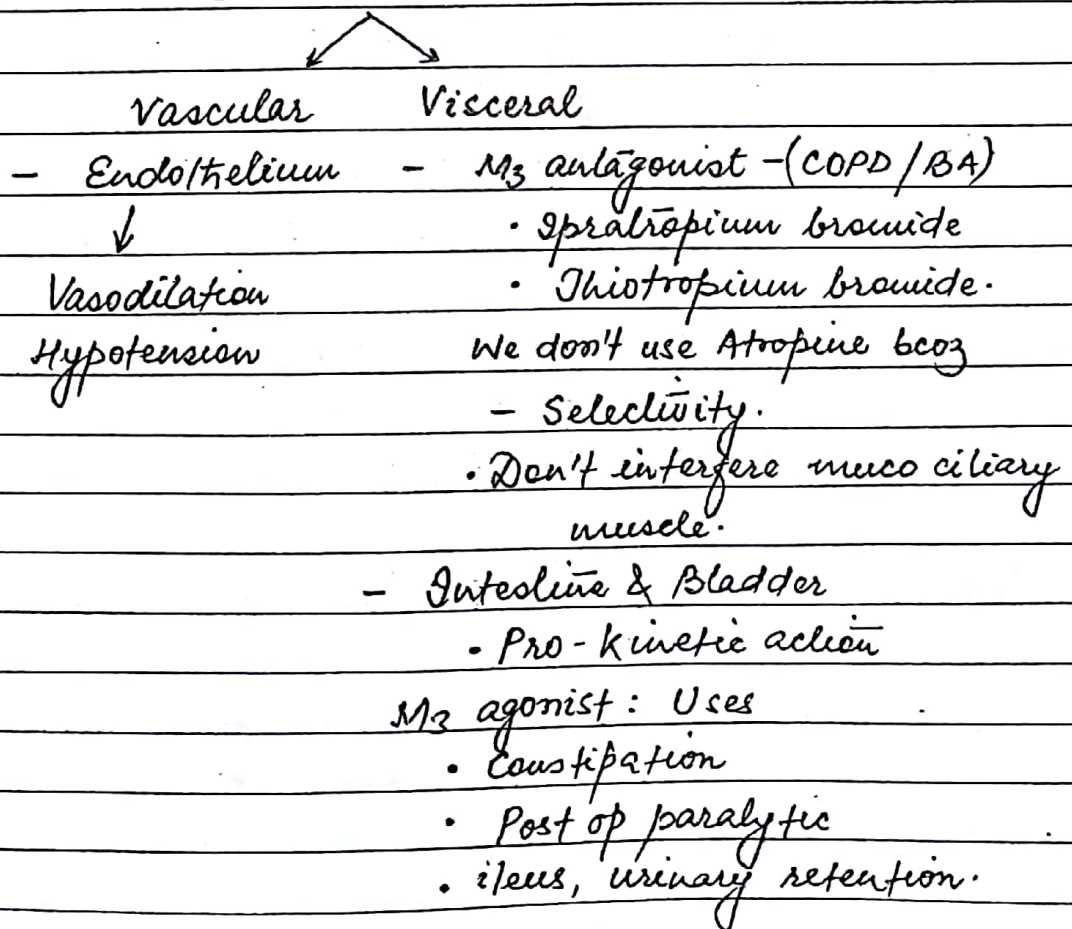
- Inhibit  $Na^+ - K^+$  ATPase test.
- Accumulate intracellular  $Ca^{2+}$  ( $\uparrow Ca^{2+}$ )
- $\uparrow$  Force of contraction
- Useful for t/t of low output CHF.

Muscarinic Receptors:

M3 Receptor - Location:

- Smooth muscle - Blood vessel (endothelium)
- Eye
- Endocrine glands.

Smooth muscle



Selective  $M_3$  agonist acting on Intestine & Bladder  
→ BETHANECHOL

Selective  $M_3$  agonist acting on GIT & Bladder

- DARIFENACIN

- SOLIFENACIN

- useful for t/t of diarrhoea &  
diarrhoeal dominant IBS.  
Overacting bladder.

Selective  $M_3$  agonist acting only on Bladder

- Vesico selective  $M_3$  agonist

• Oxybutynin

• Flavoxate

Active form →

- Tolterodine
- Fesoterodine (Prodrug)
- Trospium chloride.

Extra information on bladder:

$\beta_3$  Action - Relax detrusor - causing urinary retention

↓

MIRABEGRON ( $\beta_3$  agonist)

↳ Use - Overactive bladder.

Location of  $\beta_3$  mostly in adipose tissue

• SIBUTRAMINE ( $\beta_3$  agonist)

- Adipolysis (wt. loss)

- It is withdrawn - Ecoz Cardiotoxic.

# Nocturnal enuresis

- Imipramine (TCA)

• Anti cholinergic

DOC: Desmopressin

$V_2$  analogue - Vasopressin

Stress incontinence:

t/t → Duloxetine

- ↑ urethral tone

- also useful for t/t

- Chronic neuropathy pain

- Fibromyalgia.

- It is SNRI (Anti-depressants)



eg: Duloxetine

Venlafaxine (S/E - Sustained HTN)

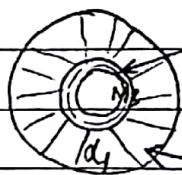
Milnacipram

Leva-milnacipram

Vilazodone

Vortioxetine } Newer drug.

$M_3$  on Eye:



sphincter muscle

Constrictor

Stimulation of  $M_3$

- Constriction of pupil  
(Miosis)

Radial muscle

Dilator

Stimulation of  $d_1$ :

- Mydriasis

↳ On Radial muscle.

$M_3$  agonist acting on eyes

- Pilocarpine

- Ecothiophate

- Organophosphorus Comp<sup>d</sup>

Irreversible cholinesterase inhibitor



$\alpha_1$  agonist acting on eyes:

- Phenylephrine

(Adrenergic agonist)

Adrenergic drugs - Only Mydriasis

Anticholinergic drugs - Mydriasis + Cycloplegia  
(loss of light reflex)

#  $\beta$ -blocker don't alter pupil size

Timolol - Use in t/t of Glaucoma.

# Oculomotor Nerve supplies constrictor muscle.  
(Circular muscle).

Causes Miosis.

Injury - Mydriasis

Even after CN III nerve injury if we use pilocarpine we will get miosis, as receptors are intact.

#  $M_3$  ~~receptor~~ agonist - Useful for glaucoma.

Pilocarpine - Useful for glaucoma by promoting drainage

Ecothiophate - s/e - Cataract.

Mydriatic anticholinergic:

Atropine (longest acting = 1wk)

Homatropine

Cyclopentolate

(M/c) Tropicamide (Fastest but shortest acting = 3-6hr)

↳ CI - Glaucoma.

Only for fundus exam - Mydriasis enough

↓  
Phenylephrine preferred

(OR)

Tropicamide.

Error of Refraction:

• Mydriasis & Cycloplegia  
DOC - Tropicamide

• In child < 5yr

• Atropine Ointment 1%

M<sub>3</sub> on exocrine glands:

M<sub>3</sub> location - Salivary gland  
Lacrimal gland  
Sweat gland.

M<sub>3</sub> agonist: Pilocarpine  
Cevimeline

Sjogren syndrome - Pilocarpine used  
Xerostomia

# Amifostine - Radio protective

↓  
Antidote for Cisplatin

↳ S/E - Nephrotoxicity.

# Radio sensitizer - Gemcitabine, Metronidazole.

Radiation Recall - Dactinomycin, Doxorubicin  
- Anticancer antibodies

Gemcitabine:

Pyrimidine anti-metabolite  
DOC - Pancreatic Cancer.

# Atropine - CI in hyperthermia

Nicotinic Receptors:

Nm & Nm

Nm:

N = Nicotinic, m = Skeletal muscle

- ① Activation of Nm causes opening of  $Na^+$  &  $Ca^{2+}$  channel.  
Entry of  $Ca^{2+}$  causes contraction of muscle.  
(Muscle depolarisation)

Ach - ↑ muscle power

So, cholinergic drugs used for Ht for Myasthenia gravis.

Skeletal muscle Relaxation (SMR):

α - Tubocurarine = Competitive antagonist.

↳ Non depolarising SMR.

For reversal - Neostigmine

̄ Atropine

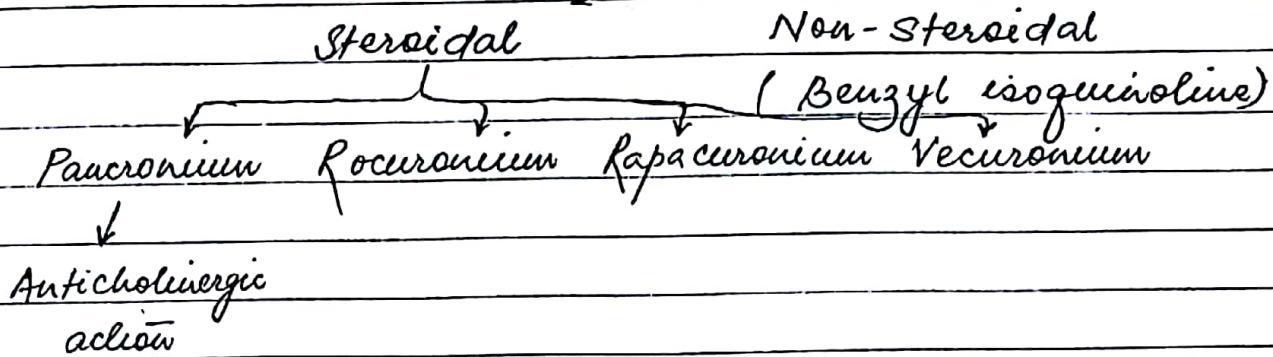
Newer drug - Sugammadex

↓

Useful for Reversal of Rocuronium & Vecuronium.

• Similar to Neostigmine

## Non-depolarizing SMR



(or) Anti Vagal.

# Glycopyrrolate: Anticholinergic agent

Useful for pre anaesthetic medication to control Secretion.

It is quaternary comp<sup>d</sup> - lipid insoluble,  
So, no CNS side effect. So it is useful  
instead of Atropine.

Rocuronium:

- Fastest acting SMR
- Alternate to Succinyl choline (Sch) for Tracheal intubation
- Least histamine releasing property.
- Severe pain during injection

Rapacuronium:

- Cause Severe Bronchospasm.

Vecuronium:

- Preferred in cardiac pts.

## Benzyl isoquinoline

Doxacurium

- Longest acting (120min)
- Most potent.

Mivacurium

- Shortest acting (15-21 min)
- Useful for day care. Sx.

Atracurium

- Undergoes Hoffman's degradation (Self metabolism)

d-Tubocurarine

- Max<sup>m</sup> Histamine Releasing
- Adverse effect
  - Bronchospasm
  - Hypotension.

Gantacurium

(5-10 min)



Newer drug.

metabolism cont

liver & kidney.

They do not

need enzyme

for degradation

- Safe in Hepatic/ Renal failure

- Produce by product



Laudanosine

(causes - Seizure)

# Cis Atracurium - Less laudanosine

Less secreting histamine.

SMR having less histamine releasing property

- Cis. Atracurium

- Rocuronium.

Depolarising SMR:

Succinyl choline (Sch):

Structurally & functionally - similar to Ach.

S/E - Muscle fasciculation

Post op. muscle pain

- Shortest acting (3-5 min)

rapidly undergo metabolism by Pseudocholine esterase.

Some people have Atypical Pseudocholine esterase

↑ action < 5 min

Lead to Sch Apnoea

T/t - Fresh blood transfusion becoz blood plasma is rich in pseudocholine esterase.

Dubucaine number:

Useful to assess whether the pt. have atypical pseudocholinesterase or normal.

Caine - Local anesthetic agent.

80% - hydrolysis - Normal Pseudocholinesterase.

<20% - hydrolysis - Atypical "

Adverse drug effect of Sch:

- Hyperkalemia (Burus), nerve injury, crush injury

- Malignant hyperthermia

- ↑ Intra ocular/gastric pressure

these who are having genetic abnormality c̄  
Ry anadine receptor.

Primaquine - Causes hemolysis only in G6PD deficiency.

Pharmacogenomic / Idiosyncrasy - Ryanodine Receptor



Occurs disease in only genetic ab<sup>o</sup> person.

T/t → Dantrolene

(Directly acting SMR)



DOC for : Malignant hypothermia  
Neuroleptic malignant Syndrome

# SMR - causes pain on injection - Rocuronium.

GA causing pain " - Propofol

Post-op muscle pain - Schw

Analgesic used during Sx causing Post-op truncal rigidity - Fentanyl, Alfentanil

T/t - Wooden chest Syndrome.

Antibiotics causing SMR:

- Aminoglycosides (Max<sup>m</sup>) - Neomycin
- Macrolides
- Quinolone
- Tetracyclines

# Aminoglycosides - Inhibit Release of ACh

Similar to Botulism toxin.

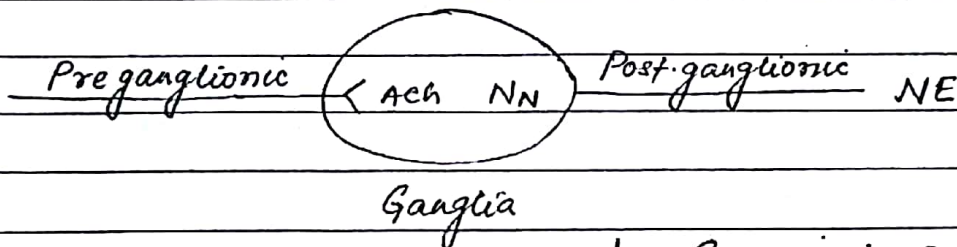
T/t - Neostigmine + Calcium.

NN :

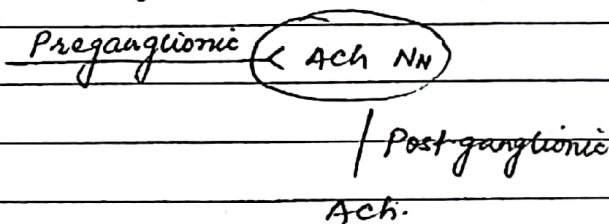
Location: Autonomic ganglia (Most)  
Adrenal medulla  
CNS

Autonomic ganglia -

Sympathetic:



Parasympathetic:



Ganglionic Blockers (NN)

- Hexamethonium
  - Trimethaphan
  - Mecamylamine → (Smoking control)
- useful to produce controlled hypotension.

Anti smoking drugs:

First line drug (therapy)

- Varenicline ( $\alpha_4\beta_2$  nicotinic agonist) - Suicidal thoughts
- Nicotine (patch, inhaler, lozenges, Chewing gum)
- Bupropion - NDRI (Norepinephrine Dopamine Reuptake Inhibitor)  
Antidepressant      Adverse drug reaction  
Weight loss      - Seizure.  
ADHD (off label)

Second line therapy:

Clonidine ( $\alpha_2$  agonist)  
Nortriptyline (TCA)



Page No.

Date: / /

## Miscellaneous:

Rimonabant

Topiramate - Antiepileptic

ADR - Weight Loss, Nephrolithiasis.

Mecamylamine

Rimonabant: Inverse agonist/ Antagonist of Cannabinoid  
1 receptor.

- Weight Loss

- Prevents craving of alcohol.

ADR - Psychiatry problems (withdraw)

ADHD (Attention deficit hyperactivity disorder):

Drug used - Amphetamine



Causes - Cardiotoxic

Addiction

Appetite Suppressant.

(Failure of growth)

## First line drugs:

- Methylphenidate (First choice)

- Atomoxetine

Ritalinic acid (Metabolite).

## Other drugs:

~~Pb.~~ Pemoline (Hepatotoxic)

Modafinil - Use: Narcolepsy

Shift worker

Obstructive Sleep apnea.

ADHD. (FDA - Unapproved)

Newer drug under Narcolepsy:  
 $H_3$  inverse agonist



Pitolisant (OR) Tiprolisant

Narcolepsy (Orphan drug status)

Drug useful for t/t of obesity:

- Sibutramine ( $\beta_3$  agonist) - Cardio toxic (Withdrawn)
- Orlistat (lipase inhibitor) - Steatorrhea
- Olestra (Sucrose polyester) - cooking medium.
- Rimonabant (Cannabinoid 1 antagonist) - Withdrawn
- Leptin (Endogenous slimming peptide)

Combination therapy:

Bupropion + Naltrexone (opioid antagonist)

Bupropion + Zonisamide (Antiepileptic)

Phenteramine + Topiramate (Antiepileptic)

(Sympathetic stimulant)

~~Crossing~~ Causing

Apetite suppressant)

Newer drug: 5HT<sub>2C</sub> agonist - LORCASERIN  
 S/E - Serotonin Syndrome.

GLP-1 → LIRAGlutide

FDA approved drug for obesity.

Extra point: Antiepileptic causing wt. loss

- Topiramate
- Zonisamide
- Felbamate

Antiepileptic causing wt. gain:

- Sodium Valproate
- Gabapentin

# Felbamate  $\leftarrow$  Hepatic failure (SE)  
Aplastic Anemia.

# Type 2 DM  $\bar{c}$  obesity — 1st line drug — Metformin  
Non-diabetic  $\bar{c}$  obesity — No Metformin.

# Antidiabetic causing:

Weight gain: — Insulin, Insulin secretagogues.  
— Sulfonyl ureas, meglitinides,  
Thiozolidindiones.

Weight loss — Pramlintide, GLP-1 agonist, SGLT-2 inhibitors.

Weight neutral — Metformin, DPP4 inhibitors.

OPC's = Organophosphorus compounds.

Page No.

Date: / /

## ANTI CHOLINESTERASE

Reversible

Irreversible

Carbamates

Acridine

OPC's

Carbamate

→ Physostigmine

→ Tacrine

→ Dyflos

→ Carbaryl

(Natural origin)

↓

→ Echothiophate

→ Propoxur

Alkaloid (plant)

Hepatotoxic

→ Parathion

(Baygon)

• Highly lipid soluble

So, not used

→ Malathion

Insecticide

in Alzheimer's

→ Diazinon

DOC: Atropine

→ Tabun

Nerve gas

poisoning

→ Sarin

or,

(Belladonna)

→ Soman

War gas.

→ Neostigmine

Pyridostigmine

Edrophonium

(Water soluble)

No CNS effect.

Neo - direct action

on NM receptor

Pyri - long acting

Orally active

Edro - Anionic site binding

• Rapid dissociation

• Used for Δ of

myasthenia gravis.

(Tensilon test

or, Ameliorative test)

- Provocative test

(done by injecting

d-Tubocurarine)

Malathion - Pediculosis (lice)

infestations

Echothiophate

Use in Glaucoma

S/E Cataract

# Aging of enzyme

Tabun (Slow)

Sarin (3-5 hrs)

Soman (2 min) - Fastest acting

t/t - Atropine + Pralidoxime

In convulsion - Diazepam

Rivastigmine } useful for t/t  
 Donepezil } of Alzheimer's ds  
 Galantamine }  
 ↓  
 deficiency of Ach.

### ■ OPC's poisoning:

Parathion, Malathion, Diazinon  
 Cholinesterase inhibitors  
 (Irreversible)

1st line DOC: Atropine (Muscarinic Blocker)



Dose & depends upon Sign & Symptoms of Atropinisation:

- HR > 100/min
- Pupil Size
- Pulmonary Secretion
- Secretion

Max<sup>m</sup> upto - 200 mg.

### Oximes:

- Cholinesterase ~~in~~ reactivators.
- Only used for t/t OPC's poisoning  
not carbamate poisoning.

eg: • Pralidoxime (1-2g; slow i.v., 15-30 min)  
 • Obidoxime (more potent)  
 • Diacetyl mono oxime (Highly lipid Soluble)  
 ↳ More CNS action

S/E - HTN

↳ T/t - Phentolamine (Non-selective α blocker)

Myasthenia Gravis (MG):

Ameliorative test

Provocative test

Definitive test → Anti Ach Receptor Radioimmuno Assay.

Confirmatory → Single fibre Electro Myography.  
(SF-EMG)

First line drug — Neostigmine  
Pyridostigmine

Others — Corticosteroids  
Thymectomy  
Plasmapheresis  
Iv Ig. } To remove  
autoantibody.

Other immunosuppressant — Azathioprine  
Cyclosporine.

Monoclonal antibody — Rituximab

↓  
Target CD20.

Remission/Exacerbation:

Rapid Recovery — Plasmapheresis  
Iv Ig.

Quinine — CI in MG  
— It is SMR  
— Used in Nocturnal leg cramps.

- Avoid Aminoglycoside in MG.

# MEMANTINE - NMDA Blocker

useful for moderate to severe Alzheimer's ds.

# Drug useful in cervical ripening - VALATHAMATE



Anticholinergic drug  
Smooth muscle relaxant.

# Diphenoxylate - Opioid

Anti diarrhoeal  
Addiction

↳ Atropine ↓ addiction of Diphenoxylate

# Glycopyrrolate - Anticholinergic

Premesothetic  
Quaternary Comp<sup>d</sup>.

# Scopolamine - Also k/A Hyosine → CNS depressant (Sedation)

↳ Used in motion sickness.

DOC: Hyosine → Narco Analysis

# 1st Gen. (H<sub>1</sub>) + (M): Promethazine

↓	↓	↓
In treating Allergic cond <sup>n</sup>	In Motion Sickness	treating EPS (Extra pyramidal Symp <sup>t</sup> )

# For Sea Sickness - Same t/t.

↳ Meclizine - 1st gen. long acting Anti-histamine.

For Mountain sickness: Acetazolamide  
(Carbonic Anhydrase Inhibitor)

Morning sickness: Doxylamine + Vit B<sub>6</sub>  
↓  
antiemetic Vitamin

Vit B<sub>6</sub> (or Pyridoxine):

- Anti-emetic
- Controls intrauterine seizure.

Stimulant of dopa decarboxylase  
C/I - Levodopa

Vit B<sub>6</sub> should not be given + levodopa.

Vit B<sub>6</sub> definitely given + Anti TB drug (Isoniazid)

↓  
To correct peripheral neuropathy.

Antidote for Vit B<sub>6</sub> - 4 deoxy pyridoxine

Folic acid -

Prophylactic - 400 µg daily in pregnancy.

Previous H/O Neural tube defect - 5mg/day.

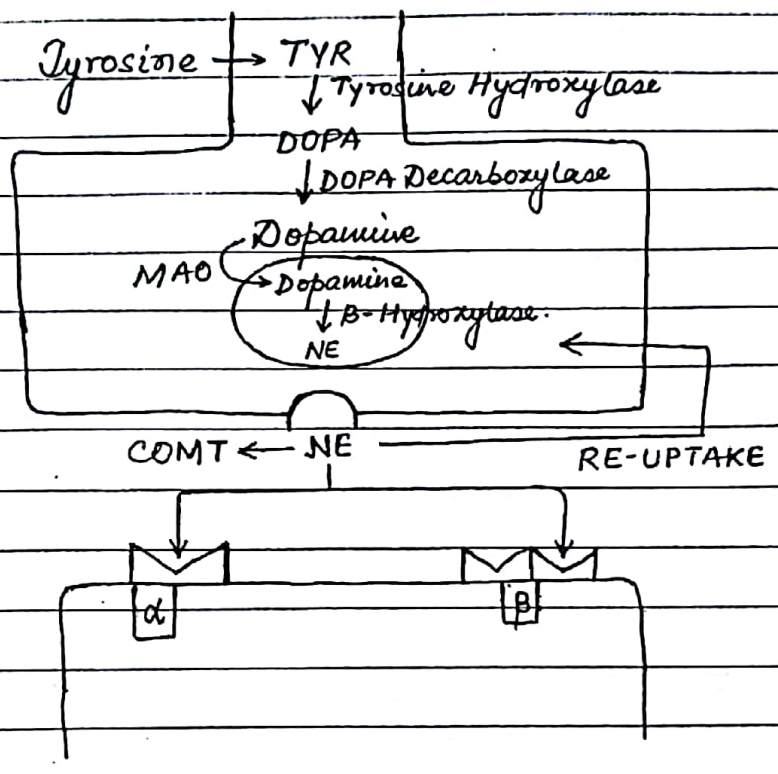


Drug having Anticholinergic activity:

- TCA's
    - Amitriptyline
    - Imipramine - Nocturnal enuresis
  - DOC: Desmopressin
  
  - Anti Psychotics
    - Thioridazine
    - Clozapine
  
  - SMR
    - Pancuronium
    - Gallamine
  
  - Class Ia Anti arrhythmic drugs.
    - Quinidine
    - Procainamide
    - Disopyramide (Highest anticholinergic property).
  
  - 1st H<sub>1</sub> Blocker
    - Promethazine
  
  - Amantidine
    - Meperidine (Pethidine)
      - ↳ opioid analgesics
      - ↳ CI in MI pain
- ↓
- Morphine is Used.

### ADRENERGIC DRUGS

#### Synthesis, storage, Release, Metabolism of NE:



# Synthesis of NE → Only in the vesicle.

Catecholamine - Dopamine  
 NE  
 Epinephrine

Monoamines - Dopamine  
 NE  
 Serotonin.

For metabolism of NE - MAO  
 COMT

- Even though NE undergoes metabolism by MAO & COMT, enzymatic degradation is not involved in termination.
- NE action is terminated by Re-uptake.
- Rate limiting enzyme of Synthesis of NE - Tyrosine Hydroxylase.
- Drug inhibiting Tyrosine hydroxylase - Alpha methyl para tyrosine (METYROSENE)
- Dopa decarboxylase inhibitor - Carbidopa Benserazide.
- Reserpine  $\rightarrow$  Anti HTN agent  
 $\rightarrow$  Vesicular uptake inhibitor.  
 - SE  $\rightarrow$  Suicidal depression.
- $\beta$ -hydroxylase blocker - Disulfiram (Used in alcoholism deaddiction)

Ethyl alcohol

$\downarrow$  Alcohol dehydrogenase

Acetaldehyde

$\downarrow$  Acetaldehyde dehydrogenase  $\leftarrow$  Disulfiram.

Acetic Acid

New drug - DROXIDOPA

( Prodrug of NE)

- Used in Neurogenic Orthostatic hypotension
- Hemodialysis induced hypotension.

BRETYLIUM : Class III drug

$K^+$ - Channel blocker.

Also called Chemical defibrillator.

Release of NE is blocked by - Bretylium  
Guanethidine.

NE Re-uptake inhibitor - SNRI, NDRI, TCA, Cocaine.

Cocaine  $\rightarrow$  One & only <sup>local</sup> anesthetic causing HTN.

- Causes mydriasis by acting on  $\alpha_1$  on the radial muscle.

Adrenergic Receptor:  $\left\{ \begin{array}{l} \alpha \\ \beta \end{array} \right.$

( Henry Ahlquist)

$\alpha$ - Receptor:  $\left\{ \begin{array}{l} \alpha_1 \rightarrow \text{post-synaptically (location).} \\ \alpha_2 \rightarrow \text{pre-synaptically} \end{array} \right.$

- $\rightarrow$  Inhibition of release of NE.
- $\rightarrow$  auto receptor for NE

$\alpha_2$  agonist:

eg: Clonidine } Centrally acting Anti HTN  
Methyldopa }  
Guanafacine }

Guana benz  
Moxonidine  
Rilmenidine

S/E - Drowsiness  
↓  
Not safe in children.  
Apraclonidine } Useful in Glaucoma.  
Brimonidine } MOA - decreases Aqueous Secretion

Tizanidine → Centrally acting SMR.

Dexmedetomidine → Used as Sedation (ICU pts) & Pre-anesthetic medication.

# Methyl dopa : DOC for t/t of HTN during pregnancy.

Hypertensive Emergency :

Labetalol ( $\beta + \alpha$  blocker)

Hydralazine ( $K^+$  channel opener)

↳ Arteriolar dilator.

Eclampsia -  $MgSO_4$ .

# Methyl dopa may cause hemolytic anemia to mother

↓

Coomb's test +ve

Drug avoided in pregnancy : ACEi (Renal & pulm agenesis)  
ARBs

Sodium nitroprusside  
(contains Cyanide)

# Apraclonidine : Specific S/E - ~~eye~~ lid lag.

Brimonidine : S/E - Anterior uveitis.

$\alpha_2$  antagonists:  $\uparrow$  NE release.

Yohimbine - Used in Hypotension & Sexual stimulation

Idazoxan

$\alpha_1$ :

location - Post synaptically.

①  $\alpha_1$  seen in vascular smooth muscle.

Action  $\rightarrow$  Vasoconstriction

$\alpha_1$  agonists:

Based on vascular action

Useful in t/t of Hypotension

Nasal congestion.

Selective  $\alpha_1$  agonists for t/t for Hypotension:

Methoxamine

Mephenteramine

Midodrine.

Selective  $\alpha_1$  agonist for t/t for Nasal congestion:

Cause Atrophic Rhinitis

(Rhinitis medicamentosa)

Naphazoline  
Oxymetazoline  
Xylometazoline.

#  $\alpha_1$  Receptor - Radial muscle of iris  $\rightarrow$  Mydriasis  
 $\hookrightarrow$  Phenylephrine

#  $\alpha_1$  Receptor seen in internal urethral sphincter

$\hookrightarrow$  Causes sphincter constriction

$\hookrightarrow$  Retention of urine.

$\alpha_1$  blocker used in BPH

# Vesico Ureteric junction  $\alpha_1$  Receptor +mt.

#  $\alpha_1$  blocker Useful in t/t of - lower ureteric calculi

#  $\alpha_1$  seen on Vas deferens of penis.  
Action  $\rightarrow$  Ejaculation.

# S/E of  $\alpha_1$  Blocker - Impairment of Ejaculation.

# Directly acting Sympathomimetic  
 $\alpha, \beta$  agonists  
Adrenaline, NA.

Indirectly acting Sympathomimetic:

Tyramine  $\rightarrow$  Act on vesicle  $\rightarrow$  Causes release of NE.  
 $\downarrow$   
Causes depletion of storage of NE  
 $\downarrow$   
Tachyphylaxis  $\rightarrow$  Rapid tolerance

# MAO inhibitors taking  $\bar{c}$  Tyramine containing food (cheese, wine, bread) causes HTN, it is called Cheese react<sup>n</sup>.

$\downarrow$   
DOC for t/t of HTN due to cheese react<sup>n</sup>: Phentolamine  
(non-selective  $\alpha$  block)

# Mixed action Sympathomimetic - EPHEDRINE

$\downarrow$   
causing hypotension  
 $\downarrow$   
Spinal anaesthesia.  
Safe in pregnancy.

Selective  $\alpha_1$  blocker:

eg: Prazosin (PDE inhibition property).  
 Doxazosin } Apoptotic action on Prostate.  
 Terazosin }

$\alpha_1$  A blockers  
 ↓  
 mainly acting on bladder.

Silodosin  
 Alfuzosin  
 Tamsulosin

Indoramine } useful in Hypertensive Emergency.  
 Urapidil }

PRAZOSIN:

- Vasodilation → on smooth muscle.

Uses - HTN

PVD

CCF

Scorpion Bite.

S/E - Postural hypotension

(1st dose hypotension)

- Impairment of ejaculation.

Selection of Prazosin as Anti-HTN:

① HTN  $\bar{c}$  dyslipidemia

② HTN  $\bar{c}$  elderly male  $\bar{c}$  BPH.

③ Can be used in diabetics  $\bar{c}$  HTN.

HTN  $\bar{c}$  dyslipidemia:

Choice - Prazosin

Anti HTN avoided - Non-selective  $\beta$ -blocker

Thiazide ~~the~~ diuretics



No problem  $\bar{c}$   $\rightarrow$  CCB, ACEi, ARB, clonidine.

HTN  $\bar{c}$  diabetics:

Choice  $\rightarrow$  ACEi = ARB  $>$  CCB

Unfavourable (avoid)  $\rightarrow$   $\beta$ -blocker  
Diuretics.

Anti-HTN causing Erectile dysfunction -

Highest risk - Diuretics (Thiazides)

High risk -  $\beta$ -blocker (Atenolol, Carvedilol,

# In BPH  $\rightarrow$  Static obstruction is overcome by  
Finasteride + Tamsulosine.

$\downarrow$  (Rapid Benefit)

It takes 3-6 months for action.

Tamsulosine overcomes dynamic obstruction.

# Pt. on Tamsulosine <sup>or</sup> may cause risk of floppy iris  
syndrome  $\rightarrow$  going for cataract.

Non-selective  $\alpha$ -blocker:

Irreversible - Phenoxybenzamine

Reversible - Tolazoline, Phentolamine.

PHENOXYBENZAMINE:

# Definitive therapy for t/t of HTN in Pheochromocytoma - Phenoxybenzamine.

# For controlling intra-operative HTN during pheochromocytoma Sx - i.v. Phentolamine  
i.v. Nitroprusside.

# Don't use Propranolol as a 1st line drug for t/t HTN due to Pheochromocytoma.

# In Pheochromocytoma Sx - ~~Drug~~ Halothane is C/I

↓  
sensitize the myocardium  
for catecholamine

↓  
Causes MI.

Phentolamine:

Use - DOC for t/t of Clonidine withdrawal HTN

DOC for t/t of HTN due to Cheeze reac<sup>n</sup>.

In intra-op HTN during Pheochromocytoma Sx

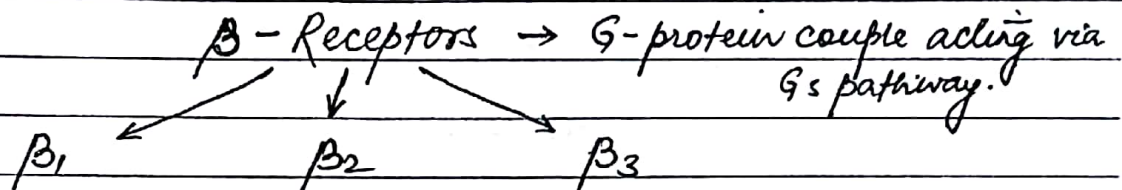
Oxime induced HTN.

Useful for t/t of Erectile dysfunction (injectable drug)

PIPE Therapy (Pharmacologically induced penile erection):

Injectable drugs used for t/t of erectile dysfunction:

- Alprostadil (PGE1 analogue)
- Phentolamine
- Papaverine (Non-selecting PDE inhibitor).



β<sub>1</sub> →

Location - Myocardium  
Kidney.

Action (Heart) → ↑ HR

↑ Force of contraction  
↑ C.O.

In kidney → Renin release.

Selective β<sub>1</sub> agonist:

Dobutamine (Synthetic Catecholamine).

# eg. of synthetic Catecholamine

- ① Isoprenaline → acting on β<sub>1</sub>, β<sub>2</sub>, β<sub>3</sub>
- ② Dopexamine → D<sub>1</sub>, β<sub>2</sub>
- ③ Dobutamine → β<sub>1</sub> (t<sub>1/2</sub> = 2 min)
- ④ Fenoldopam → D<sub>1</sub>

# Dobutamine Used in → Stress ECHO

# D<sub>1</sub> receptor seen in Renal blood vessel → Renal vasodilation

∴ Fenoldopam Used in → iv infusion  
 • HTN emergency & Renal impairment.

$\beta_2$ :  
 Location: Smooth muscle & Vascular  
 Visceral.

Stimulation of  $\beta_2$  → Vasodilation.

Visceral —

Bronchial muscle → Bronchodilation.

$\beta_2$  agonist useful for t/t of Bronchial Asthma:

Salbutamol } short acting

Terbutaline } Useful for Acute asthma.

Salmeterol

Formoterol } long acting

Indacaterol } Useful for Chronic asthma

Salbutamol:

M/C S/E — Tremors

Palpitation.

Uterus → Action → Uterine muscle relaxation.

Toxolytic — Ritordine (FDA approved)

Isoxuprine

#  $\beta_2$  agonist having anabolic action — Clenbuterol.

Phospholipase-Gg ←  $\alpha_1$   
Adenyl cyclase-Gi ←  $\alpha_2$  ] - G-Protein Couple receptor.

37

Page No.

Date: / /

## $\beta_2$ - Role on metabolism

↓                      ↓                      ↓

Carbohydrate      Potassium      Lipid  
- Hyperglycemia    - Hypokalemia    - Reducing blood cholesterol.

### Hyperkalemia:

Mild → 5.5 to 6.5 mEq/L

Moderate → 6.5 to 8.0 mEq/L

Severe → > 8.0 mEq/L

For Rapid control of potassium in Hyperkalemia (emergency) - Insulin + Glucose infusion.

For Hyperkalemia + ECG abnormalities - Calcium Gluconate.

### $\beta_3$ :

Location: Adipose tissue

Selective  $\beta_3$  agonist - SIBUTRAMINE

- lipolysis

- withdraw due to Cardiotoxic.

### MIRABEGRON:

-  $\beta_3$  agonist

- Relax detrusor

Used in - Overactive bladder.

Q Which one of the following don't have significant ~~dopaminergic~~ dopaminergic activity -

A) Dopamine ( $D_1, B_1, \alpha_1$ )      C) Fenoldopam ( $D_1$ )

B) Dobutamine ( $B_1$ )      D) Dopexamine ( $D_1, \beta_2$ )

Dopamine: has  $D_1$ ,  $\beta_1$ ,  $\alpha_1$  action.

$\downarrow$              $\downarrow$              $\downarrow$   
 $< 2 \mu\text{g/kg}$     2-5            5-10  $\mu\text{g/kg}$ .

DOC for Cardiogenic Shock - Dopamine.

Shock	T/t
Cardiogenic	NE or Dopamine
Cardiogenic $\bar{c}$ oliguria	Dopamine.
Anaphylactic	Adrenaline
Secondary	$\alpha$ -blocker
Adrenal insufficiency	Steroids

#. Blood pressure:

$BP = CO \times \text{Peripheral resistance.}$

$\downarrow$      $\downarrow$   
 SBP    DBP

Effect of Isoprenaline on BP:

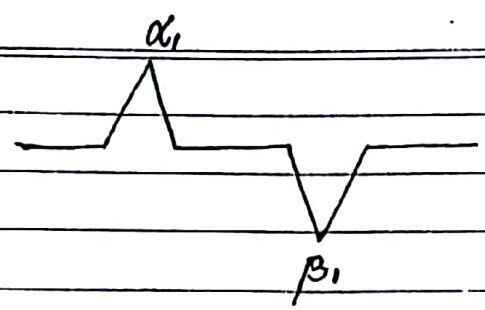
- $\beta_1$ ;  $\beta_2$ ,  $\beta_3$  action.
- No  $\alpha$  action.
- $\uparrow$  SBP;  $\downarrow$  DBP  $\rightarrow$  Reflex Tachycardia
- Wide pulse pressure.

NA:  $\alpha_1$ ,  $\alpha_2$ ,  $\beta_1$

No  $\beta_2$  action

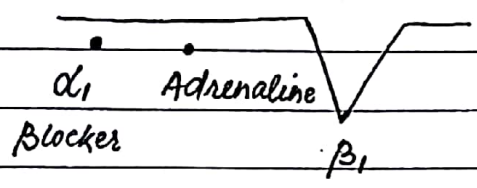
$\uparrow$  SBP;  $\uparrow$  DBP  $\rightarrow$  Reflex bradycardia

Adrenaline on BP: acting on  $\alpha_1$ ,  $\alpha_2$ ,  $\beta_1$ ,  $\beta_2$



Biphasic response of Adrenaline on BP.  
 - Adrenaline cause initial ↑ BP & later ↓ BP.

# Dale's vasomotor reversal phenomenon:



If we give  $\alpha_1$  blocker before adrenaline, adrenaline acts only on  $\beta_2$  causing fall in BP.

Q. All are lipid insoluble  $\beta$ -blocker except?

- A) Nadolol
- ~~B) Propranolol~~
- C) Atenolol
- D) Sotalol

lipid soluble  $\beta$ -blocker - Propranolol (Highly soluble)

• M/coumnest drug used for prophylaxis of migraine.

- DOC {
- Performance anxiety
  - Essential tremor
  - Akathesia

Lipid insoluble  $\beta$ -blocker - Nadolol (Most longest acting >40hrs)  
 Atenolol

Long duration of action Sotalol

No hepatic metabolism

Unsafe in Renal failure - Dose adjustment required.

## $\beta$ -blocker

Non-selective  $\beta$ -blocker: 1st generation  $\beta$ -blocker

- Drug block both  $\beta_1$  &  $\beta_2$ .

Cardioselective  $\beta$ -blocker: 2nd generation  $\beta$ -blocker  
(Predominantly blocks  $\beta_1$  blocker)

- No selective  $\beta_2$  blocker.

# 3rd generation  $\beta$ -blocker -  $\beta$ -blockers  $\bar{c}$  additional properties.

Cardioselective  $\beta$ -blocker:

Nebivolol (Most Cardioselective; Releases NO)

Vasodilation

Betaxol - Useful in Glaucoma; Safe in asthmatic.

Bisoprolol - Useful in CCF

Atenolol

Esmolol - Most ultra short acting ( $\approx 9$  min), i.v., Emergency.

Acebutolol

Metoprolol - Useful in HTN, Angina, MI, CCF.

Celiprolol

3rd generation  $\beta$ -blocker:

①  $\beta$ -Blocker having  $\alpha$  blocking property -

Labetalol -  $\beta$  &  $\alpha$  blocker

- Use  $\rightarrow$  HTN emergency in pregnancy.

- S/E  $\rightarrow$  Postural hypotension, hepatotoxic.

~~Carbi~~

Carvedilol -  $\beta$  &  $\alpha$  blocker

- Antioxidant

- USE  $\rightarrow$  in CCF.  $\rightarrow$  Bisoprolol

Metoprolol.



②  $\beta$ -blocker having NO releasing property -  
 Nebivolol  
 Nipradilol

③  $\beta$ -blocker having  $K^+$  channel opening action -  
 Tilisolol

④  $\beta$ -blocker having  $K^+$  channel blocking property -  
 Sotalol - Class III antiarrhythmic group.

# BUTOXAMINE:

- Only selective  $\beta_2$  blocker
- Used for research purpose, not for therapeutic purpose.

$\beta$ -blocker having highest uremic<sup>r</sup> stabilizing

↓

$Na^+$  channel blocking property  
 or local anesthetic action.

→ Propranolol.

$\beta$ -blocker having highest intrinsic sympathomimetic  
 → Pindolol

$\beta$  blocker having favourable effect on lipid profile  
 → Pindolol.

Antidote for  $\beta$  blocker poisoning - Glucagon.



A/c Joint National committee guidelines

First line drugs used in t/t of HTN:

- Thiazides
- ACEI
- ARB
- CCB

↳ No  $\beta$  blockers.

⑤ Useful for Portal hypertension (Prophylaxis)

↓  
Propranolol

DOC for t/t of bleeding due to esophageal varices

- OCTREOTIDE

↓

most potent vasoconstrictor

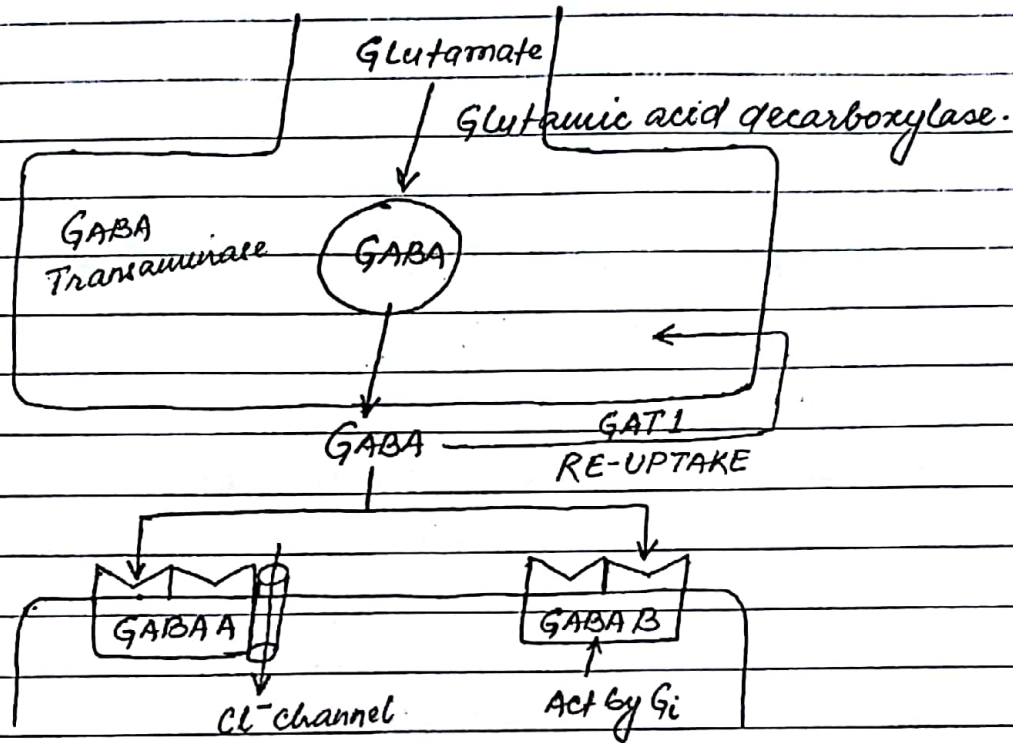
- controls bleeding

- Terlipressin -  $V_1$  agonist can be added.

DOC for prophylaxis - Propranolol, Nadolol.

## Central acting drugs

GABA:



Metabolism by - GABA transaminase.

Action of GABA: When GABA enters GABA<sub>A</sub>, Cl<sup>-</sup> channel enters causing hyperpolarization.

Drugs acting via GABA<sub>A</sub> pathway

Benzodiazepine Barbiturates.

BZD binding to BZD receptor which is made up of  $\alpha$ ,  $\gamma$  unit of GABA<sub>A</sub>.

BZD = GABA facilitatory.

↑ frequency of Cl<sup>-</sup> channel opening.

MOA of Barbiturates -

- Barbiturates binding to  $\alpha$ ,  $\beta$  units of GABA A.

Barbiturate: Low dose  $\rightarrow$  GABA facilitatory

High dose  $\rightarrow$  GABA inhibition.

$\uparrow$  duration of  $Cl^-$  channel opening.

Benzodiazepine (BZD):

Action (USE)  $\rightarrow$  Sedation

Anti-convulsion

Anti-anxiety

SMR.

Diazepam - DOC for Acute febrile seizure (Rectal Diazepam)  
Status Epilepsy (currently DOC - iv lorazepam)  
Delirium tremors.

Lorazepam - DOC for Status epilepsy.

Alcohol withdrawal: DOC: Chlordiazepoxide.  
(Delirium tremors)

Midazolam } short acting  
Remimazolam }  
 $\rightarrow$  Ultra short acting.  
 $\rightarrow$  Anaesthetic property.

Alprazolam - used in Insomnia, Anxiety disorder

Long term use of BZD - Addiction

Tolerance

Day time sleeping.

BZD safe in liver failure pt:

Temazepam  
 Oxazepam (Metabolite of Diazepam).  
 Lorazepam.

Sleep onset Insomnia:

Z compounds — Zolpidem (Most common)  
 ↳ Zopiclone  
 All are short acting — Zaleplon (Shortest)

FLUNITRAZEPAM: Date Rape drug.  
 Causes Anterograde amnesia.

KETAMINE: Also date rape drug.

BZD poisoning —

Antagonist:

Competitive antagonist — FLUMAZENIL

↓

prevent binding of BZD to  
 $\alpha, \gamma$  unit of GABA<sub>A</sub>.

- Specific antidote of BZD.
- Given i.v.
- $t_{1/2} = 60$  min

BICUCULLINE — Competitive antagonist of GABA  
 Non competitive inhibitor of BZD.

PICROTOXIN — Direct  $Cl^-$ -channel blocker.

# Inverse agonist of BZD Receptor -  $\beta$ -Carbolin

# Flumazenil used for - BZD poisoning  
 $\beta$ -carbolin poisoning  
 $Z$ -compound poisoning.

### BARBITURATES :

Long acting	Short acting	Ultrashort acting
- Primidone	- Secobarbitone	- Thiopentone Sodium
- Phenobarbitone	- Pentobarbitone	- Methohexitone.

Thiopentone sodium - Indication

- iv induction GA
- Re distribution
- Cerebro protective

Other uses - Narco analysis  
 Status epilepsy.

Methohexitone - causing convulsion.

Used in Electro convulsive therapy.

# Phenobarbitone - metabolite of Primidone.

↳ Useful in Anti convulsion in pregnancy & pediatrics.

↳ In children it causes hyperkinesia.

### General properties of Barbiturates:

- Algesic property (produce pain)
- Narrow therapeutic index. (Hence - unsafe)

↓ only  
 ∴ Used, in - Epilepsy  
 Anaesthesia

### Clinical manifestation of Barbiturates:-

- Flabby muscle
- Comatose
- Shallow & falling Respr
- Bullous eruption.

T/E:

- No specific antidote.

- Poisoning → Forced alkaline diuresis  
 Hemodialysis.

# All barbiturates are microsomal enzyme inducer.

Since powerful enzyme inducer

∴ C/I - acute intermittent porphyria.



### GABA analogues.

GABA Reuptake inhibitor: TIAGABINE

GABA Transaminase inhibitor: VIGABATRINE

SODIUM VALPROATE

Glutamic acid decarboxylase activator: VALPROATE

VIGABATRINE - DOC for infantile Spasm  
(Tuberous Sclerosis)

SE  $\left\{ \begin{array}{l} \rightarrow \text{Visual field defect} \\ \rightarrow \text{Psychosis} \end{array} \right.$

For Simple Infantile Spasm - ACTH

LEVATIRACETAM: ligand for SV2A protein



Synaptic Vesicle

- modify synaptic release of Glutamate/GABA.



Controls Seizure

New drug - GABAPENTIN } Useful in DM neuropathy pain,  
PREGABALIN } Post herpetic neuralgia.

### GANAXALONE

- Neurosteroid

- Direct  $\text{Cl}^-$  channel opener

Useful in - Absence seizure

Catamenial seizure.

GABA B (G-protein Coupled Receptor)

↳ Agonist - BACLOFEN

Antagonist - SACLOFEN

BACLOFEN - Centrally acting SMR  
Useful in - Hiccough  
Craving of alcohol.

MELATONIN:

Sleep inducing hormone  
Secreted from pineal gland.

Melatonin analogue - REMELTEON

MT1      MT2



Useful in sleep onset insomnia  
No risk of ABUSE / TOLERANCE.

TASIMELTEON - Useful in t/t sleep awake  
disorder in blind.



Melatonin analogue.

AGOMELATINE - Agonist on MT1/MT2

Antagonist on 5-HT<sub>2C</sub>

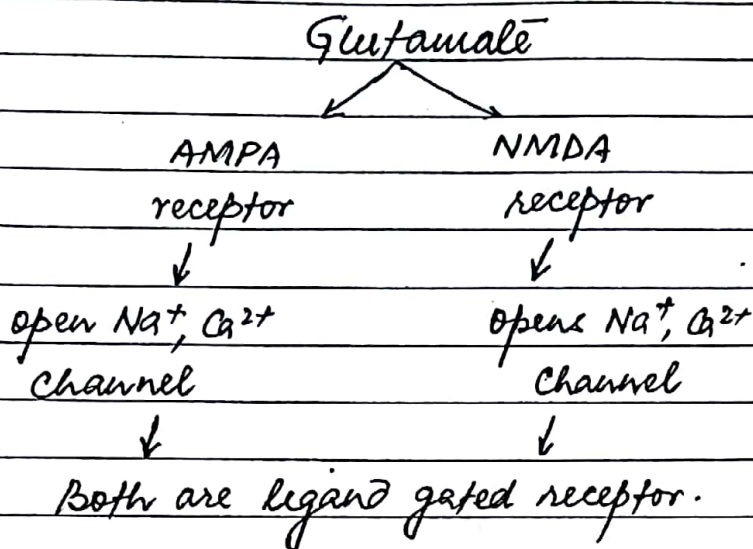
Melatonin analogue c

antidepressive property.

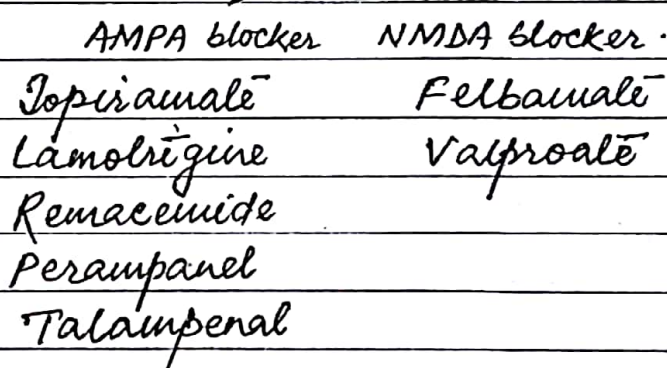
SUVOREXANT → FDA approved drug for insomnia.

ALMOREXANT → Non-selective OREXIN receptor  
antagonist.

↓  
another orexin receptor antagonist.



### # T/t of epilepsy - Glutamate antagonist



### Actions of Sodium Valproate:

- GABA agonism
- Anti glutamate
- Na<sup>+</sup> channel blocking action
- Ca<sup>2+</sup> channel blocking action
- Broad spectrum anti-epileptic.

### Lennox Gestalt Syndrome:

Rx → FELBAMATE - S/E - Hepatic failure  
Aplastic anemia.

Currently used

- ┌ VALPROATE
- ├ BZD
- └ RUFINAMIDE (Na<sup>+</sup> channel blocker)

## TOPIRAMATE :

Use → Epilepsy  
 Prophylaxis of Migraine  
 Alcohol (Anti craving)  
 Smoking ( " )  
 SE → Renal Stone  
 Wt. loss

## LAMOTRIGINE :

Useful in - Epilepsy  
 BPD depressive

Rarely cause SJS (Steven Johnson Syndrome).  
 TEN (Toxic epidermal necrolysis)

## NMDA blockers :

Anesthetic action {
 

- Ketamine : - Dissociative anesthesia
- Xenon
- N<sub>2</sub>O (laughing gas) → SE - Megaloblastic Anemia.
- Memantine → Useful in Alzheimers
- Acamprosate → GABA agonist property, Craving alcohol.
- Amantidine → Useful in Parkinsonism
- Methadone → DOC for opioid deaddiction.
- Riluzole → Useful for ALS
- Phencyclidine → Angel dust.

## Dopamine as a Neurotransmitter :

### Dopaminergic pathway :

① Meso-limbic fibre - extend upto prefrontal lobe  
secrete dopamine.

↑ dopamine - cause Psychosis

② Nigro-striatal neuron - (N) func<sup>n</sup> is to synthesise & release  
dopamine in corpus striatum.

- helps in initiation of movement.

In corpus striatum - amount of ACh & Dopamine  
balanced.

As ↑ age - adequate amount of dopamine is not  
secreted & there is ↑ in ACh activity.

Muscle rigidity occurs due to ↑ ACh.

- Hypokinesia, Tremor, ~~Rigidity~~ Rigidity.

③ Tubero infundibular fibre - extend from hypothalamus  
to anterior pituitary.

- Dopamine analogue are used for t/t of  
galactorrhoea.

- Dopamine act on D<sub>2</sub> receptor in the brain  
& causes psychosis.

- Any drug blocking D<sub>2</sub> & causing anti psychotic  
effect is called ATYPICAL ANTIPSYCHOTIC.

# Two most common SE of antipsychotic ← EPS  
Galactorrhoea.

# Levodopa & Carbidopa: long term S/E



- ① Psychosis
- ② Chorea-like movement (Dyskinesia).

PSYCHOSIS:

- Overaction of Dopamine.
- D<sub>2</sub> blockers → Conventional / Typical Antipsychotic.

Conventional / Typical Antipsychotic drugs

Phenothiazine	Butyrophenones	Thioxanthenes.
Chlorpromazine	Haloperidol	Thiothixene
Trifluoperazine	Trifluoperidol	Flupenthixol.
Thioridazine	Droperidol	
Fluphenazine	Penfluridol-LA	

# Typical antipsychotic = Neuroleptic agents.

# Most potent D<sub>2</sub> blocker / Antipsychotic = Butyrophenone



Max<sup>m</sup> EPS produced

THIORIDAZINE — S/E → Corneal pigmentation  
Cataract  
Retinal degeneration.

Most potent Antipsychotic - HALOPERIDOL



Cause Max<sup>m</sup> EPS  
Less ANS side effect.

CHLORPROMAZINE - Causes cholestatic jaundice.

Drug induced Parkinsonism:

TOC - Centrally acting Anticholinergic



Trihexyphenidyl (BENZHEXOL)

Other - Benztropine

Biperiden

Procyclidine.

PROMETHAZINE - 1st gen. antihistamine

have anticholinergic action

So, used in EPS.

Extra pyramidal Syndrome:

- ① Drug induced Parkinsonism
- ② Acute muscular dystonia: PROMETHAZINE  
BENZHAXAL
- ③ Tardive dyskinesia: No specific t/e  
Symptomatic - Valproate, Vit. E.

VALBENZAZINE (Newer drug)

- Acts by Vesicular monoamine transporter  
& inhibitor.

④ AKATHESIA - DOC: Propranolol

⑤ Malignant Neuroleptic Syndrome: DANTROLENE

↓  
directly acting SMR.

Anti-Parkinson drug:

LEVODOPA:

↳ Protein meals reduces absorption of levodopa.  
Vit-B<sub>6</sub> (Pyridoxine) should n't be given c̄  
levodopa bcoz it stimulate peripheral  
conversion.

Peripheral toxicity:

M/c S/E of levodopa - Nausea & Vomitting  
Alteration in taste sensation.

↓  
due to stimulation of D<sub>2</sub> receptor  
in CTZ.

D<sub>2</sub> receptor blocker - Domperidone  
Metaclopramide.

# Only domperidone is useful in t/t of vomiting  
due to levodopa.

# Metaclopramide is not used bcoz it crosses  
BBB & reduces efficiency of levodopa.

# Causes - Cardiac arrhythmias  
Exacerbation of angina  
- due to D<sub>1</sub>, β<sub>1</sub>, d<sub>1</sub> activation.



## LEVODOPA + CARBIDOPA

↳ Dopa decarboxylase inhibitor

Long term S/E → Abnormal choreo athetoid movement  
→ Psychosis

# Huntington's Chorea } movement disorder due to  
Tourette Syndrome } overaction of dopamine.

↓  
T/E - DOC: TETRABENAZINE

(Dopamine Depletor

Other - Chlorpromazine

Haloperidol.

# Levodopa is Precursor of melanin  
- C/I in melanoma

# Chronic therapy of levodopa may cause On & off phenomenon

↓  
dyskinesia

↓  
Severe parkinsonism

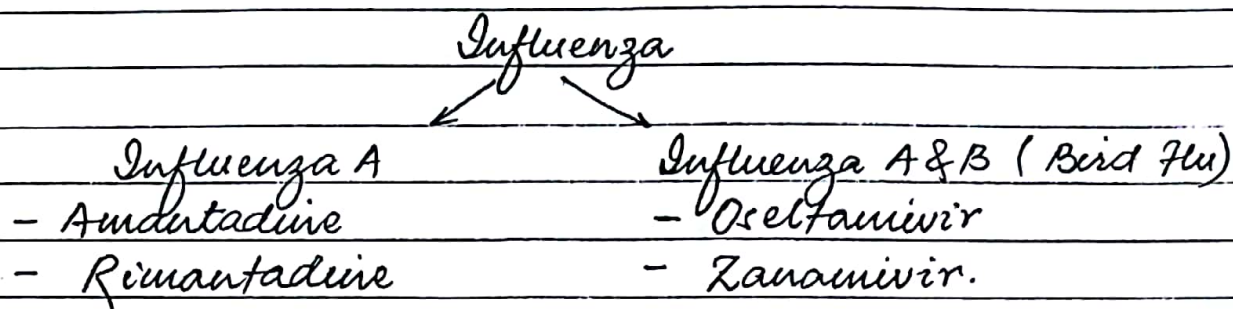
↓  
Rescue therapy

- APOMORPHINE (D<sub>4</sub>)

given S/C.

# Abrupt withdrawal of levodopa → Neuroleptic malignant Syndrome.

## AMANTADINE:



Oseltamivir — 75mg / 1 BID / 5 days — Oral  
 ↳ Prodrug — Causes Nausea & Vomiting.

Zanamivir — Intranasally — Bronchospasm

## Vaccination:

PERAMIVIR (Neuraminidase Inhibitor)

↳ IV (Intravenous)

## Amantadine:

- Anticholinergic
- Dopaminergic agonist
- NMDA antagonist.

- Useful in Parkinsonism

SE — Ankle edema

Levico reticularis. (Net like skin rashes).

# Ergot D<sub>2</sub> agonist : Bromocriptine  
Pergolide  
Cabergoline

Common S/E of these 3 drugs - Erythromelalgia.  
Cardiac valve fibrosis.

# Pergolide - Causes max<sup>m</sup> Cardiac valve fibrosis.

Other uses of Bromocriptine:

- Prolactinoma.
- Acromegaly
- Type 2 DM

Non-Ergot D<sub>2</sub> agonist: Pramipexole } m/c S/E Psychosis.  
Ropinirole }  
Rotigotine (Transdermal)

Advantage: No peripheral vasoconstriction.

[ Pramipexole ] - S/E → Compulsive shopping  
[ Ropinirole ]      Kleptomania  
                                 Sexual desire  
→ Useful for t/t of Restless leg Syndrome.

## COMT inhibitors

JALCAPONE

ENTACAPONE

Dangerous toxicity

- Rhabdomyolysis
- Severe Diarrhoea
- Hepatotoxicity.

- doesn't cross BBB.

Urine - Yellowish Orange.

## SEROTONIN (5-HT)

Source - Tryptophan

Func<sup>n</sup> of 5HT<sub>1A</sub> - Inhibition of release of Serotonin.

Autoreceptor of Serotonin.

Monoamine undergoes metabolism by Monoamine oxidase (MAO). They produce metabolite 5-hydroxyindole acetic acid.

# In Carcinoid tumour - ↑ 5-hydroxyindole acetic acid.

# Serotonin undergoes reuptake causing ↓ central serotonin.

Action of Serotonin on 5HT<sub>1B/D</sub> - Vasoconstriction

↳ SUMATRIPTAN (use - Migraine)  
(mainly 1D; min<sup>m</sup> 1B)

Action of Serotonin on 5HT<sub>2</sub> - Schizophrenia

(5HT<sub>2A/2C</sub>)  
↳ Clozapine  
Risperidone  
Olanzapine

Action of Serotonin on  $5HT_3$ : Nausea & Vomiting

$5HT_3$  antagonist - Ondansetron  
Granisetron

Action of serotonin on  $5HT_4$ : Diarrhoea.

Selective  $5HT_4$  agonist - Cisapride } withdrawn  
Mosapride } coz of  
Tegaserod } QT prolong  
-ation on ECG.

# All serotonin receptors are G-protein coupled receptor.  
except  $5HT_3$  (ligand gated receptor)

# Acute Migraine:

Main issue - Vasodilation

For t/t of Acute migraine - Vasoconstrictor



Ergot Alkaloids - Ergotamine  
 $5HT_{1B/D}$  agonist - Sumatriptan (DOC)  
Rizatriptan  
Almotriptan  
Frovatriptan  
Zolmitriptan

Care is taken for HTN & IHD in these pts.

# St. Anthony's fire → chronic treatment of ergot  
alkaloid cause peripheral vasoconstriction  
(gangrene of foot)  
Poisoning - Ergotism

## # BUTOPHANOL - Opioid

Used intranasally for Headache.

## # Drug useful for Prophylaxis of Chronic Migraine:

- ① M/c drug - Propranolol ( $\beta$ -blocker)
- ② CCB - Flunarazine  
( $\text{Na}^+$  channel blocking & Antioxidant property)
- ③ Anti-convulsant - Valproate  
Gabapentin  
Topiramate
- ④ TCA - Amitriptyline.

## ⑤ Clonidine

~~Onabotulinum toxin A~~

Onabotulinum toxin A

## ⑥ 5HT<sub>2</sub> blocker

- Pizotifen
- Cyproheptadine
  - Antihistamine + Antimuscarine
  - + Antiserotonine.

- Primary used as appetizer

- Used in Serotonin Syndrome.

- Methylsergide (Not used)

- Causes retroorbital & peritoneal fibrosis

Newer drugs - Calcitonin gene related peptide (CGRP)

- Vasodilation.

CGRP antagonist → Oliceripant - i.v.

Telcagepant - Oral

↳ Hepatotoxic

# LASMIDITAN - 5HT<sub>1F</sub> agonist

↓  
Undertrial

Atypical Antipsychotics  
(5HT<sub>2</sub> Antagonists)

- Clozapine
- Quetiapine
- Olanzapine
- Risperidone
- Lurasidone
- Ziprasidone
- Aripiprazole
- Asenapine (S/L)

- ↳ Advantages:
- Less EPS
  - Refractory Cases
  - +ve & -ve symptoms of Psychosis.

→ Not causes Metabolic Syndrome

CLOZAPINE - S/E → Agranulocytosis 0.8-1%  
(dose independent)

Seizure (10%)

Ileus (Paralytic) → Constipation  
Sialorrhoea

Metabolic syndrome.

- Pillow ~~bed~~ Syndrome  
(Wet)

- Anti-suicidal action.

QUETIAPINE - S/E - Cataract, Priapism

OLANZAPINE - USE → Mania in BPD

Adverse effect → Max<sup>m</sup> wt gain  
Max<sup>m</sup> metabolic syndrome.

**RESPERIDONE:** In addition to blocking 5HT<sub>2</sub>  
it also block D<sub>2</sub>.

- May cause EPS

**LURASIDONE:** Useful in BPD  
may also cause EPS.

**ZIPRASIDONE:** M/E & E - QT Prolongation

**ARIPIPRAZOLE:** Useful in BPD (mania)  
- Best drug among atypical  
antipsychotic

**ANXIETY DISORDER:**

- ↓ GABA activity
- ↑ 5HT activity.

**BUSPIRONE:** 5HT<sub>1A</sub> agonist

Anti anxiety agent (Chronic Anxiety)

Advantage - Non sedative

Non habit forming.

Disadvantage - Delayed in onset  
(3 to 4 wks)

For acute anxiety - Temporarily - BZD

# Performance anxiety = R<sub>x</sub>: Propranolol

Anxiety & panic attack = R<sub>x</sub>: SSRI

H<sub>1</sub> blocker: Hydroxyzine (Anti anxiety property)  
↳ 1st gen. antihistamine.

Cetirizine → Metabolite of Hydroxyzine  
↳ 2nd gen. anti-histamine.



Female Sexual Stimulant: FLIBANSERIN

↓ useful in  
HSDD - Hypoactive Sexual desire Disorder

# Deficiency of Serotonin & NE - Depression

# TCA, SNRI, NDRI → Inhibit reuptake of 5HT, NE  
SSRI → Inhibit reuptake of 5HT.

MAO-inhibitors

MAO-A

MAO-B

- involved in metabolism  
of NA & 5HT.

- Useful in depression.

- Metabolism of Dopamine

SELEGILINE

RASAGILINE

SAFINAMIDE

Selective  
MAO-A  
inhibitor

MECLOBAMIDE

CLORGILINE

Non-selective MAO inhibitors:

PHENELZINE

TRANLYCPROMINE

ISOCARBOXAZID

# Cheeze reaction = T/t: Phentolamine

## SSRI:

Fluoxetine (longest acting → 5 to 7 days)  
 Fluvoxamine - Shortest acting  
 Paroxetine  
 Citalopram  
 Escitalopram - Highly selective SSRI  
 Sertraline - Least drug interaction.

S/E of SSRI - May cause HTN

- Insomnia, Anxiety, Sexual S/E.

↓ ↘ delay in ejaculation.

∴ It is taken in morning.

↓  
 Useful in t/t of premature ejaculation.

M/c - Nausea & vomiting  
 - Diarrhoea.

## Drug interaction:

Serotonin Syndrome - SSRI + MAO inhibitor

Rx - Cyproheptadine.

↳ Primarily 5HT<sub>2</sub> antagonist

Anti H<sub>1</sub> + Ach

# FLUOXETINE: Least discontinuation Syndrome

# PAROXETINE - Wt gain

Teratogenic

Used in Premenstrual <sup>tension</sup> Syndrome (PMTS)

↓

FDA approved.

Drug interaction b/w Fluoxetine & Tamoxifen:

Tamoxifen - for anticancer activity needs activation.  
- activated w help of CYP2D6 enzyme.

Fluoxetine - CYP2D6 enzyme inhibitor.

Tamoxifen failure occurs.

# SSRI Uses:

① Depression

- juvenile depression - Fluoxetine  
Sertraline

② OCD

③ PTSD

④ Bulimia nervosa

⑤ Anxiety & panic attack.

⑥ PMTS.

DOC: SSRI: ① OCD

② PTSD

③ Anxiety & panic attack.

TCA

- Inhibit reuptake of Serotonin & NE (Non-selective)

CLOMIPRAMINE - T/E of OCD

DOXEPIN - Strong antihistaminic property

• Atopic dermatitis

• Lichen Simplex

# All TCA have antihistaminic property.

IMIPRAMINE - Strong anticholinergic activity.

- Nocturnal enuresis

DOC: Desmopressin

All TCA have anticholinergic activity.

### AMETRYPTLINE

Used in - Antidepressant

Prophylaxis of migraine

DM neuropathy pain



Gabapentin, Pregabalin

Other - Nortryline

Desipramine

Amoxapine - D<sub>2</sub> blocking action

Anti-psychotic

EPS, Galactorrhoea.

Maprotiline

Reboxetine

Adverse effect of TCA:

- All TCA having antihistaminic property

" " anticholinergic "

" " d<sub>1</sub> blocking "

- Sedation, wt gain, Seizure



∴ taken at bedtime.

- Dryness of mouth, constipation, Tachycardia  
& Retention of urine

- Postural hypotension

TCA poisoning & t/t:

Cardiac arrhythmia → Lidocaine, Bretylium, Avoid class Ia

Convulsion → Diazepam

Coma →

Metabolic acidosis → i.v. Sodium bicarbonate

- No role of dialysis in TCA poisoning  
↳ 600g large  $V_d$ .

# Anti-cholinergic

① Avoid TCA in elderly male - Aggravate Urinary Retention.

② Alzheimer's ds.

ST JOHN'S WORT:

Natural antidepressant.

# HYPERFORIN

↳ Monoamine reuptake inhibitor.

- Very powerful enzyme inducer.

↓  
Lead to OCP failure.

Anti retroviral failure.

# MIANSERIN: Presynaptic  $d_2$  inhibitor  
Useful in depression.

MIRTAZAPINE: Presynaptic  $d_2$  /  $5HT_1$  inhibitor  
Useful in depression.

- Na SSA (Noradrenergic & specific serotonergic antidepressant).

# TIANEPTIN } 5HT reuptake enhancer  
 AMINEPTIN }  
 ↓  
 Used ~~as~~ <sup>as</sup> antidepressant  
 Mechanism of action not known.

BPD (Bipolar Disorder):  
 Prophylaxis - Lithium

Acute mania - Valproate  
 Carbamazepine  
 Olanzapine  
 Aripiprazole  
 Diazepam

Depressive phase - Lamotrigine

For Rapid Cycler: DOC: Sodium Valproate  
 ↳ more than 4 episodes of mania & depression  
 in a year.

Lithium: Monovalent cation

Useful for prophylaxis of BPD.

Narrow Therapeutic Index (TDM)

Therapeutic drug monitoring  
 ↓

Monitoring plasma lithium level.

$T_{1/2} = 24 \text{ hrs.}$   
 ↓

Maintenance for BPD = 0.5-0.8 meq/L

Acute Mania = 0.8-1.2 meq/L

Toxic symptom > 1.5 meq/L

Toxicity → Hemodilysis → 4 meq/L

Adverse effect of Lithium:

LI = Leucocyte count  $\uparrow$  (Leucocytosis)

T = Tremor (M/c  $\rightarrow$  8-10 Hz)

H = Hypothyroidism (Inhibit release of  $T_3$  &  $T_4$ )

IU =  $\uparrow$  urination (polyuria = DI) (Rx: Acetazolamide)

M = Mother (Ebstein's anomaly) = Teratogen

In CVS  $\rightarrow$  T wave changes

Dermatology  $\rightarrow$  Exacerbation of psoriasis.

C/I: ① Pregnancy & Lactation

② Sick sinus syndrome.

Drug interaction b/w lithium & SMR (Succinylcholine & Pancuronium):

$\hookrightarrow$  Lithium aggravate the action of SMR.

$\hookrightarrow$  Stop lithium 1 day before Sx.

# Hyponatremia will occur in lithium toxicity.

[ Diuretic aggravate lithium toxicity.

NSAID " " " "

## Opioid Receptors.

3 imp. endogenous opioid Receptor in body

$\mu$  (Mu)

$\delta$  (Delta)

$\kappa$  (Kappa)

All opioid receptor are GPCR - via  $G_i$  pathway.

Endogenous opioid peptides:

Endorphine - more affinity toward  $\mu$

Enkephaline - " "  $\delta$

Dynorphin - " "  $\kappa$

Action of opioid:

- Due to activation of  $\mu$  &  $\delta$ .

P = Physical dependence,  $\uparrow$  Prolactin secretion

M = Miosis ~~NO Tolerance~~

C = Constipation, convulsion (M3G)

A = Analgesic

R = Resp<sup>r</sup> depression

E = Euphoria

S = Sedation

# Opioid are useful in t/t of dull pain

Continuous pain

Localised pain

Visceral pain

# Opioid (Morphine) activating Edinger Westphal nucleus (III CN) causing miosis.

↓

Only systemic Morphine cause miosis.



Action of opioid due to kappa:

D = Dysphoria

M = Miosis

A = Analgesia

R = Respr depression

D = Diuresis

S = Sedation

# Morphine having Histamine Releasing action.

↓  
Vasodilation

↓  
Shifting of pulm. fluid in systemic circulation.

↓  
It is useful for t/t of Pulm. edema.

# All the action of morphine may develop tolerance on repeated administration except - Miosis

Constipation

Convulsion

# Enkephalins may undergo metabolism by Enkephalinase.  
For the t/t of diarrhoea - Racecadotril

↓  
Enkephalinase inhibitor.

Pure agonist:

Codeine converted in morphine by CYP2D6

↑ enzyme in body.

Natural opioid - Morphine, Codeine (CYP2D6)

Semisynthetic - Diacetyl morphine (Heroin), Pholcodeine

Synthetic - Pethidine (Meperidine - Antimuscarinic,

Nor-pethidine) → Metabolite of pethidine

↓  
GI in t/t MI pain.

↳ S/E - Seizure (convulsion)

## # Pethidine & Morphine CI in Renal failure.

### Methadone:

- Longest acting opioid
- NMDA blocking property & inhibiting reuptake of NE & 5HT.
- Useful for t/t of neuropathic pain & Cancer pain
- Doc for opioid deaddiction.

### Tramadol:

- Also having property of inhibiting reuptake of 5HT & NE.

# Be careful using Methadone & Tramadol in pt. using SSRI, MAO inhibitor causing Serotonin Syndrome.

### Fentanyl: Fentanyl group.

	Fentanyl	Sufentanil	Alfentanil	Remifentanyl
Potency ↓ potent than Morphine	X100	X1000	X5	X100
Duration of action	30 min	30 min	5-10 min	3-5 min

Least potent: Pethidine & propoxyphene (1/10)

Analgesic for day care Sx: Remifentanyl.

# Fentanyl + Droperidol = Neuroleptic Analgesia

# Fentanyl + Droperidol + N<sub>2</sub>O = Neuroleptic anaesthesia.

# Fentanyl group Cause Post op trünical rigidity  
 ↓  
 (Max - Alfentanil)

# Thorax muscle rigidity = wooden chest Syndrome.

Mixed agonist - antagonist:

-  $\mu$  antagonist / Kappa agonist:

- Nalorphine (more dysphoria, not in use)
- Pentazocine (Sympathetic stimulant) C/I in MI pain
- Butorphanol (Nasal formulation)

-  $\mu$  agonist / Kappa antagonist:

• Buprenorphine

- Useful for all type of pain
- Useful for opioid withdrawal



alternate to methadone.

Pure antagonist:

Naloxone  
 Nalmefene } Intravenous

Naltrexone (Oral, long acting, Hepatotoxic)

Acute morphine poisoning:

Specific antidote - Naloxone (0.4-0.8mg)



i.v., repeated every 2-3 min.

- It blocks  $\mu$  receptor at much lower doses than those needed to block  $\kappa$  or  $\delta$  receptors.
- It promptly antagonizes

# Naltrexone → Useful to control craving for Morphine & craving for alcohol.

# For t/t of constipation due to morphine (opioid)

↓

Peripheral opioid antagonist [ ALVIMOPEN  
METHYL NALTREXONE

Newer opioid:

Peripheral kappa antagonist: ASIMADOLINE

↓

for IBS

Peripheral  $\mu$  &  $\kappa$ -agonist; delta antagonist:

ELUXADOLINE → for IBS.

Peripheral  $\kappa$ -antagonist:

NALFURAFINE → Antipyretic → CKD

# Codeine  
Dextromethorphan ] Anti-tussive opioid.

# Anti-diarrhoeal opioid:

Diphenoxylate (Atropine can be added to prevent addiction).  
Loperamide

# C/I of Morphine:

- Head injury pain (Resp<sup>r</sup> insufficiency)
- Biliary colic pain (causing constriction)
- Severe asthma. of sphincter of oddi.)

Ethyl Alcohol / Alcohol :

Deaddiction - Disulfiram like reac<sup>n</sup>

(Aldehyde dehydrogenase inhibitor)

Drug causing Disulfiram like reac<sup>n</sup>:

C = Chlorpropamide (Sulfonylurea - DM)

Cefoperazone (3rd gen. Cephalosporin)

M = Metronidazole

Prasid = Procarbazine (Anti Cancer) → Alkylated

G = Griseofulvin

T = Tinidazole

Naidu = Nitrofurantoin (Causes coffee colour urine)

# Chronic alcoholic generally suffer Thiamine deficiency.  
(Vit B<sub>1</sub>)

# Alcohol <sup>always</sup> undergoes Zero order kinetic elimination:  
Zero WATT POWER

W = Warfarin

A = Alcohol

A = Aspirin

T = Tolbutamide

T = Theophylline

P = Phenytoin

# Excretion of Alcohol - Kidney

# In acute ethanol poisoning, pt. presenting c  
hypoglycemia. T/t = Glucose + Thiamine.

Methyl alcohol:

Methyl alcohol



Formaldehyde



Formic acid (dangerous) ← Ocular damage  
Metabolic acidosis

Specific antidote for Methanol poisoning



Fomepizole

(4-Methyl pyrazole)



Acting by inhibiting Alcohol dehydrogenase.

Alternative drug - Ethanol also given.  
Hemodialysis.

Anti craving drugs for Alcohol:

- Disulfiram (DOC)
- Naltrexone (1st line drug)
- Acamprosate (2nd, NMDA blocker + GABA agonist)
- SSRI (citalopram)
- Ondansetron
- Topiramate, Baclofen (GABA agonist)
- Rimonabant, a CB1 receptor antagonist.

FAS (Fetal alcoholic syndrome):

CF - Microcephaly

Maxillofacial abnormalities

Movement disorder - Hyperkinetic

Mental retardation

Phenytoin:

Na<sup>+</sup> channel blocking antiepileptic

Fosphenytoin - Prodrug of phenytoin

Water soluble (im/slow iv)

↳ safe for

Saturation kinetics - First order → Zero order

Adverse effect:

① Acute toxicity

- On high iv. → Cardiac arrest.

- High oral → Nystagmus

Ataxia

Diplopia

Vertigo

② Chronic toxicity

- Gum hypertrophy (M/C - 30%)

↳ Due to collagen accumulation

- Blood → Megaloblastic anemia (Folic acid deficiency)

Interfere Vit K activity (Hemorrhage)

Interfere - Vit D & Calcium activity.

↳ Osteomalacia & rickets

- Hypersensitivity reac<sup>n</sup>  
↳ Pseudolymphoma.
- In female → Hirsutism
- Inhibits release of insulin from  $\beta$ -cell of pancreas - Hyperglycemia (DM)
- Teratogenicity → due to Areneoxide  
↳ C = Cleft lip & palate  
P = Hypoplastic phalanges  
M = Microcephaly.
- Extravasation of phenytoin → Purple glove syndrome.

# Phenytoin - Microsomal Enzyme inducer.

Non-epileptic uses of Phenytoin:

- Trigeminal neuralgia
- Digoxin - induced VT
- Wound healing

Carbamazepine:

DOC for Partial Seizure (Focal seizure)  
For Ht of Temporal lobe epilepsy.

Non-epileptic Uses:

DOC for Trigeminal neuralgia.

Useful for Ht mania in BPD

Carbamazepine having SIADH activity → Antidiuretic  
↳ Use in DI



# It is microsomal enzyme inducer.  
It also undergoes auto induction.

↓  
Phenobarbitone  
Carbamazepine  
Nevirapine

Sodium Valproate:

- Broad spectrum anti-epileptic.

MOA = GABA agonism property  
Anti-glutamate "  
Na<sup>+</sup> channel blocking "  
T-type CCB "

DOE for Myoclonic / Atonic / Clonic & tonic Seizure  
First line drug for Absence seizure / Lennox Gastaut  
Syndrome.

Non-epileptic uses:

- Migraine prophylaxis
- Manic in BPD (LITHIUM)
- Rapid cycler (>4 cycles/year)
- Tardive dyskinesia

# It is microsomal enzyme inhibitor

SE: V = GIT, wt. gain (Vomiting)  
AL = Alopecia / curling of hair  
P = Pancreatitis, hyperammonia  
R = Rashes  
Q = PCOD

Cleft lip &amp; palate

- A = Allergy ← Most  
 T = Teratogenic (Spinabifida / CVS problem / Orofacial / digital)  
 E = Hepatotoxicity (< 2yrs children).  
 ↓  
 t/t = Carnitine (Antioxidant)

Others Antiepileptic:

- Levacetam (SV2A)
- Magnesium Sulfate (DOC in eclampsia)
- Acetazolamide
- ACTH (Infantile spasm)

Levetiracetam - Modify synaptic release of glutamate / GABA.

Acetazolamide:

- Carbonic anhydrase inhibitor.
- Useful for Glaucoma → Taken Orally.
- Used as diuretics - acts on PCT

Use - Acute mountain sickness

Periodic paralysis

Absence seizures

Catamenial epilepsy

Rx - GANAXALONE

Absence Seizure:

- Ab<sup>o</sup> of T-type Ca<sup>2+</sup> channel (Thalamus)

Rx: T-type CCB

ETHOSUXIMIDE

SODIUM VALPROATE (1st line drug)

**TREMETHADIONE** (Withdrawn - Nephrotoxic)

↳ Hemorralopia - Day blindness.

# Anti epileptic having Carbonic anhydrase inhibiting property:

TOPIRAMATE } cause Nephrolithiasis  
ZONISAMIDE }

# RETIGABINE } Potassium channel opener  
or EZOGABINE } used for partial seizure

- causing blue colour pigmentation  
on lip & skin

↑  
(New drug)

### GENERAL PHARMACOLOGY

Pharmacokinetics (PK):

Drug absorption:

Food interfere drug absorption

eg: Milk ( $Ca^{2+}$ ) - Tetracycline

Protein meal reduces - Absorption of Levodopa.

Food enhances drug absorption

Lithium

Halofantrine

Lumefantrine

Griseofulvin

Bedaquiline

Fibrates } lowering cholesterol

} more absorbed c cholesterol diet.

- Absorption of Iron - Vit. C (Ascorbic acid)

# For a drug to absorb better - lipid soluble  
& distribution. Non-ionised.

# Acidic drug non-ionised in Acid medium.  
Basic drug non-ionised in Basic medium.

→ Aspirin  
# Acidic drug - Absorbed in stomach.  
Basic drug - Absorbed in Duodenum/Intestine.

→ Morphine  
# Strongest Acid/Alkali always seen in ionised form.

Heparin - Can't be used orally.

↓  
- Heparin ionised molecule, not cross the placenta, so not cause teratogenicity.  
DOC - for anticoagulation.

Lignocaine - For rapid absorption / onset of action  
↓  
Weak basic drug. For ↑ duration given c̄ Adrenaline.

Acidic drug poisoning -

For acidic drug poisoning if the pt. is passing acidic urine, you should alkalisise the urine.  
Urine alkalisise c̄ Sodium bicarbonate.

Alkali drug poisoning -

For the pt of alkali drug poisoning if the pt. passing alkaline urine, you should acidify the urine.  
Urine acidify c̄ Ascorbic acid

↓  
By injection Ammonium chloride.

Ion-trapping - Acidic drug (Aspirin) reached basic medium get ionised & trapped in the region.

P-glycoprotein: Permeable efflux pump.

↳ Presence of P-glycoprotein decreases the bioavailability of digoxin.

eg. of P-glycoprotein inhibitor: Quinidine

Itraconazole

Erythromycin

Amiodarone

Verapamil

Drug undergoing high first pass metabolism on orally:

Propranolol

Salbutamol

Theophylline

Verapamil

Lignocaine

Nitrates

Imipramine

# All nitrates go through extensive 1st pass metabolism except - Isosorbide <sup>mono</sup> nitrates.

# Rectally given drug absorbed via External hemorrhoidal vein - No 1st pass metabolism

If via Internal hemorrhoidal vein - 1st pass metabolism occurs.

# i.v. - 100% Bioavailability.

Henderson Hesselbach equation:

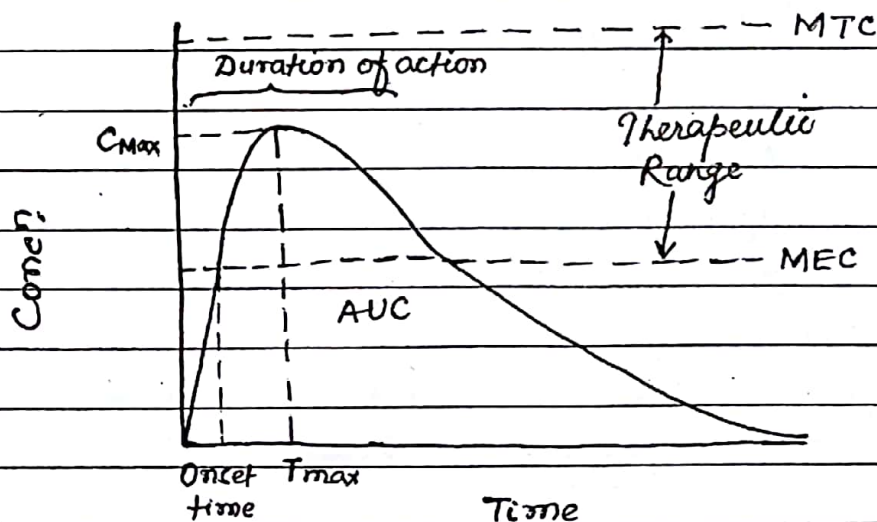
$$pKa = pH + \log \left( \frac{\text{ionized A}}{\text{unionized A}} \right)$$

If  $pKa = pH$

means, 50% drug is in ionised form  
& 50% " " unionised form

#  $pKa - pH = 1 \rightarrow 90\%$  drug in absorbed form.  
 $pKa - pH = 2 \rightarrow 99\%$  " "  
 $pKa - pH = 3 \rightarrow 99.9\%$  " "

Bioavailability curve:



$C_{max} = \text{Max}^m \text{ plasma conc}^n$

$T_{max} = \text{Time to reach } C_{max}$

$AUC = \text{Area under Curve.}$

# Same drug, same dose, same dosage form,  
 $< 20\% \rightarrow \text{Bioequivalent.}$

### Orphan drug:

- A drug useful for diagnosis/prevention & Ht of rare disease.

- eg: - Fomipizole (4-methyl pyrazole - Alcohol dehydrogenase inhibitor)  
Protamine Sulfate (Antidote of Heparin - Chemical antagonism)  
Calcitonin (1mg = 100 U of Heparin)  
Digiband (Antidote for Digoxin)  
Liothyronine (Active T<sub>3</sub> → Myxedema coma)  
↳ always given c̄ β-blocker.

Calcitonin: Useful in Hypercalcaemia

- Paget's ds
- Osteoporosis
- diagnosis for Medullary Ca Thyroid.

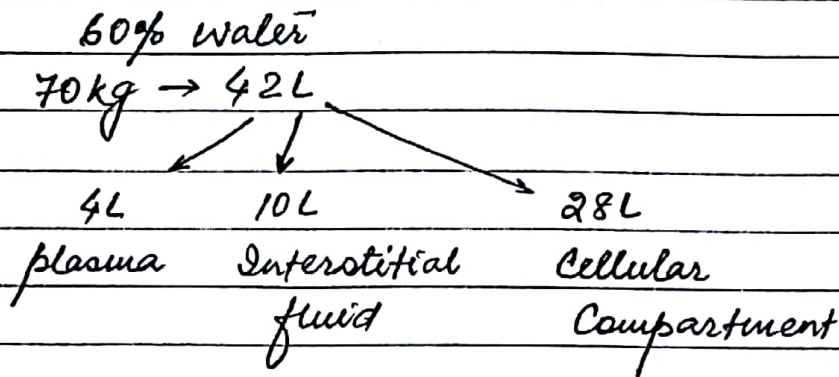
# Pitolisant / Tiprolisant: Use in Narcolepsy (Orphan drug status).

### Essential drugs:

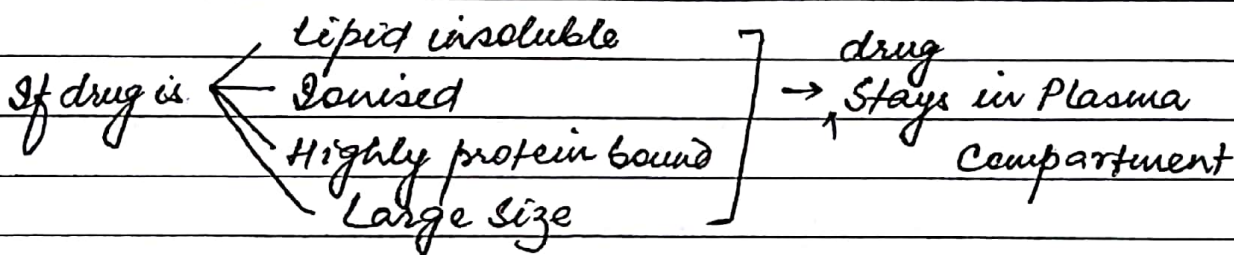
- Drug that meet health needs of the majority of population
- Affordable & Available in all area.
- Always single comp<sup>d</sup>.

Schedule H - Drug only given on prescription written by medical practitioner (Registered).

## Drug distribution:



# If a drug only in the plasma compartment, it is called as low  $V_d$ .



## - Role of Hemodialysis

# If a drug goes to cellular compartment it has high or large  $V_d$ .

↓  
Lipid soluble  
Non ionised  
Free form.

Large  $V_d$  → No role of Dialysis.



Drug can't removed by dialysis:

A = Amphetamine

V = Verapamil

O = Opioids, OPC

I = Imipramine (TCA)

D = Diazepam

Dialysis = Diazepam (BZD)

BZD - Very strong binding capacity  
can't remove by dialysis.

# Loading dose depend upon  $V_d$ .

# For drug having large  $V_d$  - for rapid action give loading dose

Volume of distribution ( $V_d$ )

$$V_d = \frac{\text{Total i.v. dose}}{\text{Plasma conc}^n / L}$$

$$\text{Loading dose} = V_d \times \text{Target plasma conc}^n.$$

$$\text{Clearance} = \frac{\text{Rate of elimination}}{(cl)} / \text{Plasma conc}^n$$

$$\text{Maintenance dose} = cl \times \text{Target plasma conc}^n.$$

$$t_{1/2} = 0.693 \times \frac{V_d}{cl}$$

### Plasma protein binding:

- Acidic drug in plasma bind  $\bar{c}$  plasma albumin.
- In nephrotic syndrome or in liver failure (hypoalbuminemia) plasma albumin conc<sup>n</sup> is low -  
Use low dose of Acidic drug.
- Basic drugs are generally bind  $\bar{c}$  Alpha<sub>1</sub> Acid Glycoprotein

### Drug displacement type of drug interaction:

eg: Warfarin displacing tolbutamide from protein binding site.

Sulphonamide displacing bilirubin from protein binding site.

### BBB:

BBB absent - Pituitary  
Pineal gland  
Area Prostruma CTZ  
Median Eminence:

Do not cross BBB - Streptomycin (Aminoglycosides)  
Neostigmine (DOC for Atropine poisoning)  
Glycopyrolate (Pre anesthetic medication)  
Dopamine

# All aminoglycosides are ionised ~~water~~ molecule, so never absorbed orally, so not given orally.  
Even though aminoglycosides not absorbed in GIT

Neomycin & Paromomycin } can given orally.

# Streptomycin - CI in pregnancy  
 bcoz it crosses placental barrier & causes permanent deafness.

Redistribution:

eg: Thiopentane Sodium

( Ultra short acting )

↳ Rapidly entering brain & rapidly comes out & distribute to liver, kidney etc.

Biotransformation ( Drug metabolism ):

Consequences of drug metabolism

① Inactivation ( more water soluble )



excreted easily.

② Active metabolite formation from an active drug

③ Activation of inactive drug.

Active metabolite from active drug:

Active drug → Active Metabolite  
 Phenacetin → Paracetamol

↳ causes Analgesic nephropathy so withdraw.

Codiene  $\xrightarrow{\text{CYP2D6}}$  Morphine

↳ In some people it is deficient.

Diazepam → Oxazepam

Spiro lactone → Canrenone.

## Activation of inactive drug

Prodrug	Active metabolite
Levodopa	Dopamine
Methyl dopa	Methyl norepinephrine
Enalapril	Enalaprilat
	↳ All ACEi are prodrug except - Captopril, Lisinopril
Dipivefrine	Epinephrine
Becampicillin	Ampicillin
Minoxidil	Minoxidil Sulphate.
Cyclophosphamide	Phosphamide mustard.

### Drug metabolism:

Non synthetic reaction (Phase I reac<sup>n</sup>):

① Oxidation (M/c Phase I reac<sup>n</sup>)

All phase I reac<sup>n</sup> taken care by microsomal enzyme - CYP450

② Reduction

③ Hydrolysis

④ Cyclization

⑤ Decyclization

### Phase II reac<sup>n</sup>:

① Glucuronidation (M/c) - Morphine

② Sulfate Conjugation

③ Glycine "

④ Glutathione " (Paracetamol metabolism)

⑤ Acetylation

⑥ Methylation

**PARACETAMOL**

PHASE I / CYP2E1

N-acetyl benzo quinone (Hepatotoxic immunogenic (NABQIN) metabolite)

Phase II / Glutathione conjugation

Inactivation

For paracetamol poisoning → N-acetyl cysteine  
Methionine.

↓  
Beoz Glutathione generator.

Chronic alcoholic → More prone for liver damage  
beoz Alcohol → CYP2E1 inducer.

# End result of phase II reac<sup>n</sup> → Inactivation.

Drug undergoes Acetylation:

- S = Sulphonamide / Dapsone.
  - H = Hydralazine
  - I = Isoniazid
  - P = Procainamide
- } may cause RA, SLE.

Methylation:

- eg: Histamine → Methylhistamine
- Noradrenaline → Adrenaline.

Microsomal enzyme:

Enzyme	Drug
CYP3A4 (M/c)	>50% of drugs

CYP2D6 (2nd)      Codeine → Morphine

Fluoxetine inhibit CYP2D6

Tamoxifen activated by CYP2D6

CYP2C9      Warfarin

CYP2C19      Omeprazole metabolism  
Clopidogrel

CYP2E1      Paracetamol - NABQIA

Clopidogrel: Anti-platelet  
Prodrug

Activated  $\bar{c}$  help of CYP2C19.

Aspirin + Clopidogrel (prodrug) -

Aspirin → Causes gastritis

t/t → Omeprazole

Omeprazole shouldn't be given  $\bar{c}$  Clopidogrel.

- Preferred PPI given  $\bar{c}$  Clopidogrel

↓  
Pantoprazole  
Rabeprazole.

## Microsomal Enzyme

### Inducers:

- G = Griseofulvin
- P = Phenytoin
- R = Rifampicin
- S = Smoking
- Cell = Carbamazepine
- Phone = Phenobarbitone

### Inhibitors

- Vit<sup>®</sup> = Valproate
- K = Ketoconazole
- Can = Cimetidine
- Cause = Ciprofloxacin
- Enzyme = Erythromycin
- Inhibition = Isoniazid (INH)
- Grape fruit

### Drug excretion:

Major source = Kidney.

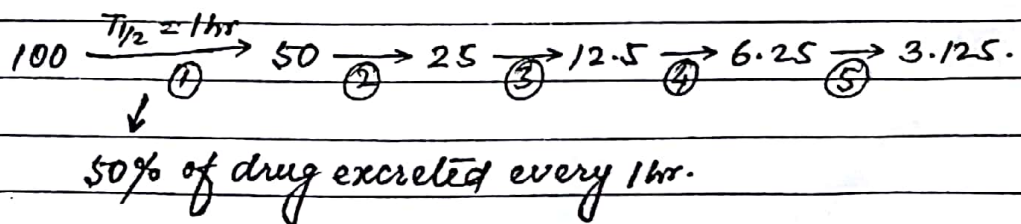
Net excretion of drug = GF + TS - Tubular reabsorption.

✓ PROBENICID - by inhibiting

prolong the action of penicillin.

### First order kinetics

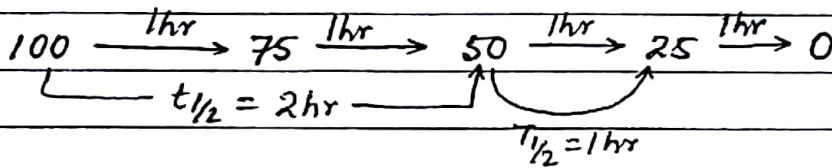
- Constant fraction of drug excreted constant interval of time.
- $T_{1/2}$  constant
- 97% drug eliminated after 5 half life.



Zero order Kinetics:

- Constant amount of drug excreted constant interval of time.
- No fixed  $T_{1/2}$ .

eg: 25 mg of drug, every 1 hr.

Common drug undergoing Zero kineticZero WAATT Power

W = Warfarin

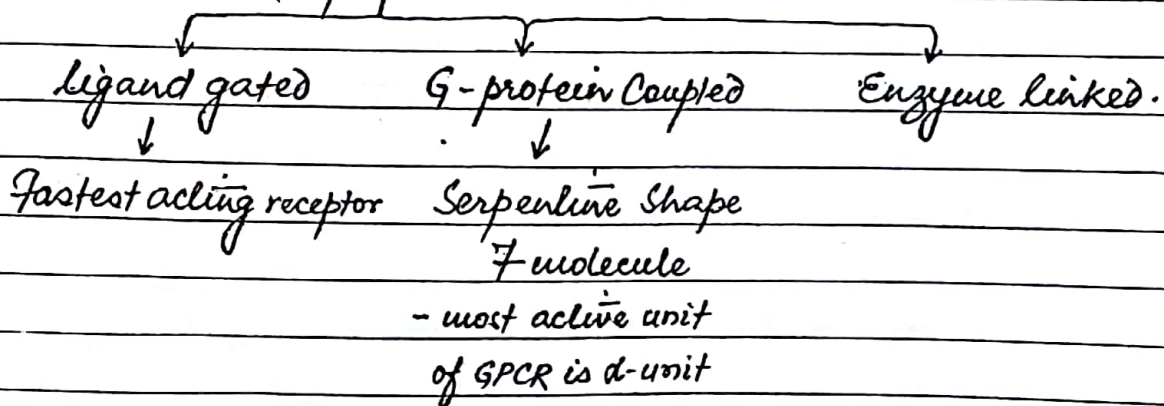
A = Alcohol

A = Aspirin

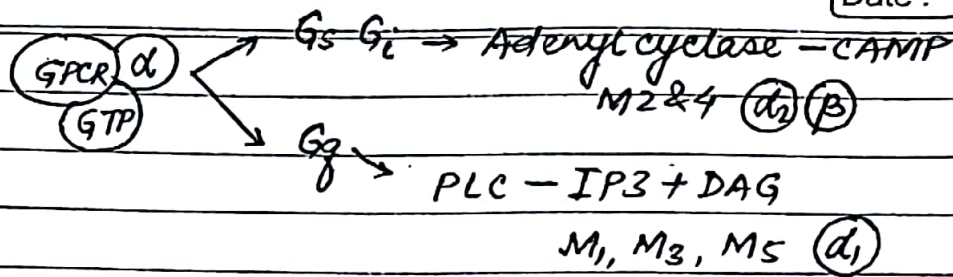
T = Tolbutamide.

T = Theophylline

Power = Phenytoin

Pharmacodynamics :-Receptor mediated MOACell memb<sup>r</sup> Receptors.





Enzyme linked receptor :

eg: Tyrosine Kinase Receptor

Insulin acting on cell memb<sup>r</sup> receptor

↓  
Activate Tyrosine kinase

↓  
Shift GLUT4 from cytoplasm to plasma memb<sup>r</sup>

↓  
Influx of glucose.

PEGVISOMENT : GH receptor blocker

Useful for t/t Acromegaly.

↑  
New drug → RUXOLITINIB : JAK enzyme inhibitor

Useful in Myelofibrosis.

TOFACITINIB : JAK 1 & 3 inhibitor

Useful in RA.

## Intracellular receptors:

Drug acting on Cytoplasmic receptor:

Steroid hormone

Vit D

Estrogen

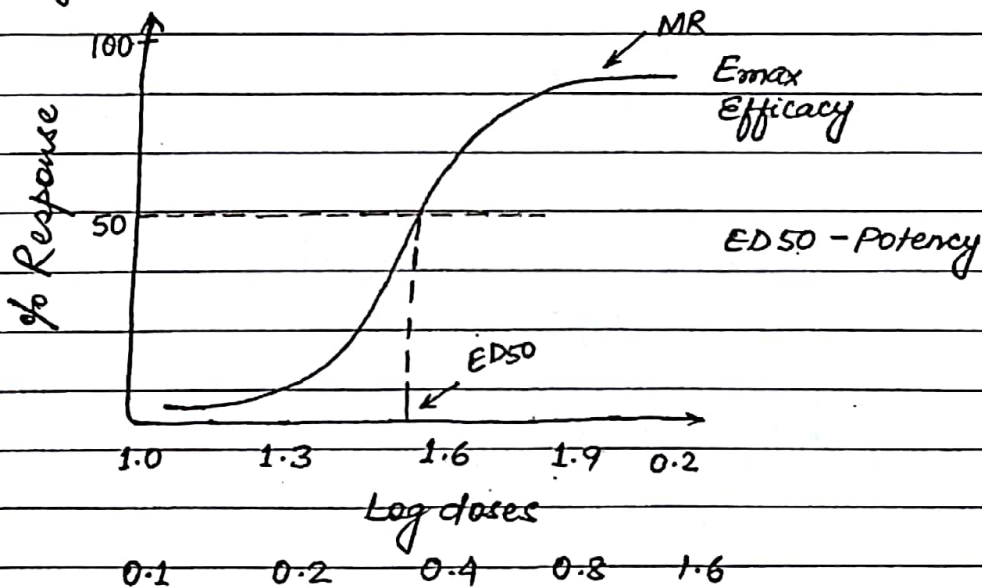
Progesterone

Testosterone.

Drug acting on nucleus:

Thyroid hormone

Log dose response curve:



Doses ( $\mu\text{g/ml}$ ) on arithmetic scale.

## Receptor Antagonism

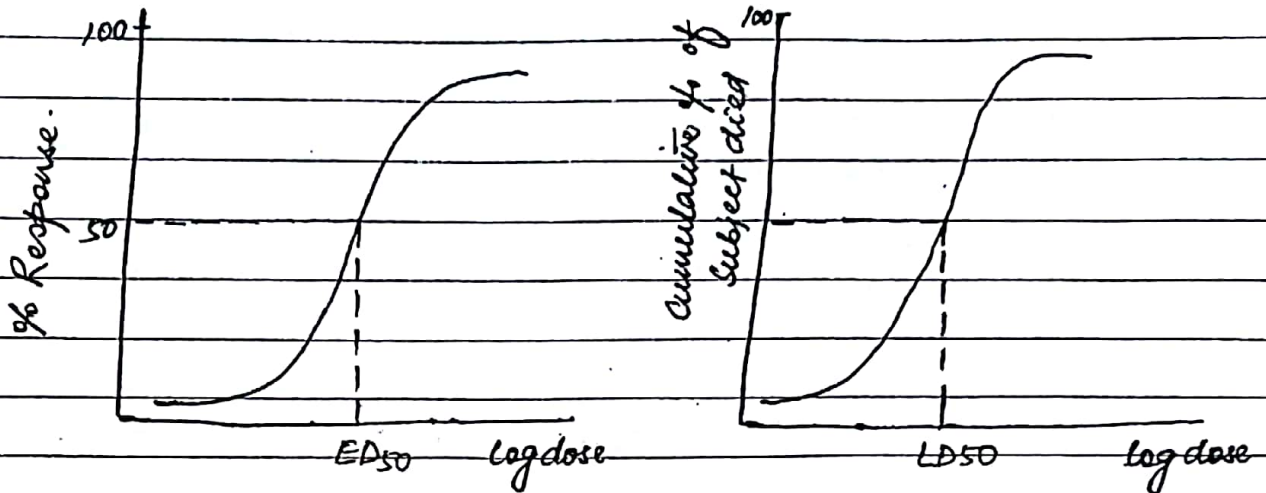
(1) In the presence of competitive antagonist DRC will be shifted parallel to right.

Efficacy  $\rightarrow$  Same; Potency  $\rightarrow \downarrow$

(2) In the presence of Non-competitive antagonism DRC will just come down

Efficacy  $\rightarrow \downarrow$ ; Potency  $\rightarrow$  Same.

## ED50 & LD50



Lower the ED50 more potent

Lower the LD50 more dangerous drug.

## Drug ~~Conf~~ Safety:

$$\text{Therapeutic index} = \frac{\text{LD50}}{\text{ED50}}$$

Theophylline  
Lithium  
Anti epileptics } Narrow  
Therapeutic  
index.

Warfarin - assessment by INR

$$\text{INR} = \frac{\text{Patient Prothrombin (PT)}}{\text{Control Prothrombin}}$$

Heparin - assessment by aPTT

LMWH - No need for monitoring

In obese pt. or Renal failure we do assessment by Anti factor Xa.

## Teratogenicity:

Preimplantation (0-2 wks)

Implantation (2-8 wks) → More teratogenicity occurs.  
↳ Organogenesis.

Growth & development (9 wks - 9 months)

① Warfarin: causing Contradi Syndrome  
(Fetal chondrodysplasia Punctata)

② Isotretinoin (Vit A) - Teratogenic

Lithium - Ebstein Anomaly  
CI in pregnancy.

### ③ THIOAMIDE:

Methimazole	}	Aplastic cutis
Carbimazole		Choanal atresia
<u>Propylthiouracil</u>		

↓  
Bcoz of strongly binding c plasma protein  
less chance of crossing placenta.

④ Alcohol - FAS (Fetal alcohol Syndrome)

⑤ Valproate - Valproate Syndrome.

⑥ ACEi - Renal agenesis

⑦ Indomethacin - Premature closure of ductus arteriosus.

⑧ Cyclophosphamide - Imperforate anus.

⑨ Busulfan & Chlorambucil (Chemotherapy)  
- Induce cleft palate

(10) Tetracycline - Bone & teeth defect. (Baby)

↓  
In mother → Fulminant hepatic failure.

So, definitely CI in pregnancy.

(11) Thalidomide - Phocomelia.

↳ Category X drug.

(12) Misoprostol - Useful for abortion

↳ Teratogenicity → Moebius Syndrome

↓  
at @ development of CN VI & VII.

(13) DES - Female → Vaginal Ca, hypoplasia  
↓ baby (in 10yrs of life) ↳ Male baby.  
if taken in pregnancy.

Drug development:

Preclinical trials - We follow guidelines

↓  
CPCSEA = Committee for the purpose of control & supervision on Experiments on Animals.

IAEC = Institutional animal ethics committee.

Clinical trial - Testing on humans.

guidelines - GCP (Good clinical practice).

HEC = Human Ethics committee.

Phase I: Pharmacokinetics studies  
Not efficacy.

Healthy volunteers (20-100)

Open label (No blinding)

- To know max<sup>m</sup> tolerable dose (MTD)

MTD - Safety & tolerability.

Anti-Cancer drug by pass Phase I.

Phase II: Therapeutic exploratory  
both efficacy & safety.

100-150 patients

Single blind

- To establish therapeutic efficacy.

- Dose ranging & ceiling effect.

Phase III: Therapeutic confirmatory.

Upto 5000 pts, from several centres

Double blind

- To confirm therapeutic efficacy.

- To establish the value of drug in relation.

Phase IV: Post marketing Surveillance.

ethical clearance is not required.

No time limits

To know rare & long term adverse effect.

### Phase 0 : Micro dosing studies.

#### Pharmacovigilance:

- Assessing, ~~monitoring~~,  
Reporting  
Monitoring  
Adverse effect.

#### # Longest acting insulin - Degludec.

##### Insulin Preparation

Fast onset & Short acting (Onset 10-20 min; duration-3-4hrs)

- Insulin Lispro  
Aspart  
Gulisine } For t/t of PP glucose.

Short acting (onset-30 min; duration → 5-8hrs)

##### Regular Insulin

↓  
made of 6 molecule (Hexamer)

↓  
dimer

it takes 30 min.  
to reach monomer status.

↓  
Monomer

given 30 min before meal.

given i.v.

Use in DKA, Hyperkalemia.

Intermediate (Onset 1-3hr; duration → 16-20hr)

NPH (Isophane Insulin) - Neutral Protamine Hagedon.

Lente Insulin (30% semilente, 70% ultralente)

→ can't mixed c̄  
other insulin

Page No.

104

Date: / /

Longer acting — Glargine (Acidic  $\Rightarrow$  pH=4)  
Detemir

Longest acting — Degludec

Adverse effect  $\leftarrow$  Hypoglycemia  
Wt. gain.

Inhalable insulin:

EXUBERA — Lack of acceptance by pts & physicians.

AFREZZA — Latest

Ultra rapid (c̄ in 15 min)

FDA approved.

[MAO]: Insulin acting on cell memb<sup>r</sup> receptor

↓  
Activate tyrosine kinase

↓  
Shifting of GLUT4 from cytoplasm to plasma memb<sup>r</sup>

↓  
Influx of Glucose.

Insulin Release:

For release of Insulin — at least 30% of  $\beta$ -cell  
are functioning.

In Type I DM — impossible to release insulin

↓  
All  $\beta$  cells are destroyed.



Sulphonyl urea

Maglitinide

- Repaglinide
- Nateglinide

Newer drugs for DM:

GLP-1 analogues:

given s/c	}	Exenatide	S/E - GIT (Nausea, Vomiting, diarrhoea) Necrotising pancreatitis Wt. loss. FDA approved - Liraglutide given for obesity.
		Liraglutide	
		Taspoglutide	
		Abiglutide	
		Dulaglutide	

- All obtained from GILA MONSTER (Salivary gland Venom).

DPP4 inhibitors: Oral

Adverse effect • Nasopharyngitis • URTI.	}	Sitagliptine → Excretion: Renal	Renal
		Saxagliptine	Renal/Hepatic
		Liraglutide	Bile
		Vildagliptine	Renal
		Alogliptine.	

# Vildagliptine: S/E - Hepatic toxicity  
pt. undergo periodic LFT.

PRAMLINTIDE: Islet Amyloid Polypeptide analog.

- ↳ given s/c
- ↳ Approved for Type 1 & 2 DM.

SGLT2 inhibitors:

Canagliflozin  
 Sertigliflozin  
 Dapagliflozin  
 Empagliflozin

Common S/E - Recurrent UTI (Bcoz Glycosuria)  
 Risk of breast/bladder CA.

C/I - In Renal failure.

Diabetes - Oral medications.

- Sulphonyl ureas
- Biguanides
- Thiazolidinediones
- Alpha-glycosidase inhibitors
- Meglitinides
- Bromocriptine
- Cholestyramin.

Sulphonylureas

1st generation:

Tolbutamide (6-12hr)

Chlorpropamide (30-60hr) - longest acting

↳ causes SIADH (dilutional hyponatremia)

2nd generation:

(Glyburide) Glibenclamide

Glipizide

Glipclazide

Glimepiride

- Cholestatic
- jaundice
- Disulfiram like react<sup>n</sup>

Glibenclamide - Safe in pregnancy.

Gliclazide - Antiplatelet, antioxidant.

M/c problem of Sulphonylurea - Hypoglycaemia  
Wt. gain.

Biguanides: Metformin

MOA = AMPK activator

↳ AMP - activated protein kinase.

Stimulates - Glucose utilisation

↓                      ↓  
Skeletal      Adipose  
muscle      tissue.

- It is insulin sensitizer.

Suppresses - Glycogenolysis  
Neoglucogenesis

# Useful in T/t of PCOD

# Renal route of excretion so C/I in Renal failure.

# Stop metformin 1 day before & 1 day after the  
Radiocontrast exposure.

# N-acetyl cysteine → t/t of Radiocontrast induced  
renal cell injury.

# Metformin Reduces ← Microvascular  
Macrovascular events.

ADR of Metformin: • GI toxicity  
• Inhibit intestinal absorption of glucose, hexose, vit B12.

Metformin causes lactic acidosis in presence of kidney, liver or cardiorespiratory failure, alcoholism.

$\alpha$ -Glucosidase inhibitors: inhibit carbohydrate digestion in small intestine.  
Acarbose  
Voglibose  
Miglitol

- Useful in PP blood glucose.

SE - Flatulence  
Abdominal distension  
Diarrhoea.

CI - in Renal failure.

Thiazolidinediones:

PPAR (Peroxisome proliferated-activated receptor)

activation-PPAR  $\alpha$

PPAR  $\beta/\gamma$

Stimulate lipoprotein lipase  
TGL (VLDL)  $\downarrow$

- Insulin Sensitizer.  
• PIOGLITAZONE

Older drugs:

Withdrawn [ Troglitazone - Hepatotoxic  
Rosiglitazone - CCF

PPAR  $\alpha$  agonist: ( $\downarrow$  TG4)

SE: Myopathy, Hepatotoxicity

- Clofibrate - Not in use (Gall stone, GB malignancy)
- Fenofibrate (Prodrug, longest  $t_{1/2}$ ,  $\downarrow$  LDL,  $\downarrow$  Plasminogen, Uricosuric action)
- Bezafibrate
- Gemfibrozil

# M/c S/E Pioglitazone - Wt. gain  
Macular edema  
Osteoporosis  
Anemia  
Bladder Ca.

Drug activating both PPAR  $\alpha$  &  $\gamma$ :  
SAROGLITAZAR

$\hookrightarrow$  Approved in t/t of Diabetes  
dyslipidemia.

Statins:

HMG CoA + Acetate  
HMG CoA reductase  $\leftarrow \ominus$  - Statins  
Mevalonic acid  
 $\downarrow$   
Cholesterol  $\downarrow$

# Statins  $\rightarrow$   $\downarrow$  Total cholesterol  
# Statins  $\rightarrow$   $\downarrow$  LDL (by upregulation of LDL receptor in liver)

S/E  $\rightarrow$  Myopathy  
Hepatotoxic  
Teratogenic

# Co-enzyme Q given c statins to control muscle weakness.

# Liver enzyme goes more than 3 times (N)  
- Stop Statins.

COLESEVELAM



Only cholesterol lowering agent in pregnancy.

#

# PCSK9 inhibitor:

ALIROCUMAB } Monoclonal antibodies  
EVOLOCUMAB } for Hypercholesteremia.

# Nicotinic acid (Vit B<sub>3</sub>) - Niacin

↓ LDL

↓ LP(a)

↑ HDL

S/E - Cutaneous flushing → (Niacin promotes the synthesis of vasodilatory PGs)



So, Aspirin added c Niacin to control flushing.

Hyperuricemia

Diabetes (causing Insulin Resistance)

Hepatotoxicity

EZETIMIBE: inhibit cholesterol absorption in intestine.

Bile acid sequestrants:

- Cholestyramine
- Colestipol
- Colesevelam

↳ approved for t/t of DM.

MIPOMERSEN: Newer drug

Given s/c Once in a week.

Useful for lowering cholesterol.

PROBUCOL: Inhibits LDL oxidation

GUGULIPID: ↓ LDL (Not use - Diarrhoea)

CETP inhibitors: (Cholesterol ester transport protein)

TOR CETRAPIB

Dalce trapib

Evace trapib

Amace trapib.

MTP inhibitor (Microsomal triglyceride transporter<sup>protein</sup> inhibitor)

LOMITAPIDE

AVASIMIBE: Inhibit conversion cholesterol to cholesterol ester.

ACAT-1 inhibitor.

## Antithyroid drugs:

Histology of thyroid gland -

Steps of Synthesis:

- ① Iodide uptake
- ② Oxidation of iodine & formation of Iodine
- ③ Organification (Iodine + Thyroglobulin)
- ④ Coupling  
 $MIT + DIT = T_3$   
 $DIT + DIT = T_4$

$T_3$  &  $T_4$   
# Stored in follicle for 3-4 days.

## THIOAMIDES:

→ Rapid control of hyperthyroidism

- Propylthiouracil (also inhibit peripheral conversion of  $T_4 \rightarrow T_3$ )
  - Carbimazole (Prodrug)
  - Methimazole
- active form

- inhibit synthesis of  $T_3$  &  $T_4$
- inhibit formation of new thyroid hormone
- lag period of 1-3 wks.

M/c S/E of Carbimazole & Methimazole: Maculopapular rash (4-6%)

Agranulocytosis (0.1-0.5%)

\* Severe hepatitis - PTU

Causing teratogenicity - Fetal aplastic cutis  
Choanal atresia.

Hepatotoxic - PTU

PTU - Used in emergency hyperthyroid crisis.

- may be safe in pregnancy



### LUGOL'S IODINE :

MOA - Inhibits release of  $T_2$  &  $T_4$  from follicle.

- Fastest acting antithyroid drug.
- Used in post-op preparation.
- Reducing vascularity.

S/E - Iodism - Acne form skin rash.

Peripheral conversion of  $T_4$  -  $T_3$  inhibitor:

$\beta$ -Blockers  
Amiodarone  
Propyl thiouracil  
Dexamethazone  
Iodate

By inhibiting 5-DE  
iodinase.

Iodide uptake inhibitor:

POTASSIUM PERCHLORATE  
THIOCYANATE

- Used in t/t of iodide induced hyperthyroidism.

Radioiodine therapy:

$I^{131} \rightarrow t_{1/2} = 8 \text{ days}$

$\hookrightarrow$  emits 2 rays  $\left\{ \begin{array}{l} \gamma \\ \beta \end{array} \right.$

Penetrating power = 0.5 - 2mm.

#  $\gamma$ -Ray useful for diagnostic purpose  
#  $\beta$ -Ray " " therapeutic "

# C/I - Pregnancy, young children, Ophthalmopathy.  
# Not useful for t/t of Medullary CA thyroid.

Newer drug for T/T of Medullary Ca thyroid:

LENVATINIB - BTC

VANDETANIB - MC

# Non-thyroid drug causing Hypothyroidism:

LITHIUM (stop release of  $T_3$  &  $T_4$  from follicle)

AMIODARONE

PROPRANOLOL ] (inhibit conversion of  $T_4 \rightarrow T_3$ )

ETHIONAMIDE ]

PAS ]

inhibit synthesis

SODIUM NITROPRUSSIDE - inhibit uptake of Iodide.

Growth Hormone Release inhibitor

- for t/t of Acromegaly

OCTREOTIDE ]

LANREOTIDE ] s/c

GH Receptor inhibitors -

PEGVISOMANT - s/c

$D_2$  analogue -

BROMOCRIPTINE ]

CABERGOLINE ]

oral

Octreotide - 40 times more potent than Somatostatin

longer acting - 12hrs.

Given (s/c) or i.v.

Never orally.

Uses - Acromegaly

Carcinoid [Diarrhoea]

AzD

Portal HTN (Bleeding esophageal varices)

S/E - Gall stone  
Vit B<sub>12</sub> deficiency (Megaloblastic anemia)  
Rarely DM also.

Dwarfism: T/E  
GH releasing factor analogue:  
SERMORELIN  
HEXARELIN  
TESAMORELIN

↳ For lipodystrophy in HIV pt.  
↓ Abdominal fat.

GH analogues

SOMATREM } also used in - AIDS related wasting  
SOMATROPIN } Turner Syndrome.  
Pituitary dwarfism.

S/E - Insulin resistance - Type 2 DM  
↑ ICT.

↳ To rule out Papilledema  
→ Fundus exam<sup>n</sup>

# Analogue of IGF + IGF binding protein 3  
MECASERMIN (S/c)

↓  
to maintain stability.

S/E - Hypoglycemia

Uterus: OXYTOCIN

- ↑ force / frequency of contraction.
- ↑ contractility to fundus & body, lower segment not contracted unlike ergometrine & methyl ergometrine.
- Useful in induction of labour.

- # Control post partum hemorrhage
- # Useful in ejection of milk.

ATOSIBAN - Oxytocin Receptor ~~Agonist~~ Antagonist.

Tocolytic of choice in heart ds -  $MgSO_4$

ZOLENDRONATE - Bisphosphonate given i.v.  
once ~~of~~ in a year

DOC for postmenopausal osteoporosis.

NATALIZUMAB - Useful for Multiple sclerosis  
given once in a month.

MIPOMERSEN - ↓ cholesterol level  
given s/c once in a week.

DALBAVANCIN - Glycopeptide  
Antibiotic  
Give once in 6-10 days.  
Single dose act 6-10 days

## Drugs for Osteoporosis

Drugs inhibit Osteoclast:

Bisphosphonates

↳ DOC: Zoledronate

Estrogen & SERM

Cinacalcet

Calcitonin

Thiazide diuretics

Denosumab - Rank L antibody.

↳ Monoclonal antibodies

Drugs promoting osteoblast:

Calcitriol (Active form of Vit D)

Androgens & Anabolic steroids

Calcium

Parathormone

(hPTH 1-34) → Teriparatide.

↓ ↳ PTH analogue

given only for 1yr (Max<sup>m</sup> 2yr)

long term therapy cause Osteosarcoma.

## STRONTIUM RANALATE

↳ Dual action ← promoting osteoblast  
inhibiting osteoclast.

## ZOLENDRONATE:

- Anti osteoclastic activity
- Interference on mevalonate pathway -  
antitumour activity (CML)
- Faster acting.
- DOC in Hypercalcaemia (osteonecrosis of jaw).
- Also used in Paget's ds.

- Less venous irritant
- Renal toxicity.

S/E - • Thrombophlebitis  
 • - During infusion Fever + chills  
 "Infusion reactn"  
 • Nephrotoxicity.  
 • Osteoporosis of jaw bone.

# M/c drug for steroid induced osteoporosis  
 - Bisphosphonate.

# Osteonecrosis of Neck of femur - S/E of steroid.

STEROIDS:

1. GLUCOCORTICOIDS:

CLASS A → Short acting (Duration < 12hrs)

Max <sup>m</sup> mineralocorticoid activity →	Glucoc	Mineralo
Hydrocortisone	1	(1)
Cortisone	0.8	0.8
(Least potent G)		

CLASS B → Intermediate acting (duration 12-36 hrs)

Prednisone	4	0.8
Prednisolone	4	0.8
Methyl prednisolone	5	0.5
Triamcinolone	5	0
Deflazacort	5-6	0

**CLASS C : Longer acting (> 36 hrs)**

Paramethasone	10	0
Betamethasone (Most potent G)	25	0
Dexamethasone (Max <sup>m</sup> G)	30	0

**Mineralocorticoids :**

• Natural.

Aldosterone	0	3000
-------------	---	------

• Synthetic

DOCA	0	20
------	---	----

Fludrocortisone	10	250
-----------------	----	-----

- # Max<sup>m</sup> glucocorticoid action - Dexamethasone
- Max<sup>m</sup> mineralocorticoid action - Aldosterone
- G<sup>c</sup> max min - Hydrocortisone
- Least potent G - Cortisone
- Most " " - Betamethasone
- Max<sup>m</sup> topical action - Triamcinolone
- Selective glucocorticoid (No mineralo) - TPDB
- Selective Mineralocorticoid (No Gluco) - DOCA

Steroid — Anti-inflammatory  
Anti cancer  
Immunosuppressive

Anti-inflammatory action of steroid  
— By inhibiting Phospholipase A<sub>2</sub>

ZILEUTON — inhibit lipoxygenase  
Not in use  
Severe hepatotoxic

NSAID — Inhibit Cyclooxygenase.

Steroid having anti-cancer activity:  
— Apoptosis of T & B cells  
— Useful for Lymphoma.

Steroid having immunosuppressive action:  
— Inhibit IL-1 & IL-6  
— Also catabolism of IgG.

Methylprednisolone — Used in pulse therapy.

ACTH

Cosyntropin — Infantile Spasm.



Medulla - Pheochromocytoma  
Adrenal cortex - Cushing Syndrome

Page No. 121

Date: / /

Drug useful for t/t of Cushing Syndrome:

Metyrapone ( $11\beta$ -hydroxylase)

Ketoconazole

Mitotane

Amino glutethamide

Trilostane

Etomidate (General anesthetic)

]- chemical adrenalectomy

PASIREOTIDE - Somatostatin analogue

useful in t/t of Cushing Syndrome.

Erectile dysfunction:

① Selective PDES blocker:

Sildenafil

Vardenafil

Tadalafil - longest acting

Avanafil

- PDES enzyme is involved in metabolism of cGMP.
- PDES blocker by blocking cGMP metabolism causes vasodilation.

Acute adverse effect - Headache

Flushing

Hypotension

Nasal congestion

Long term (chronic) therapy causes Blue vision defect.

↓  
blocking PDE6

Drug interaction b/w sildenafil & Nitrates:

Nitrates shouldn't be given c̄ sildenafil  
bc̄oz risk of severe hypotension.

Other drug for erectile dysfunction:

Amorphine (D<sub>4</sub> agonist)

Trazadone (Atypical antidepressant)

Avaptadil (VIP - Vasoactive intestinal polypeptide)

Ketanserin (Serotonin antagonist)

Naltrexone (Opioid Antagonist)

~~Ginseng~~ Ginseng

Kava

Ginkgo

Injectable therapy for Erectile dysfunction:

Alprostadil

Phentolamine

Papaverine.

Drugs useful for t/t - Premature ejaculation.

- SSRI

- PDEV inhibitors

For delayed orgasm:

Amantadine

Buspirone

Cyproheptadine.

For sexual stimulation:

- Yohimbine

Zinc

Ginkgo biloba

~~Ginseng~~ Ginseng.

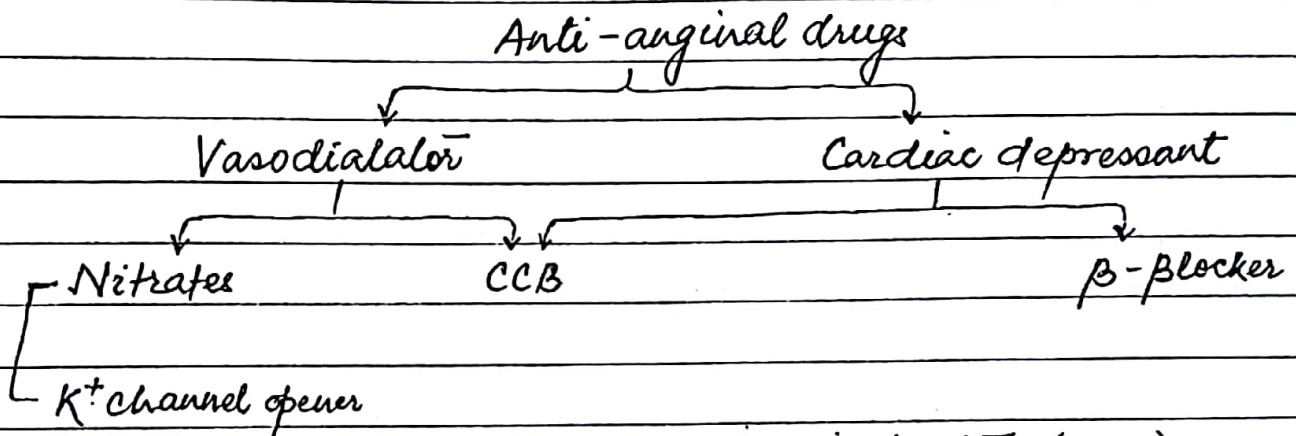
## ANTI ANGINAL DRUGS

Stable Angina

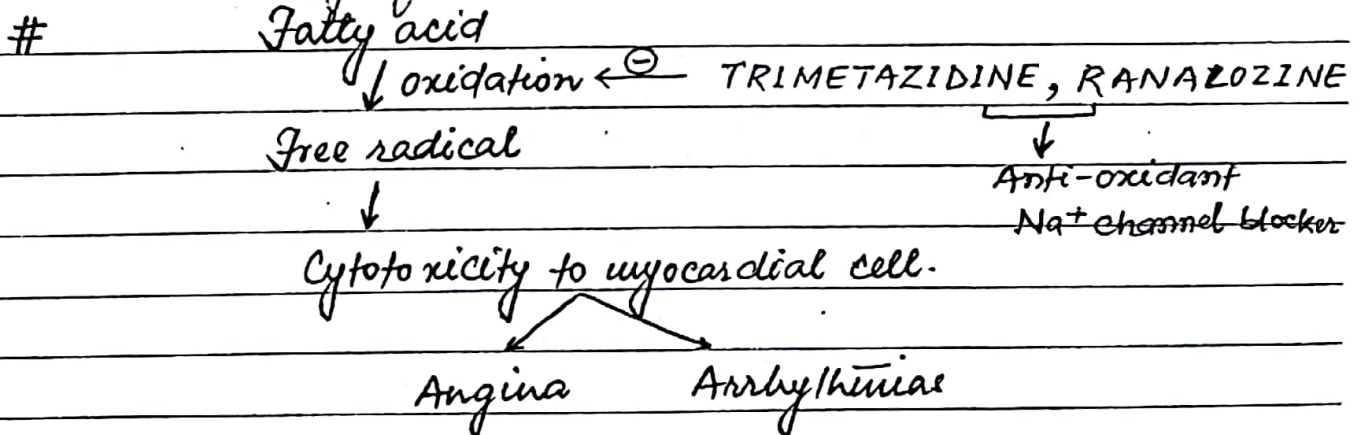
Unstable Angina

Vasospastic angina (Prinzmetal Angina) (variant angina)

Cause  $\left\{ \begin{array}{l} \text{Reduction in } O_2 \text{ supply} \\ \uparrow O_2 \text{ demand.} \end{array} \right.$



Pathway of FA oxidation inhibitors (pFox)



S/E - GI toxicity (M/C)

Thrombocytopenia

Liver dysfunction

Risk of movement disorder - C/I in Parkinsonism

QT - ~~prolongation~~ prolongation -

Excretion by Renal pathway - C/I in Renal failure

## NITRATES

Short acting	Intermediate acting	Long acting	Longest acting
• GTN	• Isosorbide	• Isosorbide	• Pentaerythritol
• Amyl Nitrite (Shortest)	dinitrate (2-3hrs)	mononitrate (6-10hrs)	tetranitrate (8-12hrs)

# For acute attack - GTN, Isosorbide dinitrate  
S/L

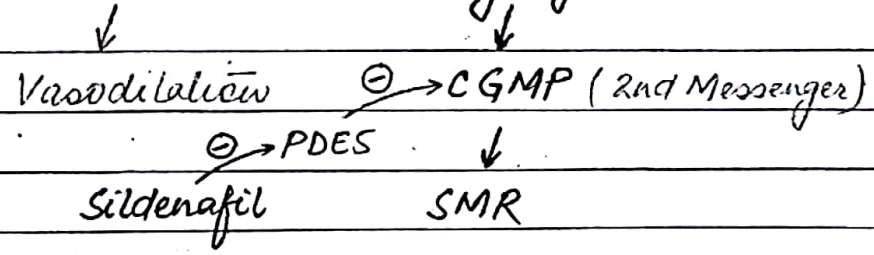
Least 1st pass metabolism - Isosorbide mononitrate.

S/L drug - Lipid soluble  
 Non ionised

Skin rash - Pentaerythritol tetranitrate

MOA of nitrates:

- Nitrates acting on Cysteine receptor, they release NO. NO activate Guanyl cyclase.



NO independent - direct Guanyl cyclase activators:

- RIOCIQUAT
- CINOCIGUAT

- Useful for t/t of Primary pulm. HTN.

# CGMP normally undergo inactivation by PDES enzyme.  
 So, PDES inhibitor = Sildenafil group of drug.

# Nitrates may get tolerance due to down regulation of receptors.

Max<sup>m</sup> Tolerance - i.v. infusion  
& Transdermal patches.

Action of Nitrates:

Visceral smooth muscle - Relaxed

- ↳ Useful for t/t of Biliary colic pain
- ↳ Useful for t/t of Achalasia cardia

Vascular smooth muscle - Vasodilator

↓  
predominantly Venodilator  
- Peripheral pooling of blood

↓  
max<sup>m</sup> ↓ in Preload.  
mild ↓ of afterload.

↓  
↓ O<sub>2</sub> demand

↓  
Reduce angina.

Uses: Cardiac uses: Angina  
MI  
CCF

Non-cardiac uses: Biliary colic pain  
Achalasia cardia  
Cyanide poisoning.  
↳ By formation of Methemoglobinemia.

ADR - Throbbing Headache (M/c)

Hypotension

→ Reflex Tachycardia (due to Sympathetic stimulation)

Tolerance

So add  $\beta$ -blocker.

Methemoglobinemia

Rashes

# Drug interaction b/w Nitrates & Sildenafil:

- Not combined together bcoz it cause severe hypotension.

Gap of 8-10 hrs should be maintained.

Sodium Nitroprusside:

- Only i.v. route

- Short acting < 10 min

Indication - Hypertensive emergency.  
Acute aortic dissection.

- Drug is sensitive to light

↳ Cover w/ black towel.

- Containing cyanide (Thiocyanate)

↓

Risk of Hypothyroidism

- C/I in pregnancy.

$\beta$ -blockers:

• ↓ Workload of cardiac.

• C/I in variant angina.

• Abrupt withdrawal ppt. angina.

•  $\beta$ -blocker + ~~GTN~~ GTN = to prevent Reflex Tachycardia.

• Controls catecholamine activity

↓

### Role of $\beta$ -blocker on MI :

- Reduces size (zone) of infarction
- Anti arrhythmic action
- Reduces mortality.

### CCB :

Chemical Type	Chemical names
Phenylalkylamines	Verapamil
Benzothiazepines	Diltiazem.
1,4-Dihydropyridines (DHP)	Nifedipine Nicardipine Nimodipine Amlodipine Nitrendipine (NO releasing property)

# Nebivolol ]  $\beta$ -blocker having NO releasing property.  
Nepradiol ]

### DHP :

Site of action - Peripheral blood vessel

↓  
Vasodilatation

- Useful for Ht of HTN & PVD.

- ↳ Maximally arterial dilatation.
- ↳ max<sup>m</sup> ↓ in PVR.

ADR → Hypotension

Reflex Tachycardia

Ankle edema (Amlodipine max<sup>m</sup> cause Ankle edema)

Constipation

→ long acting  
# Nicardipine } Approved in Hypertensive emergency.  
Clevidipine } given i.v.  
→ Short acting

Non-dihydropyridines: Verapamil  
Diltiazem.

Verapamil:

Site of action: AV node (Most imp.)  
SA node

Action → Bradycardia  
→ Anti arrhythmic agent.

Uses - Atrial Tachyarrhythmia (AT)  
SVT (Supra Ventricular Tachyarrhythmia)

ADR - Bradycardia

Block AV conduction - Prolongation of PR interval.

Ankle edema

Constipation

CI - WPW syndrome.

Diltiazem:

Uses - HTN

Angina

Arrhythmias (SVT/AT)

CCB having anti-arrhythmic property ← Verapamil } class IV  
Diltiazem } antiarrhythmic



Nimodipine: Cerebro-selective CCB

Useful for t/t of Sub-arachnoid hemorrhage (SAH)

The purpose of given Nimodipine is to prevent Reflex ischemic <sup>brain</sup> damage.

FASUDIL - Rho Kinase inhibitor

Use - SAH

~~CCB~~ PHT (Pulm. HTN)

Angina.

CCB useful in Prophylaxis of Migraine - Verapamil

Flunarizine

↓  
T-type of CCB  
Na<sup>+</sup> channel blocker  
Anti-oxidant.

K<sup>+</sup> channel openers:

Hydralazine } - Arteriolar dilator

Minoxidil } - Anti-hypertensive

Diazoxide }

Nicorandil (Anti-anginal)

Adenosine (PSVT) → DOC

Nicorandil: NO releasing property

Anti-anginal

S/E → Aphthous ulcer

Headache

### Hydralazine:

- T/t of HTN-emergency in pregnancy
- NO releasing property
- Metabolism by Acetylation

↓  
S = Sulphonamide  
H = Hydralazine  
I = Isoniazid  
P = Procainamide.

- Cause RA/SLE

### Minoxidil:

- Prodrug
- Active form → Minoxidil Sulphate.

Uses → HTN  
Alopecia

### Diazoxide:

- causing hyperglycemia by inhibiting insulin release from  $\beta$ -cell of pancreas.

Use - HTN  
Insulinoma.

↓  
Phenytoin - also inhibit release of insulin. causing  
Poor man drug for Insulinoma.

### IVABRADINE -

- Causing Bradycardia.
- $Na^+$  channel blocker (Ifunny Current)
- Reduce HR.

Two indications  $\left\{ \begin{array}{l} \text{CCF} \\ \text{Angina.} \end{array} \right.$

S/E - On chronic therapy - Causes Luminous phenomena.  
(Visual disturbance)

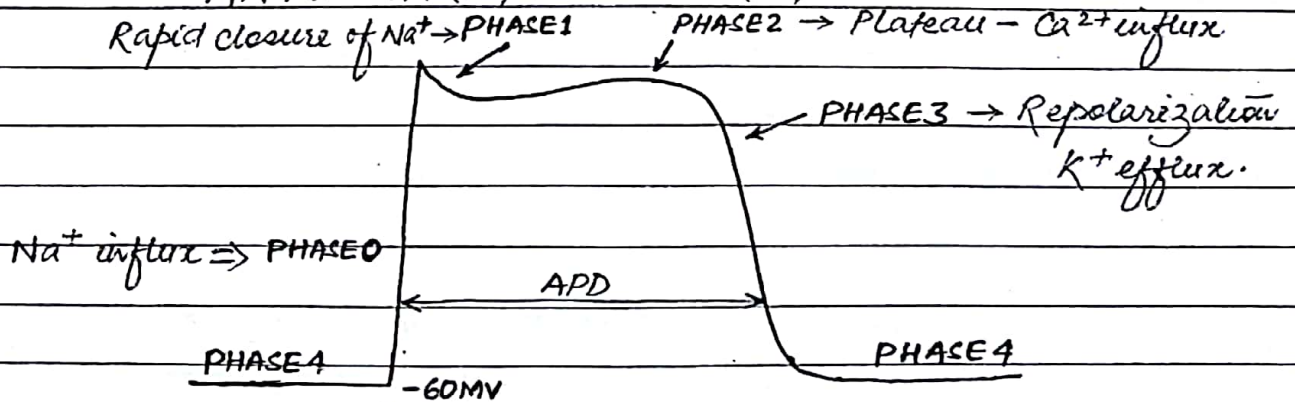
# Hemeralopia - Trimepradione (Withdrawl - due to Nephrotoxicity)  
↓  
Day blindness

Reperfusion - Thrombolysis/PTCA

Drug eluting stent:

$\left\{ \begin{array}{l} \text{SIROLIMUS (Immunosuppressant)} \\ \text{PACLITAXAL (Anti cancer drug additional} \\ \text{immunosuppressant)} \end{array} \right.$   
used  $\bar{c}$  stent to decrease rejection.

**ANTI-ARRHYTHMIC DRUGS:**



PHASE 3  $\rightarrow$  T WAVE

PHASE 2  $\rightarrow$  ST segment

PHASE 0, 1 & mid phase of 2  $\rightarrow$  QRS

APD (Action potential duration)  $\rightarrow$  QT interval.

# Any drug having  $K^+$  channel blocking property  
- Cause QT prolongation

- Class Ia & class III drug having  $K^+$  channel blocking property causing QT prolongation.

Classification: Vaughan Williams

Class I -  $Na^+$  channel blocker

↳ Class IA, IB, IC

Class II -  $\beta$ -blocker

Class III -  $K^+$  channel blocker.

Class IV - CCB

Unclassified & Miscellaneous agent



Adenosine

Atropine

Digoxin

Magnesium Sulfate

KCl

Class IA:

- Block  $Na^+$  channel +  $K^+$  channel block

- Having risk of causing QT prolongation.

Eg: Quinidine

Procainamide

Disopyramide.

} Anti vagal action

Quinidine -

Origin - Cinchona bark

↳ Symptom - ~~CI~~ Cinchonism

↓  
Tinnitus

S/E → Diarrhoea

Hypotension (Bcoz d blocking property)

Hypoglycemia (Bcoz Insulin releasing property)

SMR

Thrombocytopenia.

Drug interaction: Quinidine + Digoxin

Quinidine interfere renal excretion of Digoxin.

∴ aggravating plasma level of Digoxin

↓

∴ Digoxin toxicity.

Procainamide:

S/E - Undergo metabolism by Acetylation

SLE.

Disopyramide:

Highest anticholinergic action.

Dry mouth, constipation, Retention of urine.

↓

Not safe in elderly male  $\bar{c}$  BPH.

Class IB:

Na<sup>+</sup> block + K<sup>+</sup> opening.

- Never causes QT prolongation.

Site of action → Mainly acting on Bundle of HIS.

Rt. Bundle, Lt. Bundle & Purkinje fibres.

Used <sup>only</sup> for t/t → Ventricular arrhythmias  
(Tachycardia)

eg: Lignocaine (Lidocaine)  
Mexilitine  
Phenytoin  
Tocainide.

Mexilitine:

- Lignocaine derivative
  - Useful for t/t Ventricular arrhythmias.
  - Used for Diabetic neuropathy pain  
(Unlabeled Use)
  - Used for Phantom limb pain
- ADR - Severe Nausea & Tremor.

Phenytoin:

- Antiepileptic
- USE - t/t of Digitalis (Digoxin) induced VT

Tocainide:

Beoz of causing Agranulocytosis it is not used.


Lignocaine:

- Class IB drug
  - Never given orally beoz undergo extensive 1st pass metabolism
  - Given i.v.
  - Lipid soluble, Cross BBB.
- S/E - Convulsion  
↳ Sign - Nystagmus (1st sign)  
1st Symptom - Circum oral paraesthesia

Use - VT (Ventricular Tachycardia)  
VF (Ventricular Fibrillation)  
Digoxin induced VT (DOC: Lignocaine)

↳ # Class IB drug has no role in atrial arrhythmias

Class IC:

- $\text{Na}^+$  blocking + Negligible effect on  $\text{K}^+$  channel.
- Max<sup>m</sup> pro-arrhythmic property.
- Non commonly used.
- Only for  antiarrhythmic drug causing arrhythmia.

Flecainide (DOC: for Acute WPW)  
Encainide  
Propafenone  
Moricizine

PROPAFENONE:

- Also  $\beta$ -blocking property.

Class III:  $\text{K}^+$  channel blocker

- Prolong APD  $\rightarrow$  QT prolongation

AMIODARONE:

- Iodine containing anti-arrhythmic drug.
- Multi MOA:  $\text{K}^+$  channel blocking
- $\text{Na}^+$  channel blocking
- $\beta$ -Blocker property
- CCB property.
- $\therefore$  Broad spectrum Anti-arrhythmic.

Half life = 53 days.

USES: All type of arrhythmias  
Ventricular & Supraventricular arrhythmias.

ADR:

PLZ = Photosensitivity, Pigmentation of skin (Gray-blue)

Check = Corneal deposition (Whorl like pattern cornea)

PFT = Pulm. fibrosis, Peripheral neuropathy.

LFT = Liver damage, Pseudo alcoholic liver injury & Mallory  
Jhyline bodies.

TFT = Hypothyroidism

- due to inhibition of peripheral conversion of  $T_4 \rightarrow T_3$

Hyperthyroidism

Whorl like pattern cornea - Cornea / Verticillata  
or Vertex Keratopathy.

[ Pseudo lymphoma - Phenytoin  
Pseudo jaundice - Rifabulin

Amiodarone causing Hyperthyroidism due to:

Hypothyroidism: inhibition of peripheral conversion of  $T_4 \rightarrow T_3$ .

- Hyperthyroidism
- ① Contain Iodine  $\rightarrow$  Iodine help in synthesis of  $T_3$  &  $T_4$
  - ② Can cause inflammation of follicle.

# In each 200mg tablet there is 75mg of iodine.

Rx: Inhibit iodide trapping

- Perchlorate
- Thiocyanate.

For inflammation - Rx: Dexamethasone (steroids)



### Class III drugs:

Amiodarone

Dronedarone (No iodine)

Bretylium (Chemical defibrillator)

Sotalol

Dofetilide

New drug [ Ibutilide (FDA approved for conversion of AF-SR) - i.v.  
Vernakalant

### Class IV: CCB

Verapamil (Most potent)

Diltiazem

### Miscellaneous drugs:

#### ADENOSINE:

- Given i.v., short acting, Rapid infusion (Bolus)

Site - Close to heart.

- DOC for SVT

- It is also called Endogenous epileptic.

Antagonist - Methyl xanthine - theophylline

Agonist - Dipyridamole

Cause  $\rightarrow$  Coronary Steal Phenomenon.

For Acute SVT: i.v. Adenosine

i.v. Verapamil.

$\rightarrow$  Prefer in Asthma  $\bar{c}$  SVT.

To prevent recurrence of SVT: Oral  $\beta$ -Blocker

Oral Verapamil.

$MgSO_4$  :  
USE  $\rightarrow$  ① CNS

$\hookrightarrow$  Long QT syndrome

Congenital  $\beta$ -blocker  
(Propranolol)      Acquired.  
 $MgSO_4$

USE:  $\hookrightarrow$  Digitalis intoxication

$\downarrow$   
Hypokalemia

Hypomagnesimia  $\rightarrow$  Give  $MgSO_4$

Hypercalcemia

② Resp<sup>r</sup> System

USE: Bronchial asthma

③ GIT (laxative property)

USE: Constipation.

④ Ortho (anti-inflammatory property)

USE: Synovitis.

⑤ Obs. & Gyn.

USE: Eclampsia.

S/E - Diminished deep tendon reflex (M/C)

Rarely Resp<sup>r</sup> failure.

Safety limit - 4m Eq/L

If  $> 7m$  Eq/L  $\rightarrow$  Patellar reflex  $\downarrow$   $\uparrow$

$> 14m$  Eq/L  $\rightarrow$  Resp<sup>r</sup> failure.

Antidote - Calcium Gluconate.

ATROPINE:

- Anti-cholinergic agent.
- Causing Tachycardia.
- USE - Bradycardia or Heart Block.

DIGOXIN: Already discuss

Cardiac glycosides:

	Digoxin	Digitoxin
T <sub>1/2</sub>	40hrs	5-7 days
Route of excretion	Renal	Hepatic
Plasma conc <sup>n</sup>	0.8-1.5 ng/ml	15-30 ng/ml.

- Both have narrow therapeutic index  
 i.e. Unsafe & need monitoring.

Digoxin S/E: <sup>Non-</sup> Cardiac S/E

- Nausea & Vomitting (M/c)
- CNS depression
- Yellow vision defect (Xanthopsia)
- Gynecomastia (In male)

Cardiac S/E

- Atrial Tachyarrhythmia (AT)
- AV block

VT (Ventricular Tachycardia)

Ventricular Bigeminy (M/c)

Non-paroxysmal AT & variable AV block  
 ↳ Most characteristic arrhythmia.

For t/t digoxin induced AT — Propranolol.

# Atropine → AV Block.  
Lignocaine → VT

# No role of Hemodialysis in digoxin toxicity  
bcuz large Vd.

# Antidote for digoxin toxicity — Digibind.

↓  
Check S.  $K^+$ ,  $Mg^{2+}$ ,  $Ca^{2+}$

~~DIURETICS~~ DIURETICS.

In the PCT → Carbonic anhydrase



Reabsorption of  $\text{NaHCO}_3$  (85%)  
 Reabsorption of  $\text{NaCl}$  from urine (60%)

Thin descending limb - Absorption of  $\text{H}_2\text{O}$   
 ↳ Concentrating Segment

Thick ascending limb →  $\text{Na}^+ - \text{K}^+ - 2\text{Cl}^-$  Symporter



Absorption of  $\text{Na}^+$ ,  $\text{K}^+$ ,  $\text{Cl}^-$ ,  $\text{Ca}^{2+}$ ,  $\text{Mg}^{2+}$ .  
 (Diluting segment) (25%)

DCT →  $\text{Na}^+ - \text{Cl}^-$  Symporter



Reabsorption of  $\text{NaCl}$  (10%)  
 Reabsorption of  $\text{Ca}^{2+}$  (+PTH)  
 ↑  
 e help of

CT → Reabsorption of  $\text{NaCl}$  (e help of aldosterone) (5%)  
 Secretion of  $\text{K}^+$  &  $\text{H}^+$   
 Reabsorption of  $\text{H}_2\text{O}$  (e help of ADH)

Primary Hyperaldosteronism (Conn's Syndrome):  
 ↑↑ Aldosterone

CF - HTN

Hypokalemia

Metabolic alkalosis.

For t/t HTN →  $\text{K}^+$  sparing antidiuretic  
 ↳ Spironolactone.

## Carbonic anhydrase inhibitors:

Acetazolamide }  
Dorzolamide } Non-competitive & Reversible.  
Brinzolamide }

Site of Action - PCT

MOA - Inhibit Carbonic Anhydrase.

ADR - ① Loss of  $\text{HCO}_3^-$  }  
Metabolic acidosis.

# Acetazolamide causing Alkaliuria  
↳ So used in Alkalinisation of urine.

② Max<sup>m</sup> potassium loss.

# CA inhibitor also acting on collecting duct - it inhibit tubular secretion of  $\text{H}^+$  → so cause Metabolic acidosis & massive Hypokalemia.

# CA inhibitor are Sulpha derivative:

SE - Hypersensitivity  
Bone marrow suppression.

# C/I - liver disease (hepatic encephalopathy)

COPD

Metabolic acidosis.

Loop Diuretics: High ceiling diuretic ( $\uparrow$  dose  $\rightarrow$   $\uparrow$  diuretic action)  
 Site of action: Thick ascending loop of Henle

↓  
 MOA: Inhibiting  $\text{Na}^+ - \text{K}^+ - 2\text{Cl}^-$  symport

↓  
 Loss of  $\bullet \text{Na}^+, \text{K}^+, \text{Cl}^-, \text{Ca}^{2+}, \text{Mg}^{2+}$

Eg: Furosemide  $\rightarrow$  Vasodilatory action (USE: RF, LVF)

Bumetanide  $\rightarrow$  Most potent

Mersalyl  $\rightarrow$  Kidney damage (Not in use)

Ethacrynic acid  $\rightarrow$  Highly ototoxic (No CA enzyme inhibition)

Torsemide  $\rightarrow$  Longest  $t_{1/2}$

Role of Furosemide in Renal failure:

Furosemide promote <sup>the synthesis of</sup> vasodilatory ~~action~~ PG

↓  
 By  $\uparrow$  intrarenal blood supply

↓  
 Improving Renal failure

NSAID + Furosemide  $\rightarrow$  NSAID is not given  $\bar{c}$  Furosemide  
 in Renal failure pt. ~~by~~ bcoz it inhibit  
 synthesis of PG.

# Diuretics of choice in the presence of RF

Choice - Furosemide

ineffective - Thiazides  $\bullet$

Exception - Metolazone

OI -  $\text{K}^+$  sparing drugs.

Role of loop diuretics in heart failure:

Furosemide - Only Relief symptoms of CHF.

↓ diuretic action

Main mech<sup>n</sup>: Vasodilation

↓

Beoz of vasodilation Furosemide (i.v.) rapidly relief breathlessness in CHF.

S/E of loop diuretics:

Water loss	Electrolyte imbalance	Metabolism	Miscellaneous.
Profound ECFV Depletion	Loss of $\text{Na}^+$ , $\text{K}^+$ , $\text{Cl}^-$ , $\text{Ca}^{2+}$ , $\text{Mg}^{2+}$ ↓ Calciurea (Risk of kidney stone)	Hyperuricemia Hyperglycemia Hyperlipidemia. ↓ Exception: INDACRINONE (Ethyronic acid derivative) ↓ Uricosuric agent.	Metabolic alkalosis Ototoxicity (Irreversible) → other drugs Aminoglycosides Cisplatin Vancomycin Erythromycin

Drug interaction: Loop diuretics + Arrhythmia

- loop diuretics by causing hypokalemia & hypomagnesiemia → causing digoxin toxicity.



### Thiazide diuretics:

Site of action: DCT

- MOA: ① Inhibiting  $\text{Na}^+ - \text{Cl}^-$  Symport.  
② Promotes Reabsorption of  $\text{Ca}^{2+}$

↓  
Causing hypercalcaemia (Urine  $\text{Ca}^{2+} \downarrow$ )

↓  
Safe for Renal stones.

③ Also having antidiuretic activity.

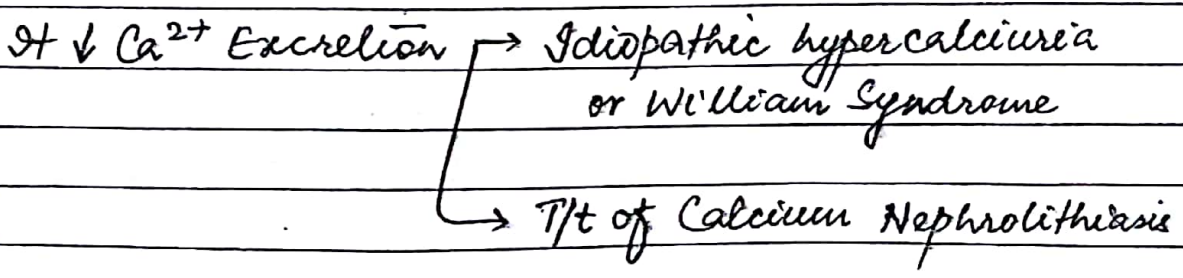
- eg: Indapamide → Vasodilatory action (No CA enzyme inhibition)  
Chlorthalidone → longest acting  
Metolazone → Useful even in severe RF.

- # A/c to JNC guidelines, the 1st line drugs are:  
Thiazides - type diuretics  
CCB  
ACE inhibitors  
ARB's

### Therapeutic effect:

- As a diuretic — ① T/t of Mild edema  
② T/t of HTN

As a anti-diuretic — T/t for Nephrogenic DI.



## Adverse effects:

Water loss	Electrolyte abnormality	Metabolism	Miscellaneous.
ECFV depletion	Hypokalemia Hyponatremia Hypercalcemia ↓	Hyperuricemia <u>Hyperglycemia</u> ↑ LDL ↓	Metabolic alkalosis Impotency (Erectile dysfunction) β-blocker also
Use in HT: Osteoporosis		Thiazide causing insulin resistance as well as inhibiting Insulin release ↓	
		HTN & Hyperlipidemia (So don't use thiazide)	

K<sup>+</sup>-sparing diuretics

Aldosterone antagonist	ENa Channel inhibitor
Spiro lactone (M/C) ←	Amiloride
Canrenone (Active metabolite)	Triamterene.
Eplerenon (No gynecomastia)	↓ Pentamidine
Drospirenone (Progesterone)	↓ Trimethoprim.
	→ Anti-microbial having ENa channel inhibitor property.

## ENaC:

- Na<sup>+</sup> from urine in CD is absorbed by ENaC.

Spiroonolactone:

MOA: One & only drug acting on interstitium.

MOA of Amiloride: Amiloride acting from lumen & blocking ENaC.

Therapeutic uses of Spiroonolactone:

↓  
Blocks Aldosterone

- ① T/t for Primary Hyperaldosteronism (Conn's)
  - ② T/t for Edema of liver cirrhosis (Ascites)
  - ③ T/t for Heart failure.
- Disease modifying HF → Spiroonolactone.

Adverse effects:

(M/C) < Hyperkalemia  
Metabolic acidosis.

Long term effect in male - Impotence  
Gynecomastia  
in female - Menstrual irregularities.

Reason of Anti androgenic action.

Drug causing Gynecomastia:

D = Digoxin

I = INH

S = Spiroonolactone

C = Cimetidine

K = Ketoconazole

O = Oestrogen/anti-androgen → Finasteroid

↓

T/t of male pattern baldness.

# Drug useful in painful Gynaecomastia - Tamoxifen.  
(DOC)

Therapeutic effect of Amiloride:



Block  $\text{Na}^+$  channels

- ① T/t of Liddle's Syndrome ( $\uparrow$  ENaC)
- ② T/t of lithium induced DI
- ③ T/t Aerosol - Cystic fibrosis. (Mech<sup>n</sup> not known)

Mannitol - Osmotic diuretics

Site - LOH & PCT

Useful for T/t of ① Glaucoma (Given i.v.)

② Cerebral edema

③ Cisplatin toxicity.

↳ Antidote - Amifostine.

Mannitol added c cisplatin to control Nephrotoxicity.

CI - Pulm. edema (LVF)

Cerebral Hemorrhage

S/E - Hyponatremia  
Headache.

## ANTIDIURETICS

- ADH (Vasopressin)

V<sub>2</sub> Receptor:Location → V<sub>2</sub> seen on medullary portion of collecting duct

Action → Water Reabsorption

• Also seen on Vascular epithelium

Action → Releasing vWF &amp; Factor VIII

Desmopressin:

- Synthetic analogue of Vasopressin acting on V<sub>2</sub>

USES: DOC for Cranial diabetes insipidus

DOC for Nocturnal Enuresis.

Useful for Hemophilia

" " Bleeding due to deficiency of vW factor.

V<sub>1</sub> Receptor:

- Seen on Vascular smooth muscle

Action → Vasoconstriction

V<sub>1</sub> analogues: Synthetic

Terlipressin } - Useful to control esophageal varices

Felypressin }

Lypressin }

↓  
DOC: Octrotide

Prophylaxis DOC: Propranolol

# Terlipressin added c̄ lignocaine to prolong the action.

Selective V<sub>2</sub> antagonist:Oral [ Lixivaptan  
Mozavaptan  
Tolvaptan ] - DOC for SIADH

Selective  $V_1$  antagonist:

Relcovaptan - Useful for HTN

Nelivaptan -  $V_{1B}$  blocker



Undergo clinical trial for  
t/t of Anxiety.

Non-selective  $V_1$  &  $V_2$  antagonist:

CONIVAPTAN ( $V_2 > V_1$ )

↳ USE: SIADH

Given i.v.

## HEMATOLOGY

Thrombolytic Agents:

MOA - Plasminogen activator → PLASMIN  
(Fibrinolysin) (Fibrinolysin)

eg:

M/C S/E

- Bleeding

Streptokinase

Urokinase

Alteplase

Reteplase

Tenecteplase

Antidote of Thrombolytic drugs:

EACA (Epsilon Aminocaproic Acid)

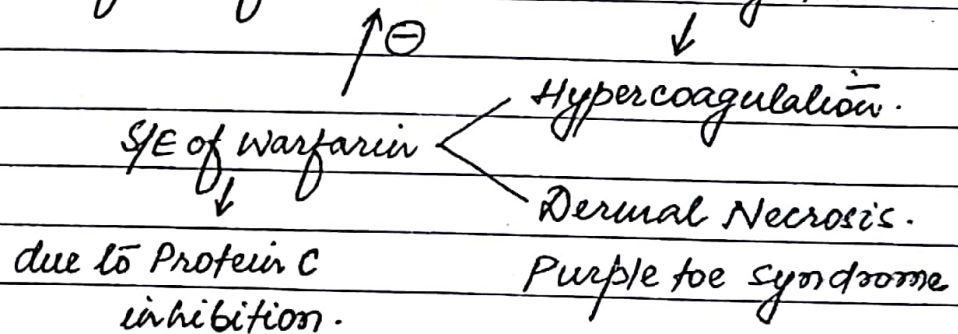
Tranexamic acid

Aprotinin.

WARFARIN: Inhibiting vitK dependent <sup>clotting</sup> factor  
( II, VII, IX, X)

Protein	Half life
Factor II	72 hrs
VII	4-6 hrs
IX	24 hrs
X	44 hrs
Protein C	8 hrs
Proteins	30 hrs.

- For full benefit of warfarin occurs, wait for 3 days.
- Not used in Acute DVT
- Useful in prophylaxis of Chronic DVT.
- Normal func<sup>n</sup> of Protein C → inhibiting Factor V & VIII



Warfarin therapy:

Narrow therapeutic index (Only INR done)

Two isomers  $\left\{ \begin{array}{l} R \\ S \text{ (Active)} \end{array} \right.$

- # CYP2C9 → involved in metabolism of Warfarin
- # Duration of action → 5 days.
- # It undergoes Zero Order Kinetic

Warfarin:  $INR = \frac{\text{Patient PT}}{\text{Control PT}}$

(N)  $\rightarrow 2-3$

Prosthetic Valve  $\rightarrow 2.5-3.5$

Long term  $\rightarrow 1.5-1.9$

C/I in Pregnancy  $\rightarrow$  Teratogenic



Contradi Syndrome

Fetal Chondrodysplasia Punctata.

Antidote of Warfarin -

Natural Vit K<sub>1</sub>  
Phytonadione

Vit K<sub>2</sub>  
Menogquinone

Vit K<sub>3</sub>  
Menadiolone

↓  
Takes about 24hrs  
to reduction INR

For immediate hemostasis - Fresh frozen plasma (FFP)

# New Oral drugs - direct IIa inhibitors

Ximelagatran (Cause severe hepatotoxicity)  
- Not used.

Dabigatran

# New oral drugs: Direct Xa inhibitors

Apixaban

Rivaroxaban

Edoxaban

Betrixaban



Injecting Anticoagulant acting Via Antithrombin III pathway:

Heparin (inhibit Xa; IIa)

LMWH (inhibit Xa)

↳ eg: Enoxaparin

Dalteparin

Tinzaparin

Nadroparin

Other injectable drugs acting via Antithrombin III but only inhibiting Xa:

Fondaparinux

Idraparinux

Idrabiotaparinux  $\xrightarrow{\text{Antidote}}$  Avidin

# Specific antidote for Heparin — Protamine Sulphate  
It is chemical antagonism.

1mg of Protamine sulfate

↓

Neutralizes 100U of Heparin.

Direct Xa inhibitor — Otamixaban

(Under trial)

Injectable — Direct Thrombin (IIa) inhibitor

Bivalent:

Hirudin

Bivalirudin

Lepirudin

Monovalent

Argatroban

(Biliary excretion)

Melagatran

- These drugs are used in pt. who developed Heparin induced Thrombocytopenia.

Adverse drug reac<sup>n</sup>:

Heparin	Warfarin
A = Alopecia	A = Alopecia
B = Bleeding	B = Bleeding
O = Osteoporosis (Supplement Ca)	O = Oral (GI intolerance)
U = Urticaria (Hypersensitivity)	U = Dermatitis
T = Thrombocytopenia	T = Teratogenicity.
Rarely Hyperkalemia	

## Monitoring:

Antiplatelet drugs (Aspirin) - Prolongs BT

Heparin (Intrinsic pathway) - Prolongs aPTT

Warfarin (Extrinsic " ) - Prolongs PT

LMWH - No need of monitoring

If monitor then Antifactor Xa



In Renal failure & Obese pt.

## ANTI PLATELETS

Drugs inhibiting synthesis of TX-A<sub>2</sub>:

Selective COX-1 inhibitor - Low Dose Aspirin  
(50mg-160mg)

Thromboxane synthase enzyme inhibitor - DEZOXIBEN

Drugs inhibiting TX-A<sub>2</sub> Receptor:

IFETROBAN

SULTROBAN

DALTROBAN

LOSARTAN (ARB having ~~an~~ Antiplatelet action)

VAPIPROST

Drugs inhibiting synthesis of TX-A<sub>2</sub> & blocking action of TX-A<sub>2</sub> receptor: Dual action  
PICOTAMIDE

Newer drug: SERATRODAST (Thromboxane A<sub>2</sub> antagonist).

ADP (P<sub>2</sub>Y<sub>12</sub>) blockers:

Ticlopidine } - Prodrug  
Clopidogrel }  
Prasugrel }  
Ticagrelor  
Cangrelor - Given i.v.

Ticlopidine - ~~Not~~ Not commonly used  
becoz thrombocytopenia & Hepatotoxicity.

Clopidogrel - Activated by CYP2C19.

# Omeprazole should n't be given c̄ Clopidogrel.  
Pantoprazol & Rabeprazol don't have drug  
interaction c̄ Clopidogrel.

Glycoprotein IIb/IIIa blocker:

Given i.v. { Abciximab - Monoclonal antibody.  
Eptifibatide  
Tirofiban

PAR1 blocker (Protease activated Receptor blocker)

Vorapaxar  
Atopaxar.

Essential Thrombocytosis:

ANAGRELIDE → Platelet maturation inhibitor.

DOC for Sickle cell Anemia — HYDROXYURIA

↓  
useful in Essential Thrombocytosis.

Drug used for T/t of CCF:

Drugs inhibiting release of Renin:

β-Blocker

Clonidine

Methyl dopa.

Renin inhibitors:

Aliskiren (FDA approved)

Renikiren

Enakiren

ACE inhibitors:

Captopril

Ramipril

Lisinopril

Fosinopril (Renal & Bile excretion)

# All ACE inhibitors are Prodrug except Captopril  
Lisinopril.

# All ACEi are having Renal excretion.

Action → Vasodilation (Equally dilates Artery & Vein)

Useful for → HTN, CCF, MI, DM, Proteinuria, Scleroderma.

↓  
Nephroprotective.

- C/I - ① Pregnancy  
 ② B/L Renal stenosis  
 ③ Severe Hyperkalemia

Bradykinin antagonist: Icatibant



Useful for angioedema & dry cough.

Hereditary angioedema:

C1-esterase inhibitor deficiency.

ICATIBANT

RUCONEST → Human Recombinant C1-esterase inhibitor

Ecallantide

Aprotinin

} Kallikrein inhibitor.

DANAZOL → Antigonadotropin  $\bar{c}$  anti-androgen action  
 (Impeded androgen)

# Sampralilat } - inhibit Vasopeptidase  
 Omapatrilat } - ACEi

Vasopeptide:

PEPTIDE

ANP

BNP

URODILANTIN

Funct<sup>n</sup>

- Natriuresis -

Diuresis

Vasodilation

Synthetic

Analogue

Carperitide

Nesiritide

Ularitide

Nesiritide:

Synthetic analogue of BNP

Action → Diuresis  
Natriuresis  
Vasodilation

Useful for t/t of CCF.

- Given iv, Never oral
- Metabolism → Vasopeptidase
- Shorter ~~life~~ half life - 20 min

S/E - Severe Hypotension

# Other name of Vasopeptidase - Neprilysin  
(Neutral endopeptidase).

Selective Vasopeptidase inhibitor:

Ecadotril

Sacubitril

Omapatrilat } - inhibit Vasopeptidase } Dual enzyme  
Sarpaprilat } - ACEi } inhibitor.

ARB's:

Losartan

Valsartan

Telmisartan

Olmisartan

Azilsartan

- Indication & C/I same as ACEi.

Losartan:

Action → Uricosuric action  
TXA<sub>2</sub> antagonism

Telmisartan

- Agonistic action on PPAR<sub>γ2</sub>  
( Peroxisome proliferator - activated receptor )  
So used in T/t of DM.

Aldosterone Antagonist:

Spiroolactone

Canrenone

Eplerenone

Drospirinone

# ACEi + Spiroolactone ⇒ Severe Hyperkalemia.

Any drug blocking RAAS pathway will cause hyperkalemia.

Other drug useful for t/t of CCF

Phosphodiesterase 3 inhibitors:

Amrinone (Inamrinone)

Milrinone

Levosimendan

Ionodilator.

→ M/c S/E - Thrombocytopenia

M/c S/E of Milrinone - Arrhythmia

Heart failure:  
 $\text{Na}^+ - \text{K}^+$  pump inhibitor: Islāroxime.

Direct myosin activator: Ome●camitū mecarbīl  
 (+ve inotropic)

Calcium sensitizer:

Pimobendan

Levosimendan (PDE-3 blocker)

Disease modifying drug /

Drug reducing mortality in CCF:

$\beta$ -Blocker (Carvedilol, Bisoprolol, Metoprolol)

ACEi

Angiotensin Receptor Blockers (ARBs)

Spironolactone

ISDN, + Hydralazine.



Isosorbide dinitrate

↳ Except these drugs, all other drugs control symptoms only in CCF.



## GIT

Drug useful for Acid peptic disease (APD):

$H_2$  Antihistamines:

Cimetidine - Least potent.

Ranitidine

Famotidine - Most potent

Roxatidine

Nizatidine

Loxalidine.

↳ Basal acid output & Nocturnal (more effective)  
So, give at Bed time.

↳ Renal excretion.

Cimetidine - Antiandrogenic

CYP enzyme inhibitor

Least potent.

PPI ( $H^+ - K^+$  ATPase inhibitors):

Short half  
life for less  
than 2hr

But acting  
for longer  
duration

Omeprazole (Metabolism by CYP2C19, CYP3A4)

Esomeprazole

Pantoprazole

Jansoprazole

Rabeprazole

→ Hit & Run drug

(Irreversible inhibition of Proton pump).

# Omeprazole not given c/ clopidogrel.

Rabeprazole

Pantoprazole

> No significant drug interaction  
(preferred c/ clopidogrel).

## Antacids:

Sodium Bicarbonate

Calcium Carbonate - shouldn't be taken w/ milk

↓  
bcz Milk alkali Syndrome.

## # GELUSIL:

Combination of Aluminium Hydroxide (Constipation)  
+ Magnesium Hydroxide (Diarrhoea)

## Ulcer protective drugs:

Sucralfate (Sucrose + Sulfated Aluminium hydroxide)

- Acts only in Acid medium (pH below 4)

- It shouldn't be combined w/ H<sub>2</sub> blocker/PPI/  
antacid.

## Bismuth

- Black stool & tongue.

CF - Renal failure.

## Ulcer healing drugs:

C. arbenclozone

↳ S/E - Displaces aldosterone from  
protein binding.

## Prokinetic drugs:

Drugs promoting GI motility.

## D<sub>2</sub> antagonist:

Domperidone

Metoclopramide

5HT<sub>4</sub> agonist:

Cisapride } - Cause QT prolongation  
 Mozapride } ∴ Withdraw  
 Tegaserod }  
 Ivosulpride }

Cholinergic agonist (M<sub>3</sub> agonist)

Beltranechol  
 Neostigmine.

5HT<sub>3</sub> blocker:

Ondansetron.

# Antibiotic having Prokinetic action: Macrolide.



acting on motilin receptor  
 of small intestine cause diarrhoea.

Among Macrolide - max<sup>m</sup> Prokinetic  
 Erythromycin

Drug used in Anti cancer / Radiation - drug induced vomiting

5HT<sub>3</sub> antagonists:

Ondansetron M/C 9/E - Headache.

Granisetron

Tropisetron

Dolasetron → QT prolongation

Palonosetron → Highly selective 5HT<sub>3</sub> antagonist  
 Long acting (T<sub>1/2</sub> = 40 hrs)

Supportive drug: For better efficacy

Ondansetron  
mixed c

→ D<sub>2</sub> blocker, BZD, Steroids



① Domperidone



Dexamethasone  
Methylprednisolone.

Antiemetic belonging to Cannabinoids

Nabilone

Dronabinol

} Antiemetic + Appetite stimulant.

2-3 days after chemotherapy → Late phase Vomiting



T/t ① Aprepitant (oral)

② Fos aprepitant (i.v.)



Neurokinin 1 antagonist

② Palonosetron.

IBS

T/t of Constipation dominant IBS:

Magnesium hydroxide

Methyl cellulose

Lactulose syrup. → Also useful for Hepatic encephalopathy.

Tegaserod } 5HT<sub>4</sub> antagonist

Prucalopride

(Lubiprostone

→ CLC-2 (Type-2 chloride channel activator)

↳ Linaclotide (Guanylate-cyclase-C activator)  
 ↳ Cystic Fibrosis transmembrane conductance regulator Activator  
 (CFTR activator)



Crofelemer - Inhibitor of CFTR  
 ↳ USE - HIV drug induced diarrhea.

Antibiotic used for t/t of constipation in IBS:

Neomycin (orally) → For t/t of Hepatic encephalopathy  
 Rifaximin → Pre-op Bowel Sterilization  
 Probiotics.

Rifaximin:

Useful for - ① IBS

② Hepatic encephalopathy

③ Traveller's diarrhea

④ Pseudomembranous colitis.

# For t/t of opioid induced constipation:

Methyl naltrexone (S/c)

Alvimopan (oral)

Diarrhea in IBS:

5HT<sub>3</sub> antagonist for t/t of diarrhoea in IBS:

Alosetron

Ramosetron

Cilansetron

Alosetron - Rarely cause dangerous problem  
 It cause Ischemic colitis

↳ So withdrawn

- But if use - give  $\bar{c}$  great caution & Informed consent.

- Only in female

Other drugs for diarrhoea:

Cholestyramine resin

Opioid for diarrhoea:

Loperamide

Diphenoxylate + Atropine  $\Rightarrow$  Control addiction.

Codeine.

For t/t Abdominal pain:

Anticholinergic drugs } muscle relaxant  
Imipramine } property.

Cholecystokinin antagonist:

Loxiglumide } Inhibits GI motility

Loxiglumide

↓

Useful for IBS (diarrhoea)

## BRONCHIAL ASTHMA.

Methyl Xanthines - Aminophylline  
Theophylline } Bronchodilator.

MAO - Adenosine antagonism - lead to seizure.  
Non-selective PDE inhibition

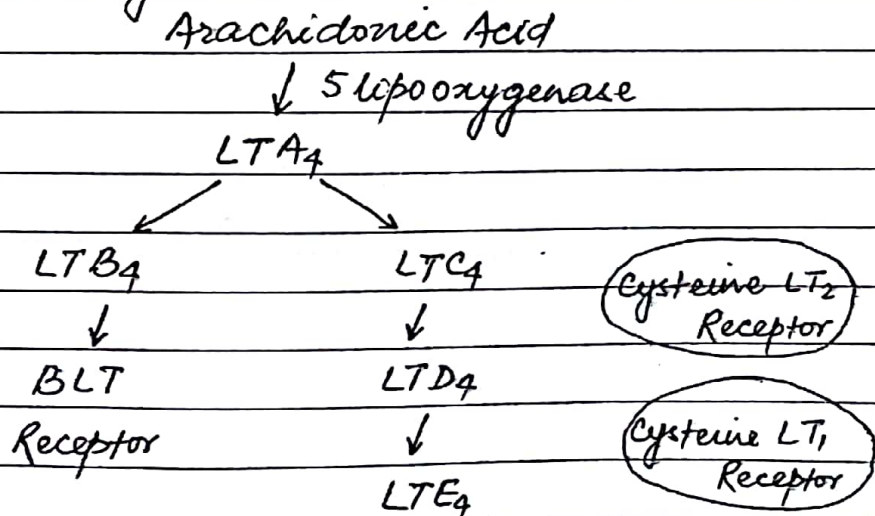
Side effect	Proposed mechanism
Nausea & Vomiting	} → PDE4 inhibition
Headaches	
Gastric discomfort	

Diuresis	} → A <sub>1</sub> receptor antagonism
Epileptic seizures	

Cardiac arrhythmias → PDE3 inhibition  
A<sub>1</sub> receptor antagonism.

# M3 Blocker } Bronchodilator  
β<sub>2</sub> agonist } M/c for acute Asthma

Leukotriene antagonists:



## Lipoxygenase Inhibitor

Zileuton

↳ Not used bcoz Hepatitis.

## Leukotriene antagonist:

Zafirlukast

Montelukast

Pranlukast

## Chronic therapy cause - Churg Strauss Syndrome



Headache

Eosinophilia

Vasculitis.



For t/t: MEPOLIZUMAB

(IL-5 antagonist)

## Mast cell stabilizers:

Sodium chromoglycate<sup>-</sup>

Nedocromil

Ketotifen (Additional Antihistaminic property)

## Monoclonal antibodies:

Omalizumab → IgE antibody agonist.

↳ s/c, Hypersensitivity.

## Newer drug - Reslizumab

Mepolizumab (IL5 antagonist)



## PDE inhibitors:

PDE inhibitors		
Methyl xanthines	PDE I, II, III, IV	Asthma
Cilomilast, Roflumilast	PDE IV	Asthma
Aprenilast	PDE IV	Active Psoriatic arthritis
Amirinone, Milrinone	PDE III	CCF
Sildenafil, Vardenafil Tadalafil ]	PDE V <del>Non-selective</del>	Erectile dysfunction
Pentoxifylline	Non-selective	PVD
Cilastazol	PDE III	PVD
Vinpocetine	PDE1, Vasodilator	Parkinson, Alzheimer's ds.

## EXPECTORANTS

## Mucolytics:

Carbocysteine  
 Methyl cysteine  
 Erdosteine  
 Bromohexane  
 Dorsane alpha  
 N-acetyl cysteine.

## Cough suppressant:

Codeine  
 Pholcodine  
 Dextromethorphan.

# Antihistamines

## 1st Generation

## 2nd Generation

→ Antihistaminic  
+ Anticholinergic action

USE: Allergic cond<sup>n</sup>  
Insect bite  
EPS  
Motion sickness

### 1st Generation drugs:

CPM (Chlorpheniramine Maleate)

Promethazine (Most sedative, Highest anticholinergic)

Diphenhydramine

Cyclizine

Mecizine (Useful for Sea sickness)

Cyproheptadine (Antihistaminic + Anticholinergic + Antiserotonergic action)

↓  
Appetizer, Useful in migraine

Cause Serotonin Syndrome.

Hydroxyzine (Antihistaminic + Anti-anxiety)  
↳ produces metabolite - Cetrizine.

Doxepin → Given topically (for itching)

↳ TCA - Atopic dermatitis, Lichen simplex

Cinnarizine (H<sub>1</sub> + M + 5HT<sub>2</sub>)

↳ Use in Vertigo

↑  
Beta-histamine (Histaminergic drug)

## 2nd Generation drugs:

Terfenadine } - Causes QT prolongation  
 Astemizole } Withdrawn  
 Metabolite Ebastine } → Still available

Fexofenadrine

Cetirizine (Metabolite of Hydroxyzine)

Levocetirizine

Azelastine (Maximum topical, nasal spray)

Mezolastine

Acrivastin

Active form. } Loratidine (longest)  
 } Desloratidine

Rupatidine (Platelet activating factor antagonist)

↓

Lexipafant } For t/t of Acute  
 Apafant } Pancreatitis

## Topical antihistamines:

Azelastine - Nasal spray

Olopatadine - Nasal spray,

↓

Ophthalmic drop.

Mast cell stabilizing Oral

Alocastadine, Epinastine - Eye drop.

H<sub>3</sub> antagonist / inverse agonist:

Pitolisant (Tirpitolisant) → Orphan drug.

↳ T/t of Narcolepsy

## Prostaglandins

PG E<sub>1</sub>:

Misoprostol:

- Useful for T/t gastric ulcer (NSAID induced)
- Used for abortion
- Teratogenicity → Mobeious Syndrome

Alprostadil

- Vasodilator
- Useful for Erectile dysfunction (Given injectable)
- Useful for maintain patency of ductus arteriosus.

PG E<sub>2</sub>:

Dinoprostone

↳ Uterine contracting agent  
Useful for abortion.

Enprostil  
Rioprostil } - Useful for t/t of Gastric ulcer.

PG F<sub>2α</sub>:

Carboprost

↳ USE: Post partum Hemorrhage (PPH)

Dinoprost

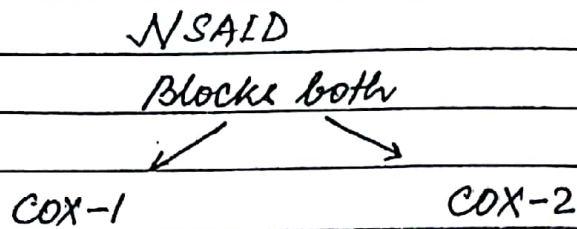
↳ USE: Uterine contracting agent for abortion.

cause  
Iris pigment  
ation } Latenoprost  
Bimatoprost  
Travoprost  
Causes  
Hypertichoses  
of eyelash } Unoprostone

- Useful for Glaucoma

↓  
By promoting drainage  
via Uveoscleral route.





Aspirin:

Analgesic  
 Anti pyretic action  
 Anti inflammatory } - All are property of all NSAID.  
 Prevent Colonic & rectal cancer

Aspirin + Nicotinic acid  $\Rightarrow$  Prevent flushing.

C/I - in t/t viral fever in children < 12yrs.

↓  
 Cause Reye's syndrome.

- Liver damage
- Encephalopathy
- Febrile illness

M/C S/E of Aspirin & other NSAID:  
 - Gastric ulcer.

Non-selective COX inhibitor

Indomethacin - Anti-inflammatory

Use: [ Frontal headache  
 Closure of ductus arteriosus  
 Batten's syndrome

Phenylbutazone

- may cause bone marrow ~~depression~~ suppression.

Ibuprofen - safe in children

Mefenamic acid - Useful in dysmenorrhoea.

Piroxicam - longest acting NSAID.

Preferable COX2 inhibitor:

- Nimmeselide
  - ↳ Cause <sup>severe</sup> hepatotoxicity in children (Unsafe)
- Nabumetone
- Etodolac
- Meloxicam

Highly selective COX-2 inhibitor:

Rofecoxib

Celecoxib

Valedecoxib

Etoricoxib

Parecoxib

Lumiracoxib.

Risk of developing HTN & CCF

COX-3 blocker

Paracetamol

Overdose ↳ Causes liver toxicity.

Other analgesic: Other than NSAID & opioids.

Ziconotide (Conotoxin)

- N type CCB

- Intrathecal given

For anti-inflammatory action of Aspirin  $\rightarrow$  300-400 mg  
aspirin required  $\hat{=}$  cause  $\uparrow$  uric acid.  
&  $>$  2 gm  $\rightarrow$  Gastric perforation.

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Nefopam - Amine uptake inhibitor  
 $\text{Na}^+$  channel blocker

Sativex - Cannabinoid  
 $\hookrightarrow$  USE - Cancer pain

~~Entonox~~ Entonox -  $\text{N}_2\text{O} + \text{O}_2$   
 $\hookrightarrow$  For painless labour.

Drug useful for t/t of Gout:

Acute Gout:

Give NSAID or, Steroids or, colchicine

Colchicine  $\rightarrow$  Acting by disruption of microtubule



Neutrophil drunken walk.

SE - Diarrhoea (Bloody)

Unsafe in RF

NSAIDs  $\rightarrow$  Naproxen  
Indomethacin  
Sulindac

# Aspirin is CI for gouty arthritis.

Drug used for chronic gout:

Xanthine oxidase inhibitor:

Allopurinol

Febuxostat.

6-Mercaptopurine.



Uricosurics:

- Probenacid (Unsafe in RF)
- Sulfipyrazone
- Benzbromarone
- Lesinurad.

Other drug having uricosuric actions are -

- Losartan
- Fenofibrate
- Amlodipine

Newer drug :-

For aggressive control of Gouty arthritis

- ↳ Give intravenously
- Rasburicase } cause Rapid metabolism
- Pegloticase } of uric acid.

Newer drug for T/t of RA:

Normal - Cytokine balance

Pro-inflammatory  
Cytokines

Anti-inflammatory  
cytokines.

TNF $\alpha$ , IL-1, IL-6

TNF $\alpha$  blocker:

Test

Immuno  
suppressant

- Infliximab (i.v)
- Etanercept (s/c)
- Adalimumab (s/c)
- Golimumab (s/c)
- Certolizumab (s/c)

Before giving TNF $\alpha$  blocker  
TB should be ruled out.  
- PPD test  
↳ Purified Protein derivative  
skin test

- All are unsafe in Hepatitis B virus infected pt.

Analogue of Interleukin 1 (IL-1) Receptor Antagonist:  
ANAKINRA

IL-6 blocker:

Tocilizumab  
Sarilumab

Newer drug - Rituximab (CD20 receptor antagonist)  
↳ Cause PML (Progressive Multifocal Leucoencephalopathy).

Targeting against  
Abatacept } CD80/86 Receptor  
Balatacept }  
↳ USE - RA

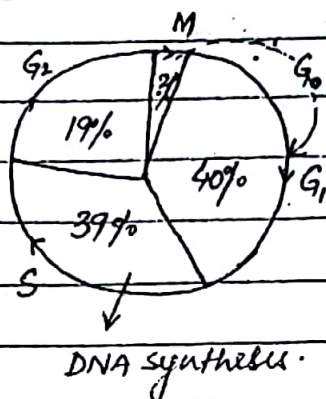
Tofacitinib - JAK 1 & 3 blocker  
↳ USE - RA.

Leflunomide

↳ Inhibit dihydro orotate dehydrogenase  
S/E - Hepatotoxic  
C/I - Pregnancy.

## ANTI CANCER DRUGS

Cell cycle:



G<sub>1</sub> (40%) → Minor development take place.

S-phase → DNA synthesis

(39%) By ← Topoisomerase II enzyme  
Folic acid, Purine, Pyrimidine.

G<sub>2</sub> (19%) → Extra development take place.

By Topoisomerase

M (2%) → Multiplication

Drugs acting on G<sub>1</sub> phase:

L-Asparaginase (enzyme)

Steroids

L-Asparaginase - Origin from E. coli (Naturally occurring)

- Useful for ALL

S/E - Hemorrhagic pancreatitis

Hypercoagulation

No significant Myelosuppression.

Thromboembolic complications.

Drugs acting on S-phase:

Anti metabolites

Epididymal phyllo toxins

eg: ETOPOSIDE

TENEPOSIDE

Drugs acting on G<sub>2</sub> phase: Topoisomerase-1 inhibitors.

Camptothecins ← IRINOTECAN - cholinergic property.  
TOPOTECAN

SE - Diarrhoea.

(dose related toxicity)

Bleomycin (Anticancer + Antibiotic)

- All anticancer + antibiotics are C-cycle non-specific  
except Bleomycin.

Drug inhibiting mitosis:

Vinca alkaloids - Vinblastin } plant origin.  
Vincristine }  
Vinorelbine }

Taxanes - Paclitaxel

Docetaxel

Cabazitaxel.

Newer drug - Ixabepilone } Useful for Breast Ca.  
Eribulin }

For HER2 +ve Breast Ca - TRASTUZUMAB

For Rx HER1 & HER2 - TK Blocker - LAPATINIB.

Newer drugs in Cancer therapy:

Tyrosine Kinase inhibitor (TKi's):

Tyrosine Kinase Receptor - EGFR (HER-1)  
 VEGFR  
 PDGFR

TKi's acting EGFR blocker:

Gefitinib } - Useful for t/t of Metastatic Small  
 Erlotinib } cell lung Ca.  
 Afatinib } → Also useful for Pancreatic Ca.

↓  
 DOC: Gemcitabine

S/E - Dysmorphic eyelashes (Erlotinib)

VEGFR blocker:

Sorafenib - Useful for RCC, HCC

Sunitinib - Useful for RCC, GIST

Lenvatinib - Useful for DTC

PDGFR blocker

Imatinib - DOC for CML

↑  
 1st gen. TKi

Useful for GIST (c-kit)

↓  
 due to alteration of c-kit - Resistance

↓ T/E of Resistance CML

DASATINIB } 2nd gen. TKi  
 NILOTINIB }

Multi-targeted TKi:

Vandetanib - Useful for Medullary Ca Thyroid.

↳ Target against EGFR & VEGFR.

Axitinib } Targeting against VEGFR & PDGFR

Pazopanib } Useful for RCC

# TRASTUZUMAB → For HER-2 +ve Breast Ca.

# LAPATINIB → Against HER-1 & 2 +ve Breast Ca.

# All the TKI are taken orally.

Common S/E - GI toxicity

(Nausea, Vomiting, Diarrhoea)

Any drug block EGFR causes HTN.

Monoclonal antibodies (MABs)

TRAS(TU)ZU(MAB) ↗

↓ ↓

Target Source

TU = Tumor Zu - Humanised

Li = lowering immunity Xi - Chimerical (Non human eg Mice)

Ci = Target circulation.

Vi = Virus.

BASILIXIMAB - Target against IL-2

ABCIXIMAB - Target against GP2B3A.

PALLIVIZUMAB - Target against RSV.

Trastuzumab -

Target against HER-2 receptor

Useful for HER-2 +ve Breast Ca.

# Most of MAB given by i.v. infusion

Specific S/E → Cardiomyopathy

Infusion reaction.

## Rituximab:

Target against CD20 on B-cell.

Useful for B-cell lymphoma

Other uses: C = CLL

H = Hemolytic anemia

I = Idiopathic thrombocytic Purpura (ITP)

N = NHL (Non-hodgkin Lymphoma)

A = Arthritis (RA)

Myasthenia Gravis.

M/C S/E - PML

## Bevacizumab: Target circulation.

Target against VEGFR

Useful for Metastatic colorectal CA (iv)

↓  
M/C → 5FU

Useful for RCC & Diabetic Retinopathy.

↓  
i.v.

↓  
Intra-vitrous

S/E - HTN

## Newer drug: RAMUCIRUMAB

- Target against VEGFR

- Useful for Gastric Cancer.

## BRENTUXIMAB

- Target against CD30 on B cell.

- Useful for Hodgkin Lymphoma.

Omalizumab - Target against IgE → USE: Bronchial Asthma (BA)

Reslizumab } - Target against ILS → USE: BA  
Mepolizumab }

Denosumab - Target against RANK-L → Osteoporosis.

Eculizumab - Target against C5 → Paroxysmal nocturnal hemoglobinuria.

Evolocumab } - Target against PCSK9 → Lipid lowering.  
Alirocumab }

Ibalizumab - Target against HIV (entry inhibitor)

### Macular degeneration (MD)

Dry type  
less blood supply

Wet type  
Age related MD (ARMD)

Drugs useful for Wet type MD:

Photodynamic therapy

VERTEPORFIN - i.v.

VEGF inhibitor:

Bevacizumab } - Intravitreal inj.  
Ranibizumab }  
pegaptanib }  
Aflibercept }



Drug for Vitreomacular degeneration:  
Ocriplasmin (Newer drug).

# Bull's eye Retinopathy - Caused by chloroquine.  
Crystalline Maculopathy - Caused by Tamoxifen.  
Field of Vision defect - Vigabatrin.  
Whorl-like pattern - Already done.

Kayser-Fleischer ring - Wilson's ds (Ceruloplasmin deficiency).

### Chelating Agents.

Metal	T/t
Copper	Penicillamine (SLE, optic Neuritis) Trientine Zinc sulphate (Safest) Potassium Sulfide
Hepatitis or cirrhosis c decompensation	Zinc
Mild - Moderate hepatic decompensation	Trientine + Zn
Neurological or Psychiatric Symptom	Tetrathiomolybdate + Zn.
For maintenance in pregnancy & children	Zinc

Metal	T/t	
Lead	BAL	C/I in Iron & Cadmium poisoning.
Arsenic	BAL	
Mercury	BAL	
Iron		Desferrioxamine Deferiprone Dexrazoxane.

# DOXORUBICIN

S/E - Cardiomyopathy  
 Antidote for Doxorubicin poisoning - Dexrazoxane.

Anti-metabolites:

Anti cancer + Immunosuppressive.

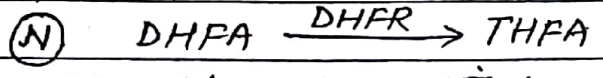
Drug acting against folic acid:

Methotrexate

Pemetrexate } - Useful for Mesothelioma  
 Trimetrexate } NSCLC

Pralatrexate - For T-cell lymphoma.

Methotrexate:



MAO: Methotrexate actively penetrate into cancer cell it inhibit DHFR, ultimately inhibiting DNA synthesis, So stop S-phase of cell cycle.

# Resistance due to alteration/mutation of DHFR.

Page No.

Date: / /

Specific antidote — Folinic acid or Leucovorin antagonist.

# Folinic acid can't be given in Renal failure.

GLUCARPIDASE — Newer drug useful for t/t of Methotrexate toxicity in a pt w/ impaired kidney func<sup>n</sup>.

USES of MTx : Anticancer :

DOC for Choriocarcinoma.

Useful for Osteosarcoma

Immunosuppressant :

RA (DMARD, low dose 7.5mg/wk)

Psoriasis

↓  
long term therapy.

C = Chorio CA

A = Abortion

N = NHL

C = Chron's ds

E = Ectopic pregnancy

R = RA.

S/E — Myelosuppression (M/C)

Alopecia

Mucosal damage. (GI toxicity)

Liver damage (on chronic therapy — In RA)

↳ Undergo LFT

Crystalluria

↳ TI — uric acid & alkalization

### Antibiotic causing Crystal

Ciprofloxacin (Alkaline)

Sulfonamide (Acidic)

Antiviral  
Causing Crystal

Indinavir → HIV  
Acyclovir

### CI of MTx - Pregnancy.

### Purine Anti metabolites:

6-Thioguanine

6-Mercaptopurine

Fludarabine } DOC: CLL  
- Useful for Hairy cell leukemia

also useful for ← Cladribine } DOC - Hairy cell leukemia

Multiple Sclerosis. Pentostatin

↳ Inhibiting Adenosine deaminase.

### 6-Mercaptopurine:

6-Mercaptopurine

↓ HGPRT enzyme.

6-Thiosinic Acid

Cause of Resistance - Deficiency of HGPRT enzyme  
(Lesch-Nyhan Syndrome)

6-MP normally undergoes inactivation (metabolism)  
by HGPRT.

If we give Xanthine oxidase inhibitor - ↑ plasma level  
of 6MP.

When we give Allopurinol ↓ 6MP  
reduce the dose 50-75% of 6MP.

INF $\alpha$  - USE: HBV, HCV  
INF $\gamma$  - USE: Ch. granulomatous ds.  
↳ Immuno stimulant.

Page No. 189  
Date: / /

Drugs useful for Multiple Sclerosis (MS):

Disease modifying drugs:

• Interferon Beta 1A & 1B

Glatiramer Acetate

Natalizumab (d4 $\beta$ 1 integrin) (iv. once in <sup>month</sup>)

Acerlizumab (anti CD20) ↓  
cause PML

Alemtuzumab (Anti CD52)

Mitoxantrone (Anti cancer + Antibiotic)

↳ Cause Cardio-toxicity.

Fingolimod (oral)

↳ Cause Bradycardia.

Dalfampridine (oral)

↳ Useful for Lambert Eaton Syndrome.

↳ Useful in MS in improving walking.

Cladribine (oral)

Teriflunomide (oral)

↳ derivative of Leflunomide

↳ Di-hydro erotate ....

↳ Useful in pregnancy & MS

Dimethyl fumarate.

<sup>meta</sup>  
Pyrimidine Antimetabolites:

Cytarabine (Cytosine arabinoside)

↳ Cause Cerebellar ataxia.

5FU

↳ M/c use - Colorectal Cancer

• Gwien  $\bar{c}$  Levamisole

Floxuridine

• Gemcitabine (DOC for Pancreatic CA)

Capacitabine. (Cause Hand foot Syndrome)



Mitomycin:

- Useful for Urinary bladder CA.



Usually Intravesical therapy: BCG  
For BCG resistance - Mitomycin  
Valrubicin

- Useful for laryngo tracheal stenosis.  
due to Antifibroblastic action.

Bleomycin:

Cell cycle specific acting on G<sub>2</sub> phase of Cell cycle.  
M/C S/E - Pulm. fibrosis.

Bleomycin hydrolase is not seen in lung.



so large accumulation of Bleomycin in lung.

Type I pneumocytes - Necrosis / destruction  
Type II " - Hyperplasia / Metaplasia.

# Anticancer drug  $\bar{c}$  No <sup>severe</sup> myelosuppression:

Vincristine → Cause Peripheral neuropathy.

Bleomycin

Asparaginase  $\xrightarrow{\text{cause}}$  Pancreatitis  
Hypercoagulation

## Alkylating agents.

Busulfan

- highly lipid soluble  
↓  
Useful for Brain Tumor
- Nitrosoureas → Lomustine
  - Semustine
  - Carbustine } Delayed Myelocuppressant.
  - Temozolamide → also for Malignant Melanoma.
  - Streptozocin (Chemical Pancreatectomy).
  - Chlorambucil (USE: CLL)
  - Cyclophosphamide, Ifosfamide
  - Melphalan (Use for Multiple myeloma)
  - Procarbazine, Dacarbazine.
  - Thiofepa
  - Mechlorethamine.
    - ↳ Cause skin Vesicant

### Procarbazine -

- Disulfiram like reac<sup>n</sup>
- Among the alkylating agent Procarbazine & Melphalan cause Secondary cancer.  
Cyclophosphamide - less Secondary cancer.
- MAO inhibitory action

### Drugs for Multiple myeloma:

Melphalan

Ihalidomide

Lenolidomide

Bortezomid (Proteasome inhibitor)

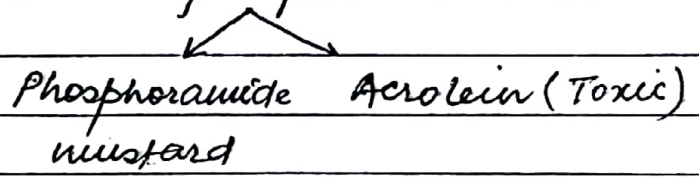
↳ DOC

- Punch out lesion.



Cyclophosphamide (Anticancer + Immunosuppressive):  
- Prodrug.

In liver it forms Aldophosphamide



DOC for Wegner's granulomatosis.

M/c s/e - Hemorrhagic cystitis

↳ Due to Acrolein

Antidote - MESNA

Supportive drug - Formalin

- N acetyl cysteine
  - Carboprost (PGF2a agonist)
- USE:

- ↳ Paracetamol poisoning
- Radiocontrast
- Nephrotoxicity
- Mucolytic

Cyclophosphamide cause { SIADH  
Cardio-toxicity.

Ifosfamide:

Active form - Acrolein

↓ Antidote  
MESNA

# Drug of choice in <sup>malignant</sup> Melanoma - LEVODOPA

Drugs for Multiple myeloma:

- Temozolamide
- BRAF V600E inhibitor - Vemurafenib  
Dabrafenib  
Trametinib

Newer drug. — Nivolumab  
Ipilimumab

Aldesleukin - IL2

↳ USE: RCC, Multiple Myeloma.

Busulfan:

Used for CML

S/E - Pulm. fibrosis

Adrenal insufficiency. (Addison's ds)

↳ Hyperpigmentation.

- # All alkylating agent action - N7 Guanine Residue
- # All " " " are cell cycle non specific.

S/E of Alkylating agent - Venoocclusive ds of Liver.

( Budd Chiari Syndrome



Minimised by DEFIBROTIDE

- Permanent sterility

Least emetogenic - Vincristin  
Chlorambucil

Page No. 195

Date: / /

Cisplatin:

Highest emetogenic

SE - Ototoxicity

Nephrotoxicity (dose limiting toxicity)

Neurotoxicity

Antidote - Amifostine.

Carboplatin:

SE - Myelosuppression

Oxaliplatin:

SE - Neurotoxicity

Pharyngeal paraesthesia.

Vincristine:

SE - Peripheral neuropathy (Sensory & motor).

SIADH

Vesicant.

Advantage - Less myelosuppressant  
less nausea.

Vinblastine:

- Myelosuppression

Taxane (Paclitaxel, Docetaxel):

- Myelosuppression

- Peripheral neuropathy (Glove & sock neuropathy)

- Allergy.

Role of hormones in Cancer:

For all premenopausal women  $\bar{c}$  ER +ve Breast cancer  
1st line choice is SERM.

If Resistance give SERD.

# For postmenopausal women  $\bar{c}$  ER +ve Breast cancer  
give Aromatase inhibitor.

#

SERM useful for t/t of Breast Ca:

Tamoxifen

Toremifen

Doloxifen

Raloxifene.

Tamoxifen -

Antagonistic action only on ER of Breast  $\rightarrow$   
Useful for t/t ER +ve Breast Ca.

Agonistic action on blood vessel

ADR - Hot flushes

Endometrial cancer

DVT

Raloxifene:

Antagonistic action on Breast → So use in Breast CA.  
 " " " Uterus

S/E - Flushing

DVT

Not cause Endometrial CA.

Aromatase Inhibitors:

Amino glutethimide (chemical adrenalectomy)

Formestane

Exemestane

Vorzole

Fadrozole

Letrozole

Anastrozole.

Extra information:

SERMs for DUB: Ormeloxifene  
 (Centchromin)

- Use as Contraceptive pill.

Twice in wk c̄ gap of four day - First  
 3 month. later once in a week.

SERMs for Dyspareunia - Ospemifene.

SERMs for induction of Ovulation - Clomiphene.

SPRM:

Ulipristal - Emergency Contraceptive (Can take 5 days after coitus)

Asoprisuol

Leuprolidone - Useful in Uterine fibroid  
Endometriosis

Prostatic Cancer:

Beoz of excess androgenic action.

Hypothalamus

(GnRH) - Pulsatile release

↓ ⊕ (60-120 min)

Pituitary

(Gonadotropins - LH/FSH)

↓ ⊕

Testis

FSH → Spermatogenesis ← Seminiferous cell.

LH → Leydig cell - Testosterone production

↓

overproduction cause Prostatic Ca.

Drugs ↓ Testosterone production:

(A) GnRH agonist (In continuous manner):

Leuprolide

Goserelin

Buserelin

Nafarelin

Desorelin

Histrelin

Triptorelin

GnRH antagonist:

Genirelix

Cetrorelix

Abarelix

Degarelix

Comparison:

Agonist  
Initial flare up  
Histamine release

Antagonist.  
No initial flare up.  
No histamine release.

↓ Testosterone cause:

Hot flush

Loss of libido

Impotence

Sarcopenia (Reduce muscle mass)

Osteoporosis

t/t → Supplement Vit D

Bisphosphonates.

Denosumab.

Drugs having histamine releasing property:

d-Tubocurarine

Morphine

Dexferioxamine

Amphotericin B

Polymyxin B

Vancomycin (Red Man Syndrome)

## Anti androgen/

Flutamide

Nilutamide

Bicalutamide

Enzalutamide

Cyproterone

Abiraterone.

## Thalidomide:

Sedative + Anti emetic

S/E - Phocomelia

CI - Pregnancy.

Category X.

- It has Anti cancer + Immune modulation property.

Indication: Multiple myeloma

ENL

Apthous ulcer

SLE.

Isomer  $\left\{ \begin{array}{l} R \text{ (Therapeutic use \& Teratogenicity)} \\ S \text{ (Sedation)} \end{array} \right.$ 

M/c S/E - Constipation

Severe peripheral sensory neuropathy.



Drug	Antidote
Methotrexate	Folinic acid
Doxorubicin	Dexamethasone
Cyclophosphamide	Mesna
Cisplatin	Amifostine
<del>Ataxia</del> Palifermin	Mucositis

Drugs useful for Ht neutropenia:

Colony stimulating Factor (CSF)

γG-CSF	GM-CSF
Filgrastim	Sargramostim
Pegfilgrastim	Molgramostim
Lenograstim	

Drug useful for Anemia:

Epoetin (Recombinant - Erythropoietin)

Darbopoietin

Peginesatide (Erythropoietin Receptor Stimulant)

Drug useful for Thrombocytopenia:

- Oprelvekin (IL-11)

- Thrombopoietin

Newer drug [ Romiplostim (TPO) for ITP → by plasma exchange.  
Eltrombopag ]  
↳ Oral

Anti-emetic useful for Anti cancer t/t:  
Already done.

Immuno suppressant:

Cyclosporin

Tacrolimus (FK506)

Sirolimus

Everolimus

Drugs inhibiting synthesis of IL-2:

Cyclosporin

Tacrolimus (FK506) > Calcineurin  
inhibitor.

↓

Both cause Nephrotoxicity

Tacrolimus > Cyclosporin

Tacrolimus - Macrolide comp<sup>d</sup>.

Common problem - Nephrotoxicity (Dose limiting).

Neurotoxicity

Hepatotoxicity

DM

Diarrhea

Alopecia

Specific SE of Cyclosporin - Hypertrophy of Gum

Hirsutism

HTN → T/t: Nifedipine.

Hyperkalemia

Hypokalemia  $\xrightarrow{\text{caused by}}$  Cisplatin  
Amphotericin B.

m-Tor blockers:

Siroliimus } -S/E - Thrombocytopenia  
Everolimus } Hyperlipidemia  
(High TGL)

Azathioprine:

Purine antimetabolite

Immunosuppressant action (CME)

No anti cancer action.

USE - RA

IBD (U. colitis)

Organ transplantation.

S/E - Myelosuppression

Azathioprine  $\xrightarrow[\text{in body}]{\text{converted}}$  6-Mercaptopurine.

↓  
Metabolism by Xanthine Oxidase.

Immunostimulants:

Cytokines

Aldesleukin (Recombinant IL2) (For RCC & MM)

Interferon  $\gamma$  (Chronic granulomatous dis).

BCG vaccine (Intra vesicle - Urinary bladder Ca)

↓  
Valrubicin, Mitomycin  
Laryngotracheal Stenosis

Levamisole (Anti helminthic property)

↳ Immuno stimulant.

IL- modulators:

Analogues of IL-1 receptor antagonist: Anakinra  
(USE - RA)

IL-3 & 4 antagonist: Pritakinra  
(USE - BA)

Analogue of IL-2: Aldeslakin  
(USE - RCC, Malignant Melanoma)

IL-2 receptor blocker: Basiliximab  
Dacizumab.

IL2 + Diphtheria toxin: Denilukin diftitor

↓

USE: Cutaneous T cell lymphoma.

↓

Histone deacetylase inhibitor  
Vorinostat

Romidepsin.

IL-5 blocker: Reslizumab (Severe eosinophilia, BA)

Mepolizumab

↳ Hypereosinophilic syndrome

Churg Strauss syndrome.

IL-6 blocker - Tocilizumab

↳ USE - RA

IL-1,6 antagonist - Steroids

Analogue for IL-11 - Oprelvekin

↳ USE - Thrombocytopenia.

IL-17 Blocker - Ixekizumab } Use: Plaque Psoriasis.  
Brodalumab

## IL 12 &amp; 23 - Ustekinumab

↳ USE - Psoriasis.

# Apafant, Lexipafant, (PAF Blocker) - For Acute Pancreatitis

Ivacaftor - For cystic fibrosis.

Imiquimod - For Chondylomata acuminata (HPV)

Alectacept - For Psoriasis

Resiquimod - For HSV

Lu-Dotatate - For Midgut endocrine tumor.

Anagrelide - For Essential Thrombocytosis

Belimumab - For SLE

Defibrotide - For Budd Chiari Syndrome.

Hydroxyurea - For Sickle cell anemia.

Olaparib - For ovarian Cancer

• Acting by Poly ADP ribose polymerase (PARP) inhibitor.

Palbociclib, Amebaciclib, Ribociclib - For Breast Cancer

↳ CDK 4/6 (cyclin dependent kinase) inhibitor

Edaravone - (Antioxidant) for ALS.

Mycophenolate mofetilale - Inhibit Inosine monophosphate  
(Immunosuppressant) dehydrogenase

Pentostatin - Inhibit Adenosine deaminase.

Vorinostat - Inhibit Histone deacetylase.

Leflunomide - Inhibit dihydro orotate dehydrogenase

Toxicity caused

Cyclosporine - Nephrotoxicity

Leflunomide - Hepatotoxicity

Sirolimus - Bone marrow suppression

Azathioprine - Hypertriglyceridemia

Muromonab - Cytokine release syndrome.

Page No. 206

Date : / /

## ANTIMICROBIAL DRUGS

Antibiotic acting by inhibiting cell wall synthesis:

N acetyl muramic acid }  
N acetyl glucosamine } N-acetyl muramic  
Acid peptidase.

Step 1:

# The first enzyme initiating cell wall synthesis  
- Alanine ligase/Racemase

↑ ⊖

Cycloserine

↳ 2nd line drug of TB

- Bacteriostatic

S/E - Psychosis.

Step 2:

Enolpyruvate transferase ← ⊖ Fosfomycin

↳ For UTI

Cause severe diarrhoea

So not in use.

Step 3:

Dephosphorylation of Bactoprenol ← ⊖ Bacitracin

↓

polypeptide group of Antibiotic

USE: Wound/ulcer healing  
(Given topically)

Step 4:

Elongation of peptide chain

↳ c help of Transglycosylase ← ⊖ Vancomycin

↓  
if Alter → VRSA

Step 5:

Cross linking of elongated peptide chain

↓  
by Transpeptidase  $\leftarrow$  Beta Lactam  
(Penicillin binding protein) (Penicillin)

↓  
If altered  $\rightarrow$  MRSA  
(Resistance)

Antibiotics acting by <sup>inhibiting</sup> protein synthesis:

Aminoglycosides & Tetracycline binding to 30s Ribosome & inhibit protein synthesis.

Drug acting on 50s Ribosome & inhibit protein synthesis:

Chloramphenicol  $\xrightarrow[\text{enzyme degradation}]{\text{Resistance due to}}$  Acetyl transferase  
Linezolid

- (M) = Macrolides
- (L) = Lincosamides (Clindamycin)
- (S) = Streptogramins.

MLS resistance  $\rightarrow$  Methylation of 50s ribosomes.

Tetracycline resistance  $\rightarrow$  Development of Efflux pump.

↓  
Tigecycline - Resistance to efflux.

Due to enzymatic degradation  $\rightarrow$  Aminoglycosides Resistance

↓  
Do not develop resistance. [ Amikacin  
Netilmicin



# All antibiotics acting by inhibiting protein synthesis are bacteriostatic exception - Aminoglycoside  
Streptogramins.

### Antibiotics

Penicillin:

Commercial source - *Penicillium chrysogenum*.

Acid Resistant: Orally.

V = Penicillin V

O = Oxacillin

D = Dicloxacillin

K = Cloxacillin

A = Ampicillin / Amoxicillin

Penicillinase resistant:

C = Cloxacillin

O = Oxacillin (hepatitis)

N = Nafcillin (Neutropenia)

D = Dicloxacillin

U

M = Methicillin (Interstitial nephritis)

$\beta$ -Lactamase inhibitor:

Clavulanic Acid + Amoxicillin

Sulbactam + Ampicillin

Tazobactam + Piperacillin

# FDC (Fixed drug combination):

Same volume of distribution

or same half life

## Extended spectrum Penicillins:

Aminopenicillins → Enteroactive

Becampicillin

Ampicillin → Causing diarrhea due to incomplete absorption.

Amoxicillin

Carboxy penicillins (Enteroactive + pseudomonas)

Carbenicillin → Cause bleeding due to disturbing platelet.

Ticarcillin

Ureidopenicillins

(Enteroactive + pseudomonas + Klebsiella)

Azlocillin

Piperacillin

Mezlocillin

# Aminopenicillins are C/I in Infectious mononucleosis  
bcz of risk of severe skin rash.

# 2nd line Anti TB C/I in HIV pt c TB: Thiacetazone

↓  
may cause Steven Johnson Syndrome

↓  
Skin Rash.

# OCP + Ampicillin → Risk of OCP failure

↓  
By interfering <sup>OCP</sup> enterohepatic circulation.

S/E of Penicillin in syphilis pt.

↓  
Jarisch herxheimer Reac<sup>n</sup>.

↓  
Secondary Syphilis

No treatment

Only symptomatic - Aspirin & Sedation.

Atypical beta lactam antibiotics:

Carbapenams:

- Imipenam

- Broadest spectrum

- Shortest acting

↳ Rapidly undergo inactivation by  
Dehydropeptidase I enzyme.

↑ ⊖

Add Cilastatin

S/E - Seizures

- Meropenam

- Ertapenam

Monobactams:

- Aztreonam

↳ No cross reactivity.

↳ Useful for Aerobic gram +ve infection.

Similar to aminoglycosides.

# For Anaerobic infection - Metronidazole

Clindamycin

↳ S/E - Pseudomembranous  
Colitis.

## Cephalosporins.

Fourth generation drugs:

Cefepime

Cefpirime

Cefclidin

Fifth generation drugs:

Ceftazidime

Ceftaroline

USE - MRSA

Community Acquired Pneumonia.

Glycopeptide Antibiotics: Vancomycin

t/t of Gm +ve infection.

Oral Vancomycin - Useful for Pseudomembranous colitis

i.v. Vancomycin - DOC for MRSA.

caused by Clostridium difficile.

Caused by 3rd gen. Cephalosporin.

Newer drug for PMC - Rifaximin

Fidaxomicin

ADR of Vancomycin: Red Man Syndrome (M/c)

Ototoxicity

Nephrotoxicity

Other Glycopeptide antibiotics:

Ticoplanin

Oritavancin

Telavancin

Dalbavancin - longest acting (6-10 days)

Drugs used for T/t MRSA/VRSA:

VRSA → Linezolid -

SE - Thrombocytopenia (M/C)

optic & peripheral neuropathy.

Also used for MDR TB.

MAO inhibitory property.

VRSA → Streptogramin

Quinupristine: Dalfopristine = 70:30.

SE - Infusion reaction

Arthralgia.

VRSA → Daptomycin

↳ causing myopathy.

VRSA → Tigecycline

given i.v. tetracycline.

Resistant to efflux

Excretion - Bile

Safe in Renal failure.

## Sulfonamides:

Sulfasalazine

In GIT split in 2 component

Sulfapyridine

5 amino salicylic acid.

Useful for RA.

Useful for ulcerative colitis

ADR - Allergy

Oligospermia (In male) → Infertility.

Topical - Sulfacetamide - For eye drop.

Silver sulfadiazine } - has anti-pseudomonal action

Mefanide

↳ CA inhibitory action

↓  
Metabolic acidosis.

useful for Fungal Keratomycosis.

# Sulfadoxine + Pyrimethamine → For T/t of Malaria.

# Toxoplasmosis:

For t/t: Sulfadiazine + Pyrimethamine  
+ Folic acid.Safest drug For t/t of Toxoplasmosis in pregnancy  
- Spiramycin (Macrolide)Cotrimaxazole: Sulfamethoxazole (400mg)  
+ Trimethoprim (80mg).Cotrimaxazole DS: Sulfamethoxazole (800mg)  
+ Trimethoprim (160mg)

DOC: Pneumocystis carinii pneumonia.

## Aminoglycosides.

For tft of TB → Streptomycin (1st line drug)  
 Kanamycin  
 Capreomycin } 2nd line drug.  
 Amikacin }

- All are ionised molecule so not absorbed via orally.

Streptomycin - DOC for Plague (mass prophylaxis)  
 Doxycycline

Also useful in - TB  
 Tularemia.

Aminoglycoside useful for Pseudomonas:

T = Tobramycin

A = Amikacin

G = Gentamycin

Among Cephalosporin  
 - Ceftazidime  
 Cefoperazone.

For severe Pseudomonas infection - TOC is combination of Cephalosporin + Aminoglycosides.  
 eg: Ceftazidime + T or A or G.

Last option for severe resistance case of Pseudomonas

↓  
 Polymyxin B.

Paramomycin -

Oral - Amoebiasis

iv - Kala azar.

Neomycin:

generally parentally

Oral - Gut sterilization

Hepatic encephalopathy.

# Aminoglycoside follow conc<sup>n</sup> dependent killing pattern  
so given OD dose.

# Beta Lactam follow time dependent killing  
so given TDS / QID.

Post antibiotic effect of Aminoglycoside:

Even though the ~~low~~ drug level is

lower than the MAC value still produce action.

# Common S/E of Aminoglycoside:

Nephrotoxicity

Ototoxicity

Neuromuscular block. (Neomycin)

Among the Aminoglycoside - Gentamycin } Highly  
Tobramycin } Nephrotoxic  
Neomycin }

Least Nephrotoxic - Streptomycin



# Max<sup>m</sup> deafness caused by - Kanamycin  
 Amikacin Max<sup>m</sup>.  
 Neomycin

Deafness 1st high frequency sound → lastly low frequency sound.  
 First damage Base of hair cell → lastly apex of hair cell.

Vestibular damage - Streptomycin  
 Gentamycin.

Equal - Tobramycin  
 Least - Netilmicin.

~~1st~~ Quinolones:

MOA: inhibits DNA Gyrase in Gram ~~+~~ -ve  
 inhibit Topoisomerase IV in Gram +ve.

Route of Excretion - Kidney.  
 ↳ So not given in Renal failure.

Excretion via liver - Prefloxacin } Used in RF  
 Trovafloxacin } (Safe)  
 Moxifloxacin }

Ciprofloxacin:

DOC for Typhoid

↳ Currently 1st line choice

- Ceftriaxone (i.v.)  
 (In children/pregnancy)  
 or in Ciprofloxacin Resistance.

Drug interaction  $\bar{c}$  theophylline:

Ciprofloxacin is microsomal enzyme inhibitor, when ~~off~~ given  $\bar{c}$  theophylline, theophyllin level  $\uparrow$  in plasma which causes convulsion/seizures.

Withdrawn Quinolones:

Trovafloxacin - liver toxicity.

Grepafloxacin - QT prolongation

Gatifloxacin - Unpredictable glucose profile.

$\rightarrow$  Only systemic use was withdrawn

Eye drop available.

Clinafloxacin - Phototoxic



available Quinolones.  $\left\{ \begin{array}{l} \text{Sparfloxacin (longest action)} \\ \text{Lomifloxacin} \end{array} \right.$

Macrolides.

Clarithromycin:

Useful for - MAC

H. pylori

Leptosy.

Azithromycin:

Useful for - MAC

Gonococci/Syphilis/Chancroid

Chlamydia

Legionella

Campylobacter jejuni

## Common S/E of Macrolides -

- GI toxicity → due to motilin
- Hearing impairment.
- Hepatitis
- Cholestatic jaundice caused by erythromycin estolate.
- ~~Erythromycin estolate~~

## Drug interaction:

- All macrolides are microsomal enzyme inhibitor

Erythromycin - Max<sup>m</sup> microsomal enzyme inhibition  
so max<sup>m</sup> drug interaction

Azithromycin - Least microsomal enzyme inhibition

# Azithromycin may cause QT prolongation.

# Erythromycin aggravates pyloric stenosis.

## Tetracycline.

## Tigecycline -

Given i.v.

Useful for MRSA/VRSA

Excreted by bile so safe in kidney failure.

## Doxycycline -

Excreted via bile, safe in RF

## Demeclocycline -

Phototoxic

Cause DI

Useful for SIADH.

Minocycline :

Used for leprosy.

↳ Rifampicin  
Ofloxacin  
Minocycline.

S/E - Vestibulo toxicity.

# All tetracycline having risk of causing elevation of ICT called Pseudotumour Cerebri.

# Outdated tetracycline may cause Fanconi's Syndrome.

# Tetracyclines are DOC for ① Rickettsial infection  
② Chlamydia infection  
③ Lymphogranuloma Venereum (LGV)

Tetracycline used as Prophylaxis of: Cholera  
Brucellosis  
Plague.

C/I in pregnancy - Fulminant hepatic failure

Baby ← Bone & teeth problem.

Most safest antibiotics in pregnancy →  $\beta$ -lactam

↓  
Cephalosporin & Penicillin > Azithromycin

### Antibiotic & Colour association:

- Grey baby - Chloramphenicol
- Yellow baby - Sulfonamide
- Red man Syndrome - Vancomycin
- Discoloured teeth - Tetracyclines.
- Coffee coloured teeth - Nitrofurantoin
- Loss of Red/green perception - Ethambutol.
- Reddish black - Clofazimine.

### Tuberculosis

#### Anti-tubercular drugs:

##### Isoniazid (INH):

- activated in the help of INH A gene & catalase peroxidase.

MOA: Inhibiting mycolic acid synthesis.

- Undergoes metabolism by acetylation.

##### S/E - Hepatotoxicity (M/C)

↳ due to formation of Acetyl hydrazine

##### Neuropathy

↳ t/t - Slow administration of Vit B6

Prophylactically - 10mg/day

Neurotoxicity - 100mg/day.

##### Memory impairment

##### Psychosis.

##### Shoulder hand syndrome

##### SLE

##### Cheese reac<sup>n</sup>.

- # It is microenzyme inhibitor.
- # Doesn't require dosage adjustment in pts w/ Renal disease.
- # Useful for prophylaxis of TB
- # Max<sup>m</sup> CSF penetration.

Isoniazid → derivative of Isoniazid.  
Used for elevating mood.

Rifampicin:

- Activated w/ help of Repo B gene.

MOA: Inhibit DNA dependent ~~RNA~~ RNA polymerase.

- Excretion via Bile & feces  
So safe in RF.

S/E - Non serious:

Reddish orange colour (Urine, Sweat & tears)  
Staining of contact lenses.

Serious:

Hepatitis  
Respiratory syndrome  
Hemolysis  
Purpura.

- # It is microsomal enzyme inducer  
pt w/ HIV Receiving antiviral drug, if we use Rifampicin for TB, HT failure occurs.  
Alternate drug → Rifabutin → Causes Pseudo jaundice.

Pyrazinamide:

- Act by inhibiting mycolic acid synthesis.

S/E - Hepatotoxicity  
Hyperuricemia

# No drug interaction bcoz Neither microsomal enzyme inducer or inhibitor.

# Undergoes renal route of excretion so need dosage adjustment in RF pt.

Ethambutol:

Bacteriostatic

MOA: Inhibiting Arabinogalactan synthesis.

S/E → • Optic neuritis

↳ loss of ability to differentiate red from green.

↳ Supplement c Hydroxycobalamin (Vit B<sub>12</sub>)

• Hyperuricemia.

Excretion → Undergo renal route of excretion

- Need dose adjustment in RF pt.

Streptomycin:

CI in pregnancy bcoz cause permanent deafness in children.

TB in liver ds pt:

Avoid - Isoniazid, Rifampicin, Pyrazinamide.

Safe - Streptomycin, Ethambutol.

TB in a Renal ds pt:

Avoid - ~~INH~~ E, P, S

Safe - R > H

Newer drug for MDR-TB:

Bedaquiline:

Inhibit mycobacterial ATP synthase.

Food ↑ Absorption.

Cross resistance  $\bar{c}$  Clofazimine

May cause QT prolongation.

↳ Cardiotoxicity.

Delamanid

Pretomanid } Inhibit Mycolic acid synthesis.

Sutemizolid - Derivative of linezolid.

Anti TB drug causing:

① Hypothyroidism - Ethionamide (also used for leprosy)

PAS

② Psychosis - INH, cycloserine.

Antibiotic useful in MAC = Azithromycin,

Clarithromycin

REC Regimen (R = Rifabutin, E = Ethambutol, C = Clarithromycin)



(3) Cross BBB - INH, Pyrizinamide, Rifampicine, Cycloserine.

(4) Uveitis - Rifabutin

Anti-leprosy drug.

- ATT drugs → Rifampicin  
Ethionamide.

Other drug → Clofazimine  
Dapsone.

Antibiotic useful for leprosy - Ofloxacin  
Minocycline  
Clarithromycin

Dapsone - Sulphonamide

Uses of Dapsone -

DOC for dermatitis herpetiformis.

# Inj. Acadapsone (i.m) one dose acting for 3 months.

SE - Allergy (M/C)

Hemolytic Anemia.

Clofazimine -

Bacteriostatic

Anti-inflammatory property.

↓

also useful for lepra reac<sup>n</sup>.

SE - Reddish black skin discoloration

Dermatological.

Lepros React<sup>n</sup>:

Type I - Cell mediated immunity to *M. leprae*.

Type IV hypersensitivity.

TOC - Prednisolone (Steroid).

Type 2 - Immune complex deposition.

Type III Hypersensitivity.

T/t - Steroids

Clofazimine

Chloroquine.

Virology.

Drugs useful for HIV:

Fusion inhibitors:

Enfuvirtide

- Given SC

SC → Injection site react<sup>n</sup>

Pneumonia (Rare)

CCR-5 inhibitor:

Maraviroc - FDA approved

Aplaviroc } under trial.

Vicriviroc }

NRTI's (Nucleoside Reverse Transcriptase inhibitor):

Zidovudine (M/C)

↳ Myelosuppressant (Macrocytic Anemia)

↳ Lipodystrophy → due to mitochondrial DNA polymerase

Didanosine

↳ Pancreatitis

Stavudine - Worst drug.

↳ S/E - Severe Neuropathy

Lactic acidosis

Lipodystrophy

Abacavir (Rule out HLA-B5701 allele, MI, safe in RF)

Zalcitabine

also useful for HBV { Lamivudine - Best drug (No serious adverse effect)

Etricitabine

Tenofovir - Causes GIT toxicity, Fanconi's Syndrome.

↳ Really a nucleotide inhibitor.

NNRTI:

1st generation:

Efavirenz

Nevirapine, NVP

Delavirdine.

2nd gen:

Etravirine

Rilpivirine.

Common S/E - Skin Rash

- Steven Johnson Syndrome

- Toxic epidermal necrolysis.

Nevirapine

↳ S/E - Hepatitis (LFT)

Efavirenz

↳ S/E - Neuropsychosis

Integrase inhibitors:

Raltegravir

Elvitegravir

Dolutegravir

} Best drug.

Protease inhibitor:

Saquinavir - Best tolerated

Indinavir - Nephrolithiasis

Nelfinavir

Ritonavir - Powerful microsomal enzyme inhibitor (CYP3A4)

↓

Called Booster.

Amprenavir

Fosamprenavir

Atazanavir → Not cause lipodystrophy.

Lopinavir.

↳ may cause intracranial hemorrhage.  
Tiplranavir } Sulfonamide  
Darunavir }

Common S/E - Hyperglycemia

Fat redistribution

Hyperlipidemia.

## # TESAMORELIN - GHRF

↳ Reduce abdominal fat in HIV c lipodystrophy.

## CROFELEMER - CFTR inhibitor

Use - HIV <sup>drug</sup> induced diarrhoea.

## Maturation inhibitor.

- Bevirimat. (Under Trial)

## # HAART/CART (Highly active anti retroviral therapy):

2 NRTI + 1 NNRTI  
NRTI + NNRTI + PI ] Triple drug therapy

↓  
To prevent drug resistance.

NACO 2011 → Zidovudine + Lamivudine + Nevirapine.

CMV (Cytomegalo Virus) → Cause Retinitis.

- Ganciclovir (DOC)

↳ M/c SE - Myelosuppression.

Valganciclovir

Fomevirsin

Foscarnet

Cidofovir

Maribavir.

Fos carnet :-

Useful for HSV (resistant to Acyclovir)

CMV (Ganciclovir resistance)

ADR - ARF

Penile ulcer.

Cidofovir - Useful for Respir papillomatosis.

Drug for Herpes simplex Virus

Acyclovir - For HSV  
ADR - <sup>Acute</sup> Renal Failure

Docosanol - Viral entry inhibitor  
given topically

Famciclovir - Prodrug  
Active form - 6-deoxy penciclovir.

Drug useful for Hep. B:  
Injections are { IFN- $\alpha$   
PEG-IFN- $\alpha$

Oral agents:

1st line - Entecavir  
Tenofovir (Anti HEV drug)

2nd line - Lamivudine

Adefovir

Telbivudine.

Drugs for HCV:

Commonly we give PEG INF $\alpha$  plus ribavirin.

Sofosbuvir - Given orally

Renal excretion

Causes Bradycardia.

Other drugs -

Telaprevir

Boceprevir

Simeprevir

Grazoprevir

~~Eltavir~~ Elbasvir

Daclatasvir

Velpatasvir

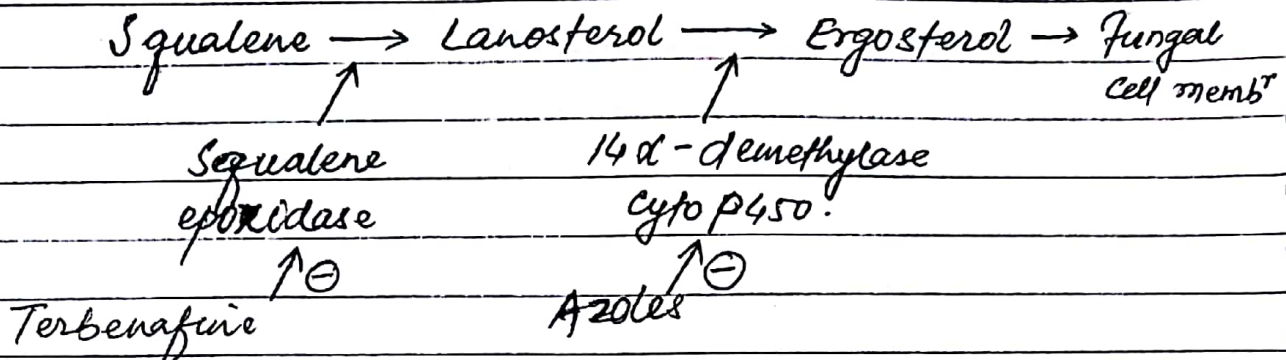
Ombitasvir

Ledipasvir

Viramidine - (Under trial)

## Antifungal drugs.

### - Membrane Active Antifungal Agents:



Polyene antibiotics -

Amphotericin B + Ergosterol  $\rightarrow$  Forms a pore in fungal cell

Act on fungal cell wall.

Destroy fungus

Amphotericin B:

Usually given as a slow iv infusion.

Very well distributed all over body /  
poorly distributed in CNS.

ADR - Infusion related reac<sup>n</sup> (Fever, chills)

Nephrotoxicity (Dose limiting toxicity).

Hypokalemia

Hypomagnesemia

Anemia

Seizure.

To avoid Nephrotoxicity - Give Hydration.

# Newer formulation: ABCD (Colloidal dispersion)

↓  
less systemic toxicity.

ABLC (Lipid complex)

Liposomal AMB (For Kalaazar)





Nikkomycin - Inhibit chitin synthesis  
Useful for Candida & Aspergillosis.

### Amoebiasis

#### Lumen Amoebiasis

- Diloxanide furoate  
(Flatulance)

- Nitazoxanide

↳ Use in Cryptosporidiosis

- Quinodochlor → Cause Subacute myelo optic neuropathy (SMA)

- Iodoquinol

- Paromomycin (oral) → i.v. for Kala-azar.

- Tetracyclines

#### Tissue

Extraintestine

Both intestine

& Extra intestine.

Extraintestine :

Chloroquine.

Both :

Metronidazole

Tinidazole

Secnidazole (Single dose) - M/c SE - Nause, Vomiting

Ornidazole

[Metallic taste]

Satranidazole (less neurological ADR)

Ementine

Dehydro ementine.

# Guinea worm : For complete removal of worm  
DOC - Niridazole.

### Helminthiasis

Trematodes	Cestodes	Nematodes.
DOC - Praziquantel	DOC - Praziquantel	DOC - Albendazole
Except - Fasciola hepatica	Except - Echinococcus granulosus	Except - Ochocerca Volvulus
↓	↓	↓
Triclabendazole	Neurocysticercosis	(Ivermectin)
Bithional	↓	Strongyloidosis
	Albendazole (hepatotoxic)	Scabies
		W. bancrofti
		↳ DEC.

### Leishmaniasis

Kala-azar	Cutaneous	Mucocutaneous
↳ For all forms	Sodium Stibogluconate	↓
(DOC) Amphotericin B (In India)	↓	↓
Hyperkalemia → Pentamidine (ENAC blocker)	Fluconazole	Amphotericin B.
Paromomycin	Metronidazole	
oral [ Miltefosine		
Sitamaquine		

## Trypanosomiasis.

African

- Sleeping sickness.

*T. gambiense*

& *T. rhodesiense*.

South American.

- Chagas disease

• *T. cruzi*

DOC - Benznidazole  
Nifurtimox.

Early haemolymphatic stage

Suramin (DOC)

Pentamidine

Late - CNS stage

Malascoprol (DOC)

Eflornithine.

## Anti-Malarial drug

Chloroquine (M/C)

↓

↳ Very large apparent  $V_d$  of 100-1000 L/kg.

Uses:

R - Rheumatoid Arthritis

E - Extra-intestinal Amebiasis

D - DLE (Discoid lupus erythematosus)

L - Leprosy react<sup>n</sup>

I - Infectious mononucleosis

P - Photosensitive react<sup>n</sup>

M - Malaria

G - Giardiasis.

- Safe in Pregnancy.

S/E → GI toxicity (Nausea & Vomiting)  
 CVS (Bradycardia, HTN)  
 Chronic therapy cause Bull's eye maculopathy.  
 Liver damage.

Mefloquine:

For t/t & prophylaxis of Malaria.  
 Long half life.  
 Single oral dose  
 S/E - Neuropsychosis.

If combine c̄ Halofen, Quinine - Risk of QT prolongation.

HALOFANTRINE, LUMEFANTRINE:

Absorption ↑ c̄ food.  
 Halofantrine - more cardiotoxic.

Lumefantrine + Artemether ⇒ ACT

# Primaquine

- Vivax curative

In G6PD deficiency → Cause hemolytic anemia.  
 C/I in pregnancy.

Artemisinin:

Artesunate	} Fast acting drug
Artemether	
Arteether	

↓  
 For extending duration of action  
 combine c̄ Mefloquine.

Indication:

Multi drug resistance Malaria  
Cerebral Malaria.

Not indicated for chemoprophylaxis of Malaria

S/E - GI toxicity (N/C)

CVS → QT prolongation, 1st degree AV block.

Hematology → Reversible Leucopenia.

WHO approved Combiviral therapies:

FDC = Artemether / lumefantrine  
Artesunate + amodiaquine  
Artesunate + SP  
Artesunate + Mefloquine } ACT's

Unsafe Antimalarial drug in Pregnancy:

Halofantrine

Tetracycline/ Doxycycline.

Primaquine