



PHARMACODYNAMICS I

MECHANISMS OF DRUG ACTION

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ILOS

➤ **Identify different targets of drug action**

Differentiate between their patterns of action; agonism versus antagonism

Elaborate on drug binding to receptors



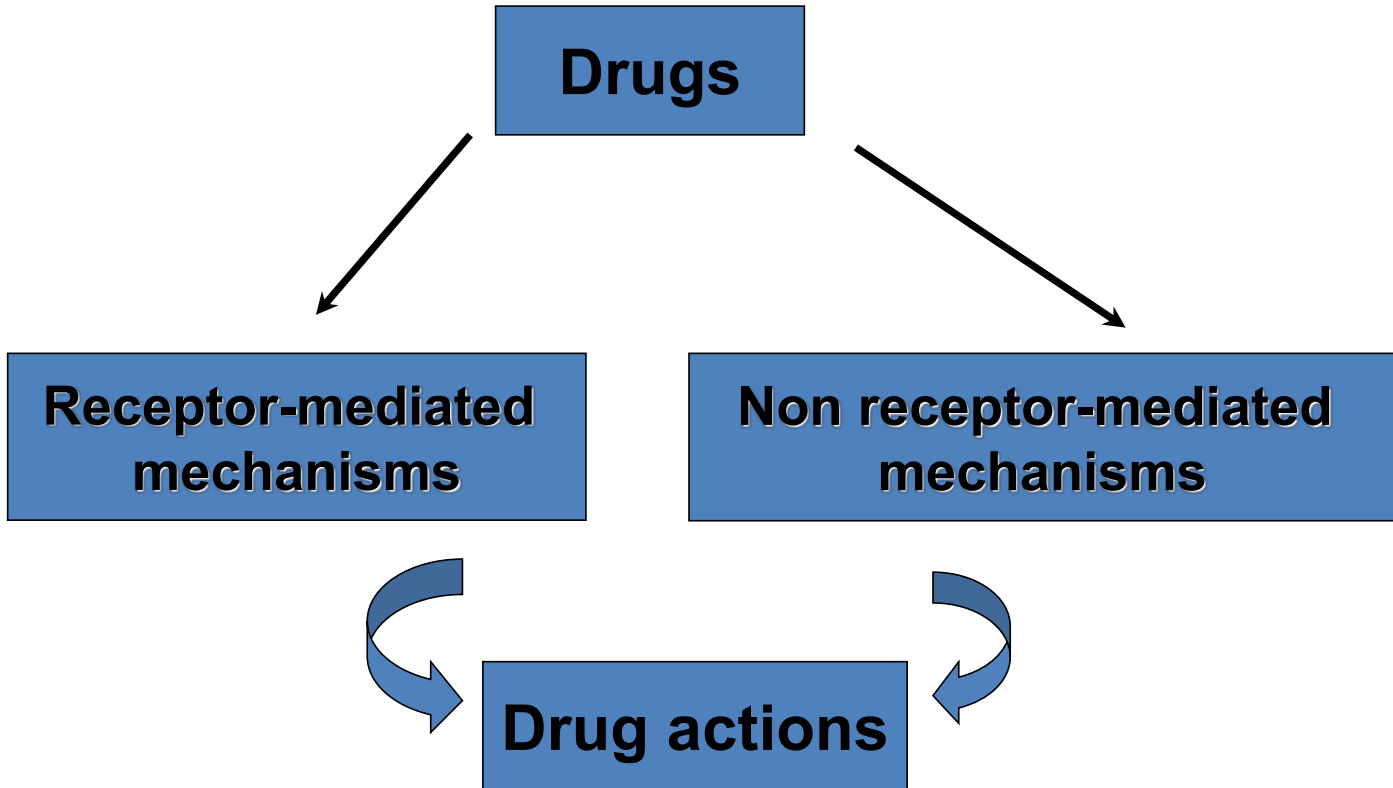
WHAT IS PHARMACODYNAMICS?

Pharmacodynamics is a branch of pharmacology that deals with the study of the biochemical and physiological effects of drugs and their mechanisms of action.

**WHAT ARE THE MECHANISMS OF DRUG
ACTION?**

How drugs produce action?

What are targets for drug binding ?



WHAT ARE THE MECHANISMS OF DRUG ACTION?

Drugs can produce their actions by one of the following mechanisms:

1) Receptor-mediated mechanisms (Binding with biomolecules):

- Receptors = Biomolecules = Targets
- Targets are mostly **protein in nature**.

1) Non receptor-mediated mechanisms

Physio-chemical properties of drugs.

Non receptor–mediated mechanisms

Drugs can produce actions by:

Chemical action

- Neutralization of gastric acidity by antacids.

Physical action

- Osmotic diuretics.

Receptor-mediated mechanisms

Drugs can produce actions by binding with biomolecules (Protein Targets)

Protein targets for drug binding

- Physiological receptors
- Enzymes
- Ion channels
- Carriers
- Structural protein

What are targets for drug binding ?

Receptors

- Is a special target macromolecule that binds the drug and mediates its pharmacological actions.

Where are receptors located?

- Cell membrane.
- Cytoplasm.
- Nucleus.

What are targets for drug binding ?

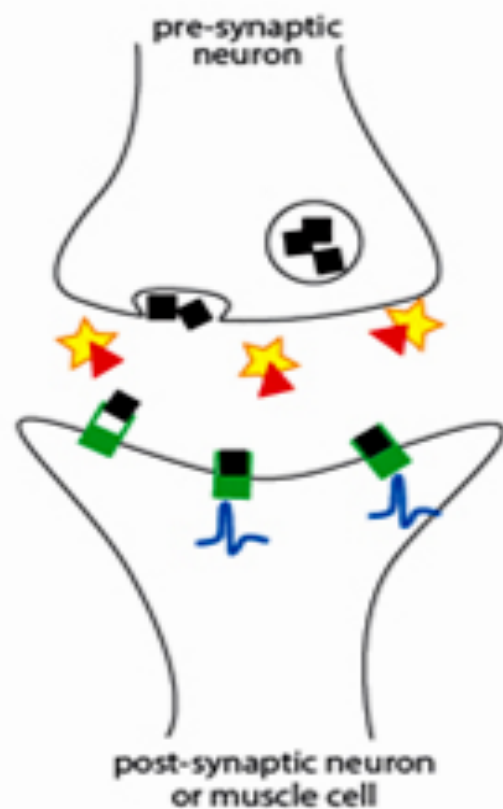
Enzymes

- The drug competes with the natural endogenous substrate for the enzyme.
- e.g. Anticholinesterases inhibit acetylcholinesterase thus producing cholinomimetic action.
- **Neostigmine reversibly** compete with **ACH** for acetylcholinesterase enzyme at motor end plate (neuromuscular junction).

ACh Esterase STOPS signaling process



- ACh
- U ACh Receptor
- ⚡ Signal transmission
- ★ ACh Esterase



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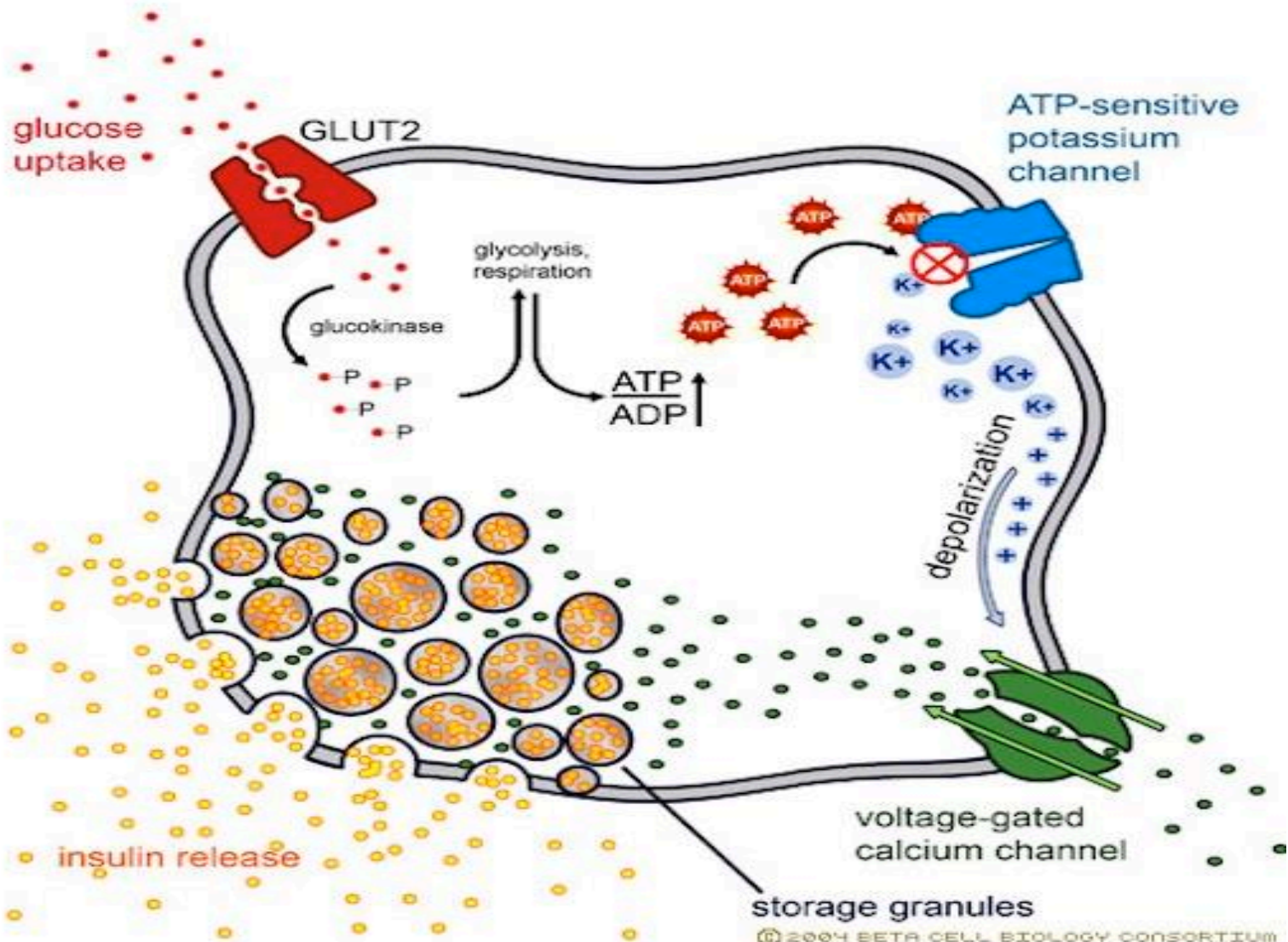
What are targets for drug binding ?

Ion channels

- Drugs bind to alter channel function **(by opening or blockade)**.
- Channels are responsible for influx or outflux of ions through cell membranes.
- They are activated by alteration in action potential.

Ion channels

- **e.g. Sulfonylurea drugs (antidiabetic drugs):**
block potassium channels in pancreatic beta cells resulting in increase in intracellular potassium & depolarization and opening of calcium channels and insulin secretion.



What are targets for drug binding ?

Carrier molecules

- Drugs bind to such molecules to alter their transport ability.
- Responsible for transport of ions and small organic molecules between intracellular compartments, through cell membranes or in extracellular fluids.

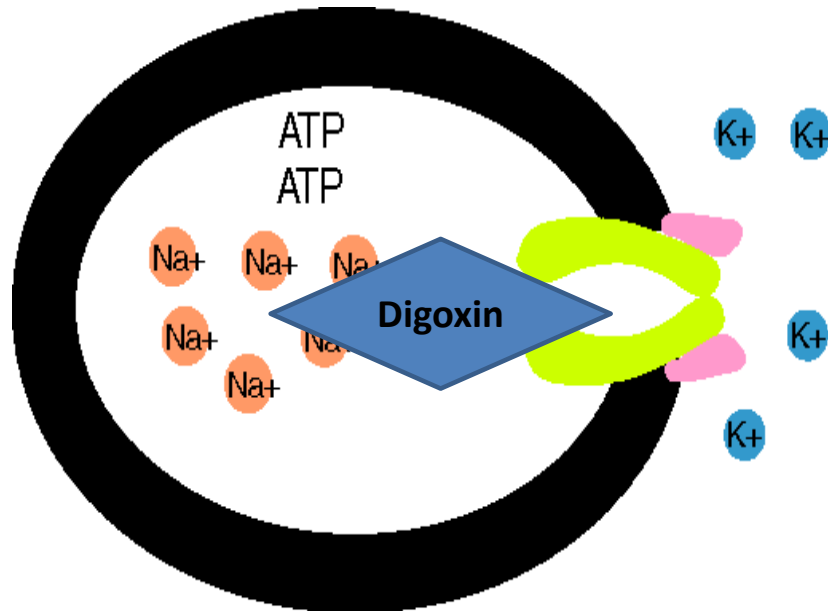
e.g. **Na pump** (Na^+/K^+ ATPase) blocked by digoxin.

e.g. **dopamine transporter** blocked by cocaine.

Carrier molecules

Digoxin: blocks Na efflux via **Na⁺/K⁺ pump (Na⁺/K⁺-ATPase)** ; used in the treatment of heart failure.

More Na⁺ in the cytosol so stronger contraction of heart muscles



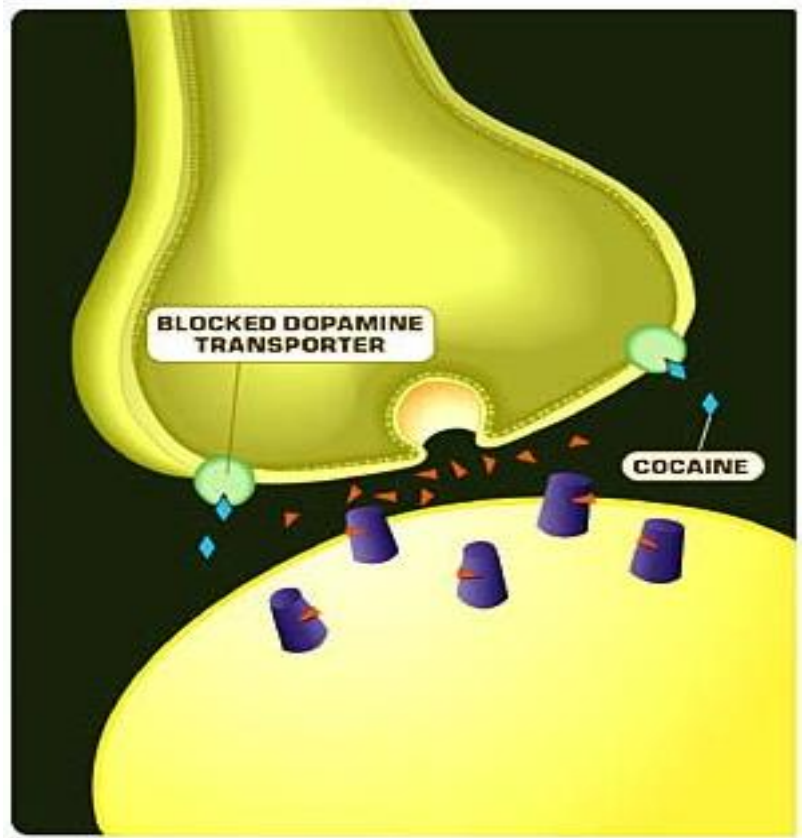
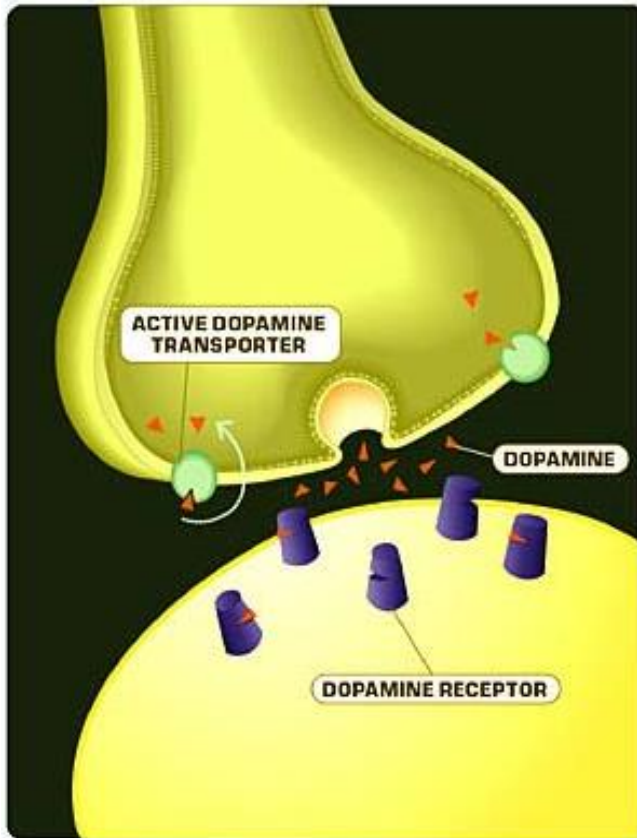
What are targets for drug binding ?

Carrier molecules

Cocaine:

- blocks transport or reuptake of (**catecholamines mainly dopamine**) at synaptic cleft.
- The dopamine transporter can no longer perform its reuptake function, and thus dopamine accumulates in the synaptic cleft producing **euphoria**.

Effect of cocaine



What are targets for drug binding ?

Structural proteins

e.g. **Tubulin** is target for drugs as **anticancer drugs** and anti gout drugs.

Tubulin is required for microtubules formation (cytoskeleton).

MICROTUBULE DESTABILIZERS

Vinca alkaloids

- Vincristine
- Vinblastine
- Vinorelbine
- Vinflunine
- Halichondrin B
- Eribulin mesylate
- Cryptophycins
- Dolastatins

Vinca binding site

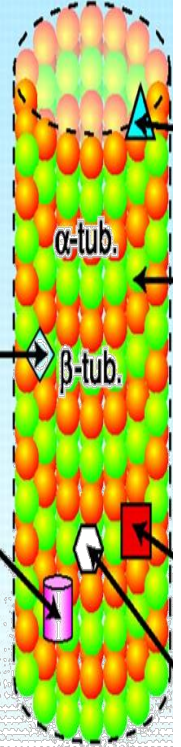
Colchicine binding site

Colchicine

2-Methoxyestradiol

Sulphonamides

Aspergillus derivatives



Tubulin Structure

Structural proteins

Vincristine

Anticancer that kills cancerous cells by inhibiting microtubule formation and cell division.

Colchicine

- used in treatment of gout
- binds to tubulin and inhibits microtubule formation, preventing neutrophil motility and decreasing inflammation

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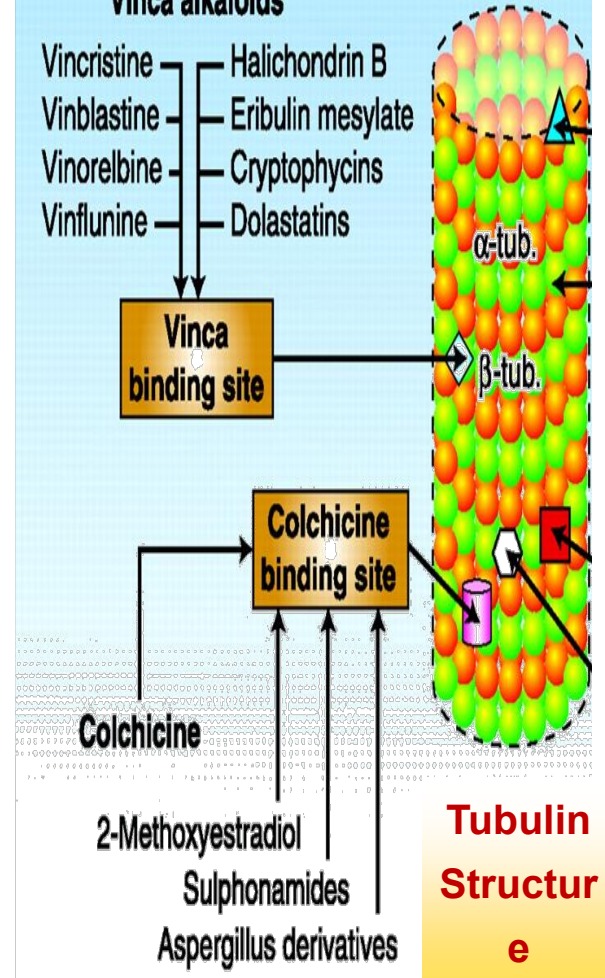
Colchicine

2-Methoxyestradiol

Sulphonamides

Aspergillus derivatives

Tubulin
Structure



What are the binding Forces between drugs and receptors?

- Ionic bond.
- Van-Dar-Waal.
- Hydrogen bond.
- Covalent bond.

Affinity

Ability of a drug to combine with the receptor.



Efficacy (Intrinsic Activity)

- Capacity of a drug receptor- complex (D-R) to produce an action.
- **E max** : is the maximal response produced by a drug

Agonist

is a drug that combines with receptor and elicit a response (has affinity and efficacy).

e.g. acetylcholine (Ach) acts upon muscarinic receptors.

Antagonist

- is a drug that combines with a receptor without producing responses.
- It blocks the action of the agonist.
- **It has affinity but no efficacy or zero efficacy.**
- It has similar chemical structure to an agonist.
- e.g. atropine block the action of Ach on muscarinic receptors.

Agonist and Antagonist



Agonist

Full agonist.

Partial agonist.

Full Agonist

A drug that combines with its specific receptor to produce maximal effect by increasing its concentration (affinity & high efficacy).

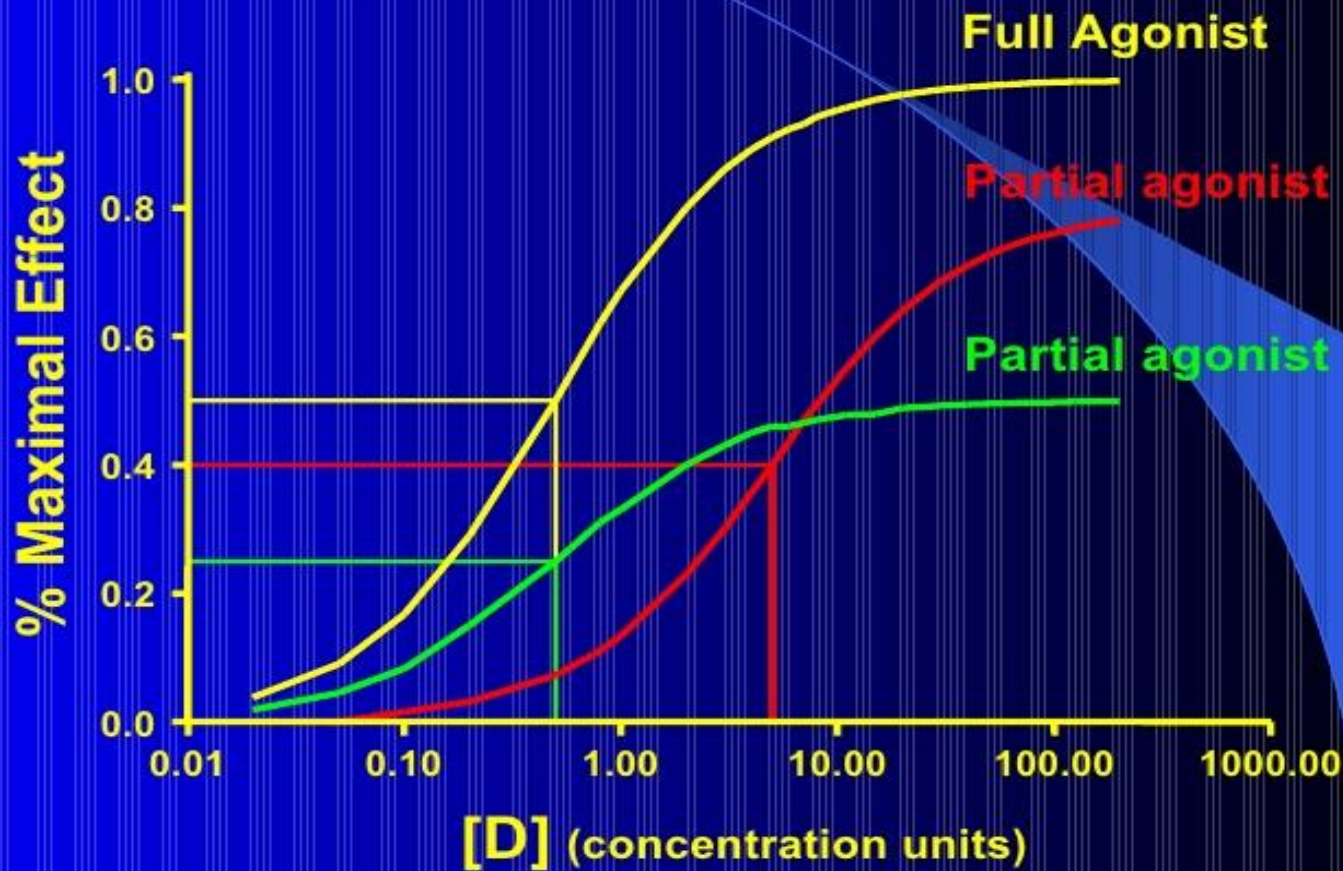
e.g. acetylcholine (Ach).

Partial Agonist

combines with its receptor & evokes a response (submaximal effect) regardless of its concentration (**affinity & partial efficacy**).

e.g. **Pindolol**: a beta blocker which is a **partial agonist**, produces less decrease in heart rate than pure antagonists such as propranolol.

Partial Agonist: Even though the drugs may combine with the same number of receptors, the magnitude they can produce may differ



TERMS DEFINITIONS



Affinity is the capacity of a drug to form a complex with the receptor(DR complex)

Efficacy(Intrinsic activity) the ability of the drug once bound to the receptor to trigger response

The value of intrinsic activity ranges from 0 to 1

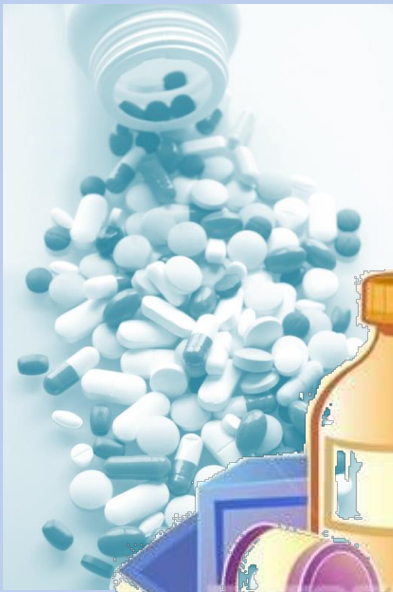
TERMS DEFINITIONS



Full agonist having a full affinity to the receptor and a maximal intrinsic activity (1) e.g. **acetylcholine**

Partial agonist having a full affinity to the receptor but with low intrinsic activity (<1) e.g. **pindolol**

Antagonist having full affinity to the receptor but no intrinsic activity(0) e.g. **atropine**



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PHARMACOLOGY