



PHARMACODYNAMICS

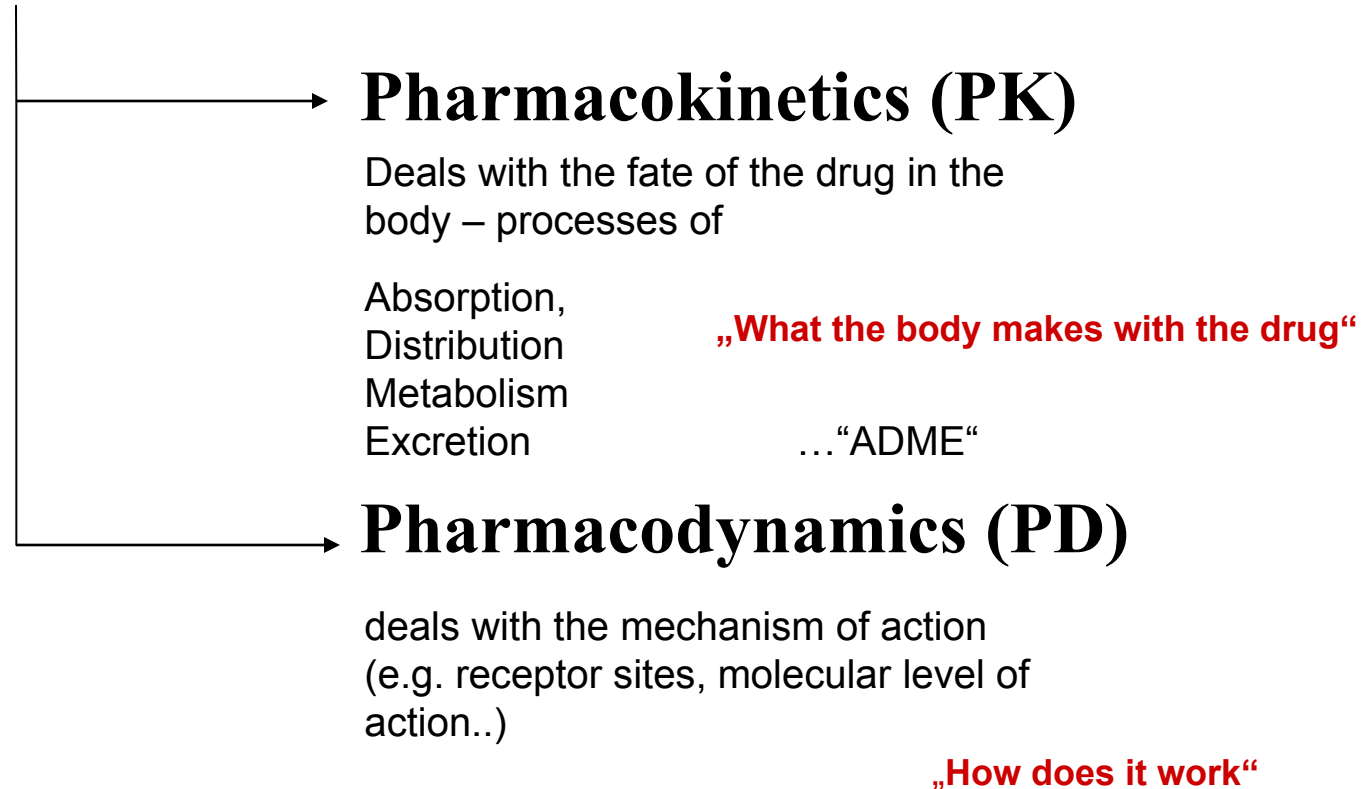
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PHARMACOLOGY



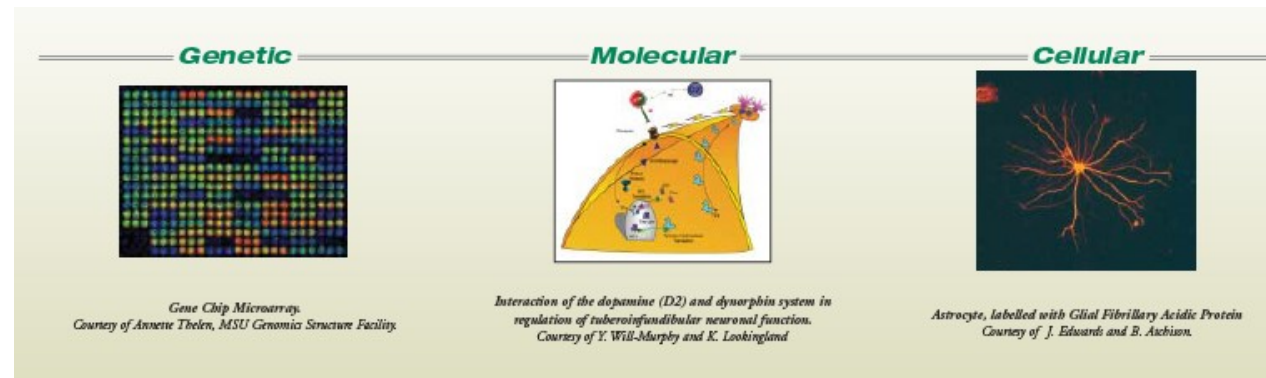


Pharmacodynamics

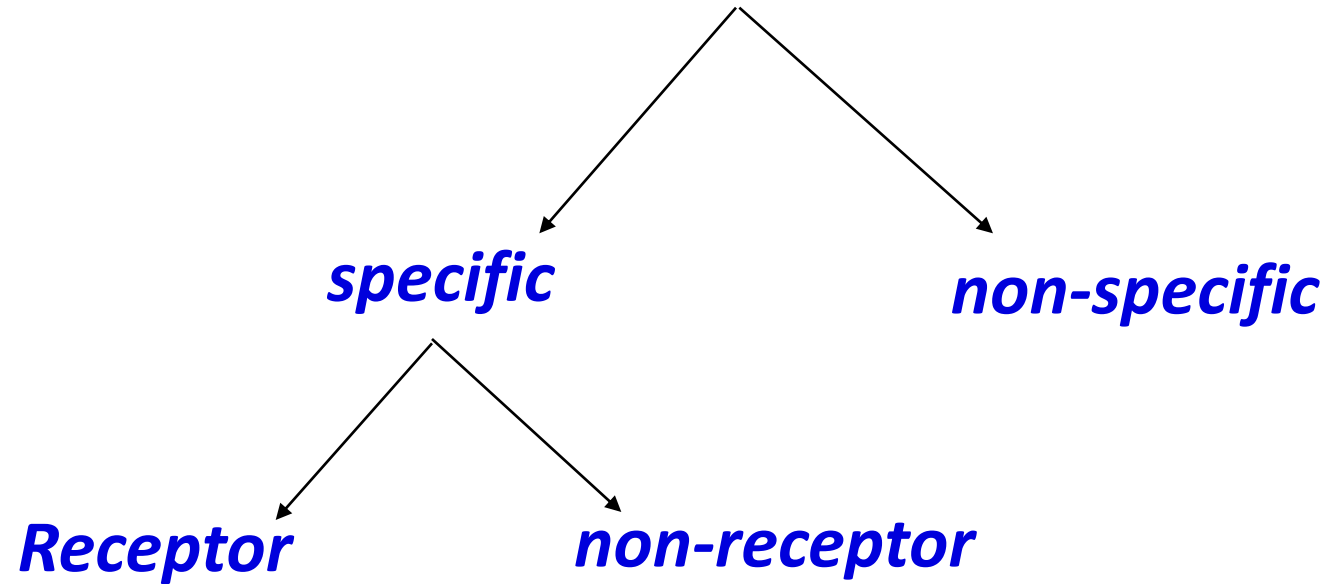
(how drugs work on the body)

- The action of a drug on the body, including receptor interactions, dose-response phenomena, and mechanisms of therapeutic and toxic action

- Main targets – cellular, molecular, genetic level...
 - Therapeutic effects
 - Adverse effects



Mechanism of drug actions





I. Non-specific drug effects

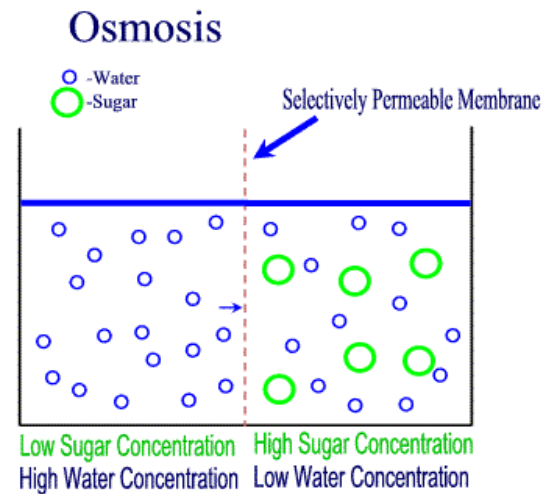
...through by the general physical-chemical properties of substances - no specific chemical and structural configuration of drugs is needed

- influencing pH
- oxidating and reducing agents
 - protein precipitation
 - adsorbents / detergents
 - chelating agents



a. based on osmotic properties

- e.g. salinic laxatives (magnesium sulphate, lactulosa)
- osmotic diuretics (mannitol)





b. influencing acid-base balance

- Antacids
 - aluminium hydroxide
 - magnesium carbonate
 - calcium carbonate
 - sodium bicarbonate
- pH modifiers (blood, urine)
 - sodium bicarbonate, ammonium chloride



c. based on oxido – reducing properties

- e.g. 3% hydrogen peroxide, boric acid, fenols
- chlorhexidine act as antiseptics



d. drugs with a large adsorption area

- intestinal adsorbents - Carbo adsorbens (activated charcoal)
- diosmectite (treatment of diarrhoea)

- bind other substances and toxins to themselves



e. surfactants and detergents

- surface active agents: carbethopendecinium bromide (and other quaternary ammonium salts) used primarily as antiseptics.
- some antibiotics (e.g. polymyxins - basic peptides) act as cationic detergents and disrupt phospholipids in bacterial membranes.



f. chelates (chelating agents)

- ethylenediaminetetraacetic acid (EDTA) is a chelating agent, it can form bonds with a metal ion
- dexrazoxane - a cyclic analog of EDTA administered with anthracyclines to prevent cardiotoxicity → Fe²⁺ ions



II. Specific drug effects

effect depends on the specific molecules configuration

➤ **most drugs act (bind) on receptors**

➤ in or on cells

➤ form tight bonds with the ligand

➤ **....on ion channels or carriers**

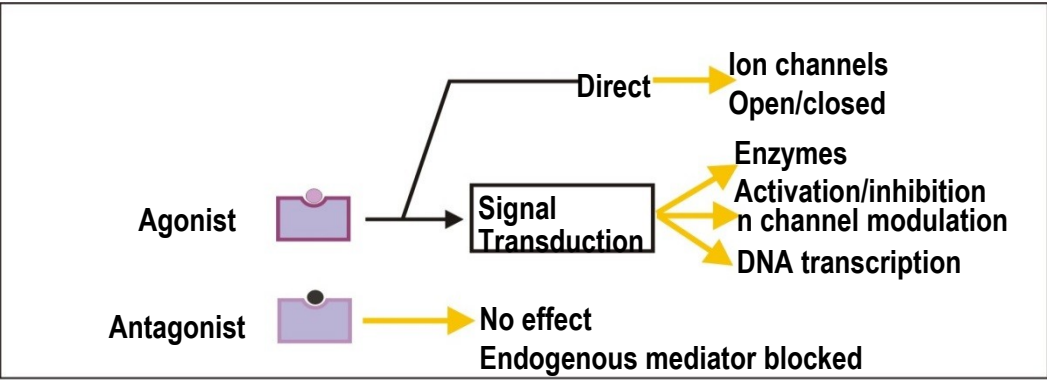


Specific drug effects

- **many drugs inhibit enzymes**
 - A very common mode of action of many drugs
 - **in the patient (ACE inhibitors)**
 - **in microbes (sulfas, penicillins)**
 - **in cancer cells (5-FU, 6-MP)**
- **some drugs bind to:**
 - **proteins (in patient, or microbes)**
 - **DNA (cyclophosphamide)**
 - **microtubules (vincristine)**



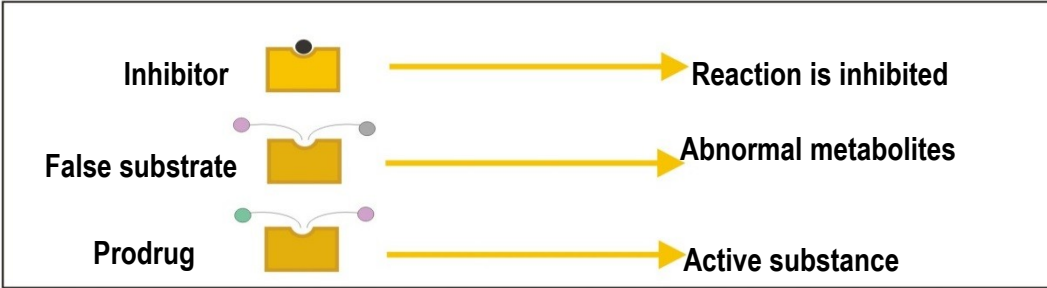
A. RECEPTORS



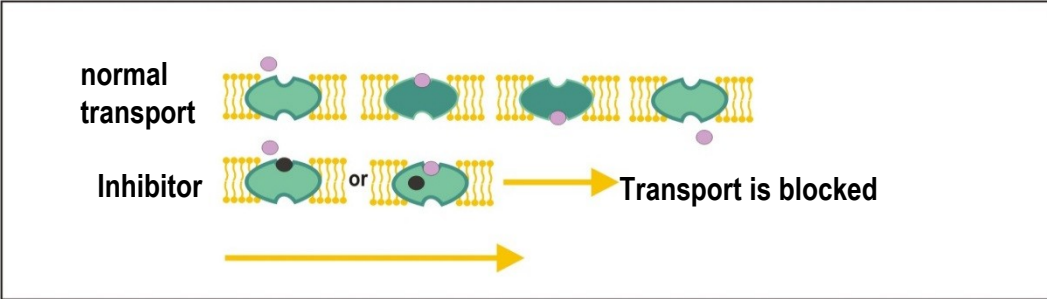
B. ION CHANNELS



C. ENZYMES



D. CARRIERS



A. Receptor – effector system



= complex of processes

extracellular signal -----> intracell. signal cascade-
-----> **effector** (own effect)

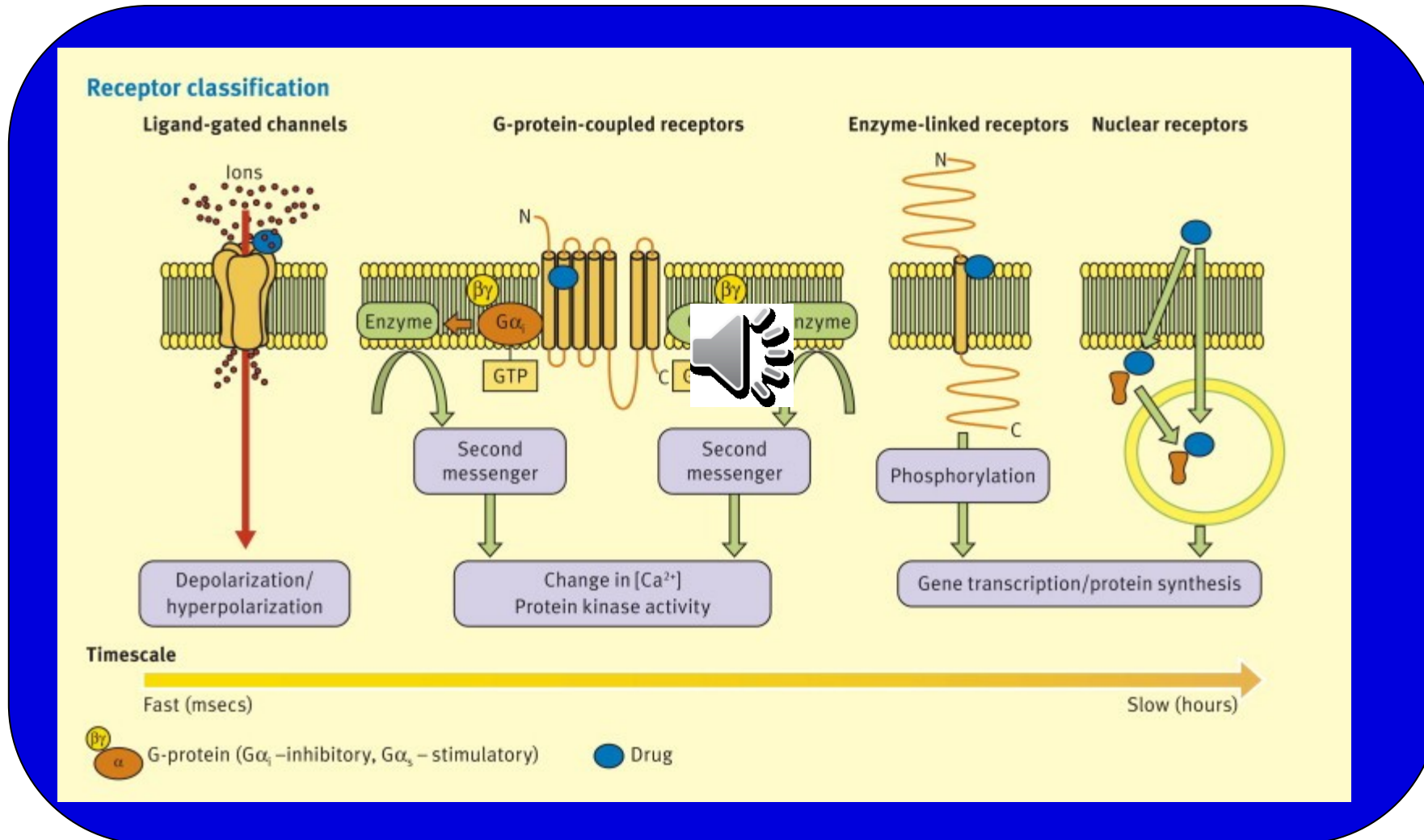
- ✓ **receptor** = protein, which interacts ligands
 - involved in signal transduction
- ✓ **effector** = enzyme, ionic channel etc. change in the activity leads to the effect of drug
- ✓ **ligand** (signal molecule) = molecule able to bind to specific receptor
 - **endogenous** - neurotransmitters, hormones
 - **exogenous** - xenobiotics, drugs

Receptor classification



Localization	Transduction	Ligands
✓ membrane	✓ metabotropic	✓ achol
✓ cytoplasm	✓ ion. channels	✓ amines
✓ organelles	✓ kinase	✓ AMA
✓ auto/heteroreceptors	✓ DNA regulating	✓ peptides

Receptor classification



4 main type of receptors



	Type 1 Receptors connected with ion channels	Type 2 G-protein coupled receptor	Type 3 Receptor tyrosin kinases	Type 4 Intracellular (nuclear) receptors
Place	Membrane	Membrane	Membrane	Intracellular
Efeator	Ion channel	Channel or enzyme	Enzyme	Gene transcription
Binding	direct	G-protein	direct	DNA mediated
Examples	Nicotin-cholinergic receptor, GABA receptor	Muscarin-cholinergic adrenoreceptors	Inzulin, growth factor, cytokin receptor	Steroids, thyroid hormon receptors
Structure	Oligomer composed by subunits surrounding center of the channel	Monomer (or dimer) containing 7 transmembrane helical domains.	Single transmembrane helical domain interconencted with extracelular kinase	Monomer structure with separate receptor and DNA binding domain



Receptor – effector system

- **Affinity**

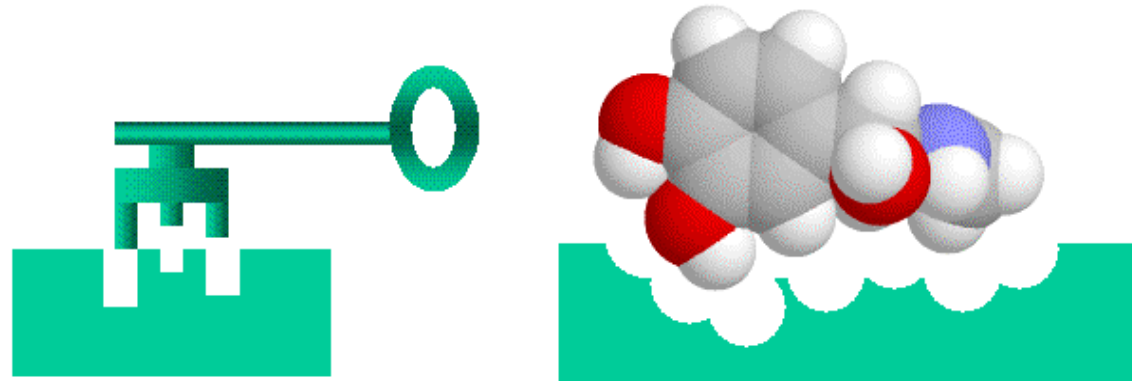
- ✓ the ability of the ligand to bind to the receptor

- **Intrinsic activity**

- ✓ ability to evoke an effect after binding to
receptor

- !!!the presence of sufficient number of receptor for the induction of pharmacological effect is essential as well as sufficient amounts of receptor ligand!!!

Receptor – effector system



Ligand classification (intrinsic activity) AGONISTS

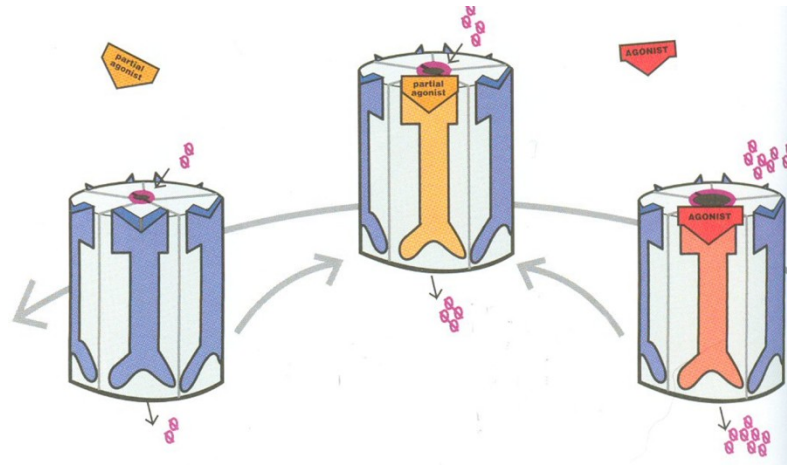


Full agonist

- IA = 1

Partial agonist

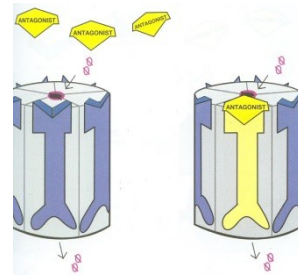
- dualist
- IA in a range from $0 <$ to > 1





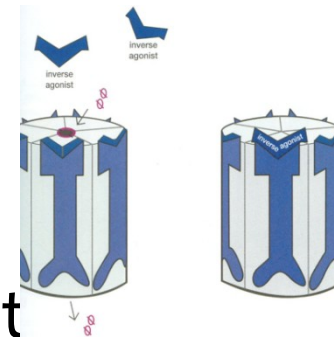
Antagonists

- ✓ $IA = 0$
- ✓ Blocks agonist binding to receptor



Inverse agonist

- ✓ $IA = -1$
- ✓ Stabilizes the receptor in the const activity



Receptor-effector system



Relation between dose and effect

Receptor Activation: Full Agonist, Partial Agonist, Antagonist

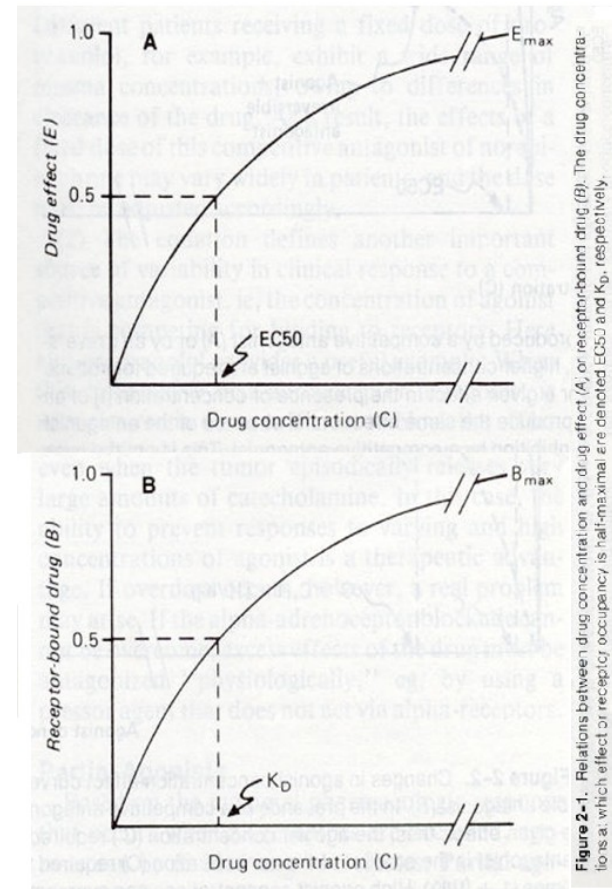
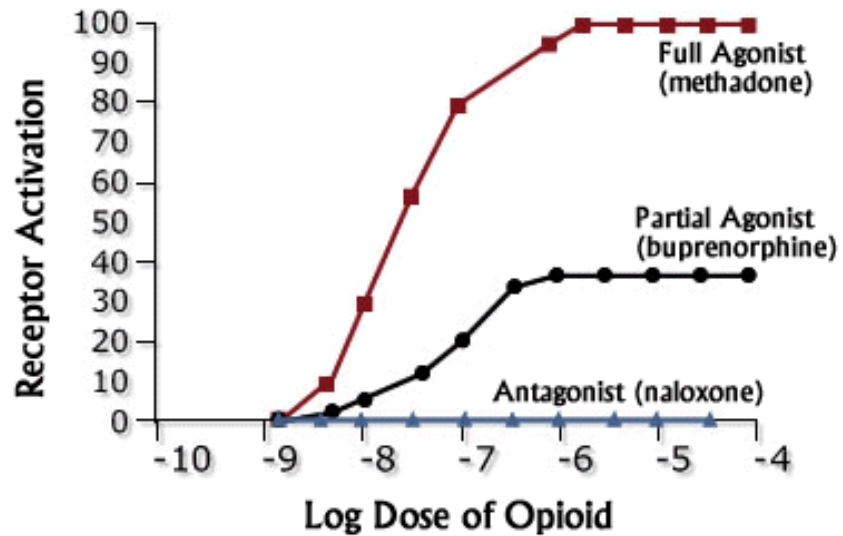
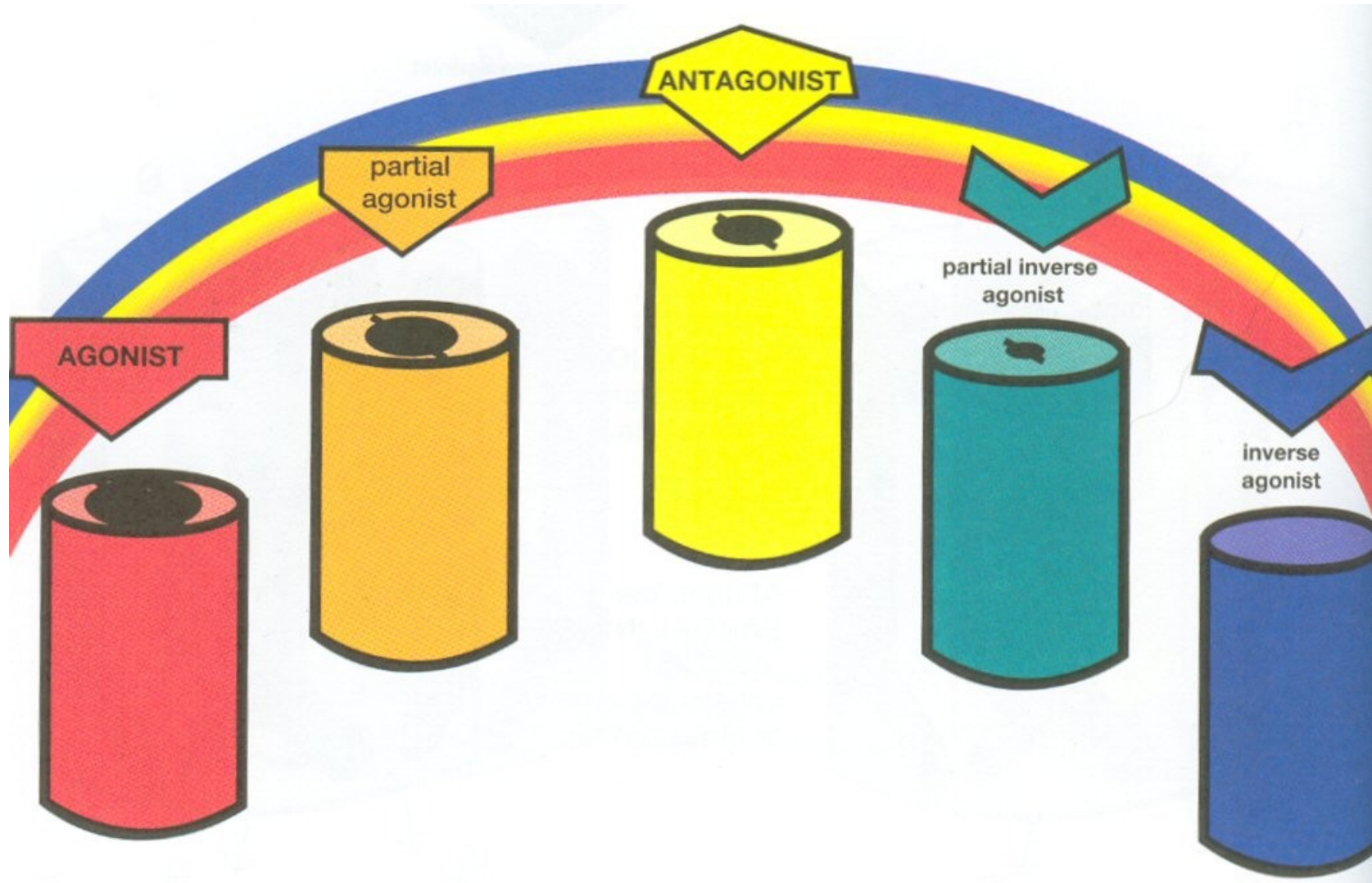


Figure 2-1. Relations between drug concentration and drug effect (A), or receptor-bound drug (B). The drug concentrations at which effect or receptor occupancy is half-maximal are denoted EC_{50} and K_D , respectively.

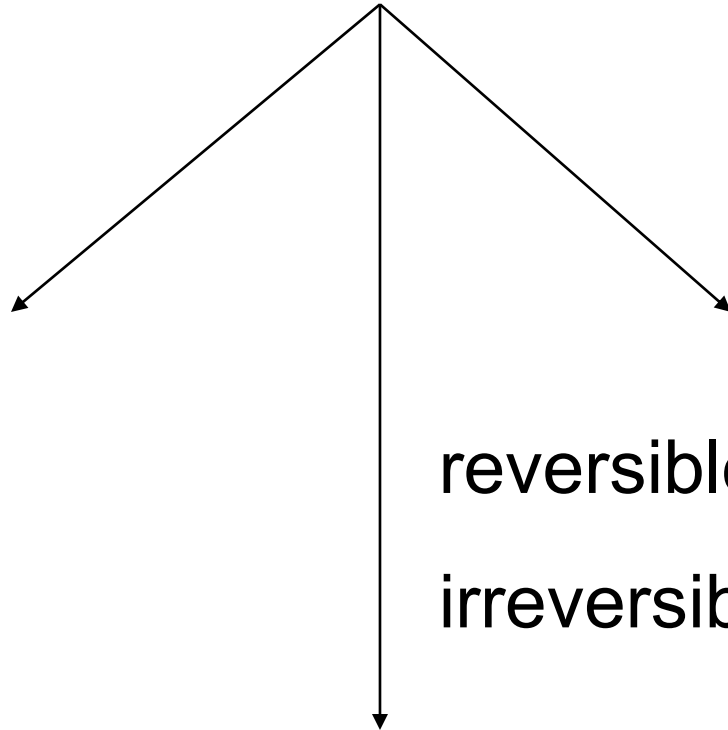
Spectrum of ligands



Antagonism



competitive
non-competitive



reversible
irreversible

at the receptor level
at the function level



Competitive

- ✓ ligands compete for the same binding site
- ✓ \uparrow c of antagonist decreases agonist effect and inversely
- ✓ the presence of antagonist increases the amounts of agonist needed to evoke the effect

Non-competitive

- ✓ allosteric antagonism
- ✓ irreversible bounds
- ✓ \uparrow c of agonist does not interrupt the effect of antagonist

M U N I
M E D



Regulation of receptor function

Regulation of receptor sensitivity and counts



Receptor desensitization

- reducing the sensitivity of the receptors after repeated agonist exposure

- **Tachyphylaxis** – acute drug „tolerance“
 - reduced sensitivity to the active substance evolving quickly (minutes) → distortion of the signal cascade
 - the reactivity of the organism returns to the original intensity after the elimination of the substance
 - Ex. of tachyphylaxis – nitrates administration, ephedrine

- **Tolerance** – reduced sensitivity to the active substance, arising from the repeated administration of the drug (days – weeks) → down-regulation, internalization of the receptors
 - to achieve the original effect required increasingly higher doses of drug
 - the original reactivity of the organism returns to a certain period of time after discontinuation of the drug
 - Ex. of tolerance – opioids administration

M U N I M E D

Regulation of receptor sensitivity and counts



Hypersensitivity

✓ increase of receptor sensitivity/counts after **chronic antagonist** exposure

Rebound phenomenon

after discontinuation of long-term administered drugs return to its original state or ↑ intensity of the original condition (hypersensitivity of receptors to endogenous ligands → up-regulation)

Example: chronic administration of β blockers



B. Non-receptor mechanism of action

Interaction with „non-receptor“ proteins

- 1. enzyme inhibition
- 2. block of ion channels
- 3. block of transporters

„non-proteins“

- binding to cellular components (ATB-ribosomes, hydroxyapatit, tubulin etc.)



1. Enzyme inhibition

Competitive or non-competitive enzyme inhibitors

- reversible
 - acetylcholinesterase – physostigmine
 - phosphodiesterase – methylxantine
- irreversible:
 - cyklooxygenaze – ASA (aspirin)
 - MAO-B – selegilin
 - aldehyddehydrogenaze – disulfiram



2. Ion channels

- Calcium channel blockers (nifedipin, isradipin...)
- Potassium channel blockers (flupirtin – selective neuronal potassium channel modulator, oral antidiabetics...)
- Natrium channel blockers – local anesthetics

3. “Carriers”



- Proton pump inhibitors (PPIs) – omeprazol
- Na^+/K^+ ATPase inhibitors – digoxin

