

# PHARMACODYNAMICS

#### **Copyright notice**

The presentation is copyrighted work created by employees of Masaryk university. Students are allowed to make copies for learning purposes only. Any unauthorised reproduction or distribution of the

presentation or individual slidesis against the law.

#### **PHARMACOLOGY**





deals with the mechanism of action (e.g. receptor sites, molecular level of action..)

"How does it work"



## **Pharmacodynamics**

## (how drugs work on the body)

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

□ Main targets – cellular, molecular, genetic level...

- Therapeutic effects
- Adverse effects









 $M \vdash 1$ 

## 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)



Low Sugar Concentration High Sugar Concentration High Water Concentration Low Water Concentration



## **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 quarternary ammonium salts) used primarily as antiseptics.
- some antibiotics (e.g. polymyxins basic peptides) act as cationic detergents and disrupt phospholipids in bacterial membranes.



 $M \vdash 1$ 

# 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**



 $M \vdash D$ 

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)

MUNI Med





## A. Receptor – effector system



= complex of processes

extracelullar 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
- Iigand (signal molecule) = molecule able to bind to specific receptor
  - endogenous neurotransmitters, hormones
  - exogenous xenobiotics, drugs

MUNI Med

## **Receptor classification**



Localization	Transduction	Ligands
✓ membrane	✓ metabotropic	✓ achol
✓ cytoplasm	✓ ion. channels	✓ amines
✓ organels	✓ kinase	✓ AMA
✓ auto/heterore	e ✓ DNA	✓ peptides
ceptors	regulating	

#### **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
Efector	lon 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

Rang and Dale Pharmacology, 2012



## **Receptor – effector system**

## □ Affinity

✓ the ability of the ligand to bind to the receptor

#### □ Instrinsic activity

✓ ability to evoke an effect after binding to receptor

Image: Image:

## **Receptor – effector system**





Ligand classification (intrinsic activity) AGONISTS



## **Full agonist**

- IA = 1

#### Partial agonist

- dualist
- IA in a range from o‹ to ›1



## Ligand classification



## Antagonists

✓ IA = 0



#### **Inverse agonist**

✓ IA = -1



 ✓ Stabilizes the receptor in the const activity

## **Receptor-effector system**



#### **Relation between dose and effect**



## **Spectrum of ligands**







# Antagonism

competitive

non-competitive

reversible

irreversible

at the receptor level at the function level

## MUNI MED co

## Antagonism



#### Competitive

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

#### Non-competitive

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



## **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"

- $\Box$  reduced sensitivity to the active substance evolving quickly (minutes)  $\rightarrow$  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

□ <u>Tolerance</u> – reduced sensitivity to the active substance, arising from the repeated

administration of the drug (days – weeks)  $\rightarrow$  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
- $\Box$  Ex. of tolerance opioids administration



✓ incerase of receptor sensitivity/counts after chronic
 anatagonist exposure

#### Rebound phenomenom

after discontinuation of long-term administered drugs return to its original state or  $\uparrow$  intensity of the original condition (hypersensitivity of receptors to endogenous ligands  $\rightarrow$  upregulation)

Example: chronic administration of  $\beta$  blockers



 $N/I \vdash I$ 

## **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
  - acetylcholinesteraze physostigmine
  - phosphodiesteraze 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<sup>+</sup>/K<sup>+</sup> ATPasa inhibitors digoxin

