

BIOMARKERS AND TOXICITY MECHANISMS 04 – Mechanisms @membranes

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Tento projekt je spolufinancován Evropským sociálním fondem a státním rozpočtem České republiky.









INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

Major mechanisms (modes of action) to be discussed in detail

- Proteins and inhibition of enzymatic activities
 Mitotic poisons & microtubule toxicity
- Membrane nonspecific toxicity (narcosis)
- Toxicity to membrane gradients
- DNA toxicity (genotoxicity)
- Complex mechanisms
 - Detoxificiation
 - defence processes as toxicity mechanisms
 - Oxidative stress redox toxicity
 - Toxicity to signal transduction
 - Ligand competition receptor mediated toxicity



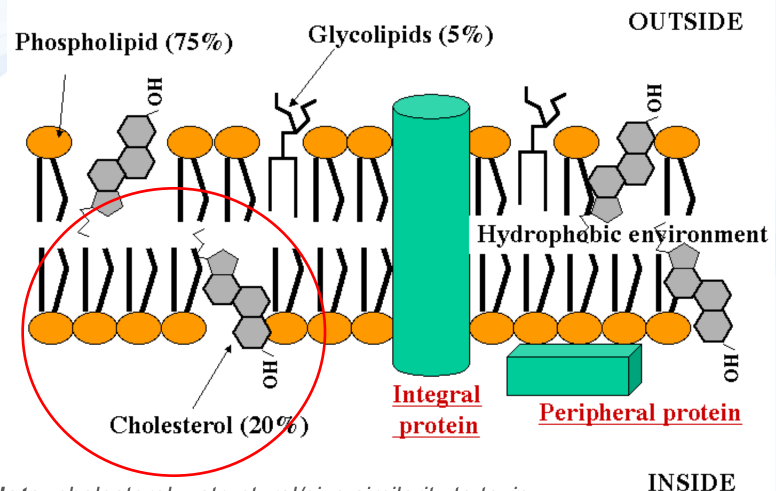
Cell membrane

Key functions for life

- Primary barrier / separation of "living" inside from "abiotic" outside
- Semipermeability for nutrients / signals
- Reception of chemical signals & regulatory molecules
- Keeping gradients necessary for life
 - H+ ATP synthesis(mitochondria / bacterial emambrane)
 - K+/Na+ neuronal signals
- Proteosynthesis (ribosomes) depends on membranes
- Many other enzymes bound to membranes (e.g. signaling, detoxification, post-translational modifications)
- Etc....

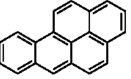


Plasma membrane



Note: cholesterol – structural/size similarity to toxic organics e.g. Benzo[a]pyrene





Nonspecific (basal, narcotic) toxicity

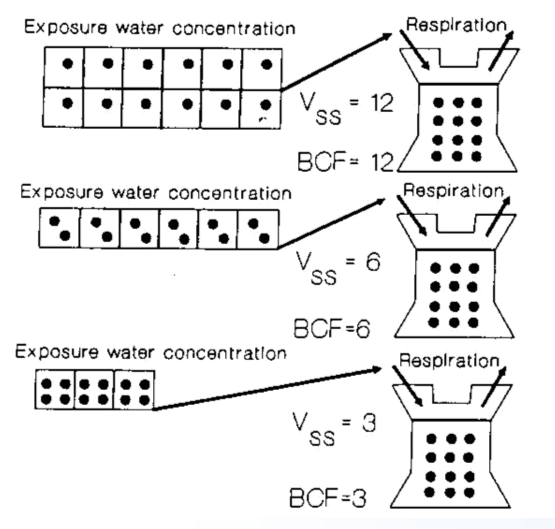
- All <u>organic</u> compounds tend to accumulate in membranes, being "narcotic" at relatively "high" concentrations
- Compounds then affect membranes
 → nonspecific disruption of fluidity
 → and/or disruption of membrane proteins
- Related to lipophilicity (Kow): tendency of compounds to accumulate in body lipids (incl. membranes)

E.g. narcotic toxicity to fish: log (1/LC50) = 0.907 . log Kow - 4.94

- The toxic effects occur at the same "molar volume" of all narcotic compounds (volume of distribution principle)



Volume of distribution principle



Centrum pro výzkun toxických látek v prostředí BCF – bioconcentration factor * Depends on hydrophobicity (i.e. Kow)

* Higher BCF

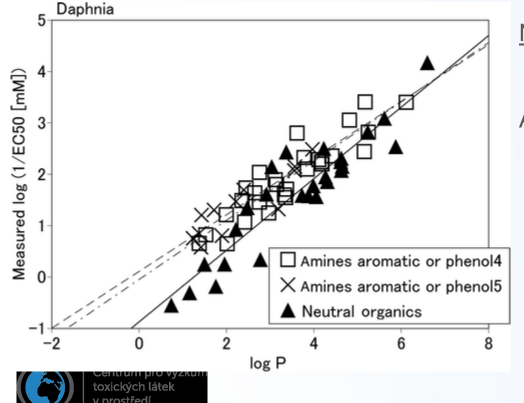
→ lower concentration is
 sufficient for bioconcentration
 to the same "tissue concentration"
 → lower external concentration
 (IC50) will induce toxic effect

* Confirmed by chemical analyses (same molar concentrations of different compounds accumulated in membranes)

Narcotic toxicity in ecotoxicology

Acute basal toxicity

Direct correlations between logKow (=logP) and EC50 for aquatic organisms (e.g. *Daphnia magna*)



Example:

Neutral organics → Nonpolar narcosis

Amines, phenols

→ Polar narcosis

(similar logP → higher toxicity, i.e. higher Values of 1/EC50 in comparison to neutral organics)

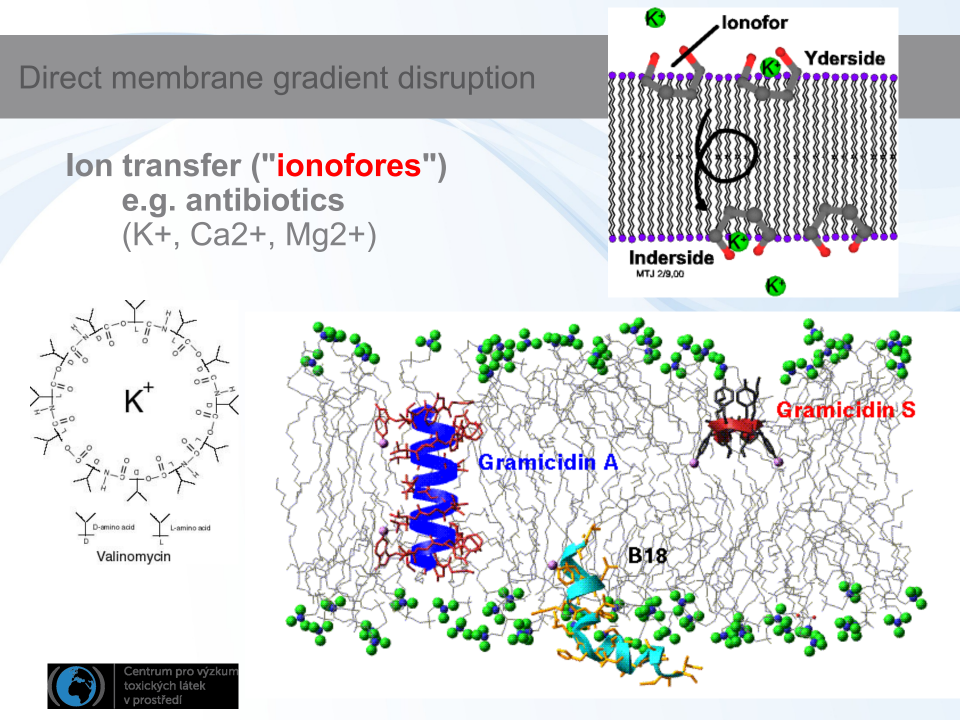
→ More specific ... In addition to membrane accumulation, direct interactions with proteins are anticipated

Toxicity to membrane gradients and transport

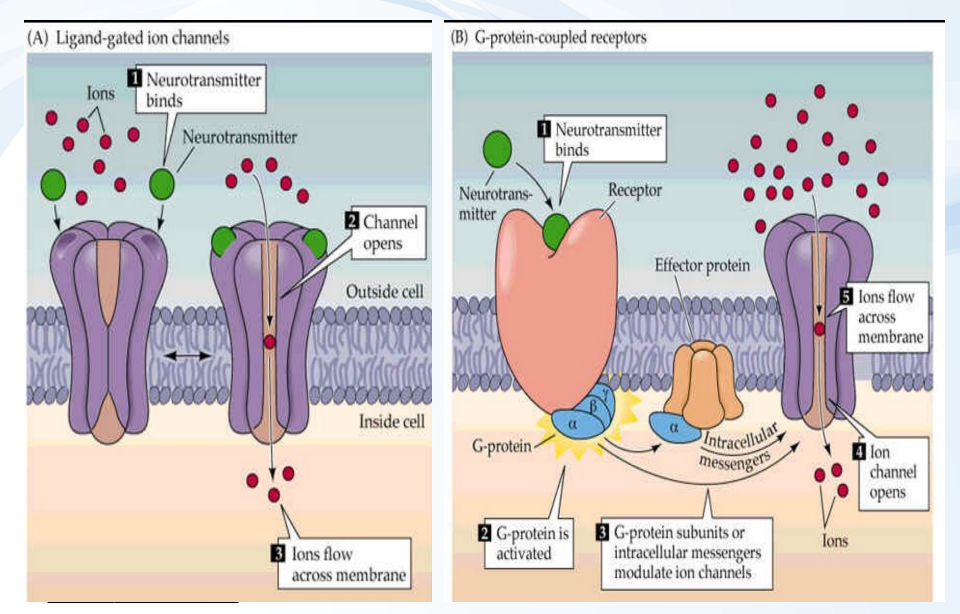
- Semipermeability of membranes and key functions

 - cytoplasmic membrane: signalling, neural cells Na+/K+ gradient
 - mitochondrial membrane: electrone flow → ATP synthesis
 - endoplasmatic reticulum Ca²⁺ signalling

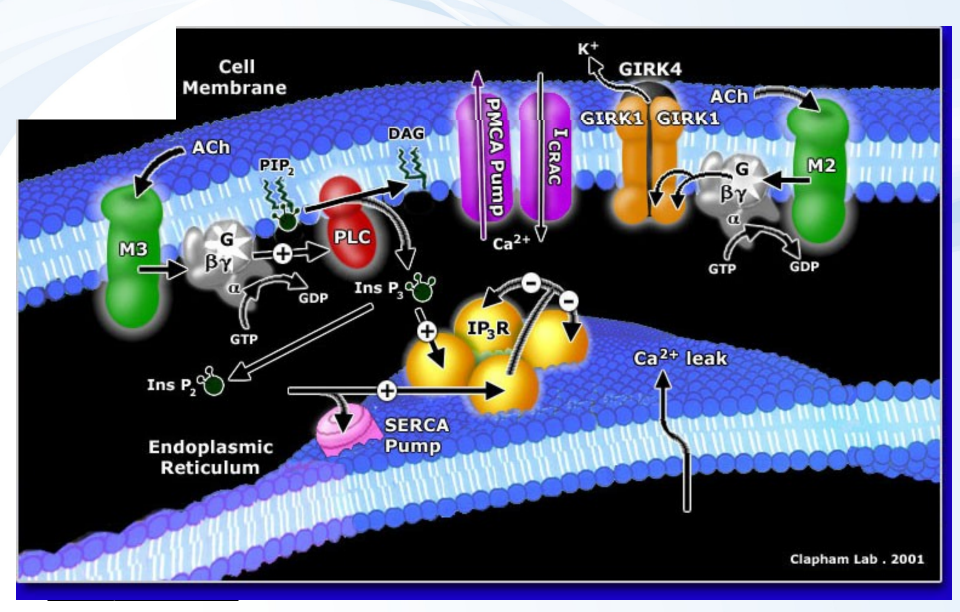




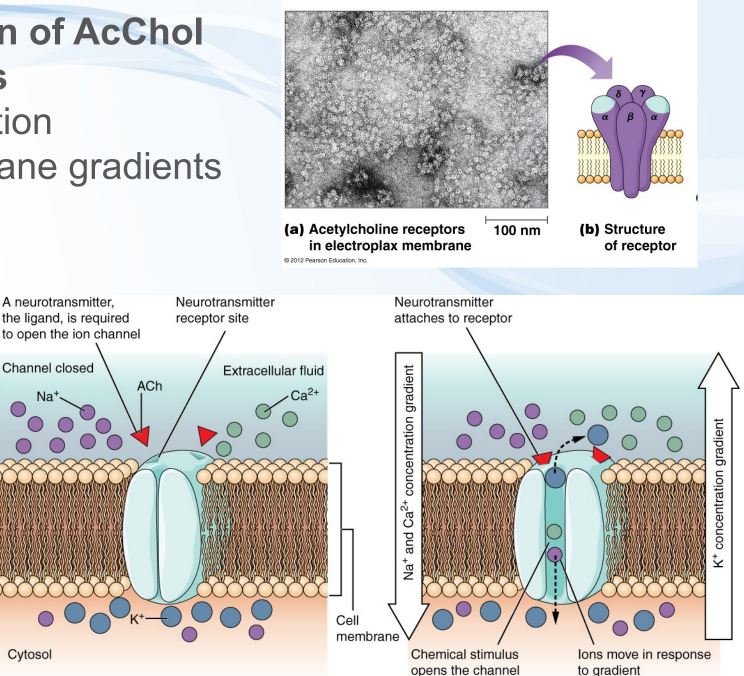
Principal types of channel activation



Various membrane channels - examples

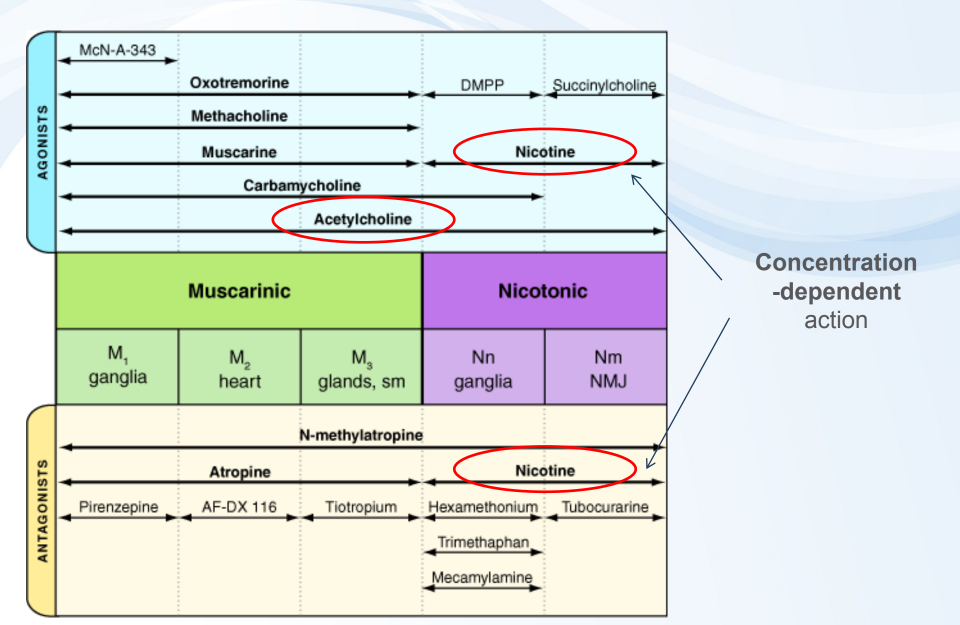


Activation of AcChol receptors → Disruption of membrane gradients



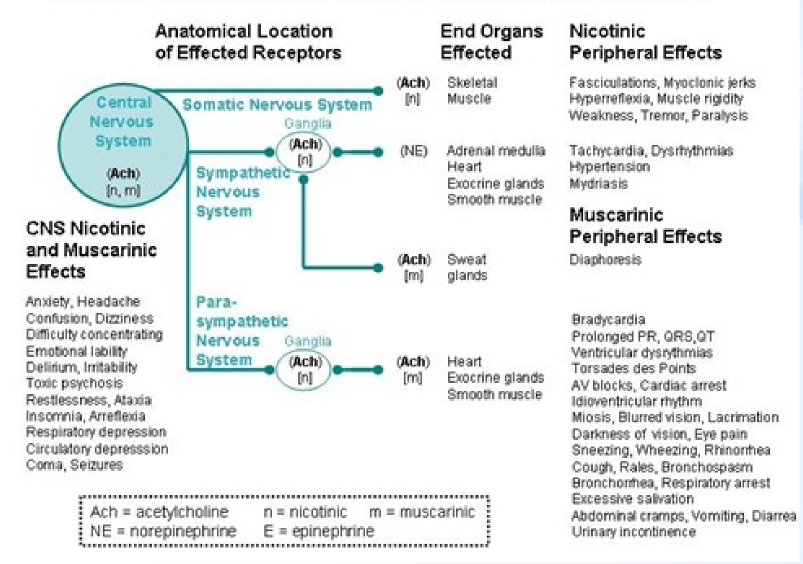


Activation / inhibition of ligand-gated channels

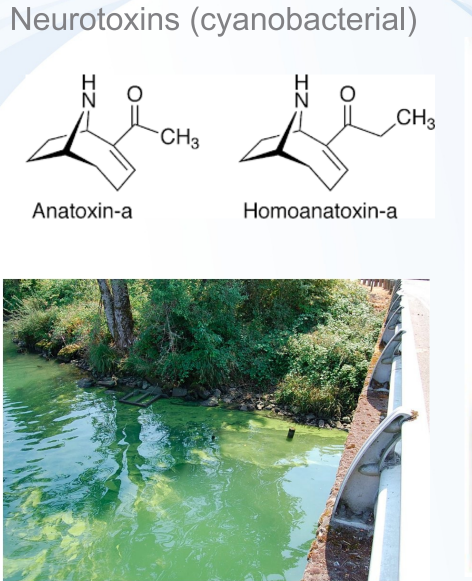


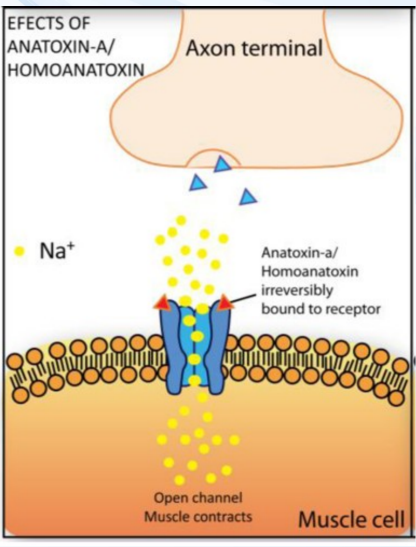
Activation / inhibition of ligand-gated channels

Nicotinic and Muscarinic Effects of Cholinesterase Inhibitors



Environmentally relevant ion channel activators



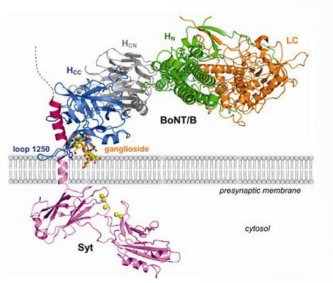


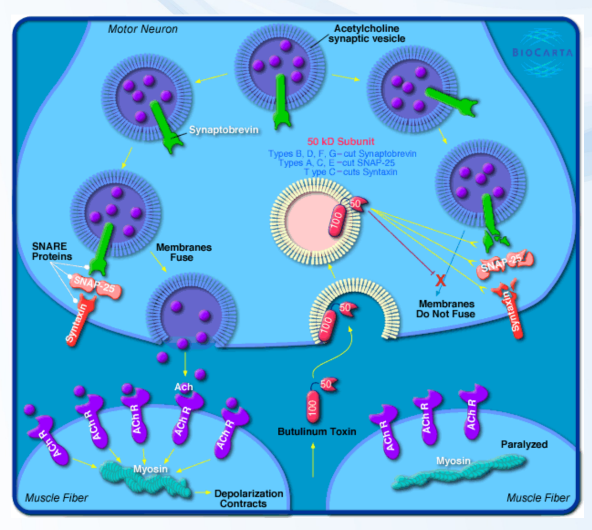
Botulinum and Tetanus toxins (Clostridium botulinum, Clostridium tetani)

Toxins = enzymes - proteases (!)

direct cleavage
of proteins involved
in vesicle formation
selective inhibition of
neutrotransmitter release

BOTULINISM → neurotoxicity (paralysis)





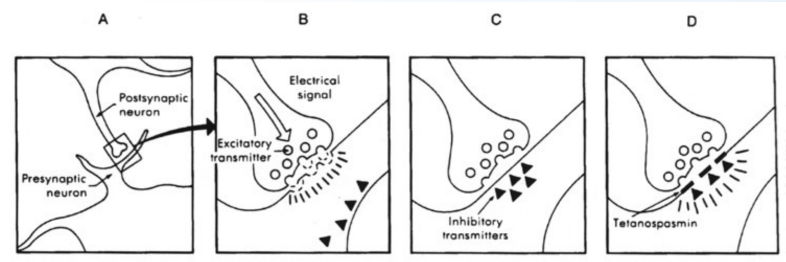
Botulinum and Tetanus toxins (Clostridium botulinum, Clostridium tetani)

TETANUS TOXIN (tetanospasmin)

blocks release of INHIBITORY NEUROTRANSMITERS (γ-aminobutyric acid (GABA) in CNS

→ neurotoxicity – permanent contraction





Gradient of $H+ \rightarrow ATP$ generation & its disruption

