

BIOMARKERS AND TOXICITY MECHANISMS 04 – Mechanisms @membranes

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









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



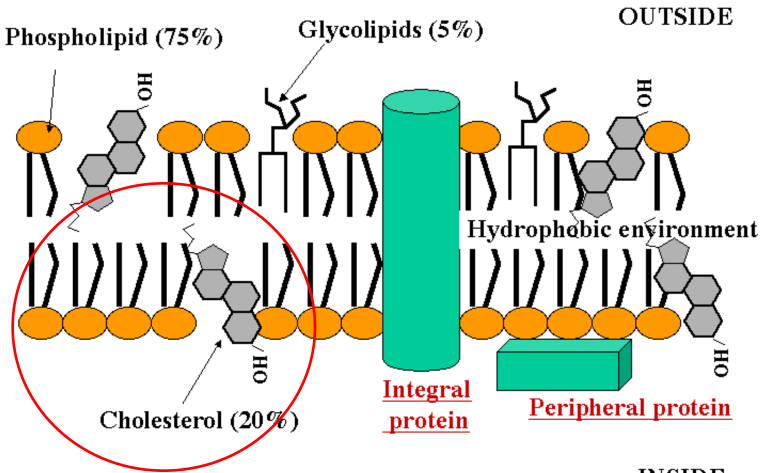
Cell membrane

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



INSIDE

Nonspecific (basal, narcotic) toxicity

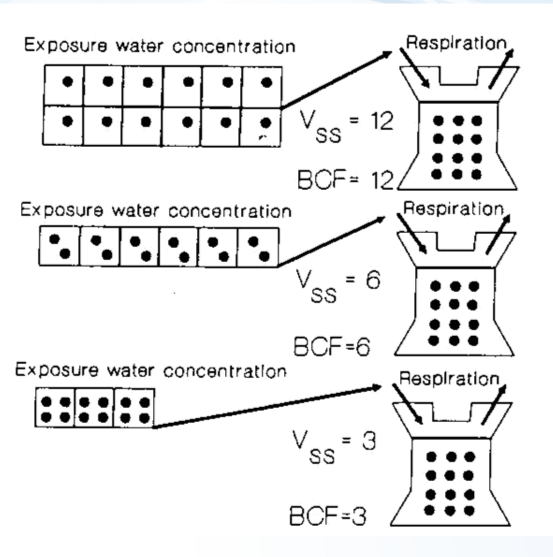
- All <u>organic</u> compounds tend to accumulate in membranes, being "narcotic" at relatively "high" concentrations
- Compounds then affect membranes
 - → nonspecific disruption of fluidity
 - → and/or disruption of membrane proteins
- Related to lipophilicity (Kow): tendency of compounds to accumulate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

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

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



Volume of distribution principle



BCF – bioconcentration factor

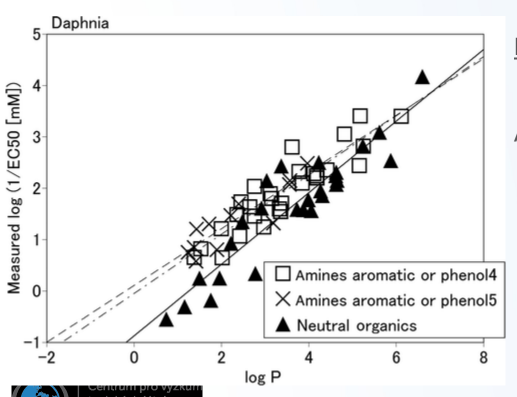
- * Depends on hydrophobicity (i.e. Kow)
- * Higher BCF
 - → lower concentration is sufficient for bioconcentration to the same "tissue concentration"
 → lower external concentration (IC50) will induce toxic effect
- * Confirmed by chemical analyses (same molar concentrations of different compounds accumulated 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

Disruption of membrane gradients



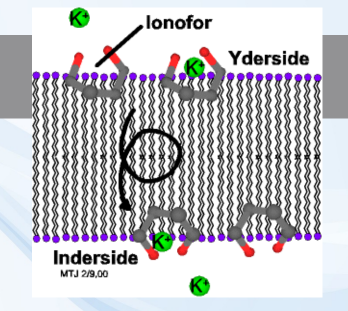
Toxicity to membrane gradients and transport

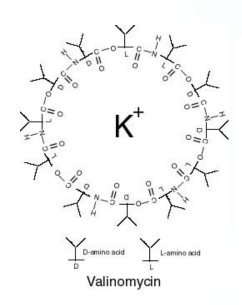
- Semipermeability is essential for membranes key functions
 - cytoplasmic membrane: signalling, neural cells Na+/K+ gradient
 - mitochondrial membrane: electrone flow → ATP synthesis
 - endoplasmatic reticulum Ca²⁺ signalling
- Disruptions can be either through nonspecific narcotic toxicity (above) or via specific effects of toxicants → discussion further

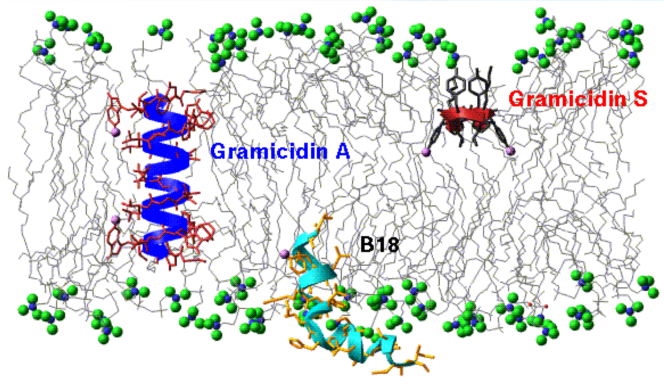


Direct membrane gradient disruption

Ion transfer ("ionofores")
e.g. antibiotics
(K+, Ca2+, Mg2+)





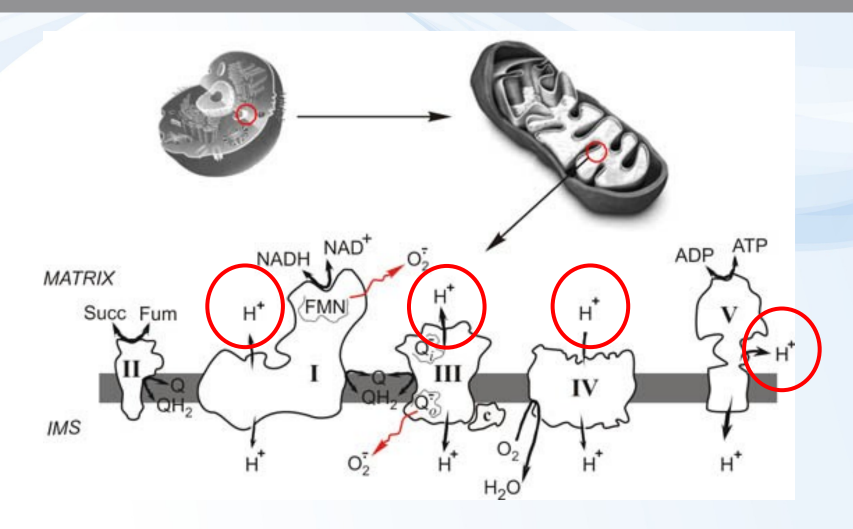




Mitochondrial membrane

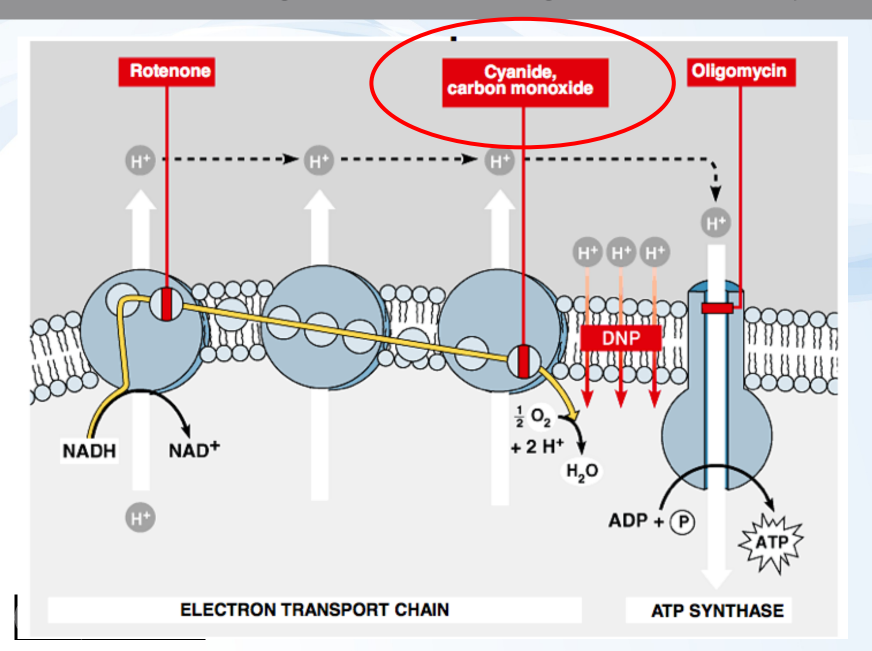


Mitochondria (= energy metabolism!) -- membrane processes: H+ formation --





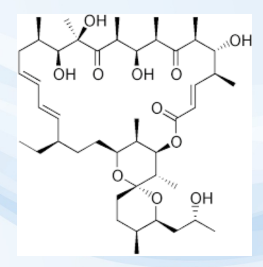
Role of membrane: gradient of H+ → ATP generation & its disruption



Gradient of H+ → ATP generation & its disruption

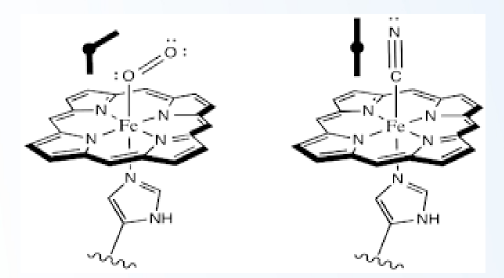
Rotenone

Oligomycin



CO (carbon monoxide)
CN (cyanide)

→ Binding to haem structures

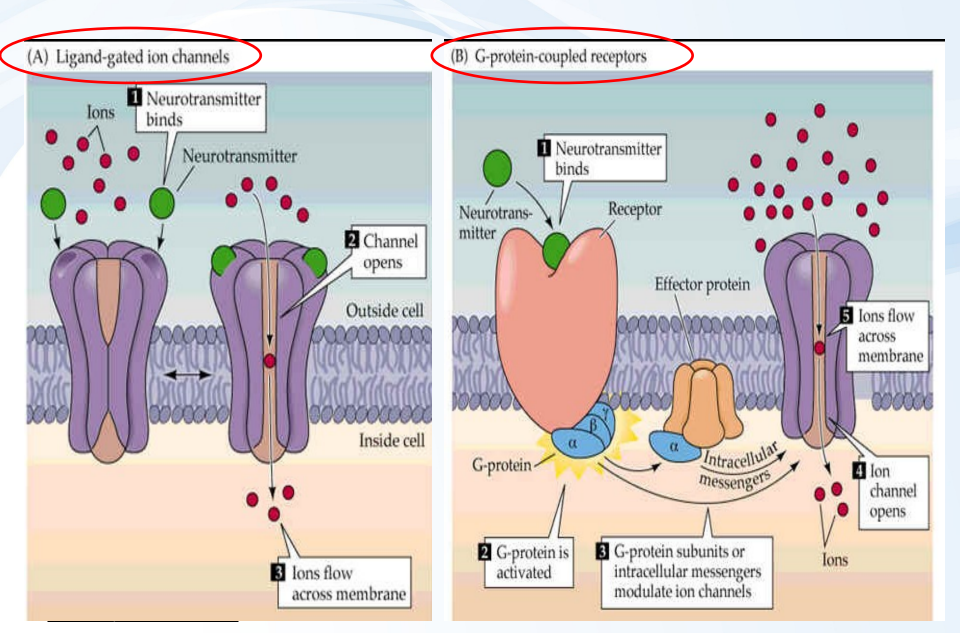




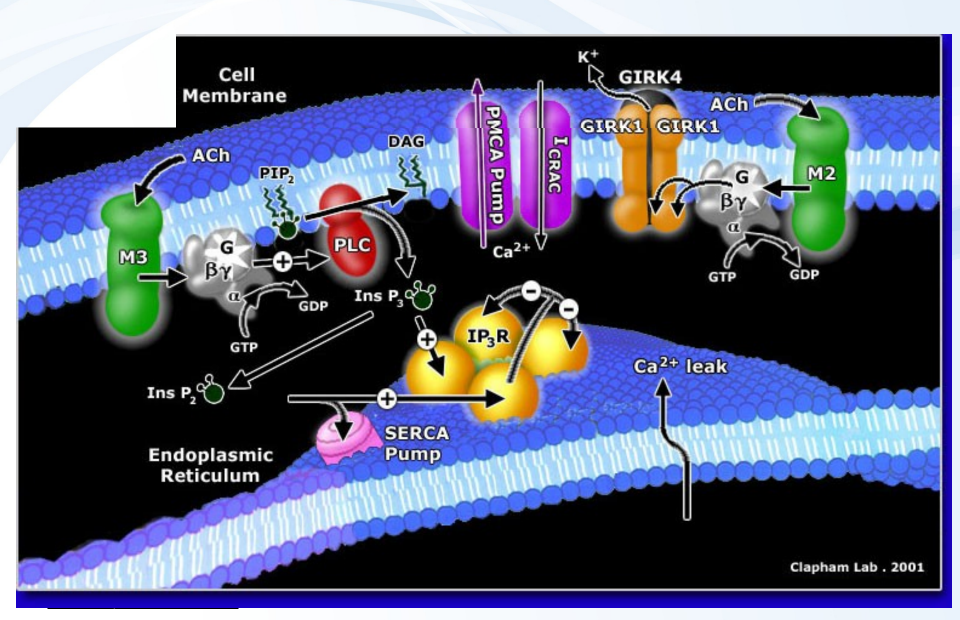
Receptors/Channels & membrane gradients



Principal types of channel activation

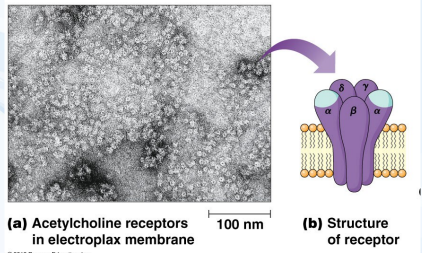


Various membrane channels - examples



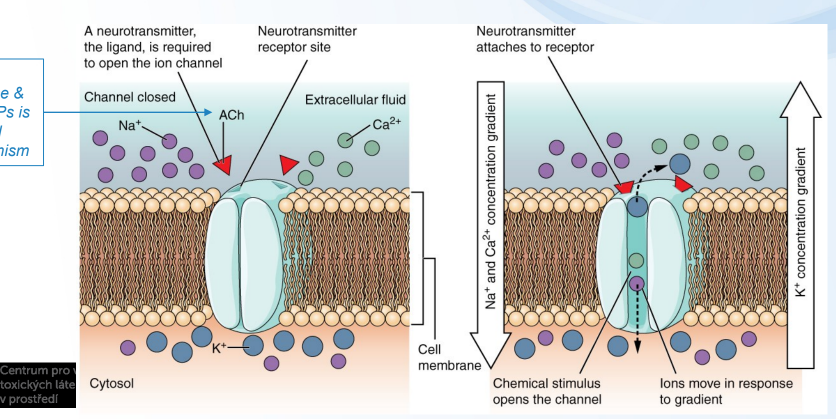
Activation of AcChol receptors

→ Disruption of membrane gradients

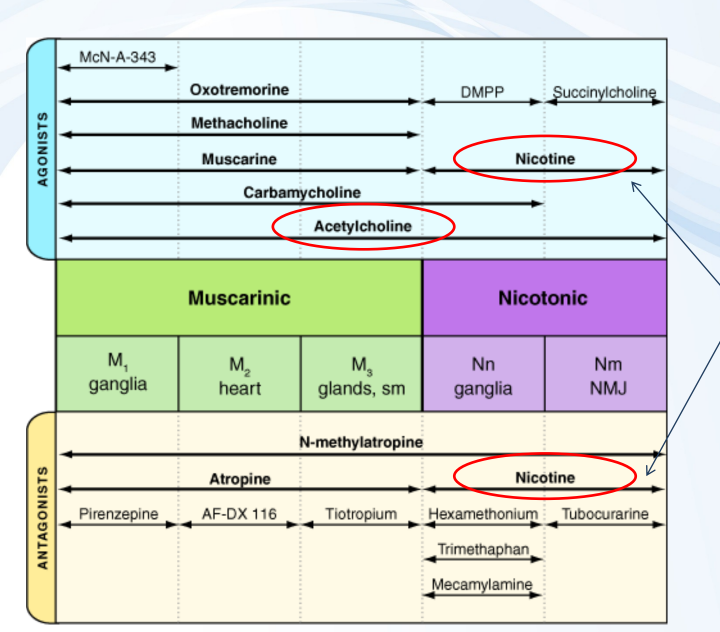


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Reminder: AcCholEsterase & inhibition by OPs is another related toxicity mechanism

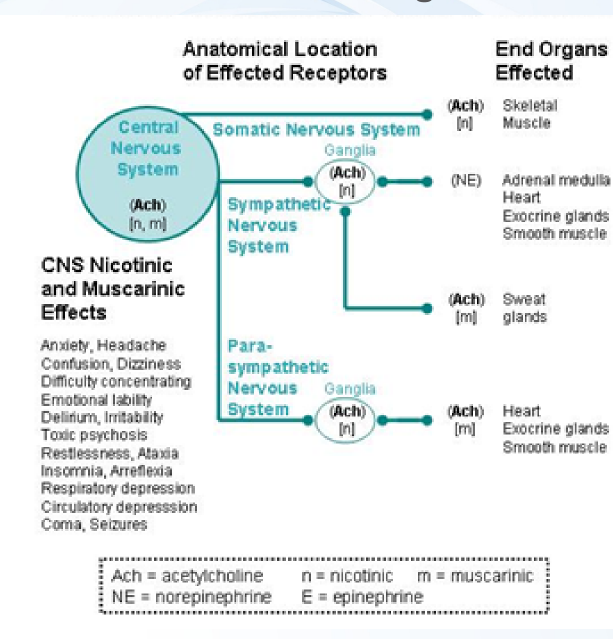


Activation / inhibition of ligand-gated channels



Agonist vs Antagonist
Concentration-dependent
action

EXAMPLE: related biological effects



Nicotinic Peripheral Effects

Fasciculations, Myoclonic jerks Hyperreflexia, Muscle rigidity Weakness, Tremor, Paralysis

Tachycardia, Dysrhythmias Hypertension Mydriasis

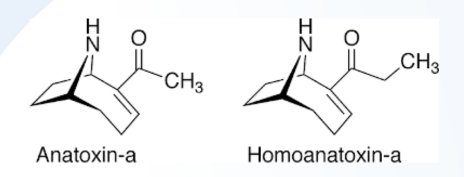
Muscarinic Peripheral Effects

Diaphoresis

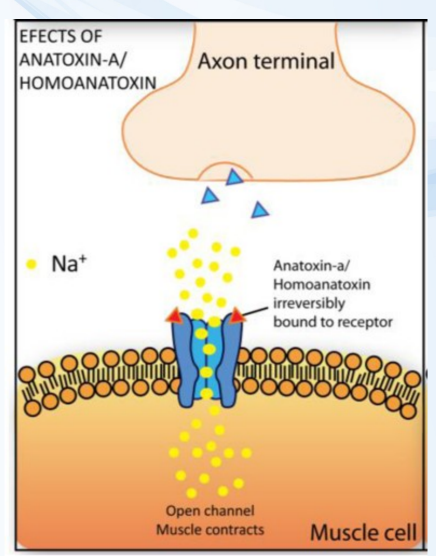
Bradycardia
Prolonged PR, QRS,QT
Ventricular dysrythmias
Torsades des Points
AV blocks, Cardiac arrest
Idioventricular rhythm
Miosis, Blurred vision, Lacrimation
Darkness of vision, Eye pain
Sneezing, Wheezing, Rhinorrhea
Cough, Rales, Bronchospasm
Bronchorrhea, Respiratory arrest
Excessive salivation
Abdominal cramps, Vomiting, Diarrea
Urinary incontinence

Environmentally relevant toxins - ion channel activators

Neurotoxins (cyanobacterial)



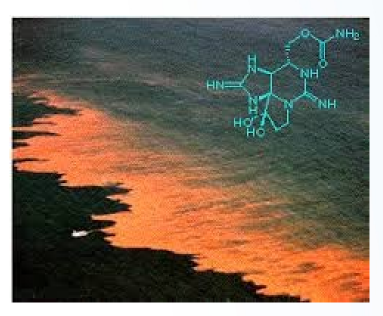


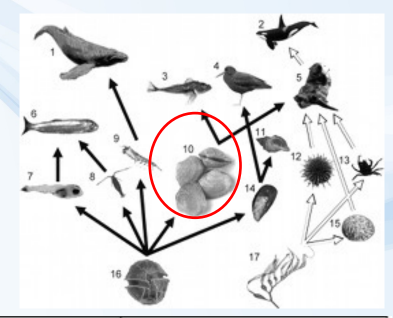


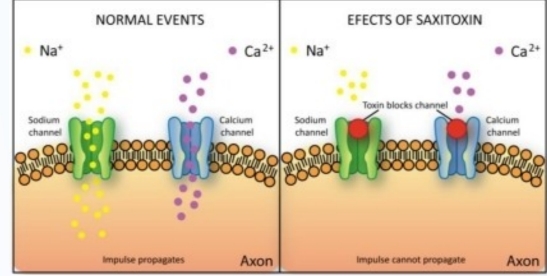
Environmentally relevant toxins - ion channel activators

SAXITOXINS

- Produced by dinoflagelates and cyanobacteria
- (toxic blooms, "red tides")









Roles of membranes in the <u>release</u> of neurotransmitters



Botulinum and Tetanus toxins

(Clostridium botulinum, Clostridium tetani)

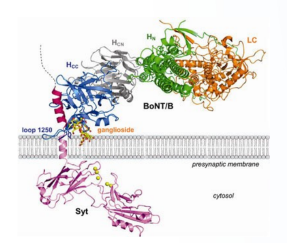
((Complex MoA - mediated through both proteins and membranes))

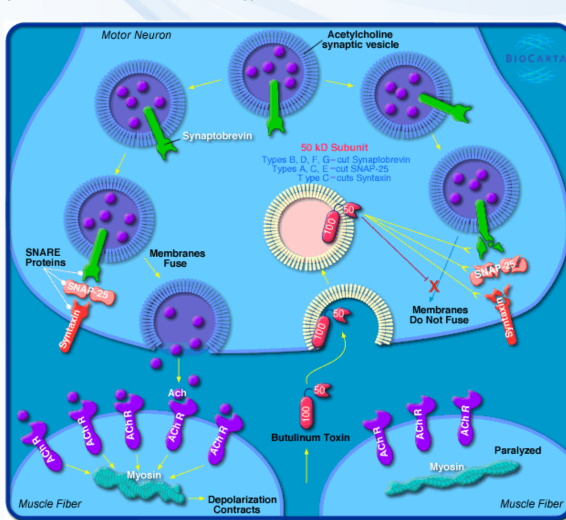
Toxins = enzymes - proteases (!)

- 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 muscle contraction



