

# BIOMARKERS AND TOXICITY MECHANISMS 08 – Mechanisms Signalling and regulation

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









### Cell communication & regulation: a target for toxicants

... especially sensitively regulated processes are highly susceptible to toxicants

→ toxicity to REGULATIONS & SIGNALLING

### Hierarchy in signalling

- **systems**: neuronal ←→ endocrine
- cell-to-cell

  hormonal & neuronal signal transmission
  contact channels
- intracellular signal transduction

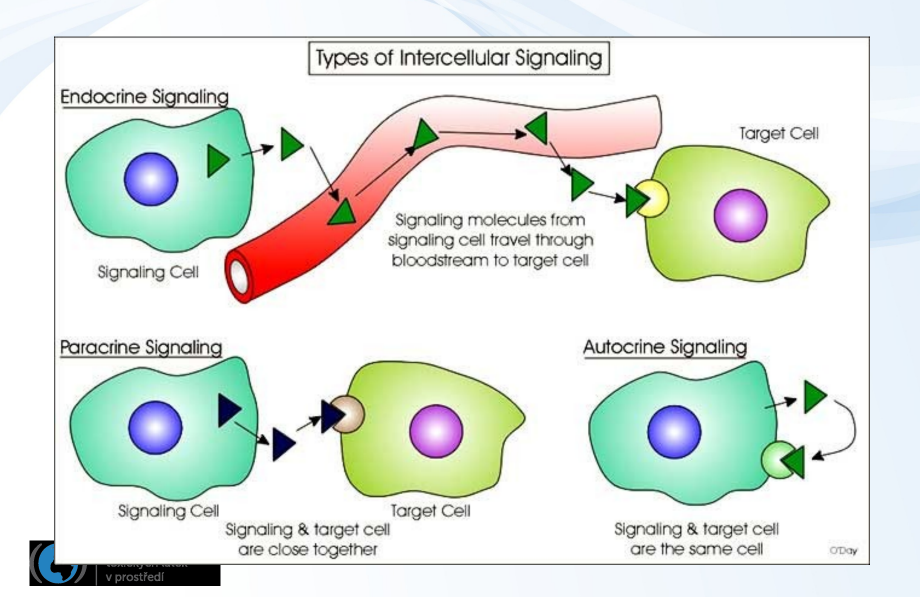


### INTER-cellular signals

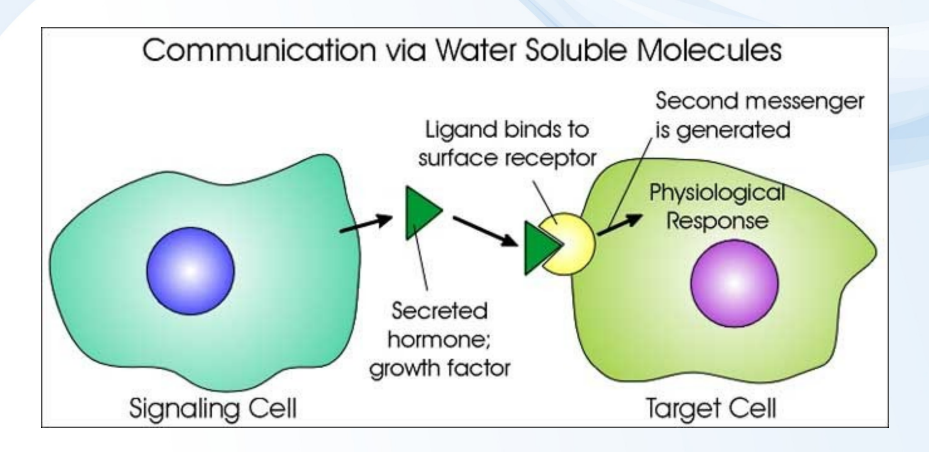
Overview



### Cell to cell communication & regulation: a target for toxicants

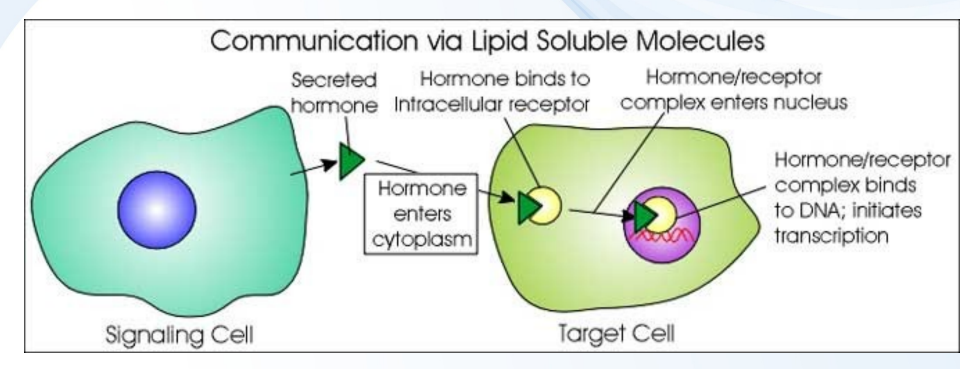


### Cell to cell communication (1)



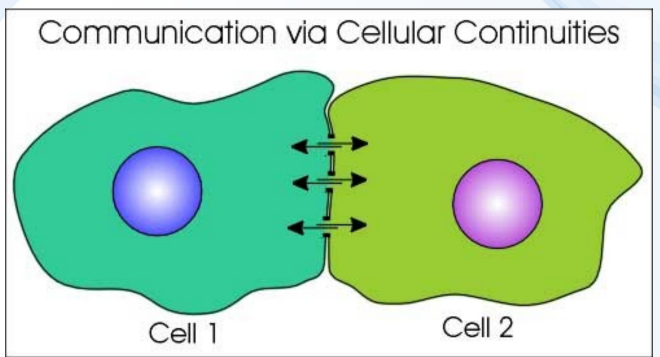


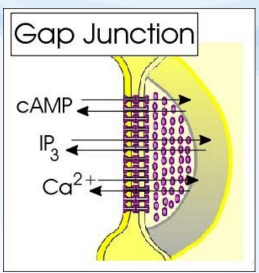
### Cell to cell communication (2)





### Cell to cell communication (3)



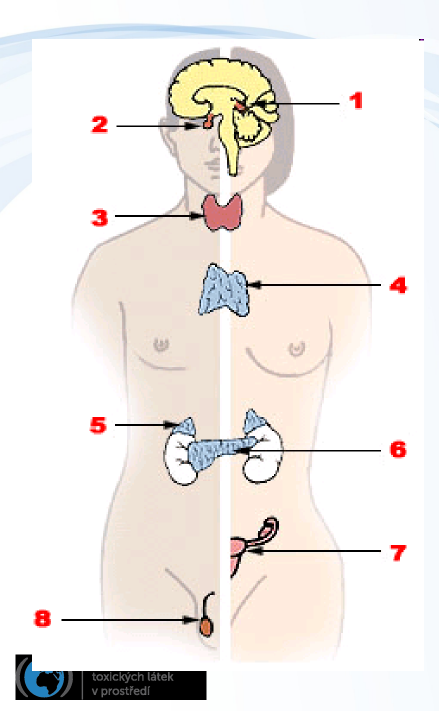




### INTER-cellular signals

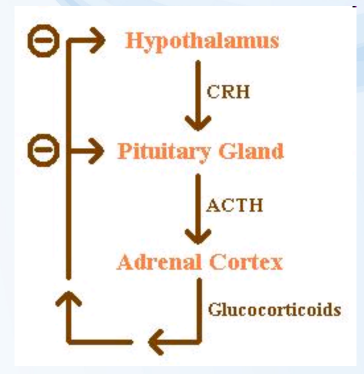
### Hormones





### **Endocrine system:**

1. Pineal gland, 2. Pituitary gland, 3. Thyroid gland, 4. Thymus, 5. Adrenal gland, 6. Pancreas, 7. Ovary, 8. Testis



**Example:** feedback loop

### **FUNCTIONS OF HORMONES**

- \* stimulation or inhibition of growth
- \* mood swings
- \* induction or suppression of apoptosis (programmed cell death)
- \* activation or inhibition of the immune system
- \* regulation of metabolism
- \* preparation for fighting, fleeing, mating ...
- \* preparation for a new phase of life (puberty, caring for offspring, and menopause)
- \* control of the reproductive cycle .... etc.



Chemicals interfering with various hormonal functions

→ diverse impacts (effects)



### System regulation = HORMONES & ENDOCRINE SYSTEM

### **FATE OF HORMONES: target for toxicants**

Toxic compounds can affect "hormone signalling" at various levels (highligted):

- 1. Biosynthesis of a particular hormone in a particular tissue
- 2. Storage and **secretion** of the hormone
- 3. **Transport** of the hormone to the target cell(s)
- 4. **Recognition of the hormone** by an associated cell membrane or intracellular receptor protein.
- 5. Relay and <u>amplification of the received hormonal signal</u> via a signal transduction process -> cellular response.
- 6. The reaction of the target cells is recognized by the original hormone-producing cells (negative feedback loop)
  - 7. **Degradation and metabolism** of the hormone

More details will be discussed in the lectures dedicated to nuclear receptors



### Toxicity to hormone regulation = ENDOCRINE DISRUPTION

### **ED & EDCs (endocrine disrupting compounds)**

= major problem in environmental toxicology

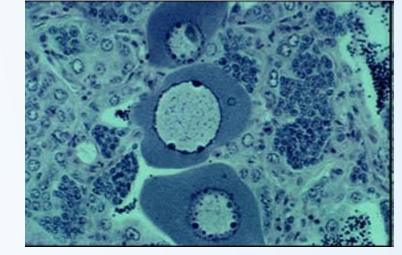
Effects at all levels of hormonal action have been demonstrated

- → synthesis, transport, site of action ....
- Multiple effects due to ED (! Not only "xenoestrogenicity" & feminization)
  - → immunotoxicity, developmental toxicity

(ED - WILL ALSO BE DISCUSSED FURTHER)

### Example of ED - Intersex roach testis

containing both oocytes and spermatozoa, caused by exposure to environmental oestrogens





### Types of hormones in vertebrates

### **Amine-derived hormones**

structure: derivatives of the amino acids tyrosine and tryptophan. Examples - catecholamines and thyroxine.

(small molecules - similar to organic toxicants → TOXIC EFFECTS)

#### Adrenalin

### Thyroxin

Norepinephrine



### Types of hormones in vertebrates

### **Peptide hormones**

structure: chains of amino acids.

- small peptides: TRH and vasopressin;
- <u>large proteins</u>: insulin, growth hormone, luteinizing hormone, follicle-stimulating hormone and thyroid-stimulating hormone etc.

Large molecules; receptors on surfaces of the cells (Interactions with toxic chemicals less likely)

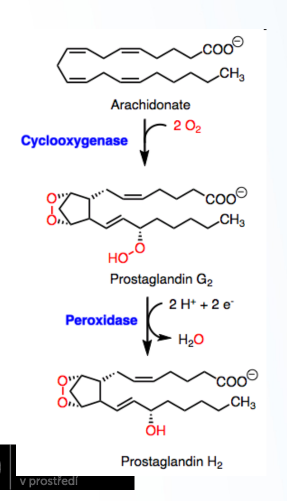
Example - insulin

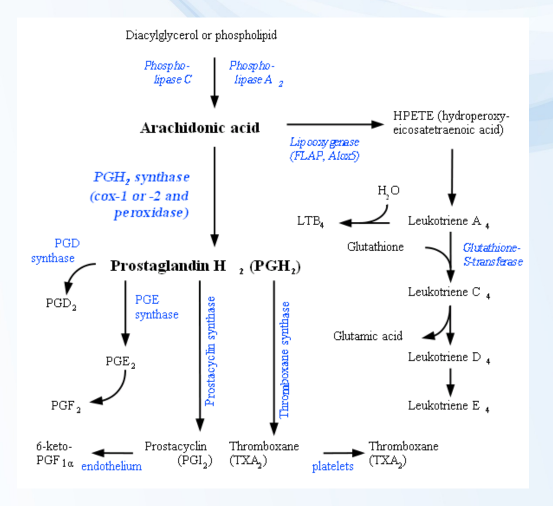




### Types of hormones (signal molecules) in vertebrates

### Lipid derived "hormones" (1) - from linoleic acid, arachidonic acid - prostaglandins





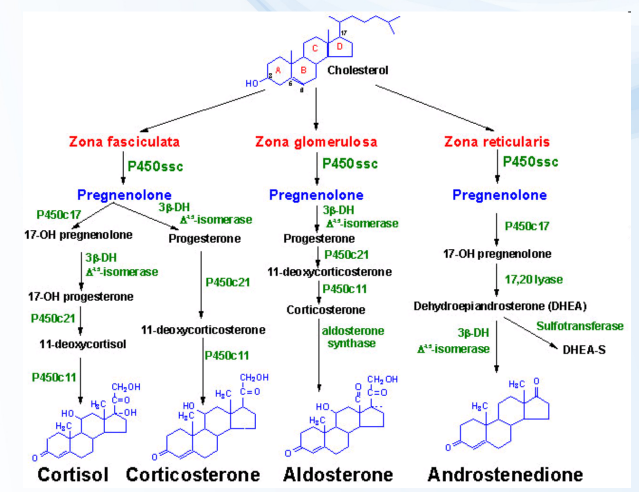
### Types of hormones in vertebrates

### Lipid derived hormones 2 - steroid hormones

- \* Small molecules similar to organic toxicants:
- → several compounds interfere with steroid hormones → toxicity !!!

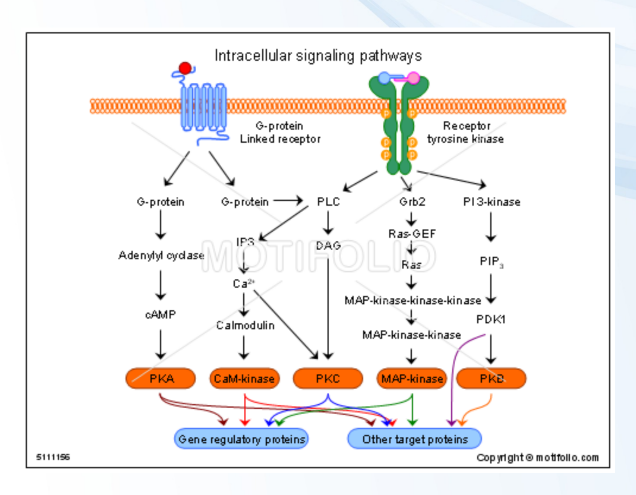
Derived from cholesterol

Examples: testosterone, cortisol, estradiol ...





### **INTRACELLULAR** signals





### Intracellular signal transduction: target of toxicants

- Regulation of cell life = control of major cell functions
  - metabolism
  - proliferation
  - differentiation
  - death (apoptosis)
- Regulation controlled by complex signalling
  - "network" of general pathways
  - similar in all cells / different cell-specific effects



### Intracellular signal transduction: target of toxicants

- Consequences of signalling disruption
  - unwanted changes in "homeostatic" rates among proliferation / differentiation / apoptosis
  - → cell transformation (carcinogenicity)
  - → embryotoxicity
  - → immunotoxicity
  - → reproduction toxicity
    - .... and other chronic types of toxicity



### Signal transduction - principles

### Two major signalling processes

protein-(de)phosphorylation

ProteinKinases - PKs, ProteinPhosphatases - PPases

- secondary messengers

cAMP / IP3, PIP2, DAG, Ca2+, AA

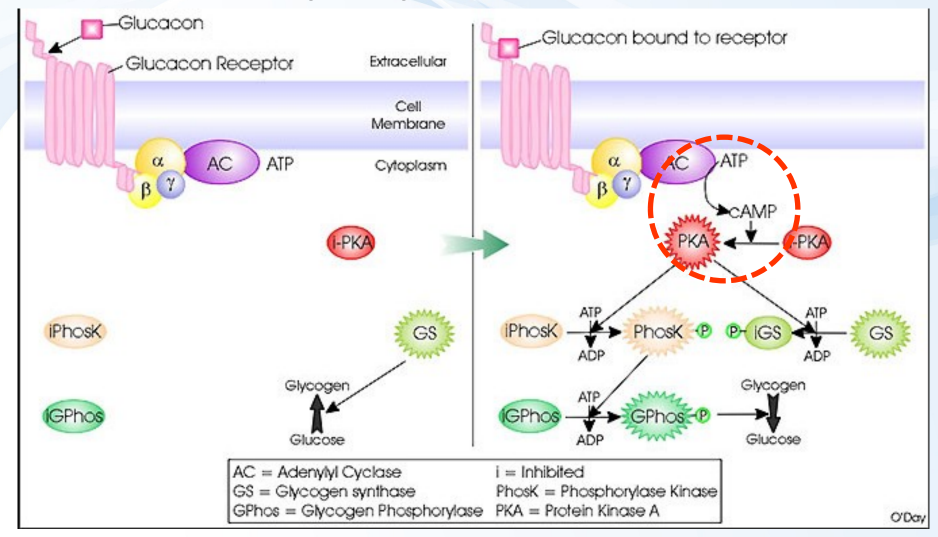
### Three major types of signalling

- 1: Membrane receptors (G-protein, kinases)
  - → activation of protein kinase A (PKA): major messenger: cAMP, <u>MAPKs</u>
- 2: Membrane receptors
  - → activation of membrane lipases → and later proteinkinase C IP3, PIP2, DAG, Ca2+, AA
- 3: Cytoplasmic (nuclear) receptors (discussed in detail in other sections)

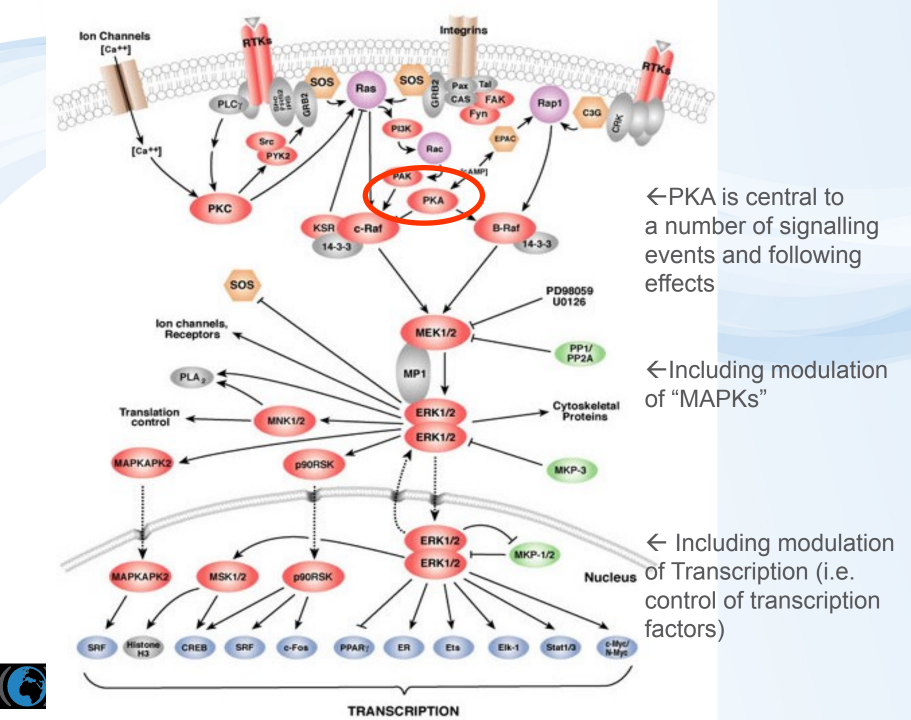


### Signalling mechanism 1

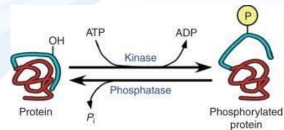
→ Activation of adenylate cyclase → cAMP → PKA



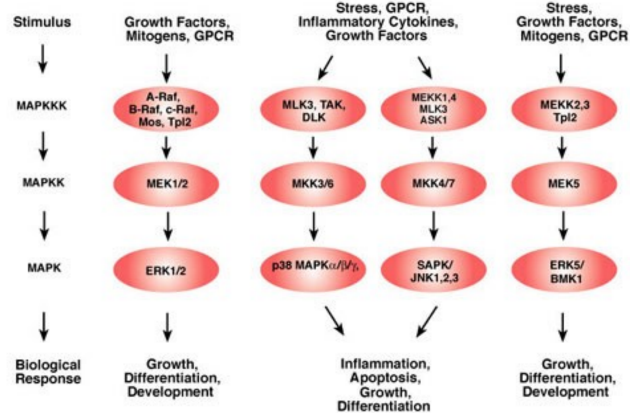




### Mitogen Activated Protein Kinases (MAPKs) & dependent effects



#### MAPK signaling cascades

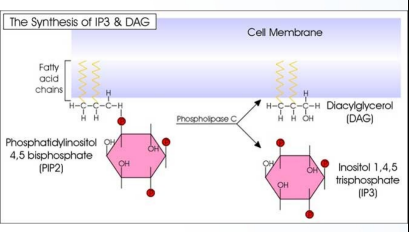


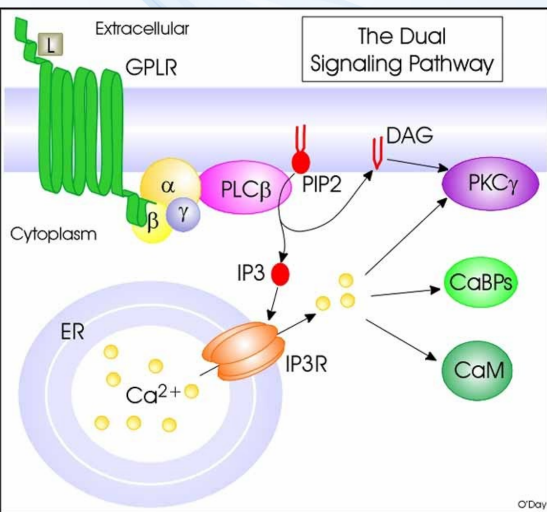


### Signalling mechanism 2

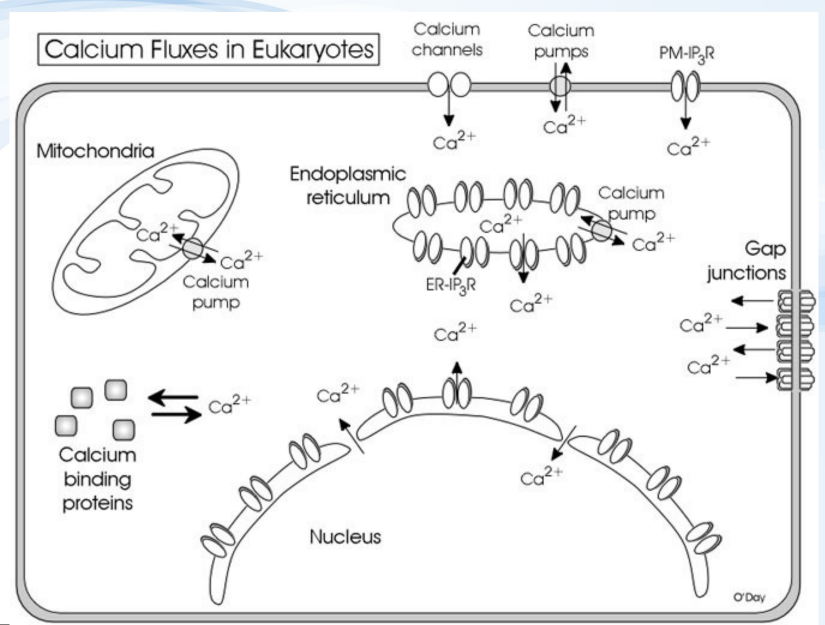
Activation of Phospholipase C

- → release of PIPs → DAG → PKC / arachidonic acid
- + IP3 → activation of Ca<sup>2+</sup> signalling











### Different signalling crosstalks → networks

Some Signaling Pathways Leading to Gene Regulation

Transcription Factors

NEAT

Nuclear Factor of Activated

T-cells

(SRF)

SerumResponseFactor

(CREB)

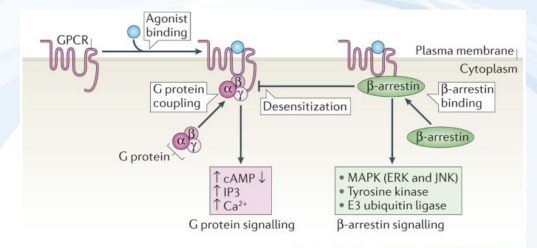
cAMP Response
 Element Binding
 protein

Ca<sup>2+</sup> MAPKs DAG CAMP CaM PKA **PKC** CaMKs I, II, etc. CREB NFAT SRF PP ] SRF)~P CREB NFAT Activation of Gene Regulation

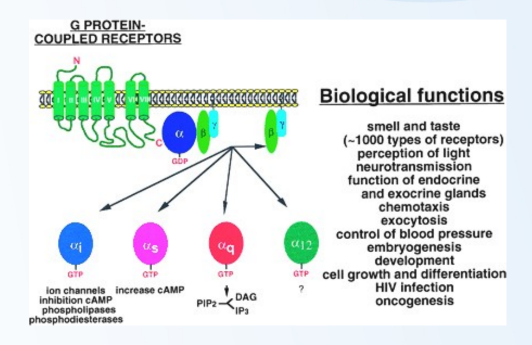
O'Day

### G-proteins & G-protein coupled receptors - GPCRs

### Involved in many functions → triggering multiple downstream events & networks



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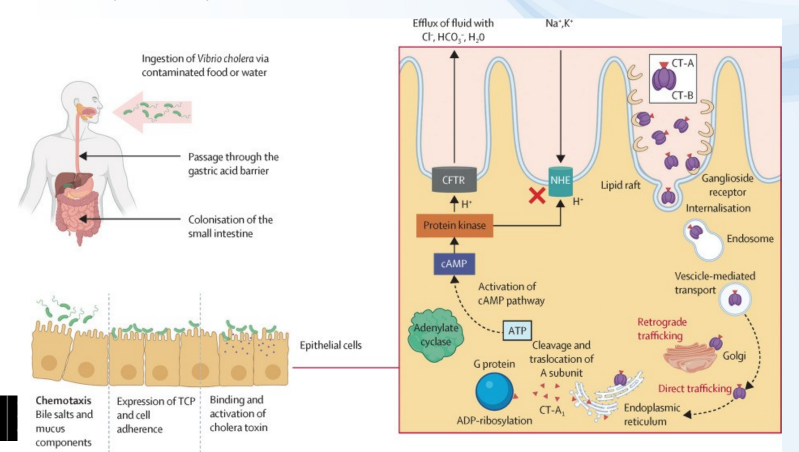


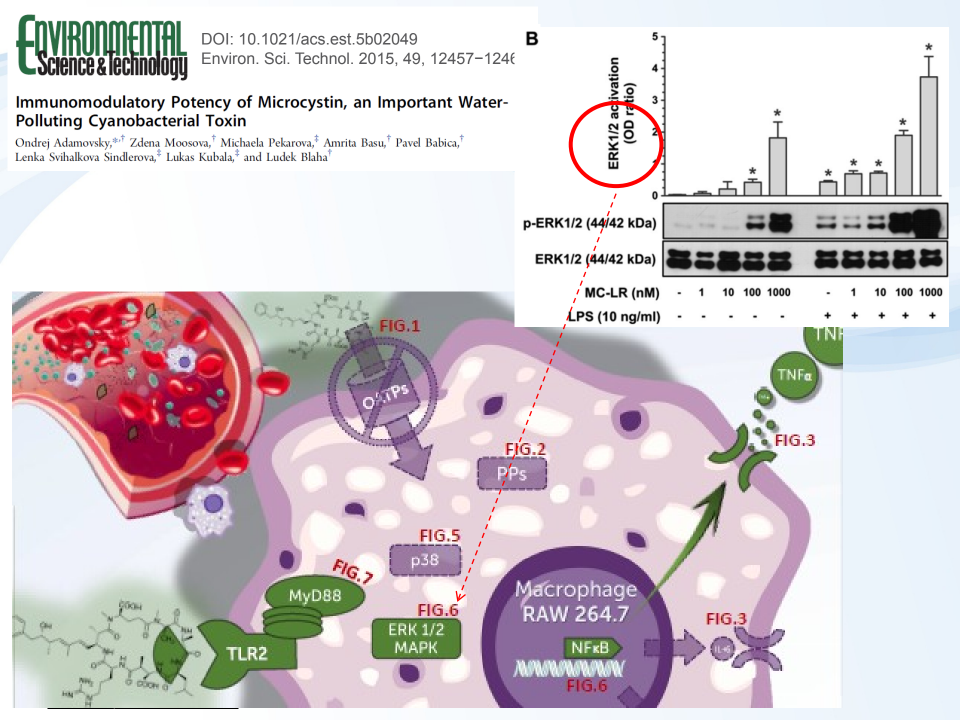
### Disruption of intracellular signaling - EXAMPLES

### Cholera toxin (from Vibrio cholerae)

### CT acts as adenylate cyclase enzyme

- → increasing cAMP levels
- → TOXICITY (diarrhea)





## Example: Lipopolysaccharides & exogenous agents inducing immune pathologies – allergies, auto-immune diseases

→ hyperactivation of intracellular signals → immunotoxicity

