

# BIOMARKERS AND TOXICITY MECHANISMS 13 – BIOMARKERS of EFFECTS

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









INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

## In vivo biomarkers of effects / response

**Behavioral and Clinical biomarkers** Pathology **Clinical chemistry and hematology Enzymatic changes** Gene and protein expression biomarkers **Detoxification and oxidative stress** markers



# Behavioral and clinical biomarkers



# Examples of behavioral biomarkers

Table 7.4 Effect of some agricultural chemicals on behavioural parameters of the rainbow trout

Chemical	LD <sub>50</sub> (96hr)	Swimming capacity	Swimming activity	Strike frequency	Daphnia consumed	% consuming daphnia	% survival from predation
Carbaryl	1.95	0.1-1	0.1–1	>1	0.1–1	0.1–1	<0.01
Chlordane	0.042	>0.02	0.002-0.02	0.002-0.02	0.002-0.02	0.0020.02	0.002-0.02
DEF	0.66	0.05–0.1	0.005-0.05	0.005-0.05	< 0.005	0.005-0.05	0.005-0.05
2,4-DMA	100	5–50	5-50	5-50	5 -50	0.5-5	5-50
Methyl parathion	3.7	>0.1	< 0.01	0.01-0.1	<0.1	0.01-0.1	0.01-0.1
Pentachlorophenol	0.052	>0.02	0.002-0.02	0.002-0.02	0.0002-0.002	>0.02	0.002-0.02

DEF: tributyl phosphorotrithioate

2,4-DMA: 2,4-dichlorophenoxyacetic acid After Little *et al.* (1990).

# Concentrations affecting behaviour: often lower than LD50 → early markers of (lethal) toxicity



# Behavioral and clinical "biomarkers"

# Interpretation

: are these really biomarkers ?
 (effects already demonstrated in vivo?)
 = biomarkers of existing serious stress / intoxication

# **Parameters evaluated**

- body weight
- food consumption
- fitness & welness



# (Histo)pathology biomarkers



# Pathology

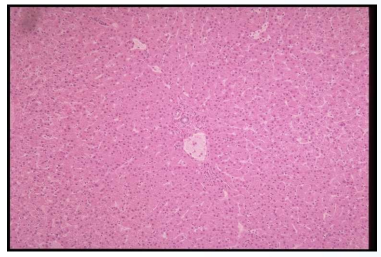
(-) Destructive methods, Time consuming, Professional requirements(+) High relevance – organ/tissue changes

## 1) microscopy of internal organs

A) observations of non-specific changes in internal organs

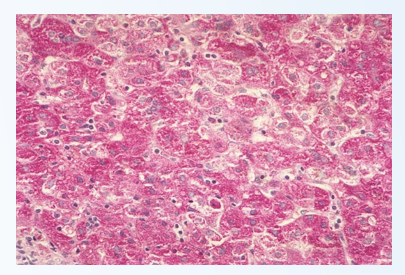
B) specific <u>changes</u>, e.g.

in liver (dioxin-like POPs, cyanobacterial toxins ..) intersex / imposex formation (xenoestrogenicity)





Centrum pro výzkum toxických látek v prostředí

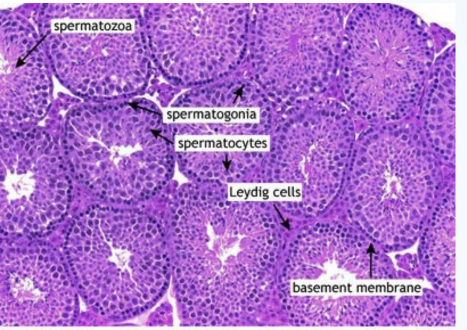


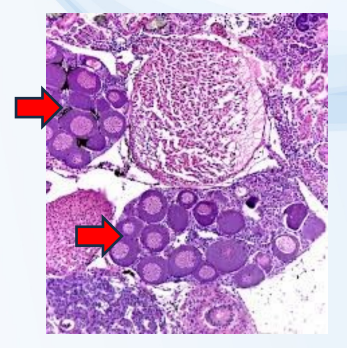
Example: Liver damage by cyanobacterial toxins microcystins

## Endocrine disruption: Intersex microscopy

## **Testicular tissue**

# Oocytes within testis





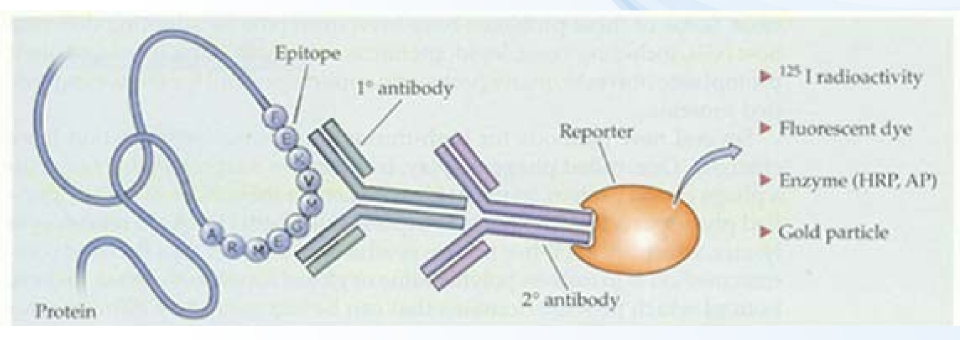
C Deltagen Inc.



## 2) immunohistochemistry & microscopy

- : determination of "specific" changes in tissues
- : Fluorescein (FITC) labeled antibodies (Ab) applications

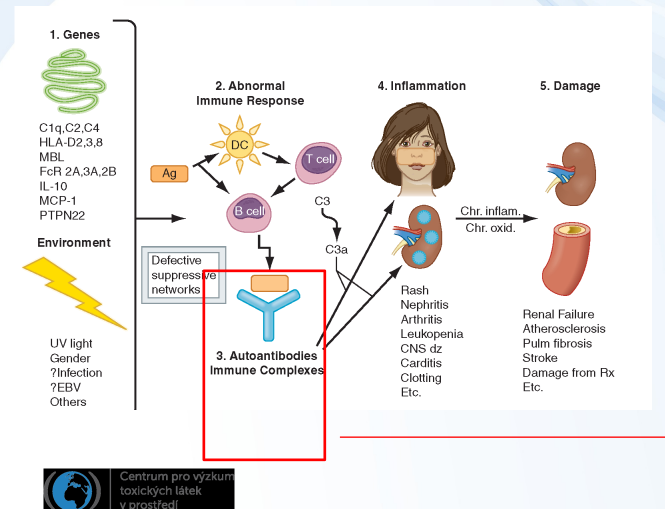
Example → toxicant induced autoimmunity: anti-nuclear Ab (ANA test)





# 2) immunohistochemistry & microscopy anti-nuclear Ab (ANA test)

## Systemic lupus

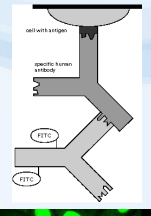


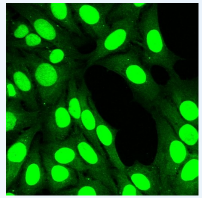
#### ANA test

\* Determination of antibodies in patient blood acting against "nuclei" proteins (ANA)

: target: permeated liver cells on slide

- : application of blood (Ab)
- : visualization (secondary Ab)

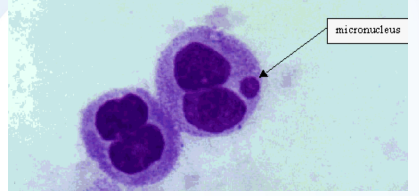




# Pathology

## 3) Nuclear DNA damage characterization 3.1. micronuclei (MN) evaluation by microscopy

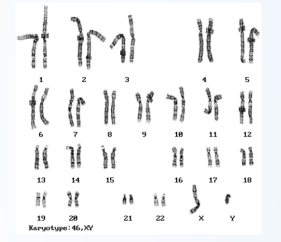
: **example:** MNs in blood lymphocytes of hospital workers (exposed to anticancer drugs – they are often carcinogenic)



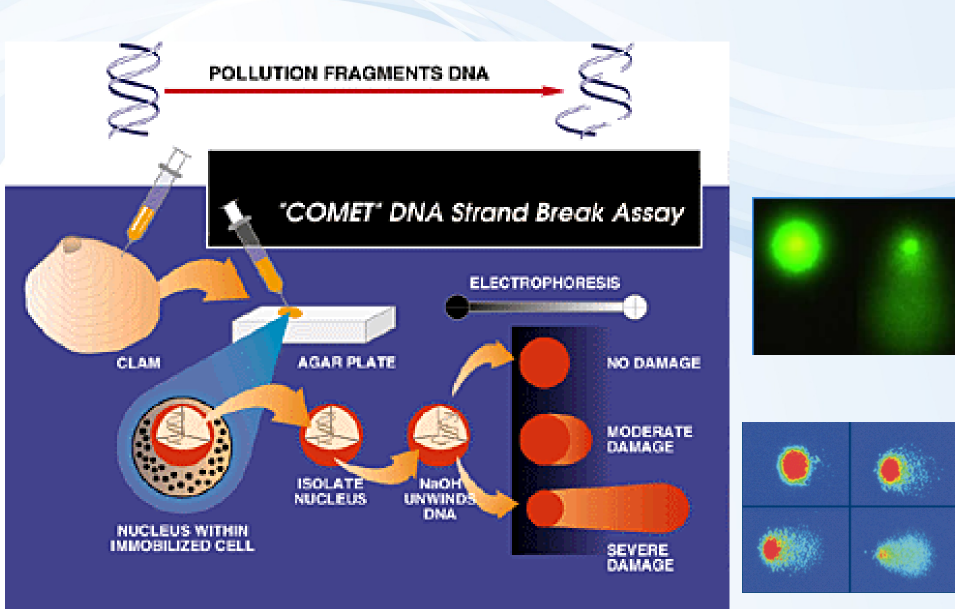
## 3.2 chromosomal abnormalities

karyotype biomarkers (human genetic disorders)

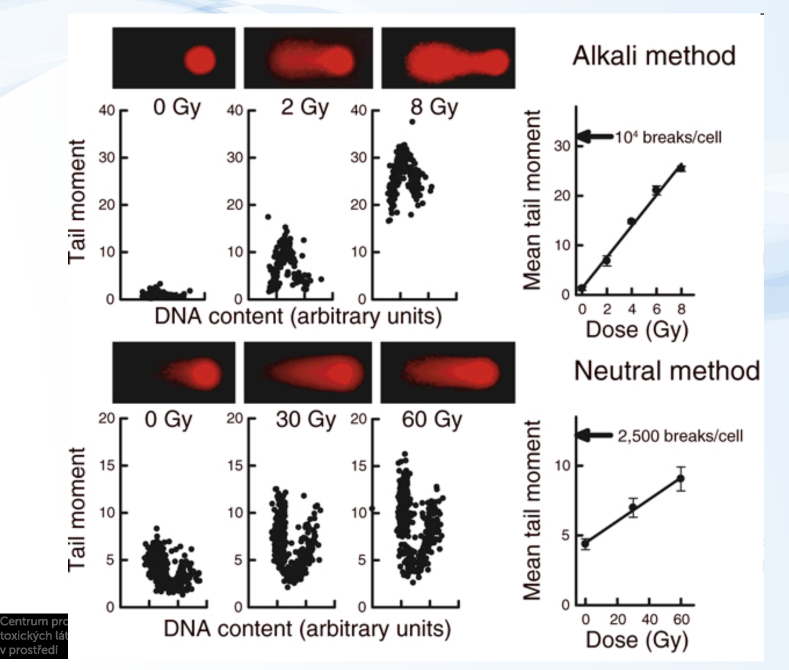




3) Nuclear DNA damage characterization 3.3.COMET ASSAY



## Example results - Comet assay vs. radiation



# Standard clinical chemistry & hematology biomarkers



# **Clinical chemistry & hematology**

# Non-destructive (BLOOD, URINE sampling)

# Multipe parameters can be measured

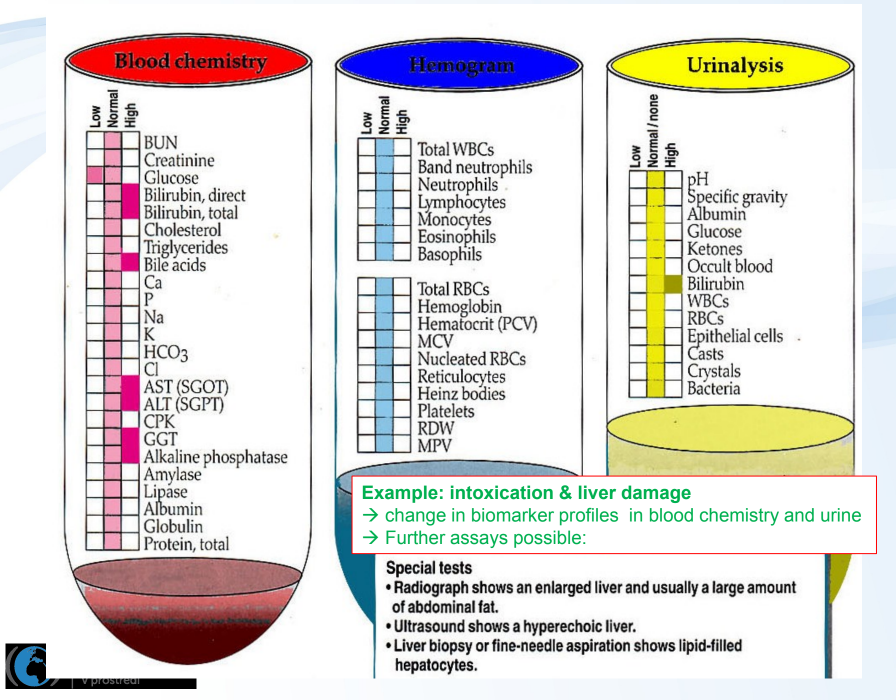
- responses to various types of stresses (including toxic stress)
- "normal" value ranges known for humans, rats and few other species (*limited use as biomarkers in other organisms*)

## **Blood** analyses

- chemistry and biochemistry
- cells (hemogram)
- **Urine analyses** 
  - chemistry, cells, bacteria etc.







## Methods in clinical chemistry

**Methods:** 

- automatic biochemical and hematological analyzers
- different "analytes": various principles of methods (see example  $\rightarrow$ )



# Methods in clinical chemistry

# Example

- determination of enzymatic activities in blood
- interpretation: tissue/organ-specific damage damage

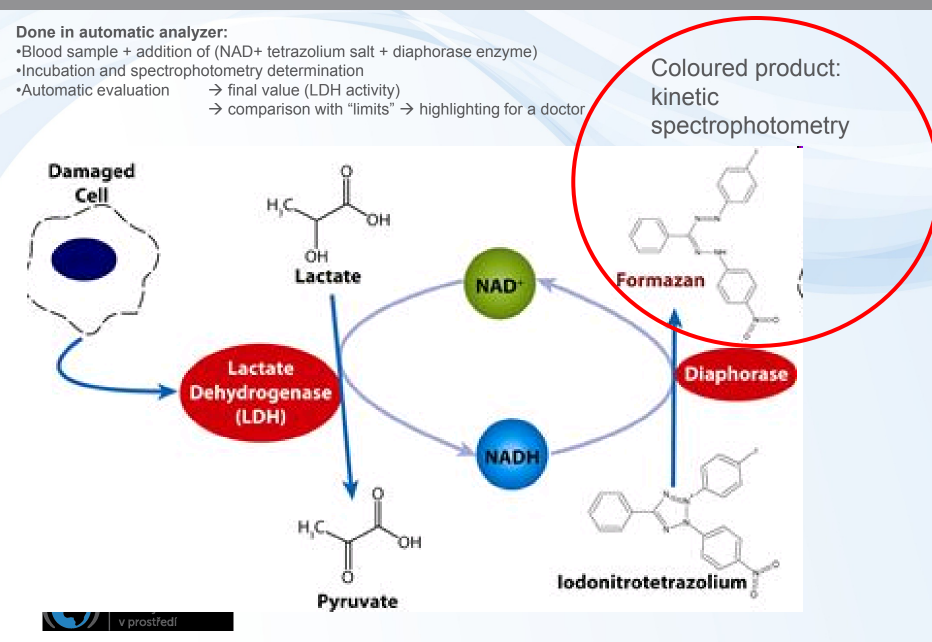
# Examples (toxicological studies)

- Liver damage (toxicants, POPs, alcohol)
  - **AST** (Aspartate aminotransferase),
  - ALT (Alanine aminotransferase) in blood
- General damage in cell (tissue non-specific)
  - LDH lactate dehydrogenase
- Muscle damage:
  - **creatine kinase** in serum (isozymes tissue specific muscle vs heart);

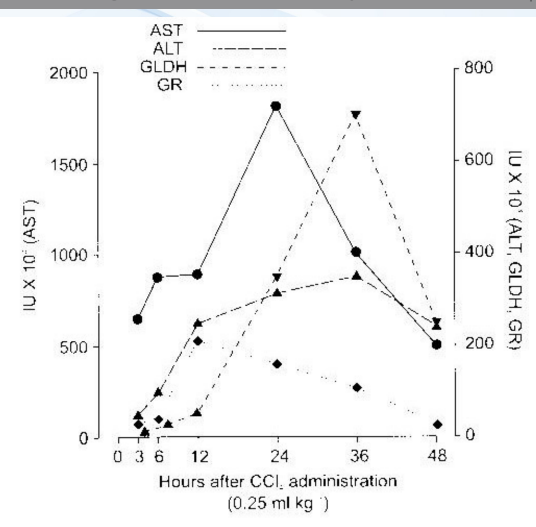
## Other enzyme biomarkers → see further



# Methods in clinical chemistry: example LDH analysis



Example – changes in rat serum enzymes after CCl<sub>4</sub> exposure



**Figure 3** Serum enzyme levels in rats following dosing with carbon tetrachloride (CCl<sub>4</sub>, 0.25 ml kg<sup>-1</sup>). Redrawn from Zimmerman (1978).



Cell damage (Liver) enzyme activity (LDH) is also highly variable and species-specific

#### Table 6.2 Effects of pollutants on LDH

+ Quail
+ Starling
= Redstart
= Redstart
+ Quail
+ Starling
– Fish
(Ophiocephalus)
+ Rat

+ Rat

+ Ouail

- Carp

+ Carp

+ Carp

+ Quail

+ Fish

+ Starling

+ Chicken

+ Chicken

= Brook trout

= Brook trout

= Brook trout

(Notopterus)

+ Starling

Methylparathion Phosmethylan Methidathion

#### Metals

PHAHs

DDE

DDT

**PCBs** 

Endrin

OPs

Photomirex

Malathion

Cadmium chloride Copper sulphate Lead nitrate Mercuric chloride

#### Methylmercury

#### Others



Centrum pro výzkum Oil toxických látek Paraquat v prostředí = Striped mullet + Carp Dieter (1974) Dieter (1975) Karlsson *et al.* (1974)

Dieter (1974) Dieter (1975) Sharma *et al.* (1979)

Chu et al. (1981)

Dragomirescu *et al.* (1975) Dieter (1974) Dieter (1975) Dragomirescu *et al.* (1975) Somlyay *et al.* (1989)

Asztalos et al. (1990)

Christensen *et al.* (1977) Dragomirescu *et al.* (1975) Christensen *et al.* (1977) Dieter (1974) Christensen *et al.* (1977) Verma and Chand (1986)

Dieter (1975)

Chambers et al. (1979) Asztalos et al. (1990)

# Biomarkers: Changes in enzyme activities



# **Enzymatic changes**

**Biomarkers reflecting "enzyme changes":** 

EXAMPLES - inhibitions of specific enzymes (as also discussed earlier during the class: MoA) AcChE (organo-phosphates) Proteinphosphatases (microcystins)

(+) Rapid enzymatic assays, specific responses(-) Some ~ EXPOSURE biomarkers



# Reminder: AcChE inhibition mechanism

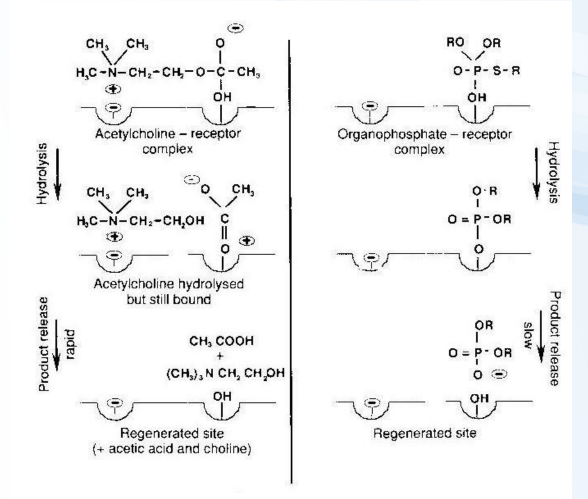


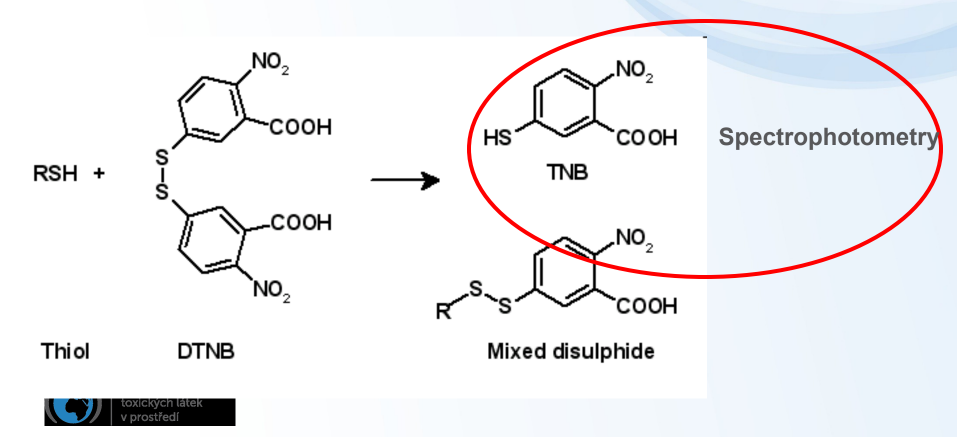
Figure 2.2 Mode of action of inhibition of acetylcholinesterase.



# AcChE assessment

Model Substrate (butyryl-thio-choline, acetyl-thio-choline)

- cleaved by AcChE → formation of free –SH groups
- reaction of SH with thiol reactive probe = Ellman's reagent (DTNB)
- → DTNB-S-choline: yellow colour (spectrophotometry A420)



## Changes in AcChE in birds after exposure to organophosphates

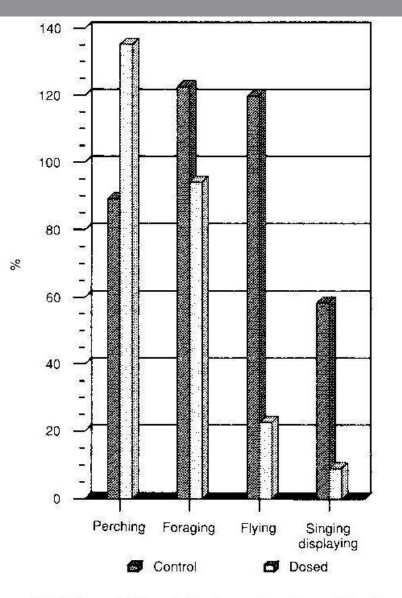


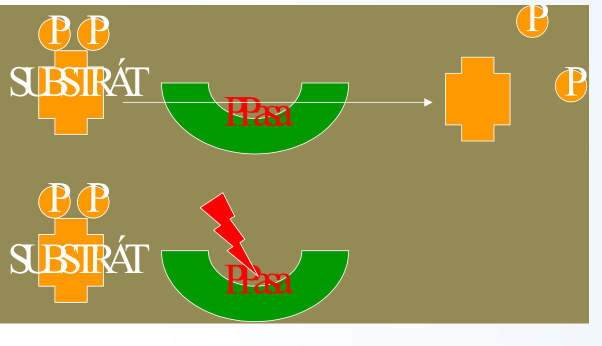


Figure 2.4 Effect of OP on behaviour of starlings. After Grue and Shipley (1981).

# Proteinphosphatase (PPase) inhibition assay

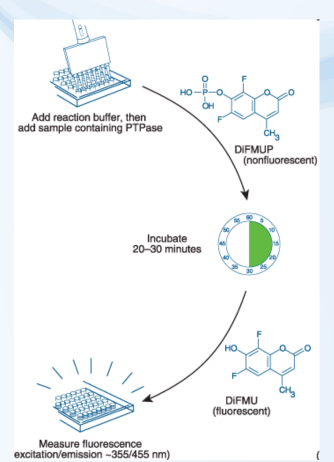
# Model substrates cleaved by PPase

<sup>32</sup>P-labelled protein
 → free <sup>32</sup>P radioactivity
 6,8-difluoro-4-methylumbelliferyl phosphate
 → fluorescence





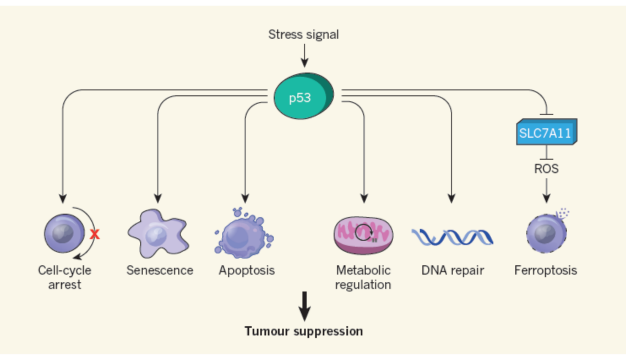
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# Biomarkers – assessing gene and protein expressions / levels



# Protein modulation: toxic response at several levels



Toxicants induce various changes in the cell ...

- ... many of these changes result in
- ightarrow activation / deactivation of specific genes
- $\rightarrow$  modulated gene expression
- $\rightarrow$  modulated protein levels
  - ... and protein activities



Nature (2015) vol 520, p. 37

# How to measure gene and protein modulations?

## **Traditional methods of QUANTIFICATION at different levels**

- mRNA levels
  - PCR / quantitative RT-PCR
- protein levels
  - electrophoresis and Western-(immuno)blotting
  - ELISA techniques
- induced protein enzymatic activities associated with elevated protein levels
  - eg. enzymatic activity

New types of complex techniques: "omics" → also will be discussed later

## Examples of targeted protein biomarkers – discussed further $\rightarrow$

specific protein markers of disease / e.g. cancer

heat shock proteins (hsp90, hsp60, hsp 70, ubiquitin)

metalothioneins

endocrine disruption biomarkers

- Vitellogenin(-like) Vtg proteins in male
- Aromatase
- Induction of detoxification enzymes CY
- CYP450 / EROD
  - GST



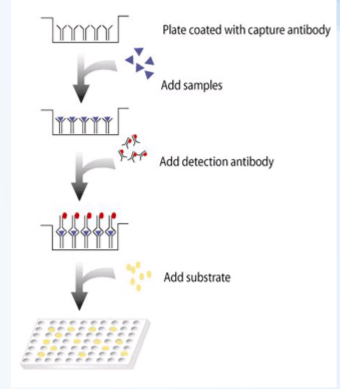
## **ESTABLISHED PROTEIN MARKERS – determination in blood**

## **Tumor genes and tumor markers**

- cancer genes ras, myc e.g. metastasing bowel cancer
- $\alpha$ -fetoprotein (AFP) elevated during fetus development AND e.g. liver cancers
- tumor suppressor genes (e.g. *p53) indicate better prognosis for certain cancers*
- PSA prostate-specific antigen: prostate cancer in males (over 50 years of age)

# Methods of determination in practice: ELISA

(enzyme linked immunosorbent assays)





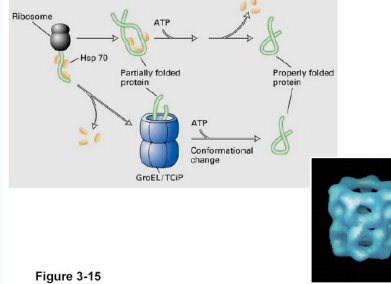
# Heat Shock Proteins (hsp)

# General stress = synthesis of new proteins

- ~ equilibrium and homeostasis buffering
  - temperature (cold / heat) → proteins assuring cryo-preservation
  - salinity & metals  $\rightarrow$  ion buffering
  - organic xenobiotics  $\rightarrow$  detoxication

# New proteins must be folded to their 3D stucture by activity of "CHAPERONES"

Chaperons = hsp90, hsp60, hsp 70 ~ 60-90 kD molecular weight kD

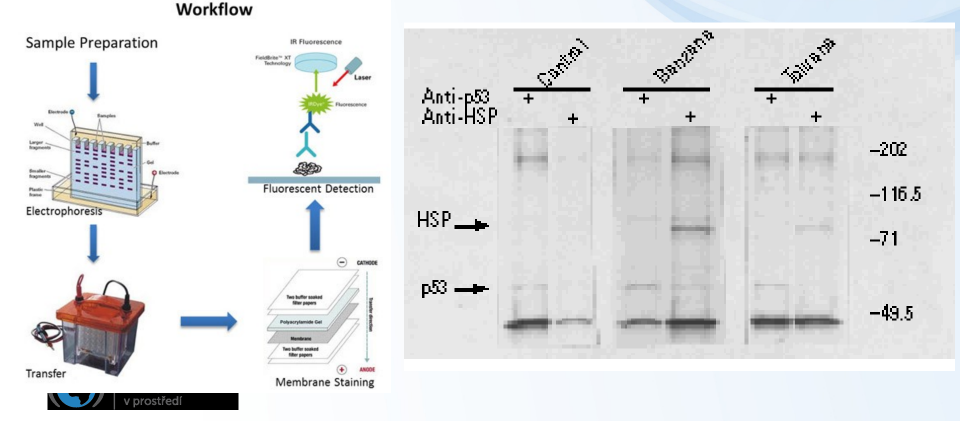




# HSP determination - example

## HSP = GENERAL STRESS biomarker, non-specific

- phylogenetically conserved (similar genes in most of the organisms)
- structural similarity  $\rightarrow$  easy determination:
  - electrophoresis + immunoblotting (Western blotting)



# Metalothioneins (MTs, MT-like proteins)

## Low MW proteins (6-10 kD) rich of Cystein (-SH)

- detected in numerous eukaryotic organisms
- induced in the presence of metals or less specific stress (low O2, T)
- long halflife (~ 25 days)
- binding of divalent metals (Zn, Cd, Hg)  $\rightarrow$  exposure elimination
- natural function (?) regulation of essencial metals in cells

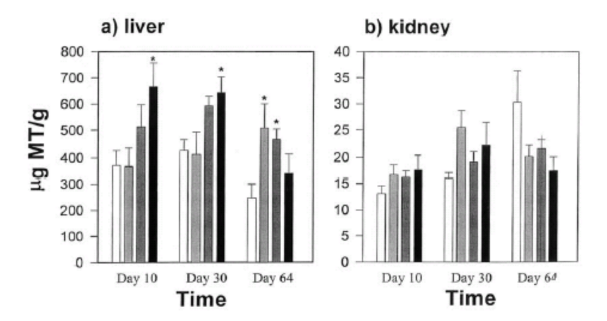


Fig. 2. Metallothionein (MT) concentrations in the (a) livers and (b) kidneys of lake whitefish fed a control diet and three As contaminated diets for 10, 30, and 64 days. Data are expressed as mean ( $\pm$ S.E.). Asterisk denotes mean is significantly different from the control at that duration (P < 0.05). See Fig. 1 for an explanation of histogram shading.

# Protein biomarkers of estrogenicity / ED-like effects

## ER = transcription factor controling number of target genes

Target genes of ER = biomarkers of estrogenicity

Major examples

Vitellogenin
Aromatase - CYP19A



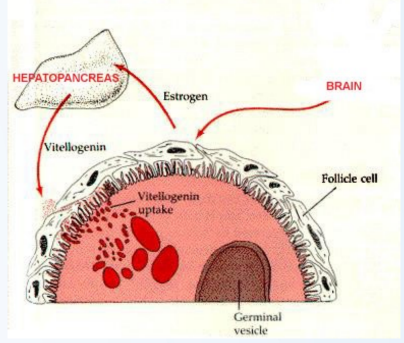
# Vitellogenin (Vtg)

Precursor of yolk proteins, phospho-protein ("energy" rich) → egg formations (females) **at oviparous vertebrates** 

Synthesized in liver and distributed via blood / haemolymph

## **Xenoestrogens & other endocrine disruptors**

- $\rightarrow$  increased levels or early production in FEMALES
- $\rightarrow$  production de novo in **MALES**



(A) VITELLOGENESIS AND OOCYTE DIFFERENTIATION



## Vitellogenin (Vtg) assessment

## 1) ELISA in exposed organisms (F/M) or in vitro

(-) specific antibodies are necessary for each species (low crossreactivity of Abs)

## 2) "Vitelin-like proteins"

- total amount of "alkali-labile" phosphate in haemolymph (mussels)

- alkaline extraction of P from sample  $\rightarrow$  spectrophotometric determination

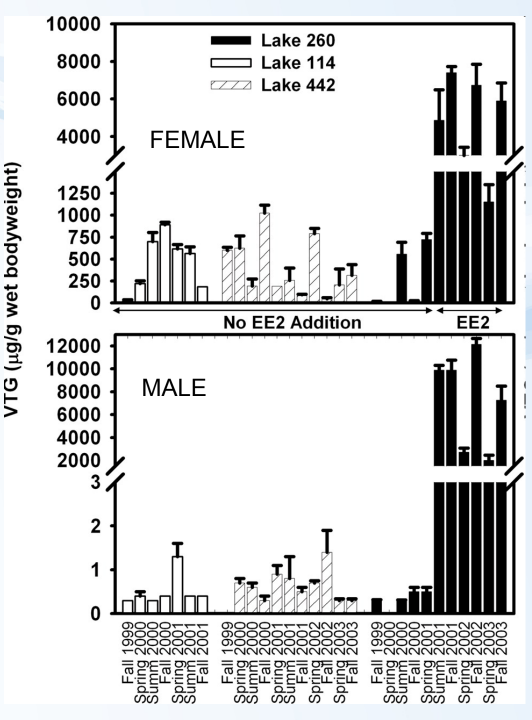


### Vitellogenin in fish

### Kidd et al. (2007) PNAS

Fig. 1. Mean SE (n = 4-7) VTG concentrations in whole-body homogenates of male (*Lower*) and female (*Upper*) fathead minnow captured in 1999–2003 from reference Lakes 114 and 442 and from Lake 260 before and during additions of 5–6 ng·L<sup>-1</sup> of EE2 (low catches of fish in Lake 260 in 2004 and 2005 did not allow for these analyses in the latter 2 years of the study).





## Vitelin-like proteins in mussels

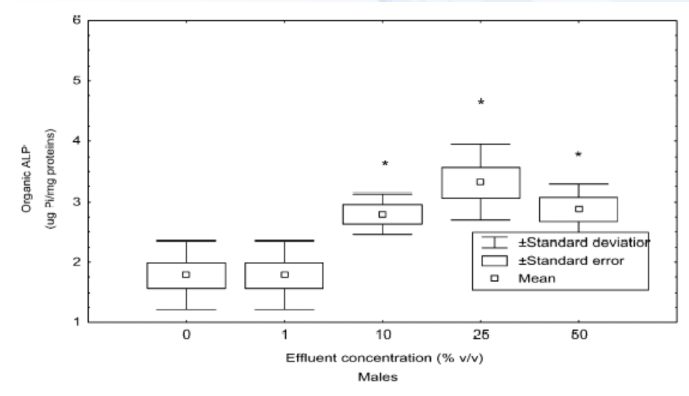
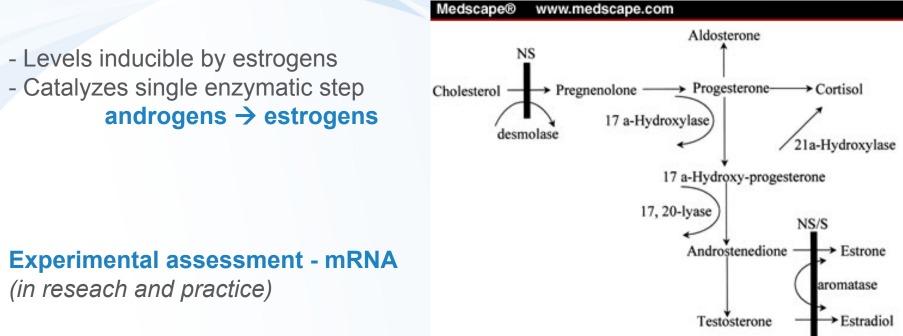


Fig. 4. Induction of Vg by exposure to a municipal effluent. Mussels were exposed for 96 h to a municipal effluent at 15°C. They were then collected for Vg and sex determinations. The asterisk (\*) indicates significant difference at P < 0.05.



# Aromatase (CYP19A)



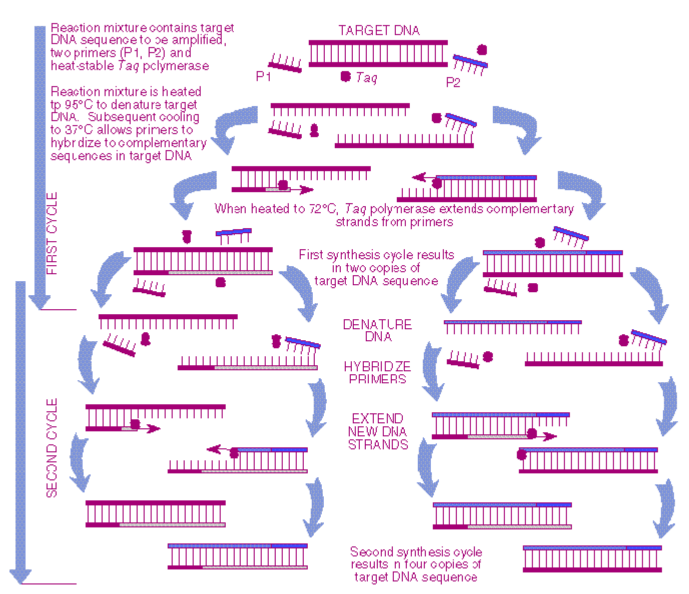
1. PCR / Quantitative-Real-Time-PCR

### 2. GM-organisms (zebrafish): reporter gene with GFP

Green Fluorescence Protein under the control of aromatase promoter



### **DNA Amplification Using Polymerase Chain Reaction**







# Visualization of PCR product

## 1) Electrophoresis (qualitative)

### Intercalation dyes – e.g. **ethidium bromide**

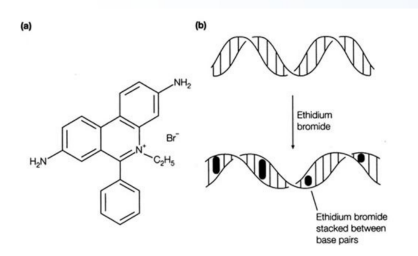
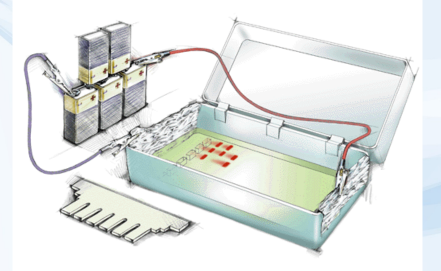


Fig. 3. (a) Ethidium bromide; (b) the process of intercalation, illustrating the lengthening and untwisting of the DNA helix.



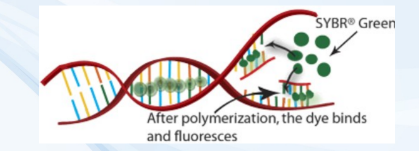


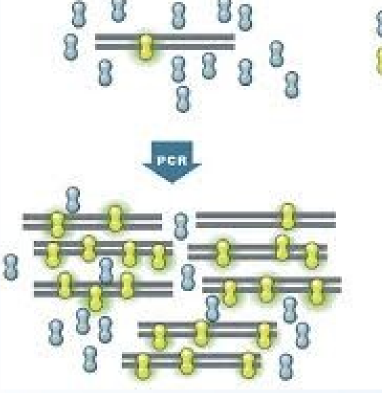


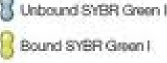
# Visualization of PCR product

### 2a) Real-time (quantitative) SYBR GREEN dye

→ more DNA synthesized,
 more fluorescent dye incorporated
 → Higher fluorescence





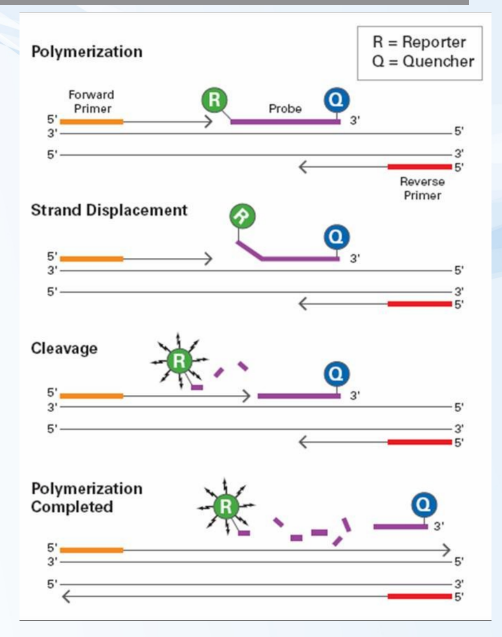




# Visualization of PCR product

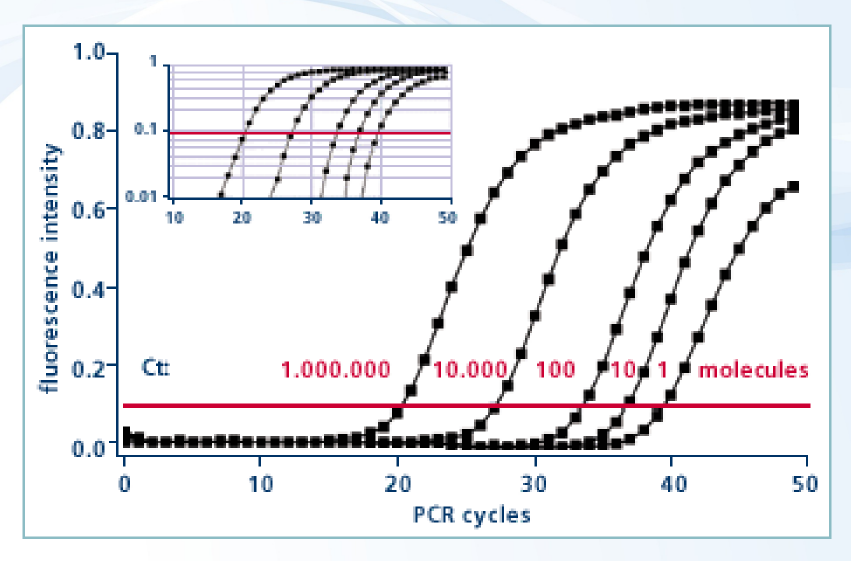
### 2b) Real-time (quantitative) TaqMan probes

(more DNA replications more fluorescent dye released)



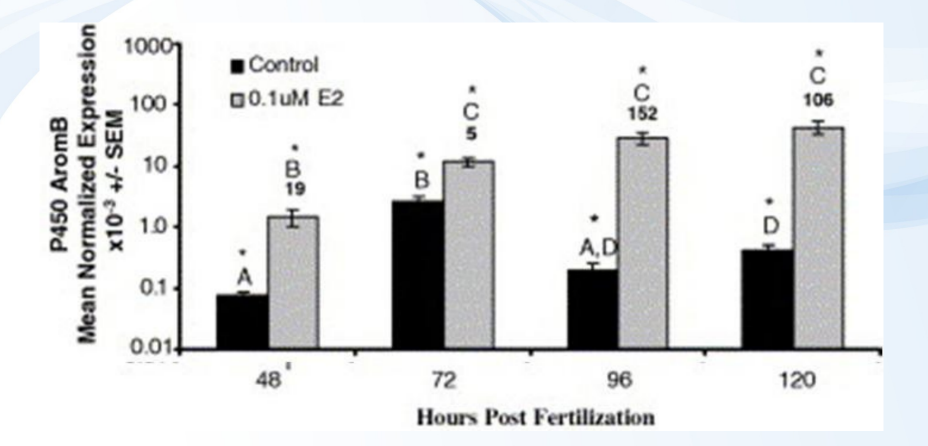


## "Quantitative" determination of PCR product





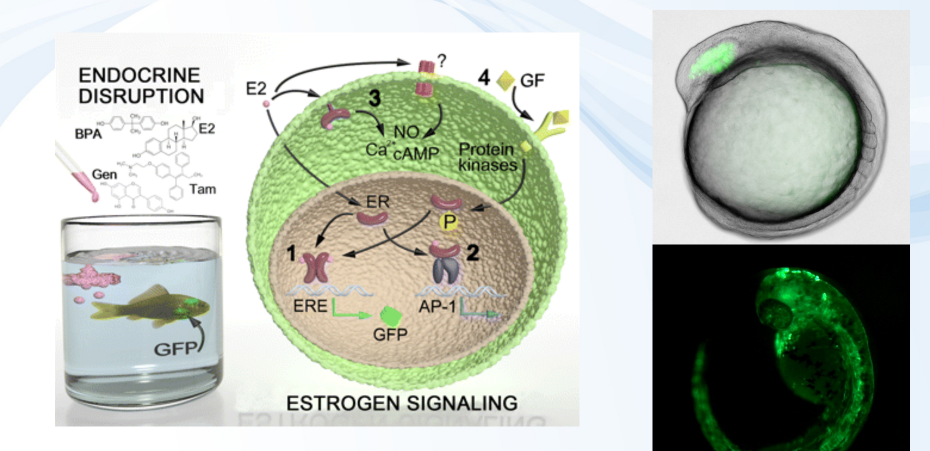
## qPCR determination of the aromatase gene in Zebrafish



http://dx.doi.org/10.1016/j.ygcen.2005.12.010,



## GFP-reporter for estrogens in zebrafish embryo

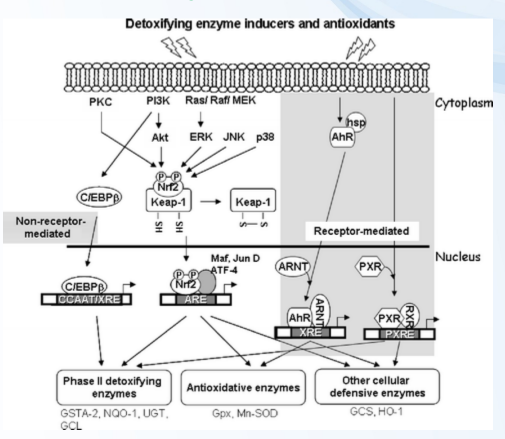


http://endo.endojournals.org/content/152/7/2542.full



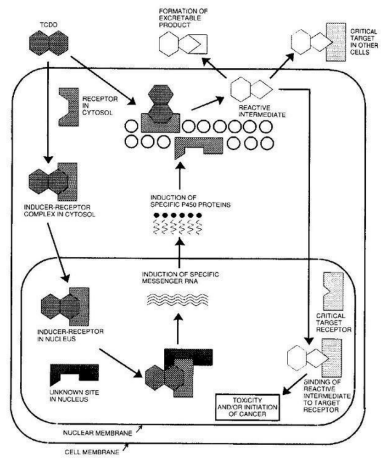
## DETOXIFICATION / ANTIOXIDANT DEFENCES

Inductions of detoxication & oxidative stress enzymes (hepatopancreas / liver / blood) MFO - CYP classes - EROD / MROD / BROD Phase II enzymes (GSTs) Glutathion metabolism enzymes (GPx, GRs)





# MFO (CYPs) - reminder







#### Table 5.1 Classification of P450s

Nomenclature	Induced by/specificity	
P4501	Polycyclic aromatic, TCDD	
P450II	Phenobarbital-inducible family*	
P450IIA	Specific for testosterone hydroxylase	
P450IIB	PB inducible	
P450IIC	PB inducible	
P450IID	Specific for debrisoquine 4-hydroxylase	
P450IIE	Ethanol inducible	
P450III	Steroid inducible	
P450IV	Specific to lauric acid w-hydroxylation	
P450XI	Located in mitochondrion	
P450XIA		
P450XIB		
P450XVII	Formation of steroid 17-hydroxylases	
P450XIX	Involved in synthesis of oestrogens	
P450XX1	Formation of steroid 21-hydroxylases	
P450LI	Plant/yeast	
P450CI	Prokaryote	

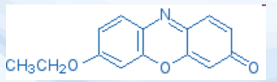
\* PB-inducible genes largely confined to P450IIB and C. After Nebert and Gonzalez (1987).

# Assessment of CYPs (MFO) - "EROD"

## Determination of CYP1A1 activity "EROD" - EthoxyResorufin-O-Deethylase activity

Substrate: Ethoxyresorufin

- : Oxidation by CYP1A1 (MFOs)
- → Fluorescence (easy determination)



## **EROD** = sensitive biomarker of organic pollution (exposure & effects)

- : AhR-activating compounds (PCDD/Fs, PCBs, PAHs)
- : often used in environmental studies

Use of other substrates: assessment of other CYPs BROD – butoxy-ROD (CYP3A), MROD, PROD ...



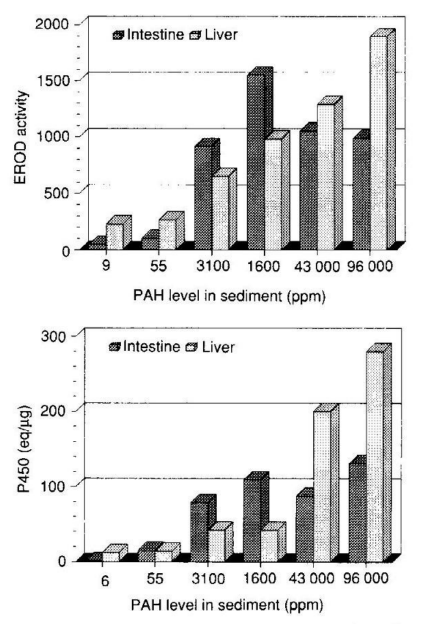
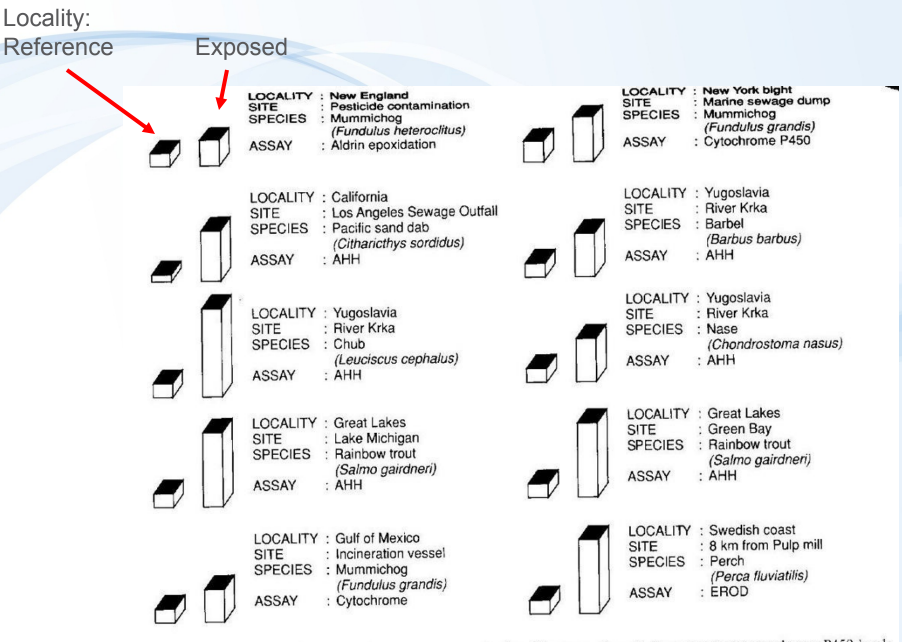


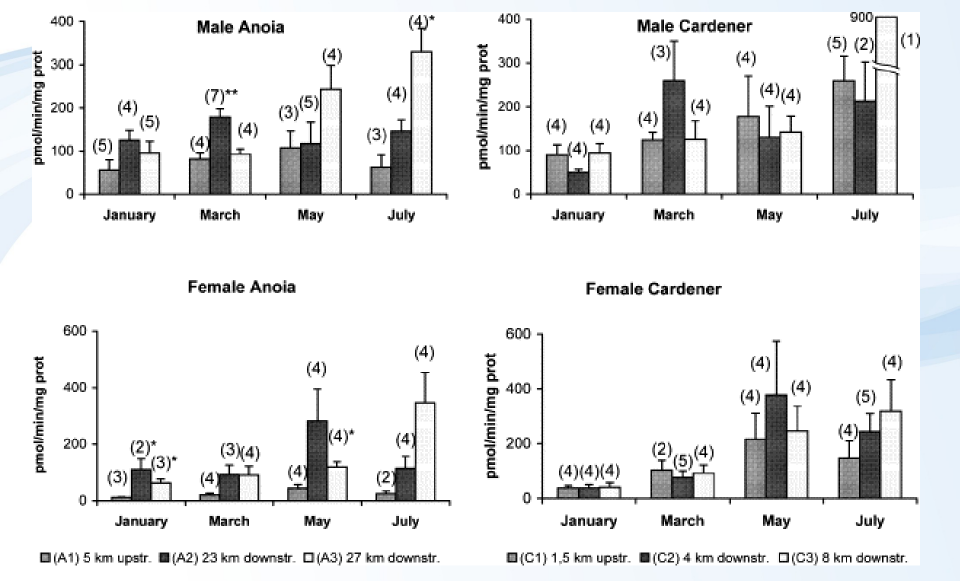


Figure 5.6 Relationship of sediment concentration of PAHs to EROD activity in liver and intestine of spot. After Van Veld *et al.* (1990).



Centru toxický v prost

Figure 5.5 MFO changes in fish exposed to organic contamination. The proportion of either enzyme or cytochrome P450 levels letected at reference (short towers) and experimental sites (long towers) is presented in schematic form. All differences between reference and experimental sites were statistically significant (P < 0.05 or better). Payne *et al.* (1987).



EROD variation on male and female carp from the Anoia and Cardener tributaries – seasonal variability & response at contaminated localities



MFO responses (EROD) are strongly **species specific** & not always related to clinical signs

Table 3.3 Comparison of the effects of PCB congeners on the reproduction of mink and rats

C	122 - Kr	
PCB congener	Mink	Rat
2,4,2',4'-TCB	Clinically normal No change in cytochrome P450	Clinically normal No change in cytochrome P450
(	No induction of MFO	Some induction of MFO enzymes
	Severe anorexia and diarrhoea	Clinically normal
	Increase of cytochrome P450	Increase in cytochrome P450
	No induction of MFO enzymes	Induction of MFO enzymes

After Gillette et al. (1987a).



# MFO-responses (EROD) depends on animal size and metabolism rate

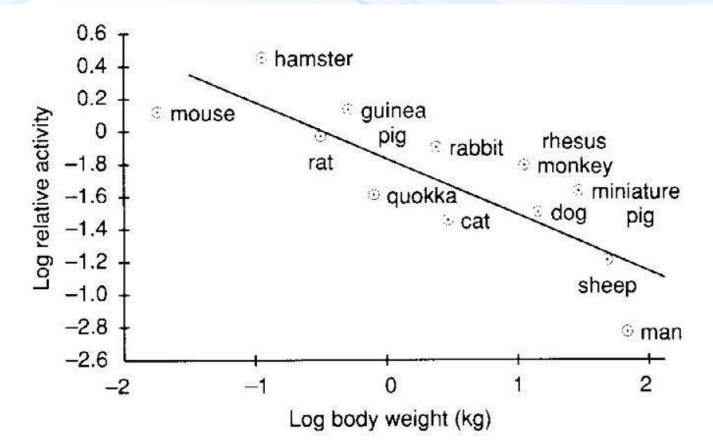


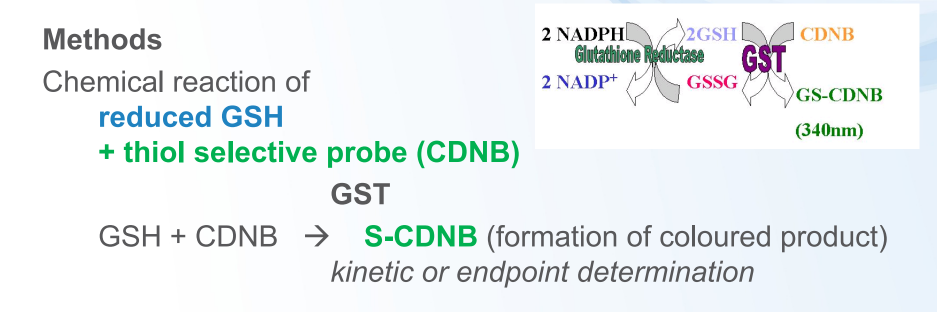
Figure 5.3 Relationship of body weight to MFO activity in mammals. Walker (1978 and 1980).



# Phase II conjugation enzymes - GSTs

## GSTs

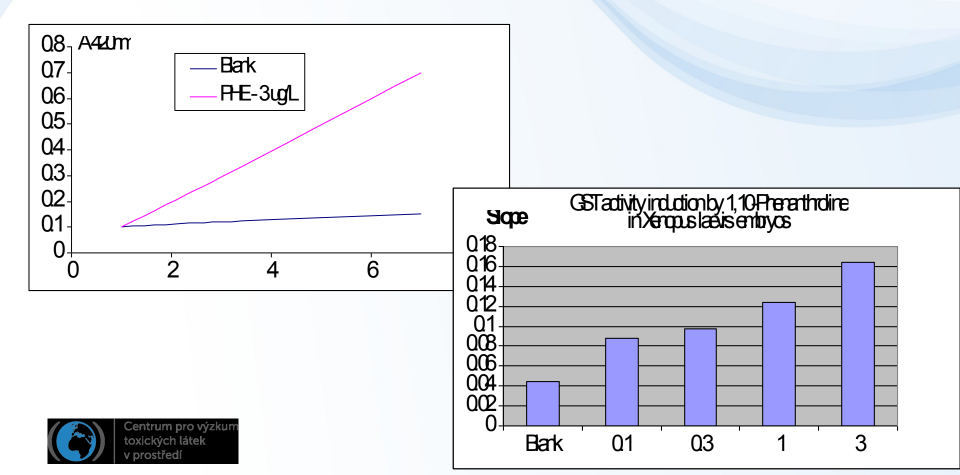
soluble and membrane (endoplasmic reticulum) variants: activities can be measured in cytoplasm or ER microsomes





# GST activity determination: example

Kinetic assessment of GSTs stress → Induction of GSTs faster reaction = increasing slope of the kinetics



# Biomarkers of oxidative stress



# **Oxidative stress markers**

Several parameters respond to oxidative stress

- : **enzymes** detoxification, antioxidants: GPx, GR, GSTs) .. - enzymatic activities (see elsewhere)
- : antioxidants e.g. GSH (discussed further), vitamin E
- : markers of oxidative damage
  - membranes: MDA (discussed further)
  - DNA: 80H-dG
  - (see at DNA damage / adducts-exposure biomarkers)
  - proteins: oxidized forms (carbonyls)



# **Oxidative stress markers**

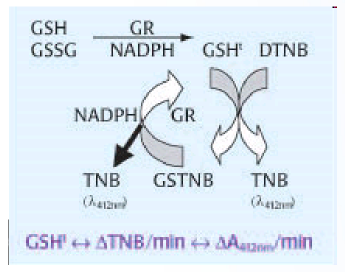
## GSH

- antioxidant (scavenger of ROS) & reactive molecules
- conjugation molecules for detoxication
- probable intracellular regulatory molecule (? apoptosis ?)

### Total glutathione = reduced GSH + oxidized GSSG

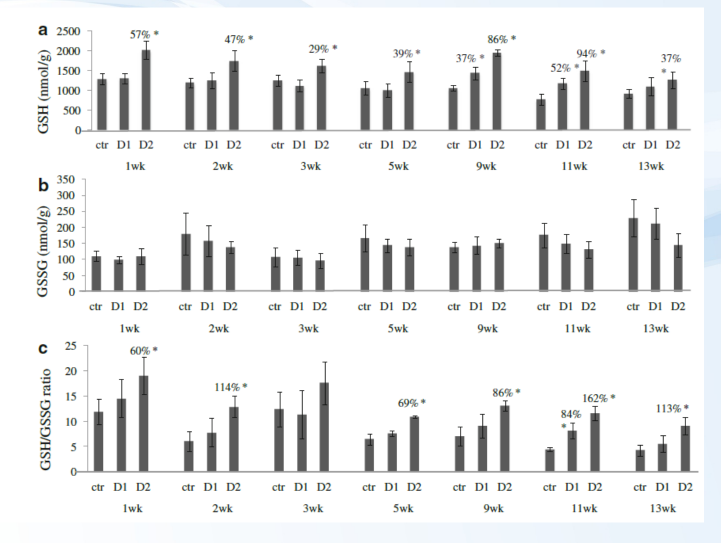
Method of determination (thiol selective probe DTNB)GSH + Ellman s reagent (DTNB)→ Reduced GSHGSH + GSH-reductase + DTNB→ Total GSH

Total – Reduced = Oxidized



# Example - GSH modulation by toxic nanoparticles

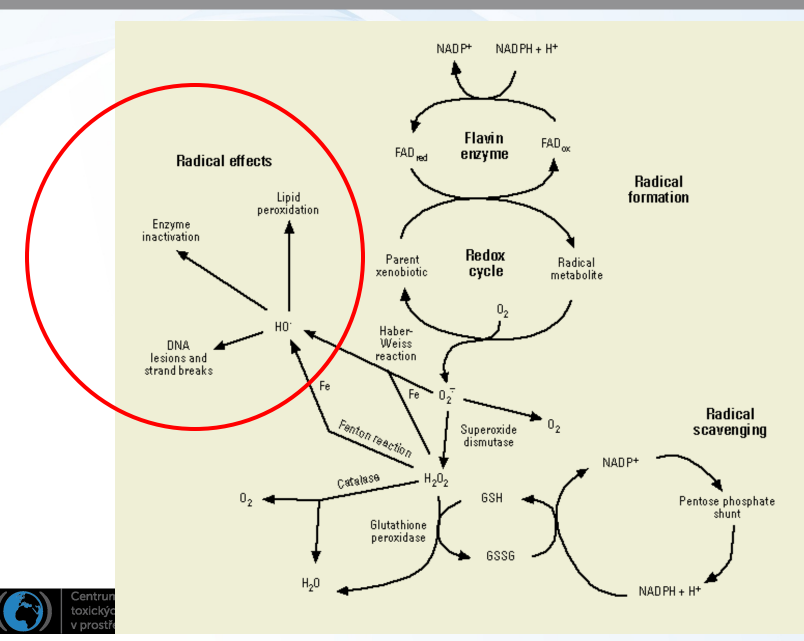
Fig. 6 Content of GSH (a), content of GSSG (b), and GSH/ GSSG ratio (c) in lung of mice after chronic exposure (1–13 weeks) to CdO nanoparticles at dose 1 (*D1*) and dose 2 (*D2*). *Numbers with asterisk* (\*) in the graph indicate significant differences compared to the control variant within the respective week (p < 0.05; N=5animals)





Bláhová et al. 2014 Anal Bional Chem 406:5867–5876

# Markers of oxidative DAMAGE

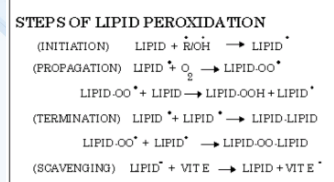


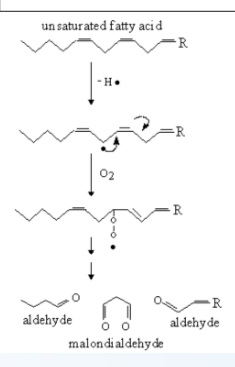
## Lipid peroxidation $\rightarrow$ Malondialdehyde (MDA)

### MDA – malondialdehyde

product of lipid peroxidation







## Malondialdehyde (MDA) determination

## MDA – formed from oxidized membrane phospholipids

- : determination:
  - HPLC (instrumental)
  - TBARS (spectrophotometric) method

## **TBARS – ThioBarbituric Acid Reactive Species**

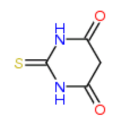
- : less specific than HPLC
- : easy determination (spectrophotometry)

## Method:

- 1) sample extract (with MDA)
- 2) add TBA
- 3) boil (cca 30 / 90 C)

## → formation of red/violet coloured product

4) determination by spectrophotometry (A 540 nm)



TBA



## MDA modulation - examples

### Effects of nanoFeOxide particles on MDA in fish

### Induction of MDA (TBARS) by carbamazepine (and protection by antioxidants)

