

# BIOMARKERS AND TOXICITY MECHANISMS 13 - BIOMARKERS of EFFECTS

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









#### In vivo biomarkers of effects / response

Behavioral and Clinical biomarkers
Pathology
Clinical chemistry and hematology
Enzymatic changes
Gene and protein expression biomarkers
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



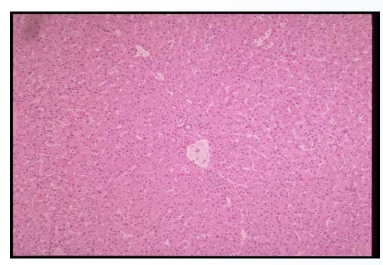
## (Histo)pathology biomarkers



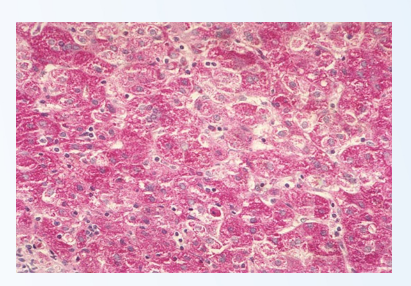
#### Pathology

- (-) 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 **changes**, e.g.

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



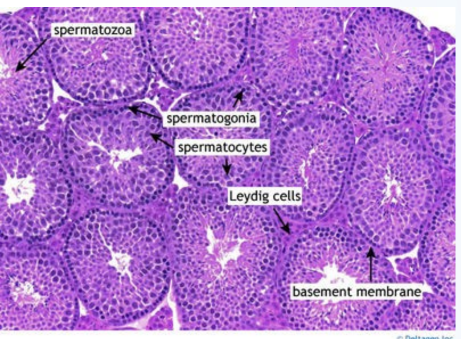




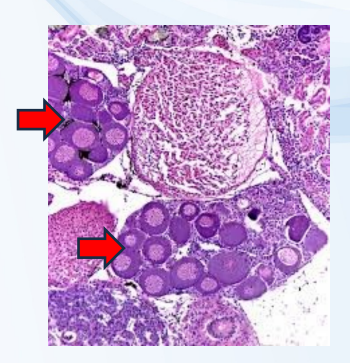
Example: Liver damage by cyanobacterial toxins microcystins

#### Endocrine disruption: Intersex microscopy

#### Testicular tissue



#### Oocytes within testis





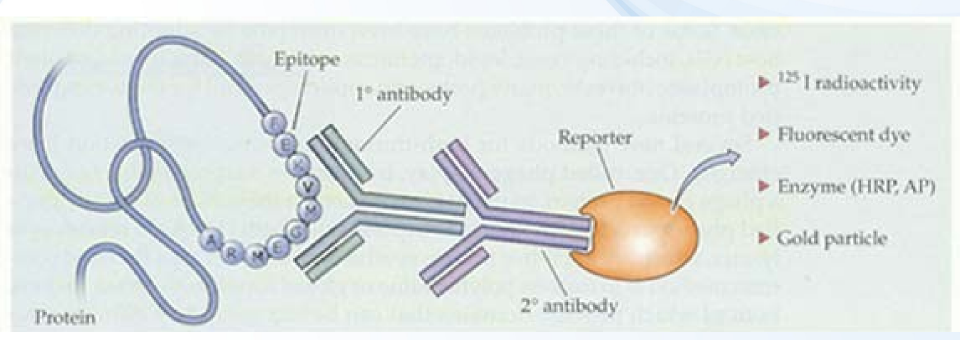


#### 2) immunohistochemistry & microscopy

: determination of "specific" changes in tissues

: Fluorescein (FITC) - labeled antibodies (Ab) applications

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

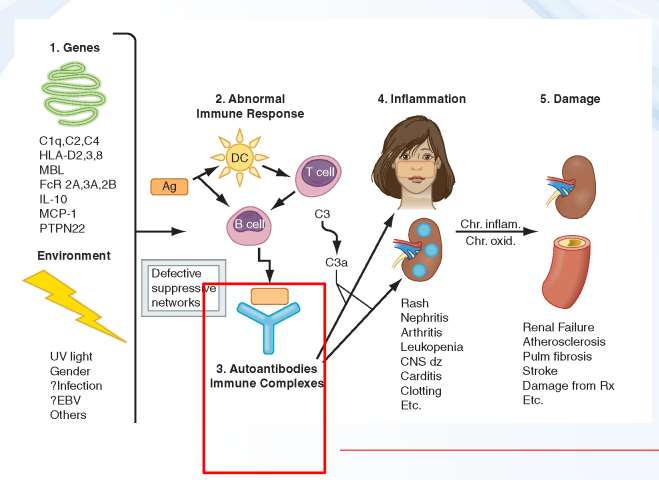




### 2) immunohistochemistry & microscopy anti-nuclear Ab (ANA test)

#### Systemic lupus

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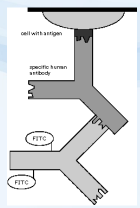
#### **ANA** test

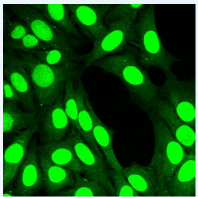
\* Determination of antibodies in patient blood acting against "nuclei" proteins (ANA)

: target: permeated liver cells on slide

: application of blood (Ab)

: visualization (secondary Ab)



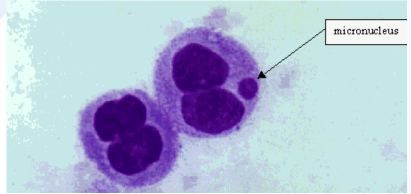


#### Pathology

#### 3) Nuclear DNA damage characterization

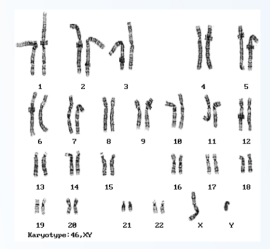
#### 3.1. micronuclei (MN) evaluation by microscopy

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



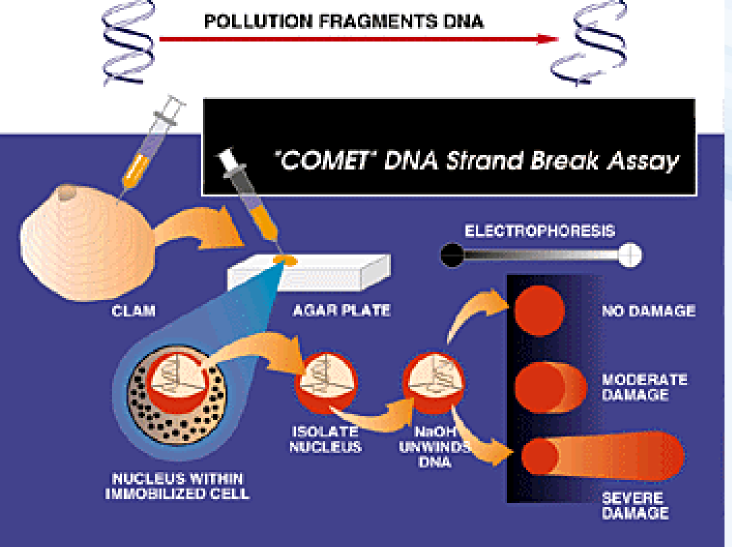
#### 3.2 chromosomal abnormalities

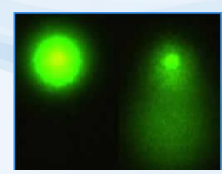
karyotype biomarkers (human genetic disorders)

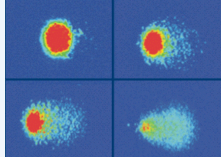




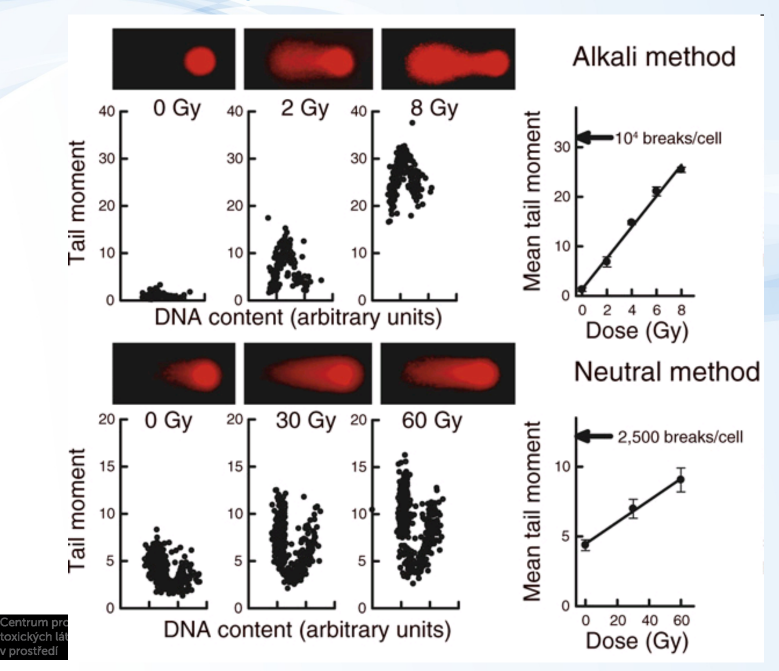
### 3) Nuclear DNA damage characterization 3.3.COMET ASSAY







#### **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**

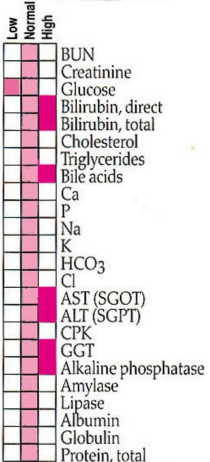
- chemistry, cells, bacteria etc.



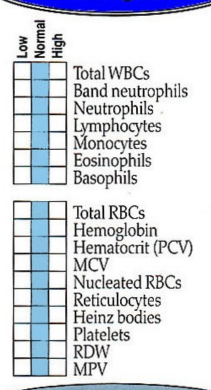




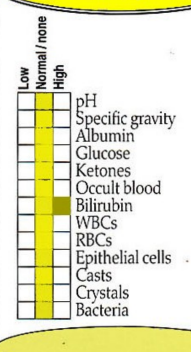
### Blood chemistry



#### Hemogram



#### Urinalysis



#### Example: intoxication & liver damage

- → change in biomarker profiles in blood chemistry and urine
- → Further assays possible:

#### Special tests

- Radiograph shows an enlarged liver and usually a large amount of abdominal fat.
- · Ultrasound shows a hyperechoic liver.
- Liver biopsy or fine-needle aspiration shows lipid-filled hepatocytes.



#### Methods in clinical chemistry

#### Methods:

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



#### Methods in clinical chemistry

#### **Example**

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

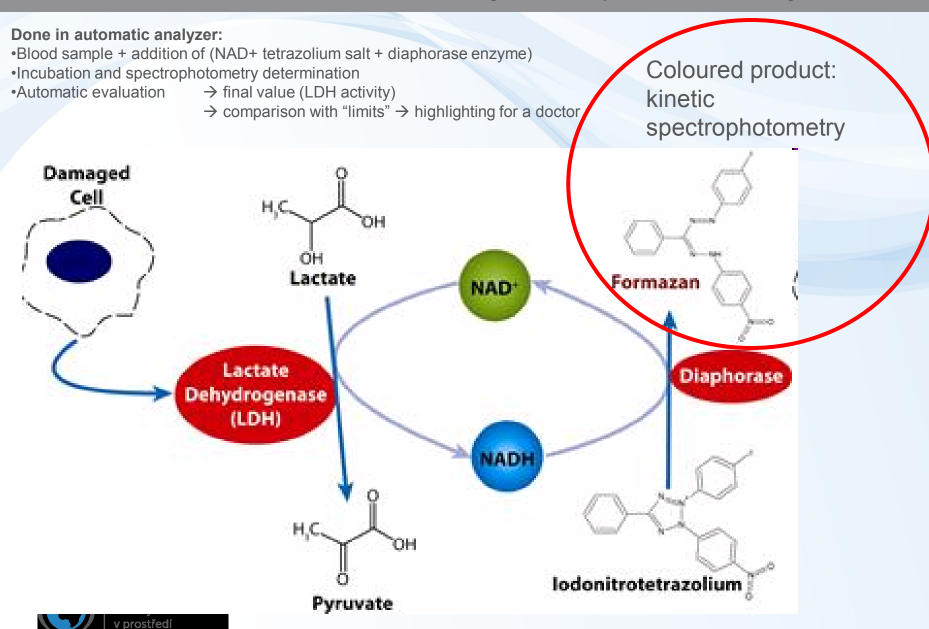
#### Examples (toxicological studies)

- 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);

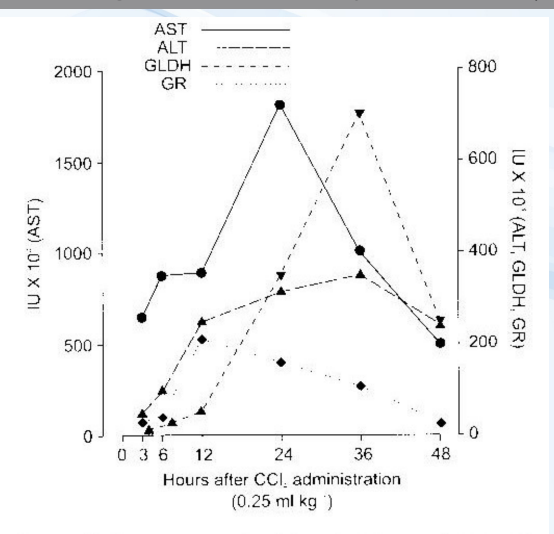
#### Other enzyme biomarkers → see further



#### Methods in clinical chemistry: example LDH analysis



#### Example – changes in rat serum enzymes after CCI<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).



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

Table 6.2 Effects of pollutants on LDH

PHAHs			
DDE	+ Quail	Dieter (1974)	
	+ Starling	Dieter (1975)	
DDT	= Redstart	Karlsson <i>et al.</i> (1974)	
PCBs	= Redstart	1 un (1574)	
	+ Quail	Dieter (1974)	
	+ Starling	Dieter (1975)	
Endrin	- Fish	Sharma et al. (1979)	
	(Ophiocephalus)		
Photomirex	+ Rat	Chu et al. (1981)	
OPs			
Malathion	+ Rat	Dragomirescu et al. (1975)	
	+ Quail	Dieter (1974)	
	+ Starling	Dieter (1975)	
	– Carp	Dragomirescu et al. (1975)	
Methylparathion	+ Chicken	Somlyay et al. (1989)	
Phosmethylan	+ Chicken	3-3 (13-03)	
Methidathion	+ Carp	Asztalos et al. (1990)	
Metals			
Cadmium chloride	= Brook trout	Christensen et al. (1977)	
Copper sulphate	+ Carp	Dragomirescu et al. (1975)	
Lead nitrate	= Brook trout	Christensen et al. (1977)	
Mercuric chloride	+ Quail	Dieter (1974)	
	= Brook trout	Christensen et al. (1977)	
	+ Fish	Verma and Chand (1986)	
	(Notopterus)	mid Official (1700)	
Methylmercury	+ Starling	Dieter (1975)	
Others			
Oil	= Striped mullet	Chambers et al. (1979)	
Paraquat	+ Carp	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

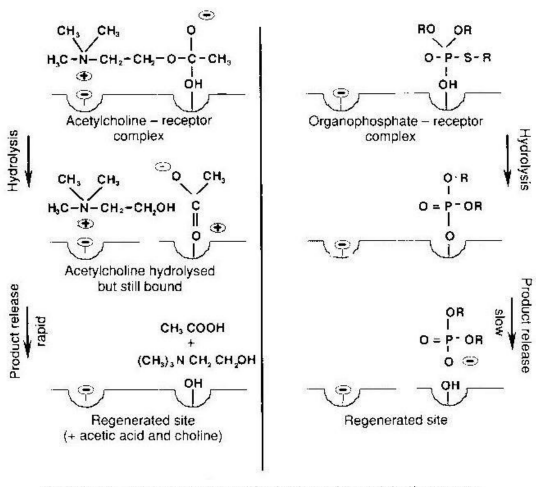


Figure 2.2 Mode of action of inhibition of acetylcholinesterase.



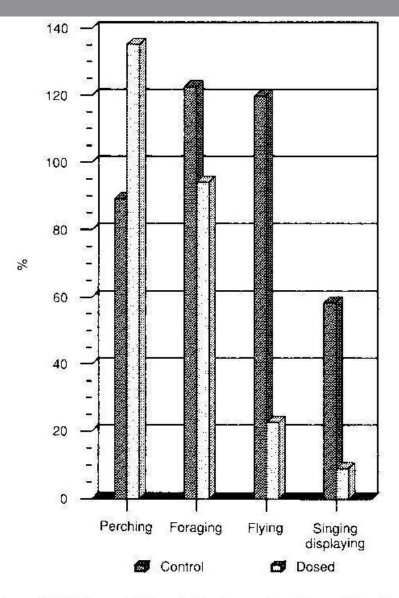
#### AcChE assessment

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

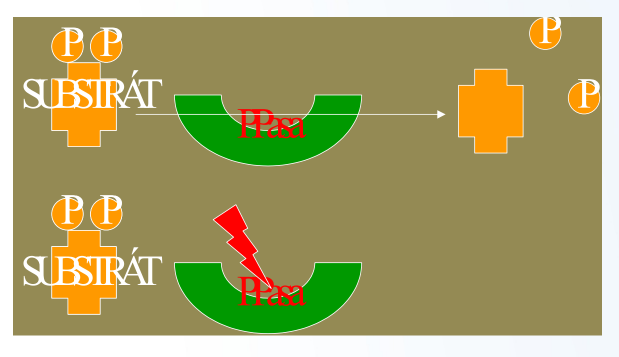
#### Model substrates cleaved by PPase

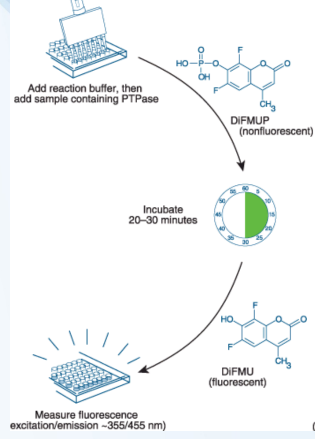
<sup>32</sup>P-labelled protein

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→ free <sup>32</sup>P radioactivity

6,8-difluoro-4-methylumbelliferyl phosphate → fluorescence





# Biomarkers – assessing gene and protein expressions / levels



#### 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
  - activities of induced enzymes

**New types of complex techniques:** "omics" → discussed later

#### Examples of targeted protein biomarkers – discussed further →

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 - CYP450 / EROD

- GST



#### PROTEIN MARKERS OF DISEASE – determination in blood

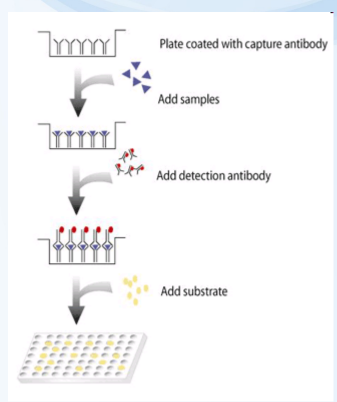
#### **Example: Tumor genes and tumor markers**

- 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 → ion buffering
  - organic xenobiotics → detoxication

#### New proteins must be folded to their 3D stucture

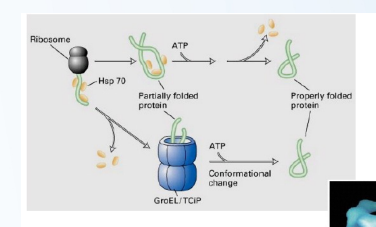
by activity of "CHAPERONES"

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

Methods of determination:

ELISA, Western Blotting

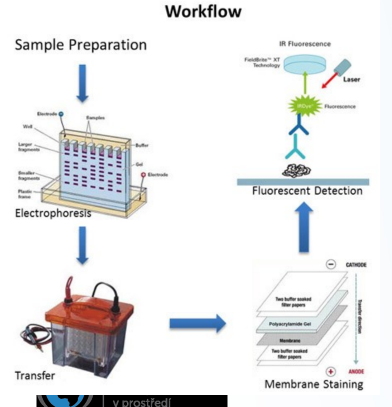


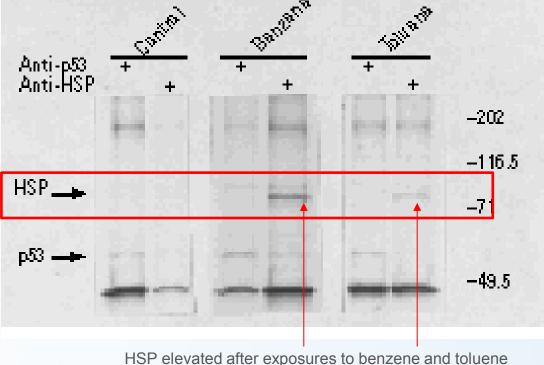


#### HSP determination - example

#### **HSP = GENERAL STRESS biomarker, non-specific**

- phylogenetically conserved (similar genes in most of the organisms)
- structural similarity → 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) → exposure elimination
- assessment ... similar to other whole proteins, i.e. ELISA, Western blotting

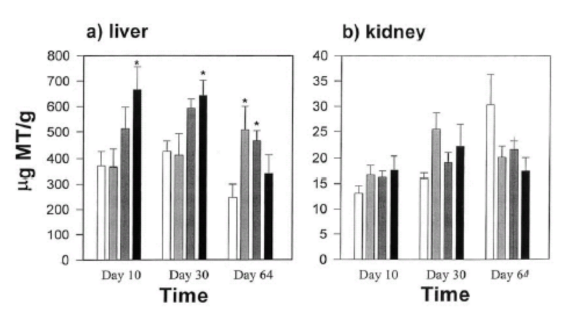


Fig. 2. Metallothionein (MT) concentrations in the (a) livers and (b) kidneys of lake whitefish fed a control diet and three As contaminated diets for 10, 30, and 64 days. Data are expressed as mean ( $\pm$ S.E.). Asterisk denotes mean is significantly different from the control at that duration (P 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

## Protein biomarkers of estrogenicity

**ER** = transcription factor controling number of target genes

Target genes of ER = biomarkers of estrogenicity

Major examples

- Vitellogenin
- Aromatase CYP19A



#### Vitellogenin (Vtg)

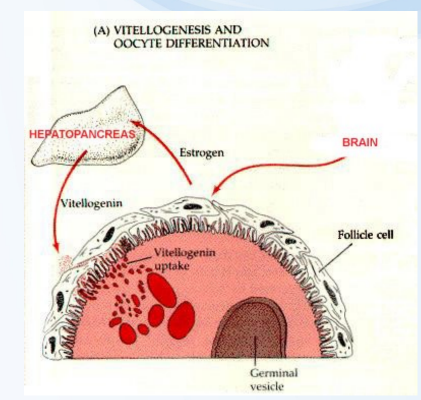
Precursor of yolk proteins, phospho-protein ("energy" rich)

→ egg formations (females) at oviparous animals

Synthesized in liver and distributed via blood / haemolymph

#### Xenoestrogens & other endocrine disruptors

- → increased levels or early production in FEMALES
- → production de novo in MALES





#### Vitellogenin (Vtg) assessment



## Vitellogenin in fish

# **ELISA** in exposed males and females

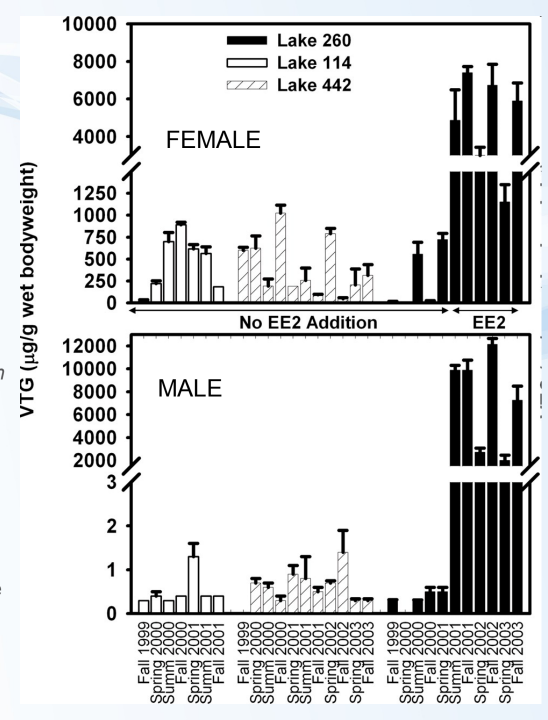
(-) specific antibodies are necessary for each species (low crossreactivity of Abs)

#### Kidd et al. (2007) PNAS

Ethinylestradiol → collapse of fish population

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).





## Aromatase (CYP19A)

Medscape®

- Levels inducible by estrogens
- Catalyzes single enzymatic step androgens → estrogens

## **Experimental assessment**

(in reseach and practice)

1. PCR / Quantitative-Real-Time-PCR

Cholesterol

Pregnenolone

Progesterone

Cortisol

17 a-Hydroxylase

17 a-Hydroxy-progesterone

17, 20-lyase

NS/S

Androstenedione

Testosterone

Aldosterone

Cortisol

NS/S

Estrone

aromatase

Estradiol

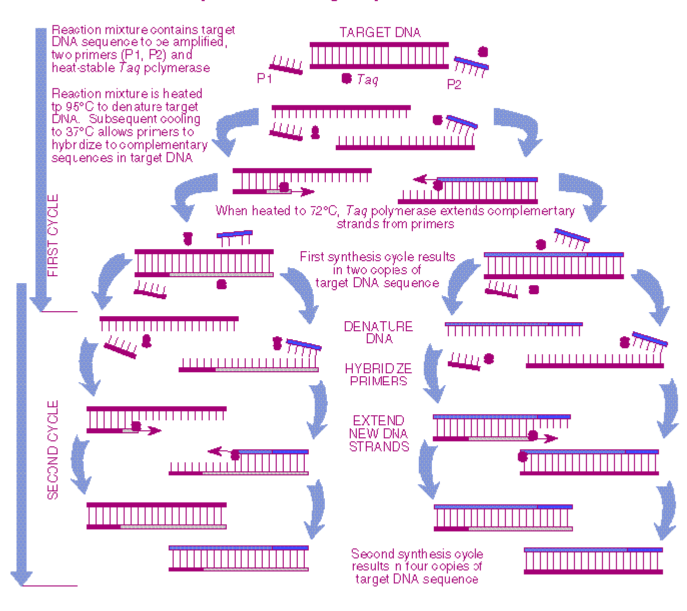
www.medscape.com

2. GM-organisms (zebrafish): reporter gene with GFP
Green Fluorescence Protein under the control of aromatase promoter

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# PCR principle

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



# Visualization of PCR product

## 1) Electrophoresis (qualitative)

## Intercalation dyes

- e.g. **ethidium bromide** 

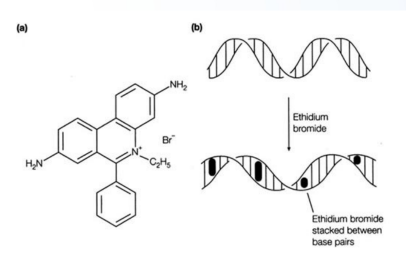
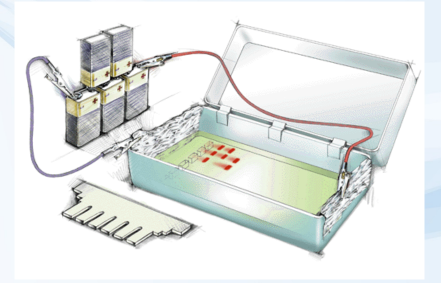
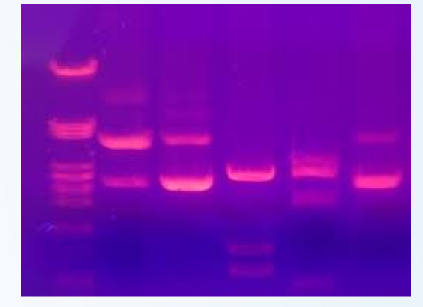


Fig. 3. (a) Ethidium bromide; (b) the process of intercalation, illustrating the lengthening and untwisting of the DNA helix.



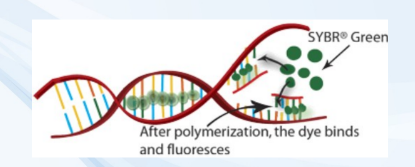


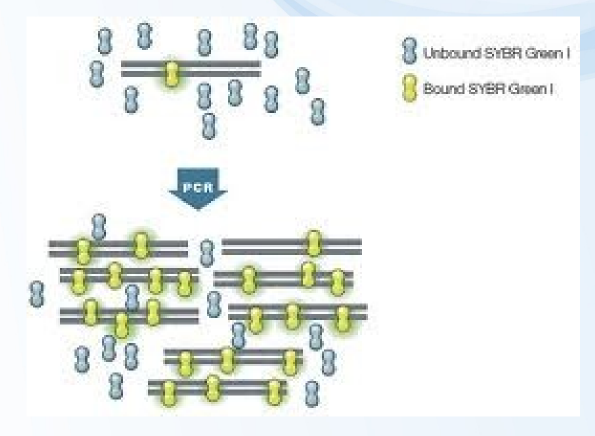


# Visualization of PCR product

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

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



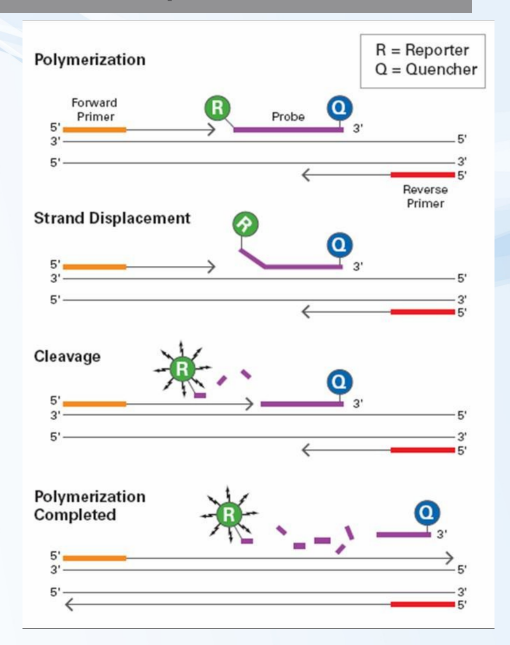




## Visualization of PCR product

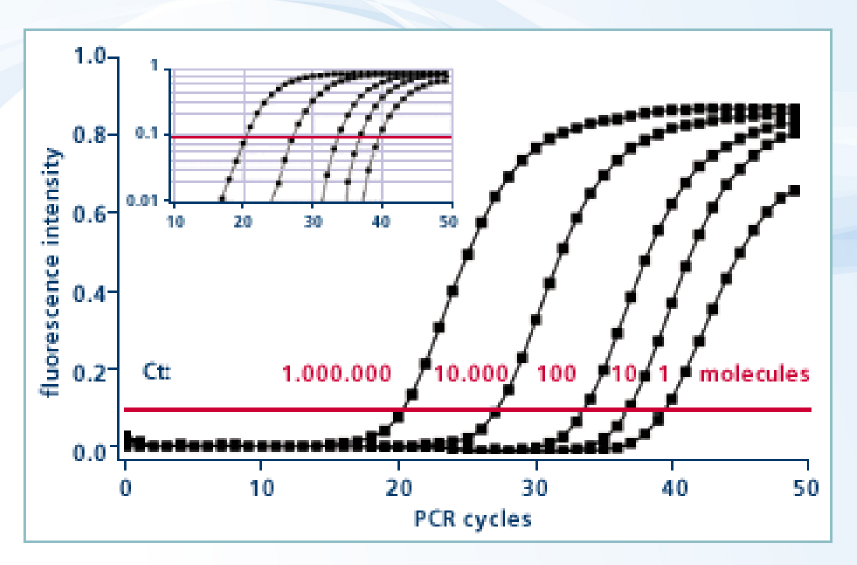
# 2b) Real-time (quantitative) TaqMan probes

(more DNA replications more fluorescent dye released)



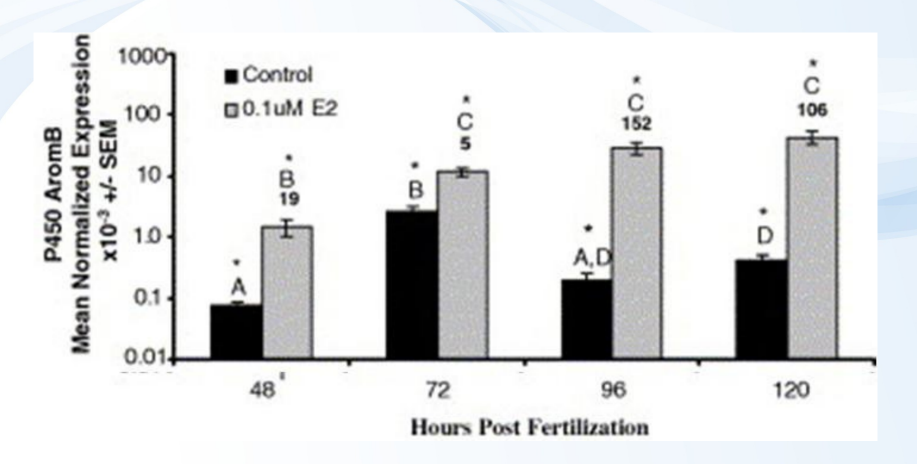


## "Quantitative" determination of PCR product





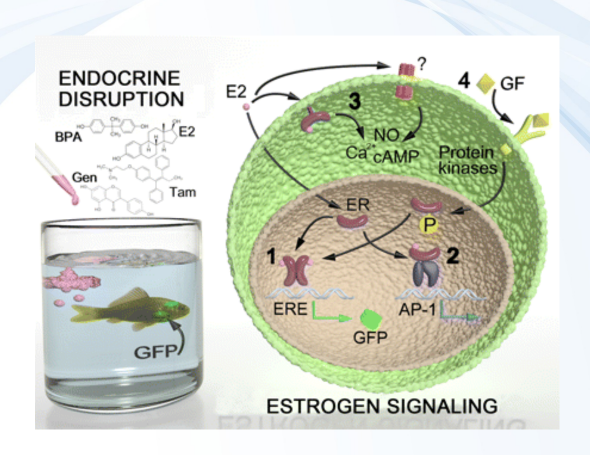
## qPCR determination of the aromatase gene in Zebrafish

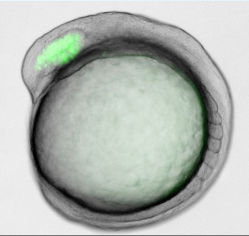


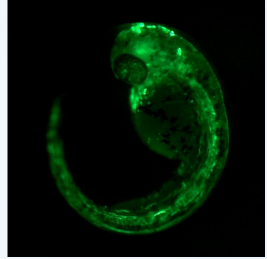
http://dx.doi.org/10.1016/j.ygcen.2005.12.010,



## GFP-reporter for estrogens in zebrafish embryo







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



## DETOXIFICATION / ANTIOXIDANT DEFENCES

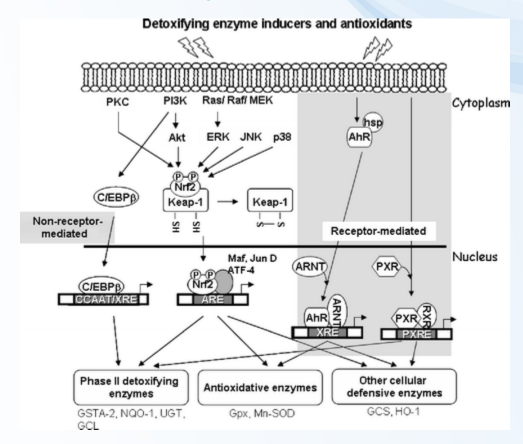
## Inductions of detoxication & oxidative stress enzymes

(hepatopancreas / liver / blood)

MFO - CYP classes - EROD / MROD / BROD

Phase II enzymes (GSTs)

Glutathion metabolism enzymes (GPx, GRs)





# MFO (CYPs) - reminder

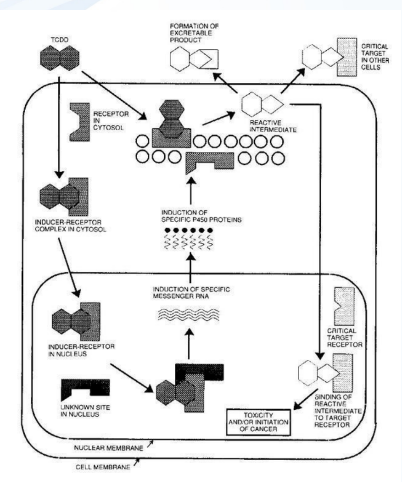


Figure 5.1 Diagram of MFO system. Nebert and Gonzalez (1987).



Table 5.1 Classification of P450s

Nomenclature	Induced by/specificity
P450I	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
P450XXI	Formation of steroid 21-hydroxylases
P450LI	Plant/yeast
P450CI	Prokaryote

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

## Assessment of CYPs - "EROD"

## **Determination of CYP1A1 activity**

## "EROD" - EthoxyResorufin-O-Deethylase activity

Substrate: Ethoxyresorufin

: Oxidation by CYP1A1

→ Fluorescence (easy determination)

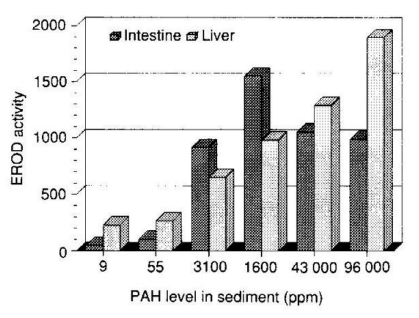
## **EROD** = sensitive biomarker of organic pollution (exposure & effects)

: 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 ...





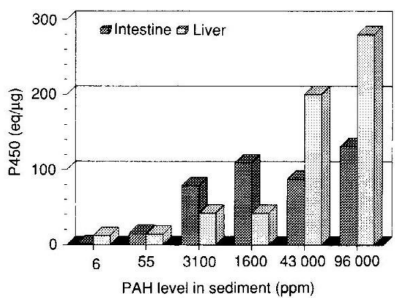
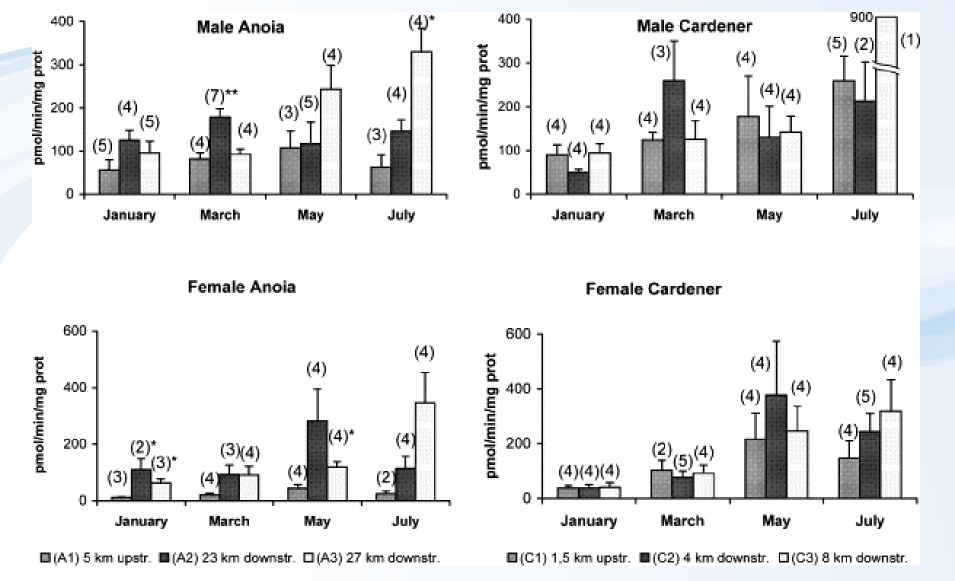


Figure 5.6 Relationship of sediment concentration of PAHs to EROD activity in liver and intestine of spot. After Van Veld et al. (1990).





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



## Phase II conjugation enzymes - GSTs

## Glutathione-S-transferases (GSTs)

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

#### **Methods**

Chemical reaction of

reduced GSH

+ thiol selective probe (CDNB)

**GST** 

GSH + CDNB

 $\rightarrow$ 

**GS-CDNB** (formation of yellow product) kinetic or endpoint determination



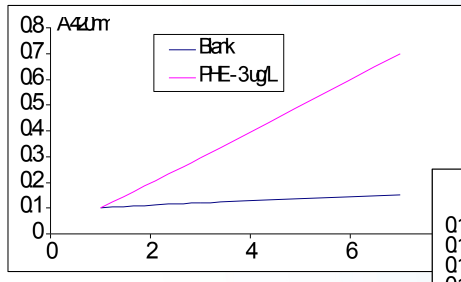


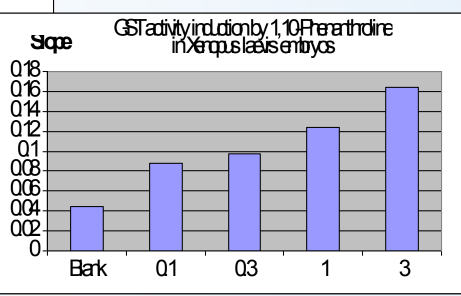
# GST activity determination: example

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 (notes on methods e.g. GST- 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)
  - proteins: oxidized forms (carbonyls)



## Oxidative stress markers

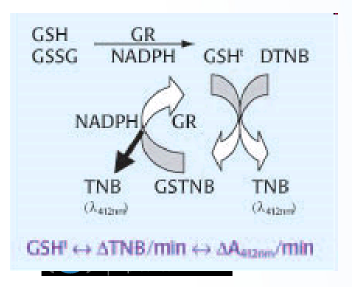
## **GSH**

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

## Total glutathione = reduced GSH + oxidized GSSG

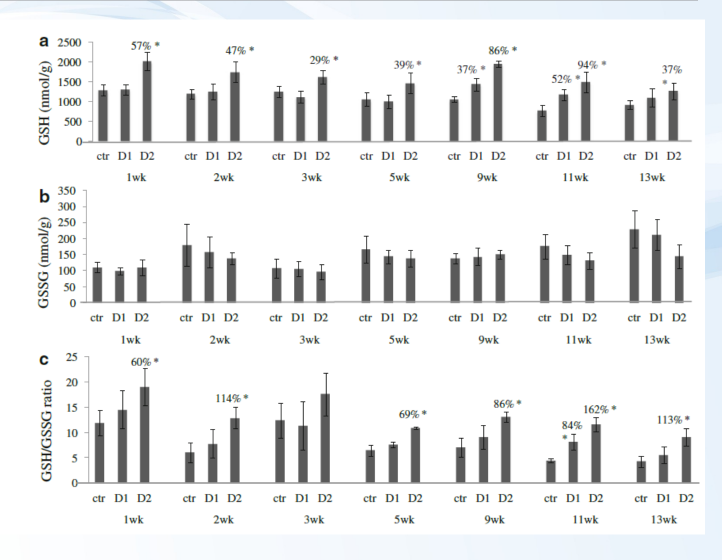
#### Methods of determination of GSH

- Spectrophotometry: GSH + Ellman s reagent (DTNB) → Reduced GSH
- LC-MS/MS selective assessment of GSH and GSSG



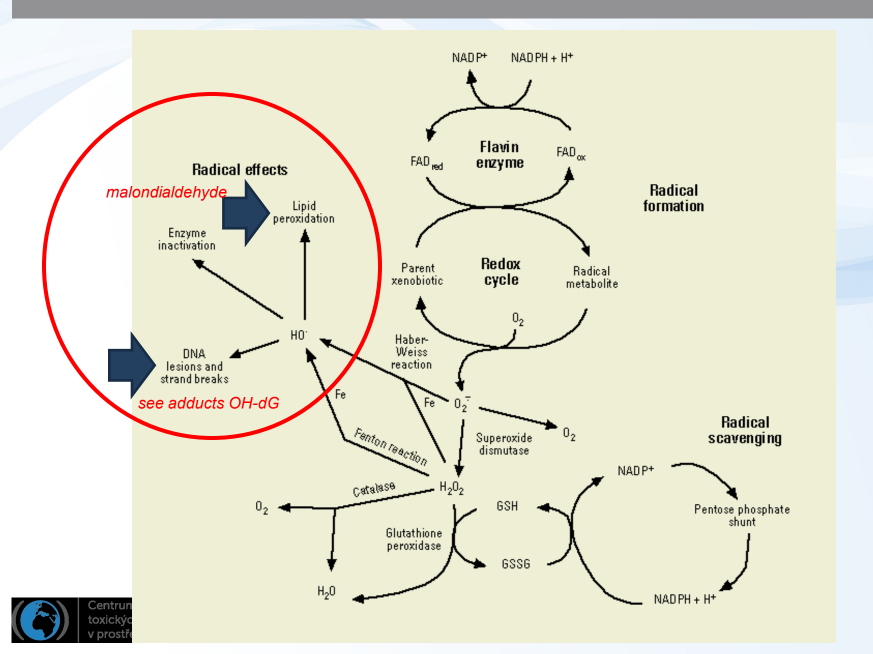
## Example - GSH modulation by toxic nanoparticles

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*=5 animals)





## Markers of oxidative DAMAGE

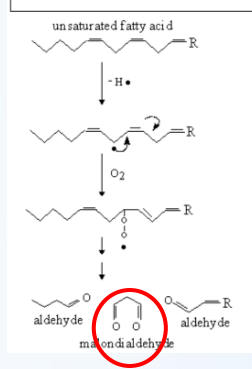


## Lipid peroxidation → Malondialdehyde (MDA)

#### MDA – malondialdehyde

product of lipid peroxidation

#### STEPS OF LIPID PEROXIDATION





## Malondialdehyde (MDA) determination

#### MDA – formed from oxidized membrane phospholipids

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

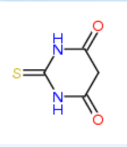
## **TBARS – ThioBarbituric Acid Reactive Species**

: 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)



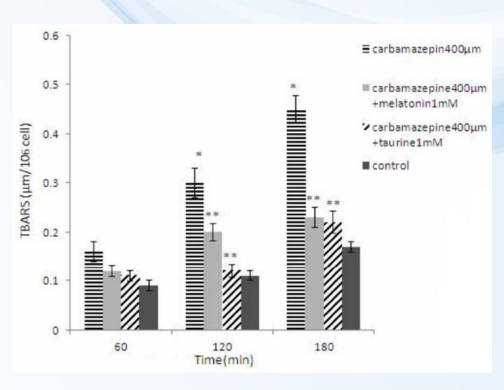
**TBA** 

## MDA modulation - examples

Effects of nanoFeOxide particles on MDA in fish

2.0 -1.8 1.6 MDA content (nmol-mg<sup>-1</sup> 14 1.2 10-0.8-0.6

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





50

100

200

nano-Fe<sub>1</sub>O<sub>2</sub> concentration (mg·L<sup>1</sup>)

400

800

1600

0.4

0.2 -

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