

# BIOMARKERS AND TOXICITY MECHANISMS 12 - BIOMARKERS of EXPOSURE and SUSCEPTIBILITY

Luděk Bláha, PřF MU, RECETOX www.recetox.cz

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









### Biomarkers of Exposure

#### Biomarkers of internal and effective dose

depends on toxicokinetics

#### Biomarkers of internal dose (short / long term)

- examples: Cd in urine, DDE in fat tissues
  - should be easy to sample (urine, breath)
  - instrumental analytical methods (analyses of toxicant)

#### Biomarkers of effective dose

- the chemical interacted with the biological target
  - → analyses of ADDUCTS

Two types of adducts: selective and non-selective



#### SELECTIVE ADDUCTS OF TOXICANTS with BIOMOLECULES

#### **SELECTIVE = CHEMICAL-SPECIFIC**

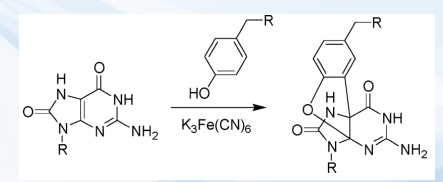
Adducts with DNA

<u>styrene</u>-oxide-O6-guanine N7-guanyl-<u>aflatoxin</u> B1

Hemoglobin-pesticides adduct

#### **Methods of analyses:**

- analytical chemistry
  - extraction from biological sample
  - chemical determination by HPLC or GC



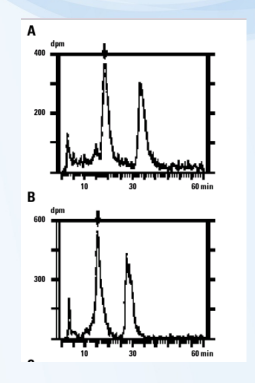
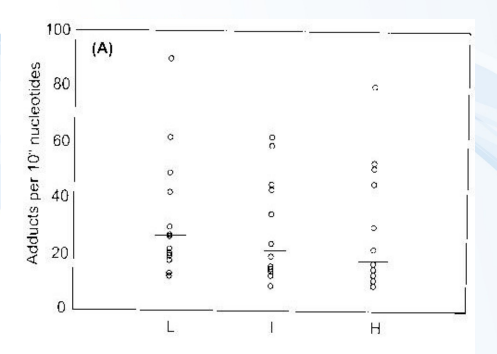




 Table 1
 Reported human haemoglobin adduct levels for various xenobiotics

Chemical (type of exposure)	Adduct/analyte	Method	Adduct level (nmol g - haemoglobin)
N, N-Dimethylformamide (occupational)	3-Methyl-5-isopropylhydantoin	Hydrolysis; GC-MS	75–1000 (exposed) 4–12 (control)
Epichlorohydrin (occupational)	N- (2, 3-Dihydroxypropyl)valine	Modified Edman; GC-MS	0.020 (exposed smokers) 0.007 (exposed non-smokers) 0.013 (control smokers) 0.007 (control non-smokers)
Acetaminophen (drug overdose)	3-(Cystein-S-yl)acetaminophen	Immunoassay	100-4100
PAHs (occupational)	BPDF-Hb	Spectrofluorimetry	0.005-0.139
Ethylene oxide (occupational)	N- Hydroxyethylvaline	Modified Edman; GC-MS	5–20 (exposed) 0.1–0.5 (control smokers) 0.01–0.1 (control non-smokers)
Ethene (occupational)	N- Hydroxyethylvaline	Modified Edman; GC-MS	0.02
Propylene oxide (occupational)	N- Hydroxypropylvaline	Modified Edman; GC-MS	0.05-3.5 (exposed) < 0.02 (unexposed)
Acrylonitrile (smoking)	N- Cyanoethylvaline	Modified Edman; GC-MS	0.09
NNK (smoking)	4- Hydroxy-1-(3-pyridyl) butan-1-one	Hydrolysis; GC-MS	0.0015 (smokers) 0.0005 (non-smokers)
4-ABP (smoking)	4-ABP-cysteine	Hydrolysis; GC-MS	0.00025–0.0025 (smokers) 0.00005–0.0005 (non-smokers)
Acrylamide (occupational, smoking)	N- (2-Carbamoylethyl)valine	Modified Edman; GC-MS	9.5 (production workers) 0.054 (laboratory workers) 0.116 (smokers) 0.031 (non-smokers)
Butadiene (occupational)	N- (2,3,4-Trihydroxybutyl)valine	Modified Edman; GC-MS	0.010-0.014 (exposed) 0.002-0.003 (control)
Styrene (occupational)	2-Phenylethanol	Cleavage with Raney nickel, GC-MS	3.7–8.0 (exposed) 2.0–8.6 (control)



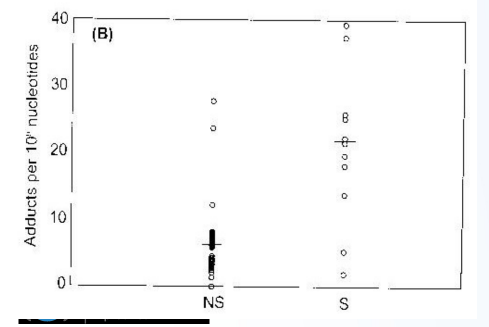
#### PAH-DNA adducts

PAH (polycyclic aromatic hydrocarbons)

\* often high variability

\* may have difficult interpretation

Occup. exposure (Low / Intermed. / High)



Occupational
Non-exposed (NS)
vs.
Exposed (S)

#### Non-selective adducts

 binding with macromolecules (DNA, proteins) with no further information on the structure of actual adduct (i.e. causative agent not clear)

### Typical nonselective biomarker methods

- <sup>32</sup>P-postlabelling assay
- oxidized DNA: 8-hydroxy-2'-deoxyguanosine



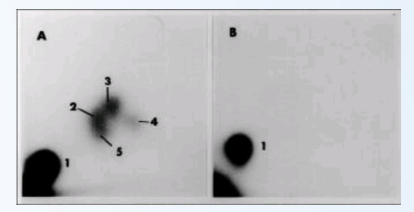
# <sup>32</sup>P-postlabelling assay principle

- Digestion of NA
- •Enzymatic labelling with 32P (kinase)
- •TLC or HPLC analyses of products

# TLC result (thin layer chromatography)

A - 2-5 = various adducts

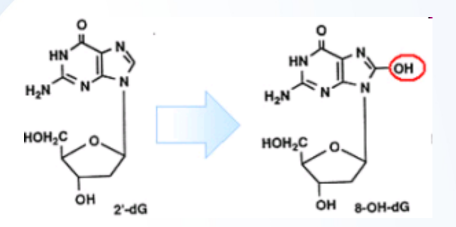
B - controls



#### 8-hydroxy-2 -deoxyguanosine analysis

#### Oxidative damage to DNA

- many causes → 8-OH-dG is the most common marker of DNA oxidation



#### **Analysis: analytical chemistry methods**

- HPLC
- immunochemistry (ELISA)

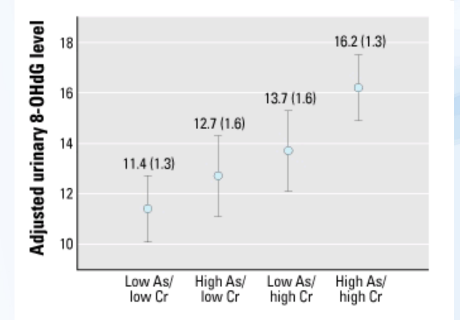
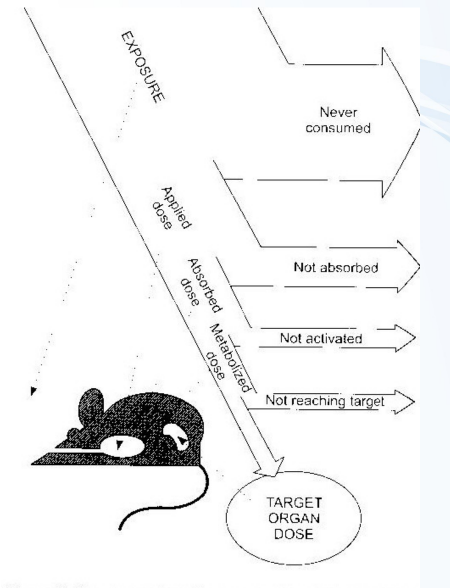


Figure 1. Adjusted urinary 8-OHdG level (ng/mg creatinine) by urinary arsenic and urinary chromium concentrations. Values shown are mean  $\pm$  SE. Cut points were determined according to medians (arsenic, 7.7 µg/g creatinine; chromium, 2.0 µg/g creatinine) of urinary creatinine-adjusted levels among all subjects.



## Biomarkers of susceptibility





**Figure 2** Representation of the relationships between ambient exposure and critical target dose and the progressive decrease in effective exposure due to various biological barriers. Source: *Low-Dose Extrapolation of Cancer Risks: Issues and Perspectives*, p. 188. Used with permission. c. 1995 International Life Sciences Institute, Washington, DC, U.S.A.

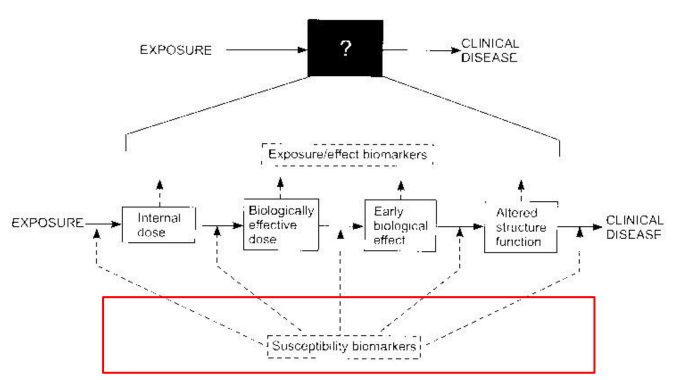
#### **Toxicokinetics**

determines susceptibility of an individual at various levels

# → Biomarkers of susceptibility

Will the individual be sensitive? Will patient respond to a drug?

#### Importance of susceptibility biomarkers



**Figure 1** The biomarker paradigm linking exposure with disease and showing expansion of the classical epidemiological 'black box' to reveal discrete mechanistic stages. Reprinted with permission from *Environ, Sci. Technol.* (1997) **31**, pp. 1837–1848. Copyright 1997 American Chemical Society.



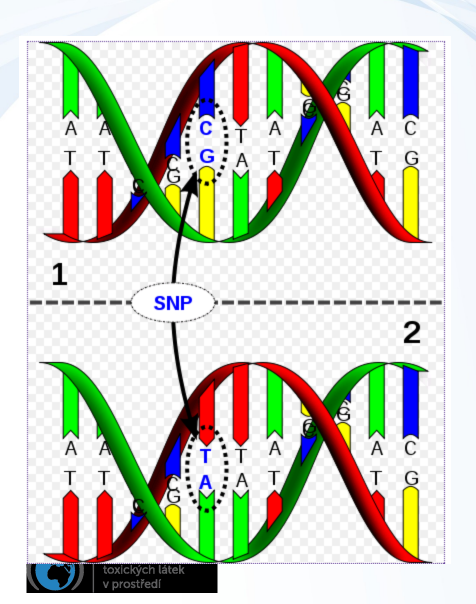
#### Biomarkers of susceptibility

#### Susceptibility depends on genotype and metabolism

- genetic polymorphism in detoxification enzymes
- variability in specific isoenzymes
- → susceptibility to "activate" toxicants: example: N-acetylation of arylamines NAT2
  - → susceptibility to genotoxins
- → family cancers
- → susceptibility to drugs (including anticancer drugs)



# Example: genetic polymorphism SNPs - single nucleotide polymorphism



#### **SNPs**

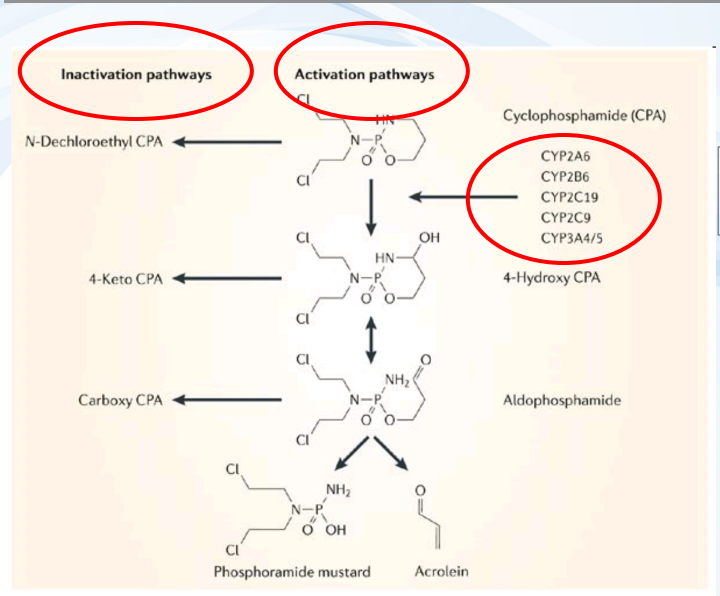
- → affects protein functions
- → in specific cases (see example) some SNPs identified

#### → PERSONALIZED MEDICINE

To identify SNP as a biomarker

Many genotypes (from many individuals) must be sequenced and compared with phenotype (e.g. responsiveness to certain drug)

#### Cyclophosphamide (anticancer drug) and its toxicity





Copyright © 2006 Nature Publishing Group Nature Reviews | Cancer

#### Example: genetic polymorphism

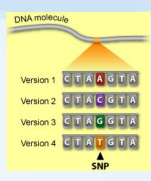
#### **CYP450 Enzymes and Polymorphisms**

	_	-	-	_	
/	b	_	_	١.	А
C	n	0	С	ш	9)
`	c				7

Diagnostics

Enzyme	Fraction of drug metabolism	Major polymorphisms
CYP3A4	40-45%	Rare
CYP2D6	20-30%	'2xn, '4, '10, '17, '41
CYP2C9	10%	<b>'2, '3</b>
CYP2C19	5%	<b>'2, '3</b>
CYP1A2	5%	'1K
CYP2B6	2-4%	-
CYP2E1	2-4%	-
CYP2A6	2%	<b>'4, '9</b>
CYP2C8	1%	*3
CYP3A5	<1%	.3

Alleles known to be involved in polymorphism



The CYP 2D6 gene is extremely polymorphic with more than 70 allelic variants described so far <sup>1</sup>

Ingelman-Sundberg, TRENDS in Pharmacological Sciences, Vol. 25 No.4 April 2004 <sup>1</sup> Dahl, Clin. Pharmacokinet 2002; 41 (7); 453-470

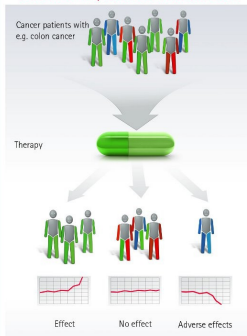
AMPLI@HIP

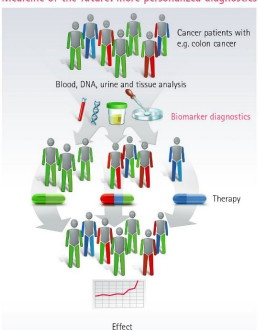


#### Personalized medicine

#### Personalized medicine: tailored treatments

Medicine of the present: one treatment fits all Medicine of the future: more personalized diagnostics



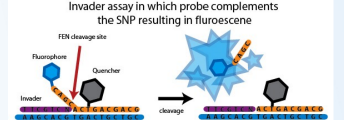


Different people respond differently to the same therapy: while one treatment brings about the desired success in one group of patients with e.g. colon cancer, it does not change the condition of other groups at all, or even leads to adverse effects (left). The reason: the genetic makeup and metabolic profile of each individual patient influences the effect of a drug. Personalized medicine takes these individual patterns of cellular and metabolic products into account in the diagnostic phase: biomarker diagnostics separates patients into groups with similar characteristics, and provides information on the best individual treatment. This should enable all patients to benefit from their own, "personal" therapy.



#### SNP diagnostics:

- 1) DNA isolation
- 2) Multiplication of specific gene eg. CYP
- 3) SNP identification
- ... Molecular biology methods such as
- \* NA sequencing
- \* Probe pairing ... number of variants



Invader assay in which probe mismatches at the SNP location preventing cleavage from occuring

Fluorophore

Quencher

Invader

A G G A C G G A C G A

