

# Glutathione Conjugation

- I. Glutathione Metabolism, Homeostasis
- II. GSH Conjugation Reactions
- III. Glutathione S-Transferases
- IV. Reactions of GSH Conjugates

### Suggested Reading

Bernard Testa, Stefanie D. Krämer. The Biochemistry of Drug Metabolism - An Introduction : Part 4. Reactions of Conjugation and Their Enzymes. Chemistry and Biodiversity, vol. 5(11): 2171-2336 (2008)

Methods in Enzymology vol 401(2005). GST and  $\gamma$ -glutamyltranspeptidases

Frova, Glutathione transferases in the genomics era. Biomolecular Engineering 2006 vol 23:149-169.

Hayes JD, Flanagan JU, Jowsey IR. Glutathione Transferases. Annu Rev Pharmacol Toxicol. 2004

Sheehan D, Meade G, Foley VM, Dowd CA. Structure, function and evolution of glutathione transferases: implications for classification of non-mammalian members of an ancient enzyme superfamily. Biochem J. 2001 Nov 15;360(Pt 1):1-16

Townsend DM, Tew KD. The role of glutathione-S-transferase in anti-cancer drug resistance. Oncogene. 2003 Oct 20;22(47):7369-75

Dekant W. Chemical-induced nephrotoxicity mediated by glutathione S-conjugate formation. Toxicol Lett. 2001 Oct 15;124(1-3):21-36

van Bladeren PJ. Glutathione conjugation as a bioactivation reaction. Chem Biol Interact. 2000 Dec 1;129(1-2):61-76

Anders MW. Glutathione-dependent bioactivation of haloalkanes and haloalkenes. Drug Metab Rev. 2004 Oct;36(3-4):583-9.

# I. Glutathione Metabolism and Homeostasis

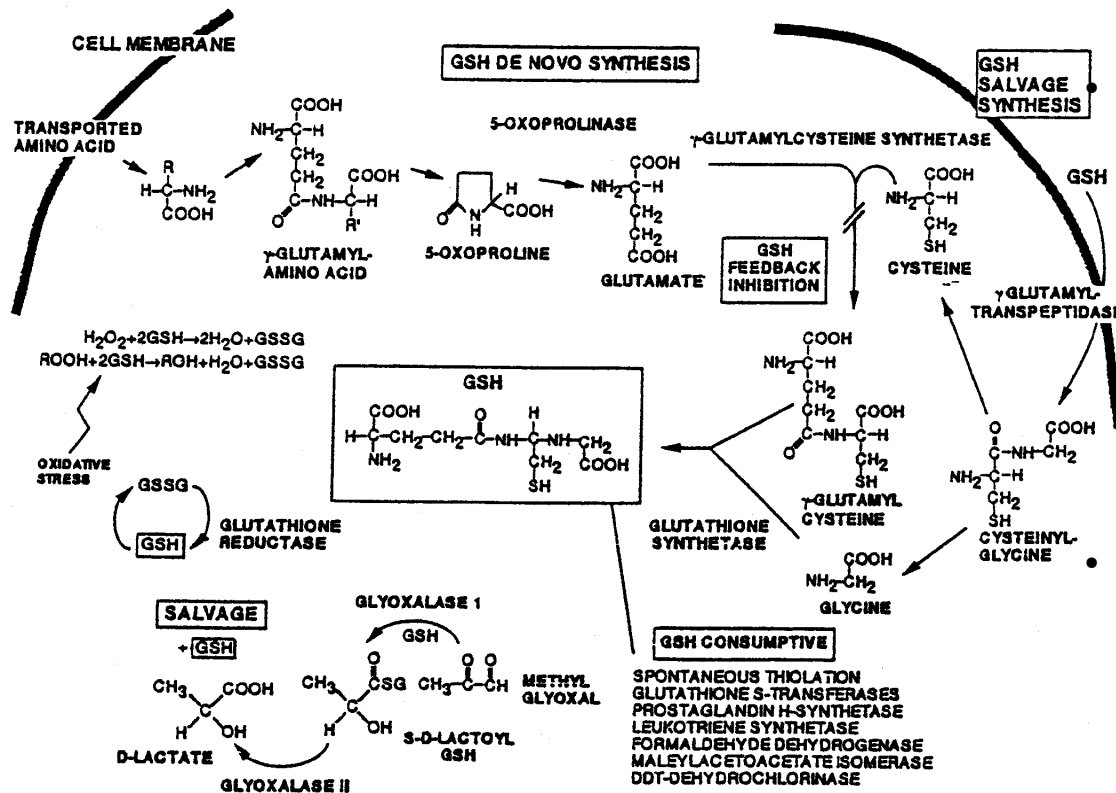


Fig. 1. Summary schema of interrelated GSH-producing and -utilizing pathways. GSSG, reduced glutathione; DDT, bis(*p*-chlorophenyl)-trichloroethane.

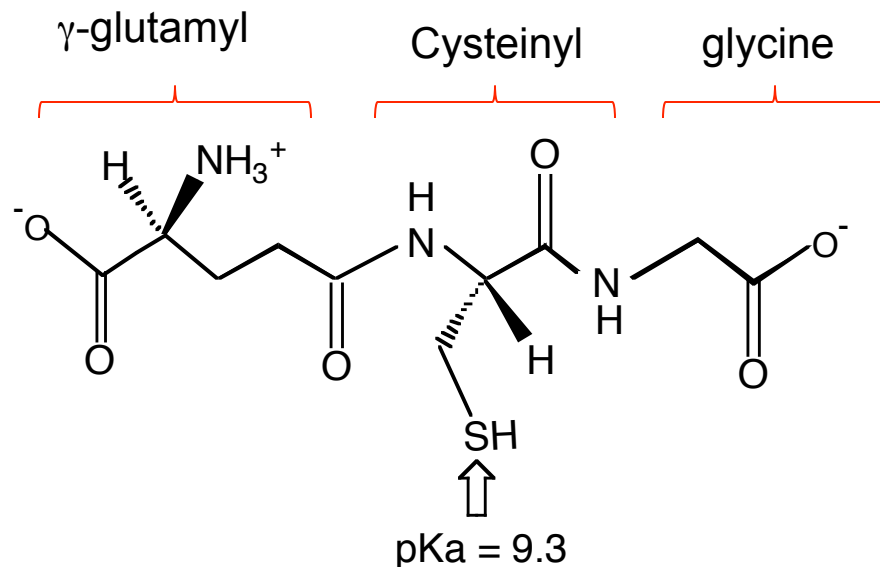
important in:

- xenobiotic deactivation,
- regulating 'redox' state of cells, maintenance of reduced cysteines,
- general oxidative stress responses

tissue distribution is variable, but generally high (1-15 mM). In hepatocytes there are two pools of GSH:

mitochondrial pool ~ 20%  
cytosolic pool ~ 80%

# I. Glutathione: Chemical Properties



- Nucleophilic cysteine-chemical strategy is to have a good nucleophile on a polar, water-soluble, co-factor. In contrast to UDPGA and PAPS, **electrophilic drugs react.**
- Redox active cysteine
- Unusual peptide linkage; γ-glutamyl-cys-gly

# I. Glutathione Metabolism & Homeostasis

- Enzymatic Reactions of GSH



Detoxification of peroxides, control of oxidative stress



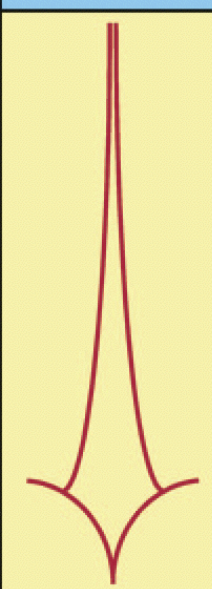
Maintenance of GSH levels



Protein folding, protein regulation

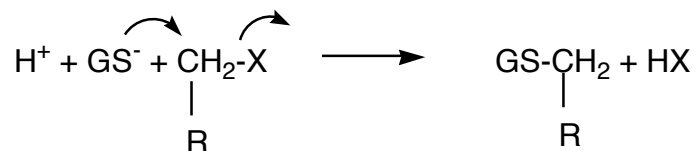
**Glutathione S-transferases:** detoxification

## II. GSH Reactions

A ranking of nucleophilic and electrophilic sites in bio(macro)molecules		
Nucleophilic sites	Increasing hardness	Electrophilic sites
Thiol groups (R-SH) in Cys residues in proteins and peptides ( <i>e.g.</i> , GSH)		Aldehydes, polarized double bonds
Thioether groups (R-S-R') in Met residues		Epoxides, strained lactones, alkyl sulfates, alkyl halides
Amino groups in Arg, Lys, and His residues		Arylcarbonium ions
Amino groups of bases in RNA and DNA		Benzylic carbonium ions, nitrenium ions
Oxygen atoms (=O) of purines and pyrimidines		Alkylcarbonium ions
Phosphate oxygen (P=O) of RNA and DNA		

GSH is a 'soft' nucleophile due to the polarizability of Sulfur. GSH 'prefers' soft electrophiles.

## II. GSH Reactions



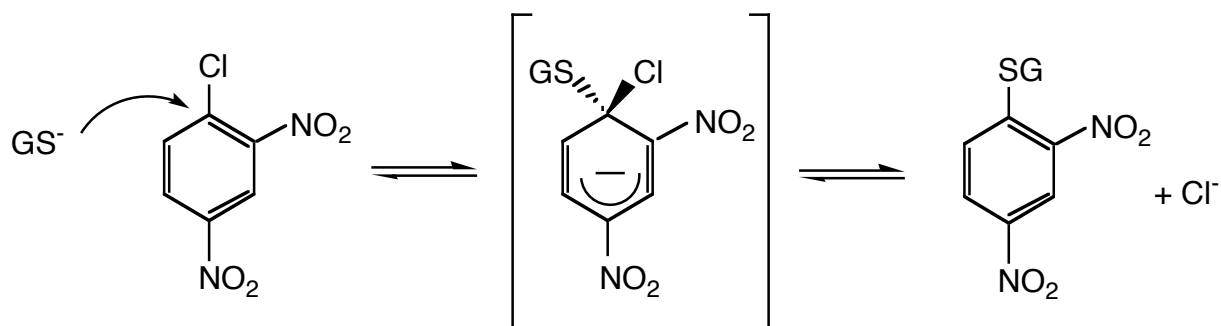
R = alkyl, aryl, benzylic, allylic

X = Br, Cl, I,  $\text{OSO}_3^-$ ,  $\text{OSO}_2\text{R}$ ,  $\text{OPO(OR)}_2$

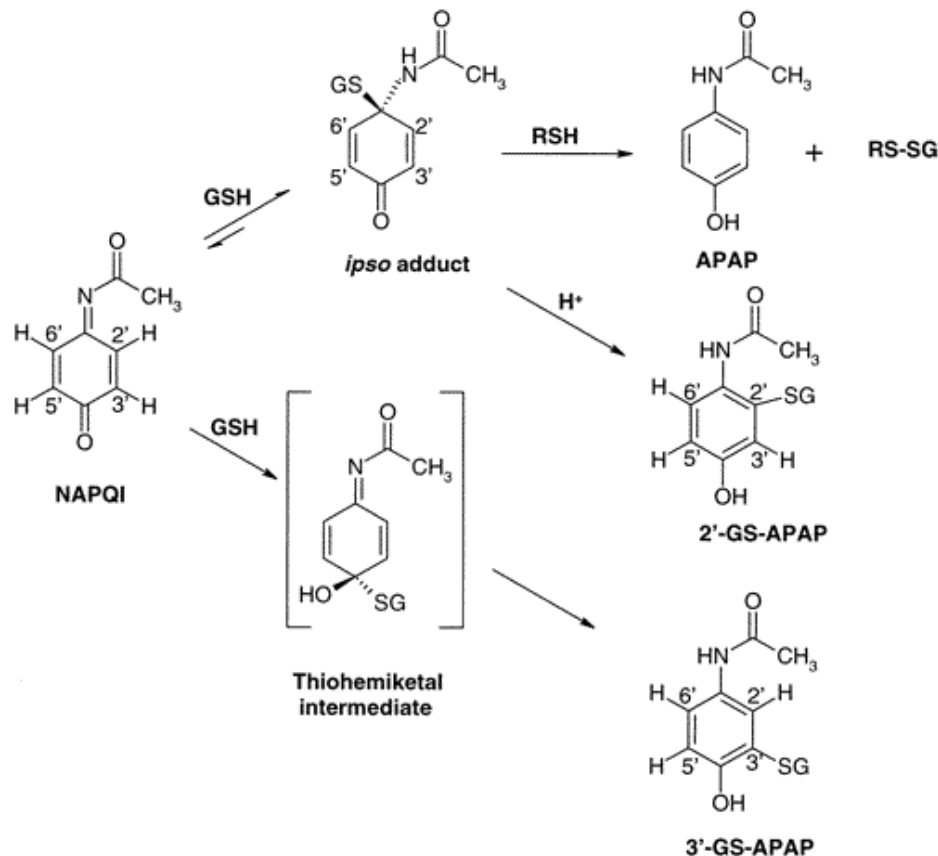
### A. nucleophilic substitutions and nucleophilic aromatic substitutions

the 'universal' GST substrate

1-chloro-2,4-dinitrobenzene (CDNB)



## II. GSH Reactions



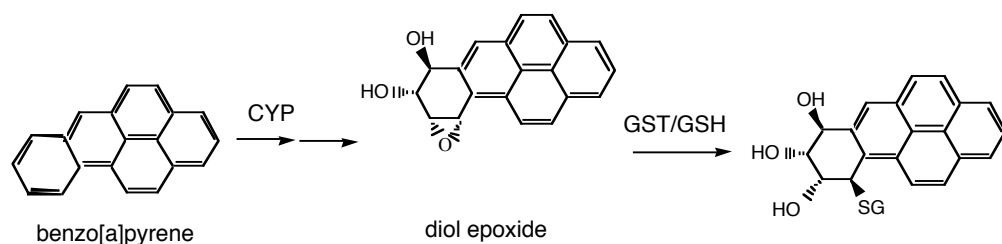
B. nucleophilic additions  
ketones, aldehydes,  
esters, nitriles,  $\alpha,\beta$   
unsaturated Michael  
acceptors.

e.g. NAPQI

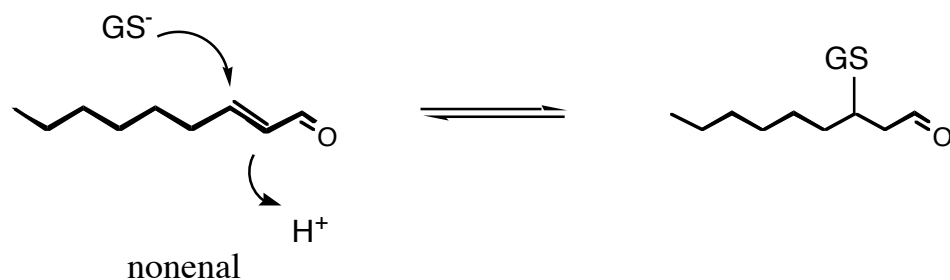


## II. GSH Reactions

### B. Nucleophilic Additions (cont'd)



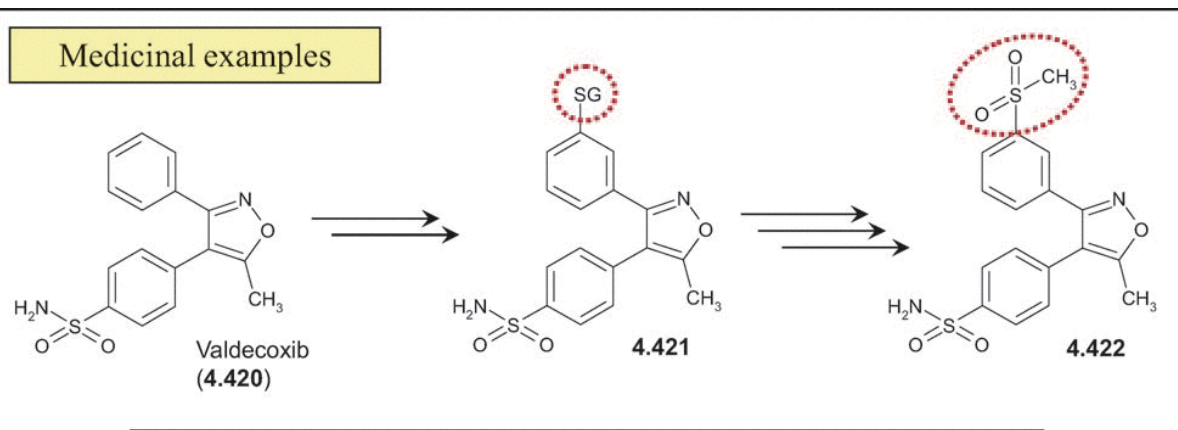
e.g. epoxides



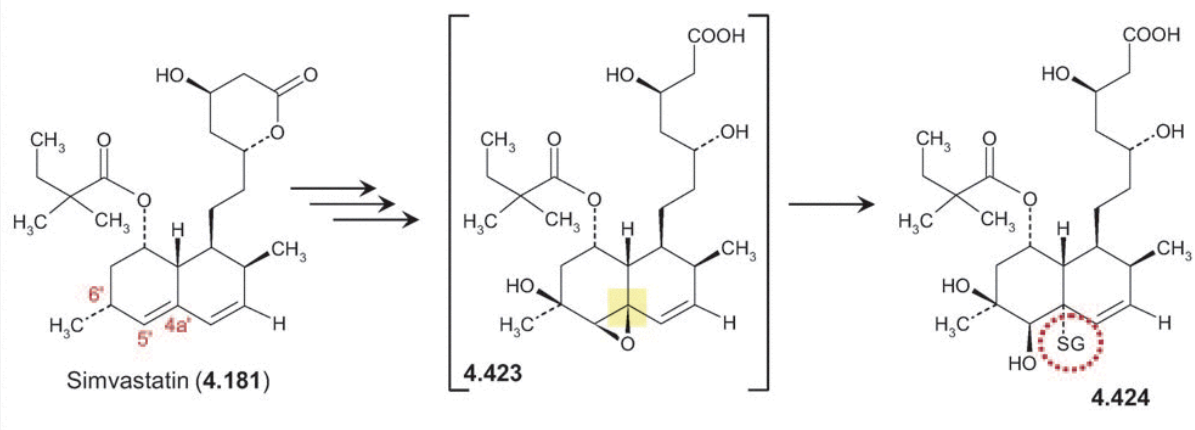
e.g.  $\alpha,\beta$ -unsaturated  
aldehydes

## II. Reactions: Examples of GSH addition to Drug Epoxide Metabolites

Epoxidation,  
followed by  
GSH attack  
and  
dehydration



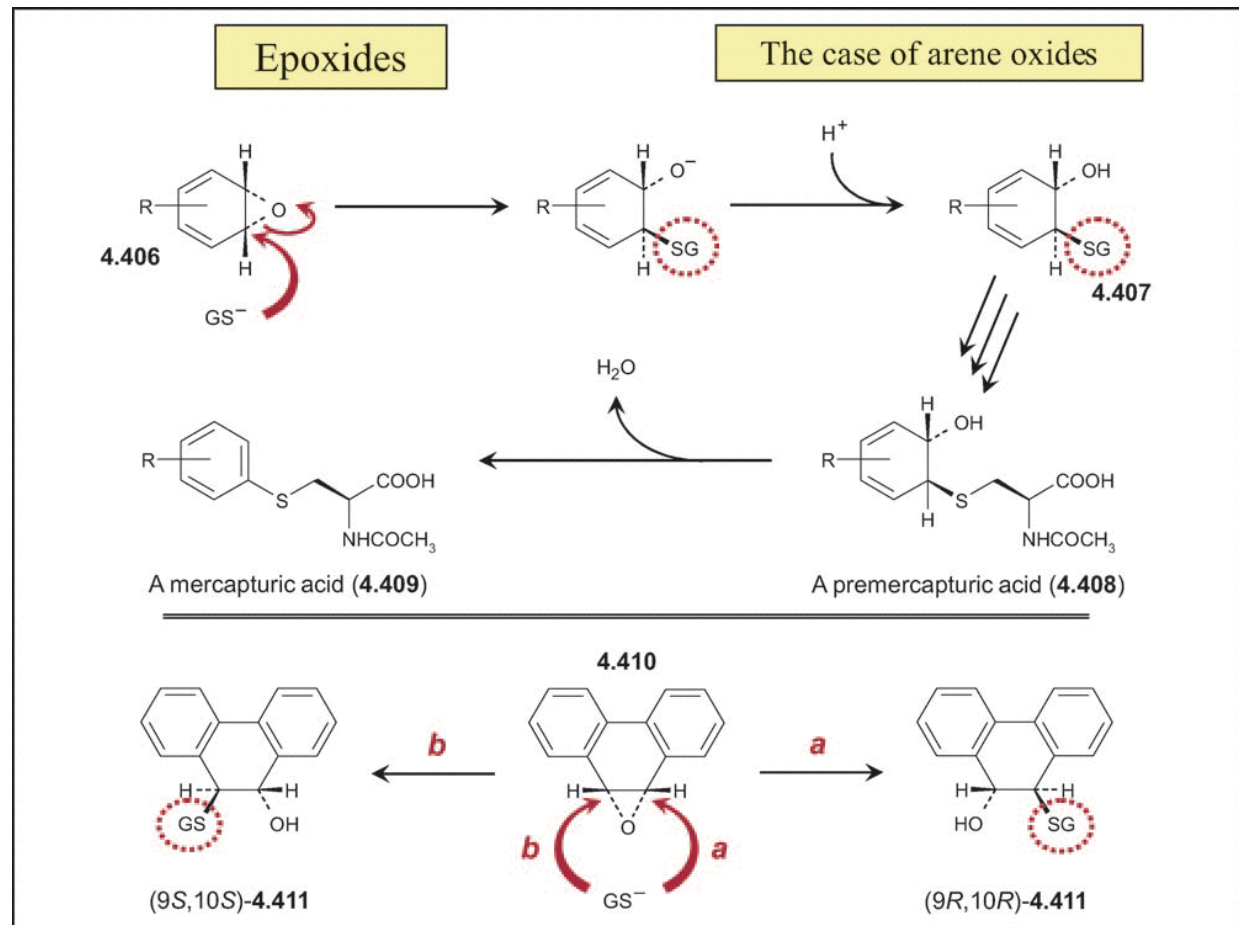
Epoxidation,  
followed by  
GSH attack  
but no  
dehydration



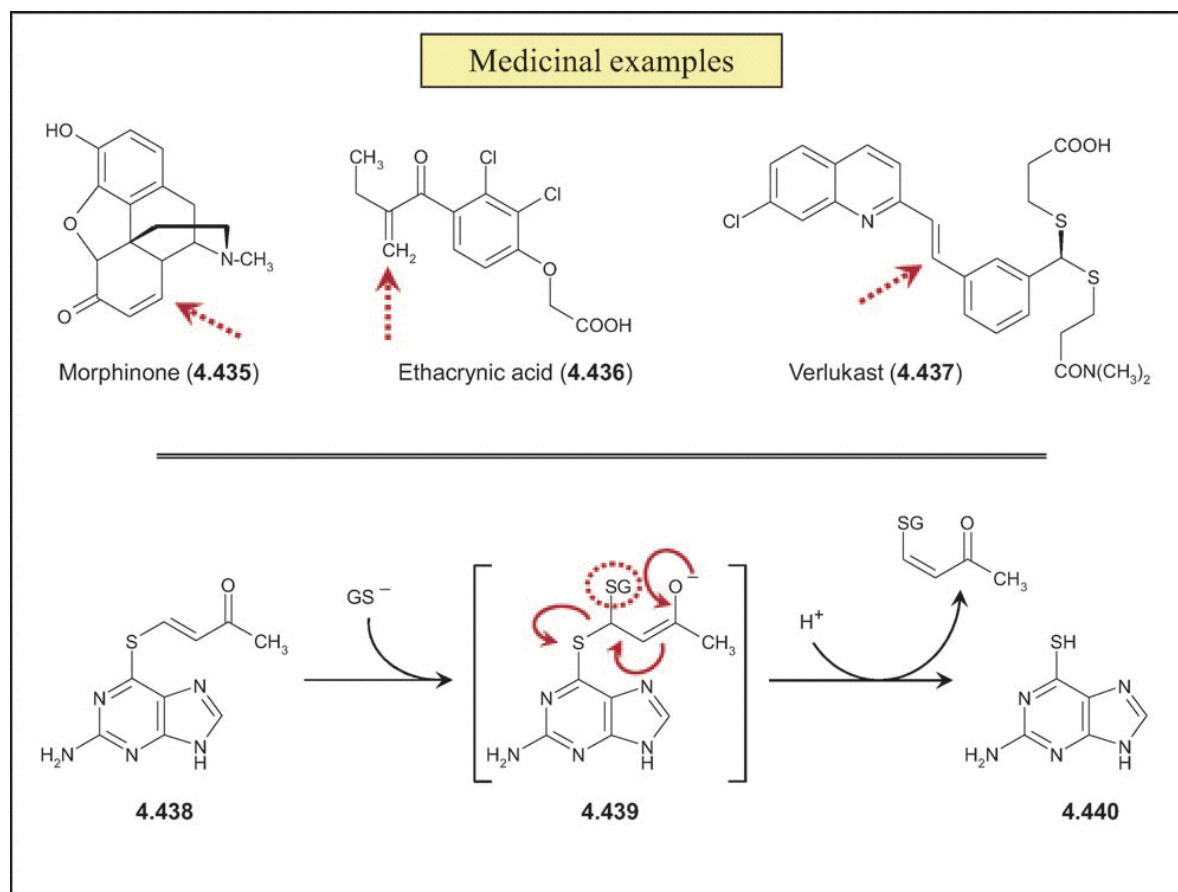
## II. GSH Reactions Addition to Epoxides: Mechanistic Aspects

$S_N2$ -type, 'inversion' of stereochemical configuration. And here, aromatization drives dehydration, after mercapturic acid pathway.

Regio- and stereoselectivity of addition depends on the GST isoform.



## II. GSH Reactions: Polarized Alkenes, Alkenals



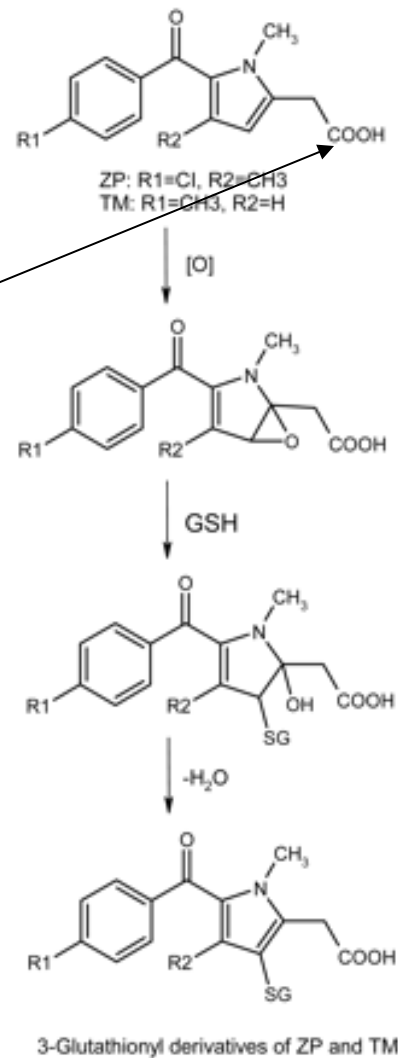
Addition-elimination has been used as a Glutathione-dependent strategy for pro-drug activation.

## II. GSH Reactions: Trapping 'bioactivated drugs'

Contrast to glucuronidation:

Zomepirac, Tolmetin are 'toxic' NSAIDs.

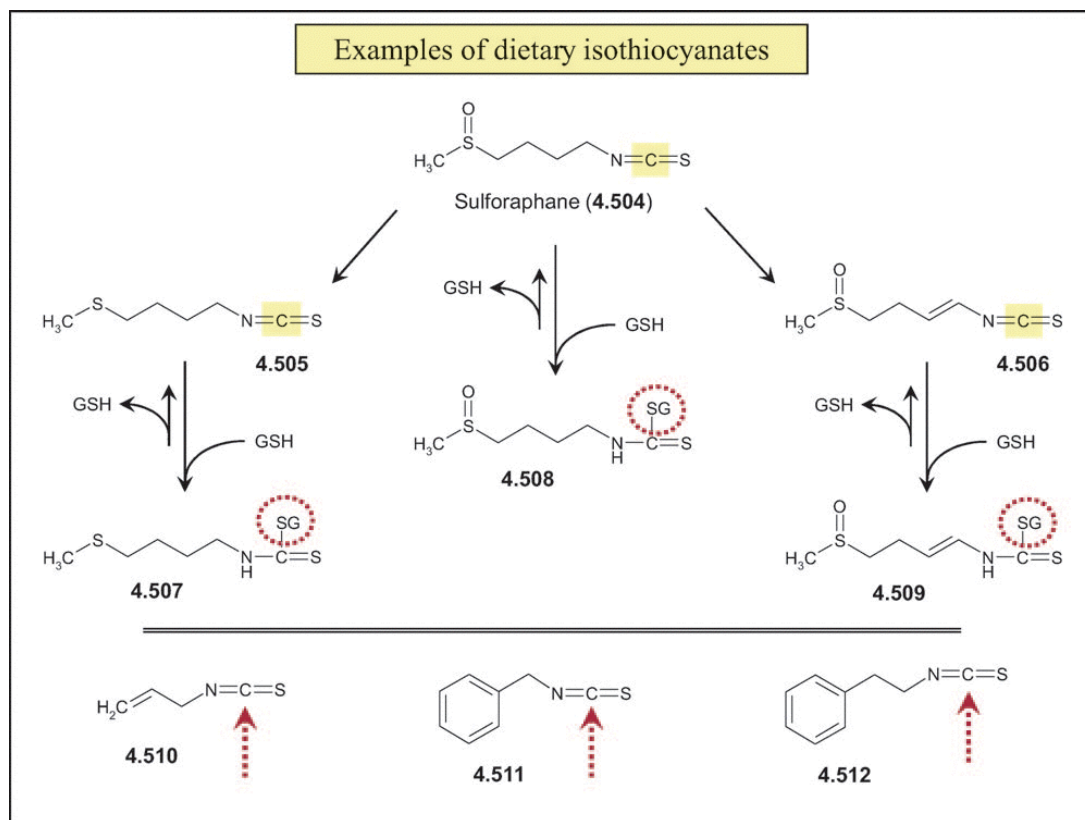
Dogma has been: acyl glucuronide adducts proteins.



- Epoxidation - metabolic activation of zomepirac.
  - CYP-dependent epoxidation, followed by GSH addition.
- epoxidation: alternative possible mechanism for toxicity of zomepirac, vs. acyl glucuronide formation.

Chen et al., 2005 DMD 34:151-155.

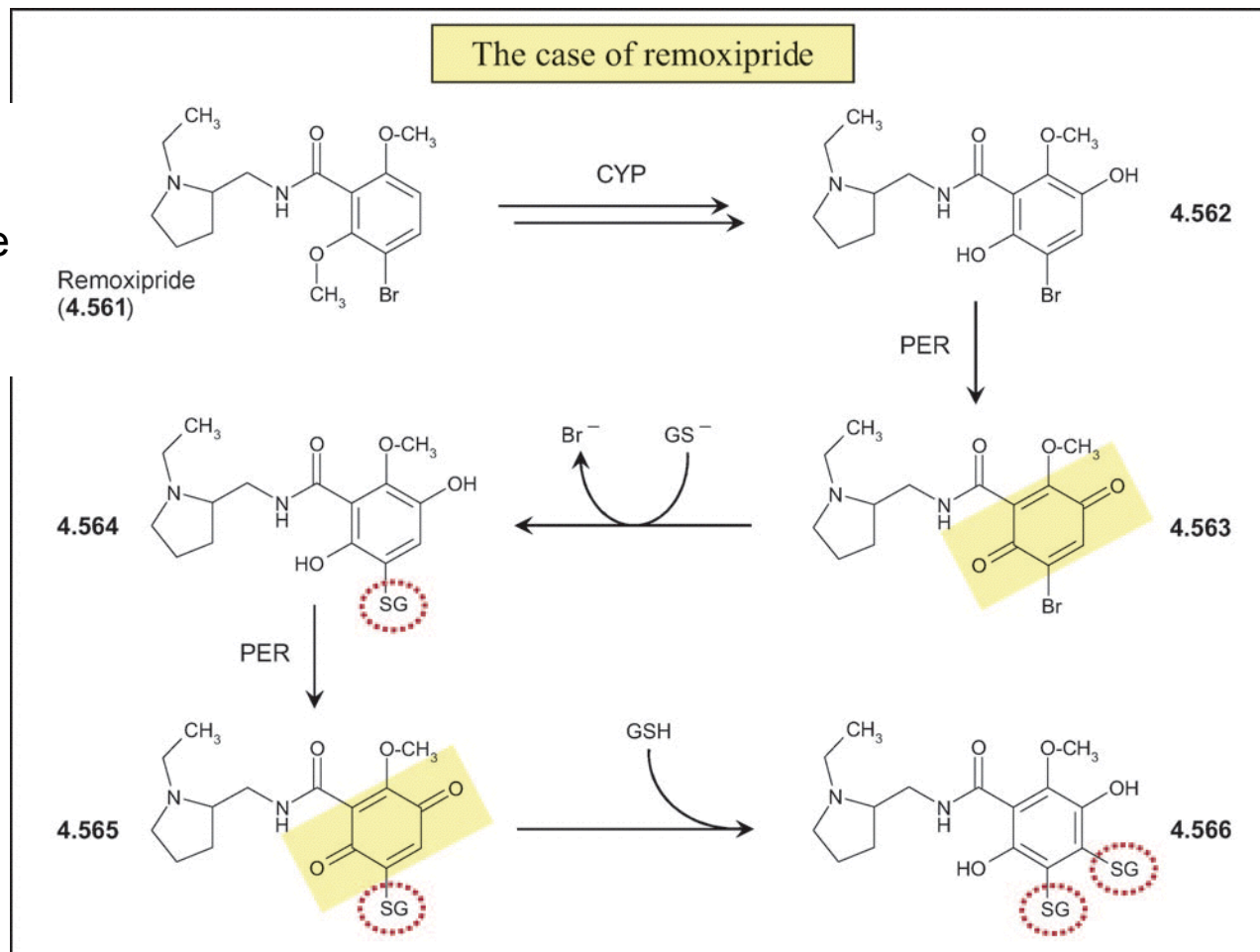
## II. GSH Reactions: Addition to Isocyanates and Isothiocyanates



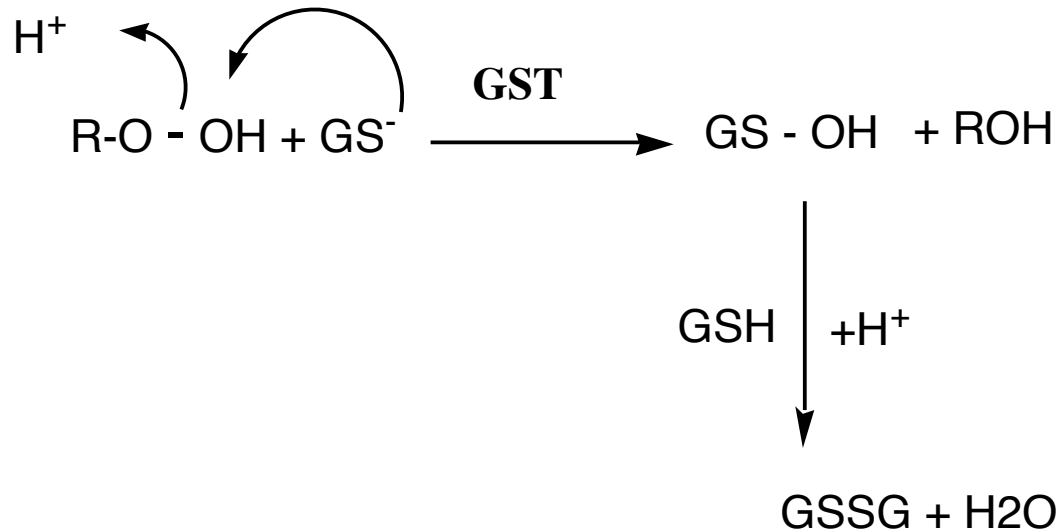
- Found in cruciferous vegetables.
- Isothiocyanates used in industrial processes. Bhopal disaster, 1984 release of methylisocyanate. CCN=C=O
- Induce GSTs.
- **Reversible**: can serve to transport drug/toxin as the GSH conjugate.

## II. Multiple GSH Reactions on a Single Parent Drug

Antipsychotic,  
removed from  
the market due  
to hepatic  
toxicity.

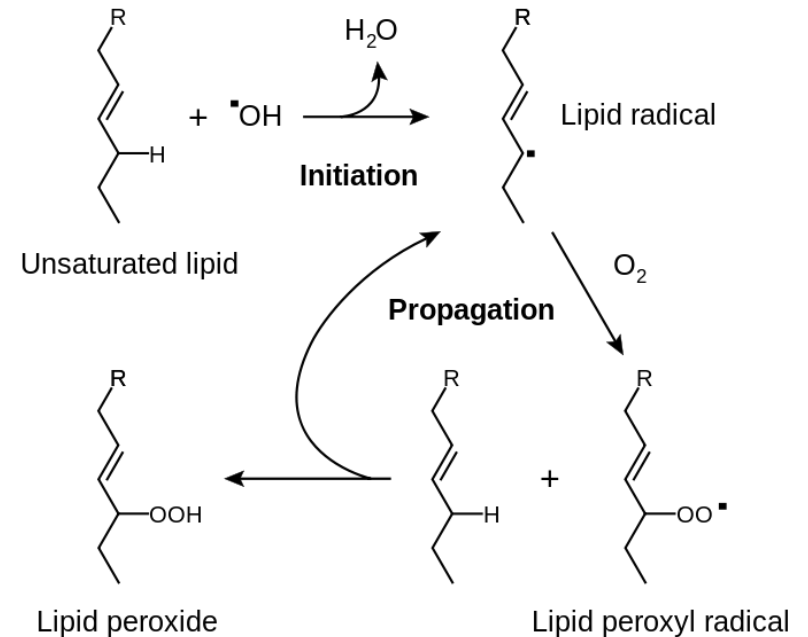


## II. GSH Reactions



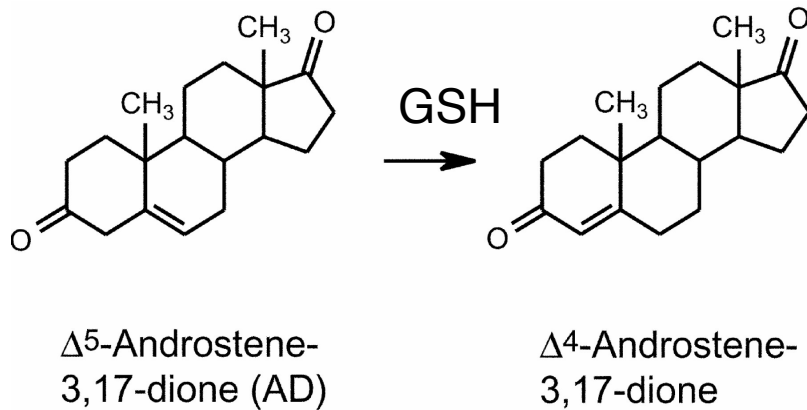
**C. Reduction of peroxides**

e.g. lipid peroxides





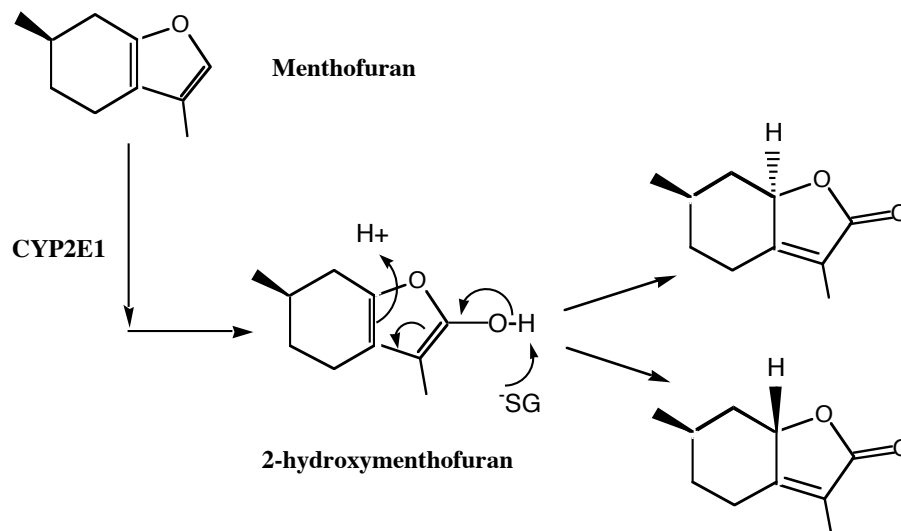
## II. GSH Reactions



### D. Double Bond Isomerization

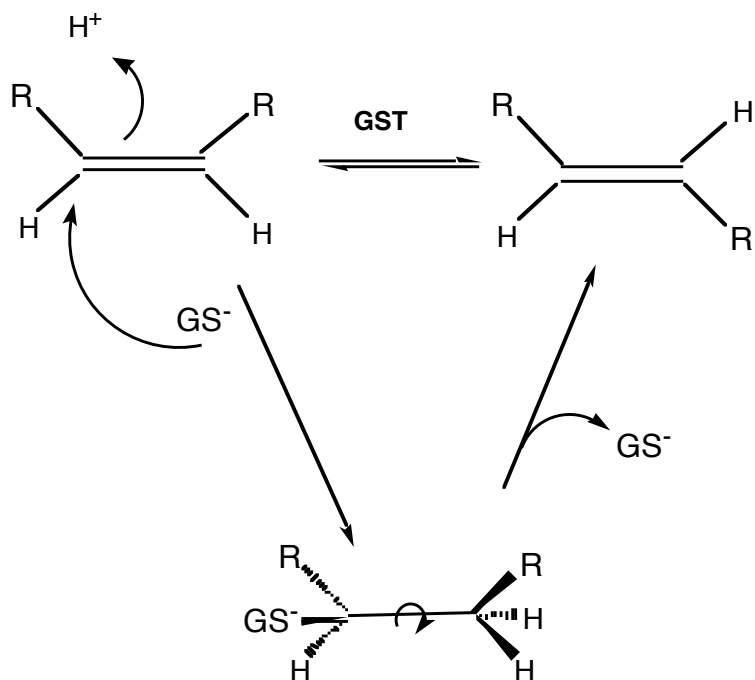
GS<sup>-</sup> is base catalyst (C-4 protons), no net consumption of GSH

e.g. positional isomerization of androstenedione



e.g. hydroxymenthofuran/ a sort of keto-enol isomerization

## II. GSH Reactions



### D. Double Bond Isomerization

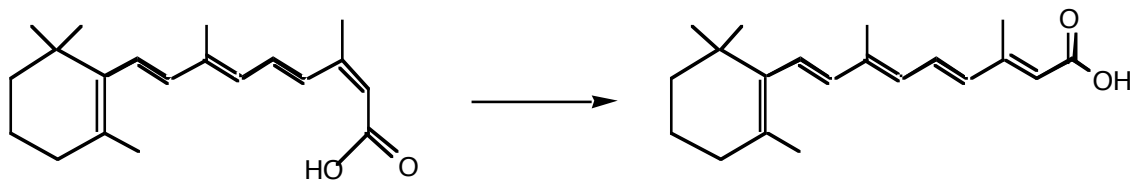
cis-trans isomerization

e.g. maleate  $\rightleftharpoons$  fumarate

prostaglandins

atypical example:

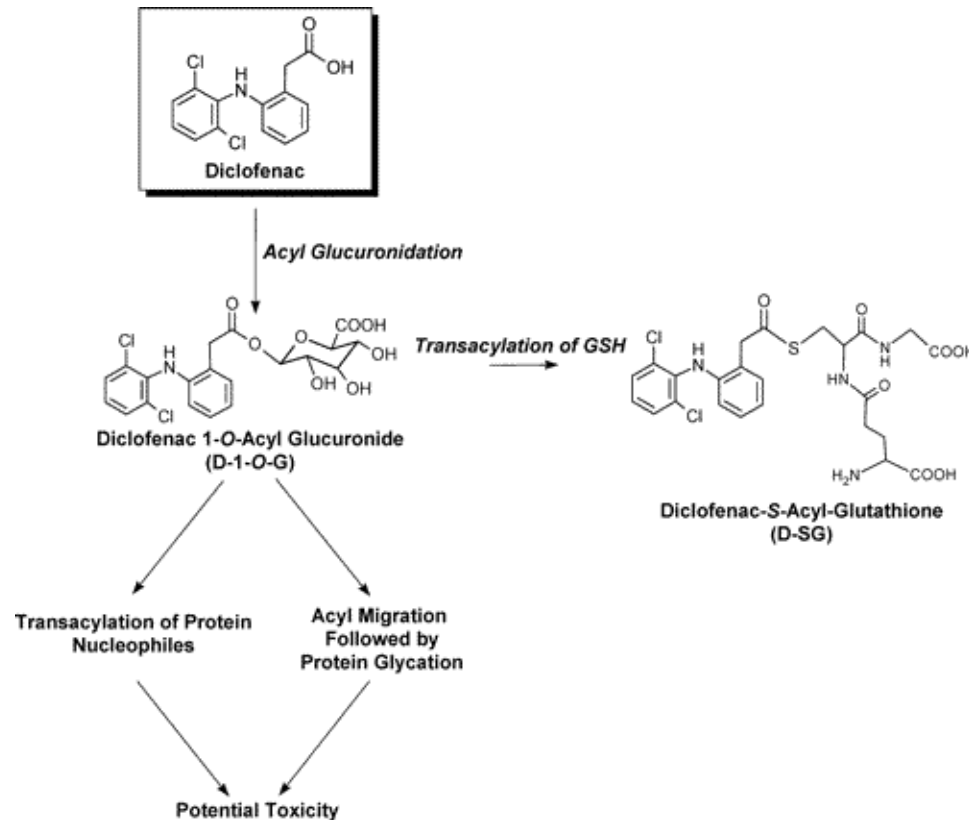
Retinoic acid isomerization  
catalyzed by GSTP1-1 does  
**not** require GSH.



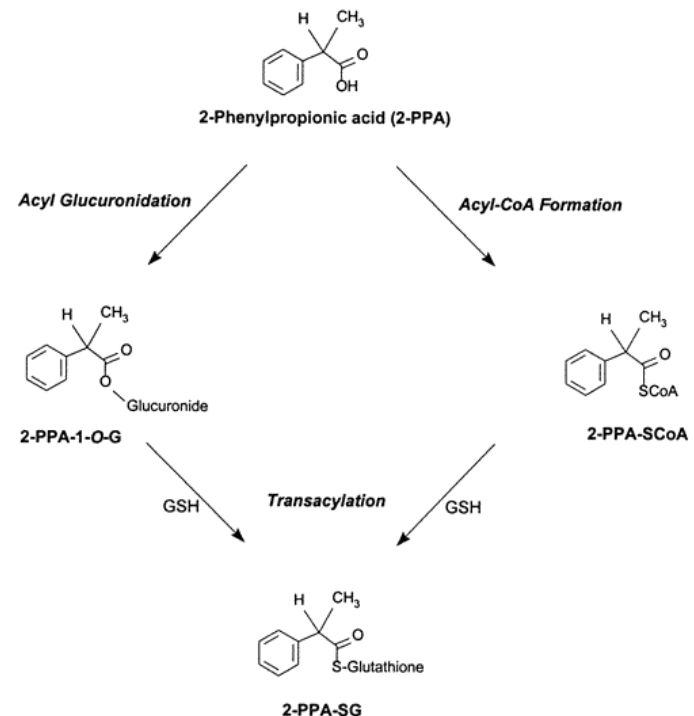
13-cis retinoic acid

all-trans retinoic acid

## II. GSH Reactions



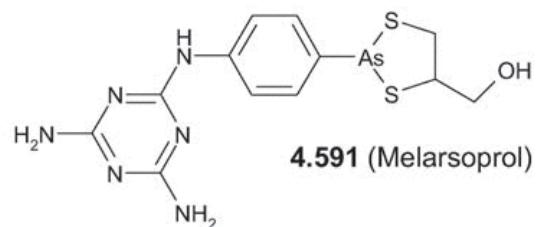
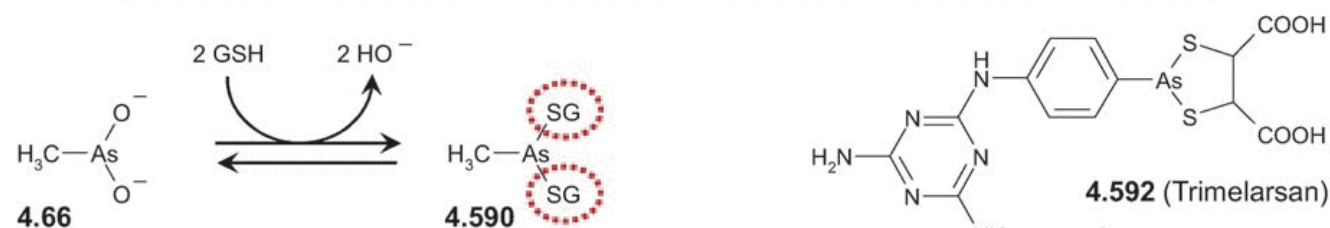
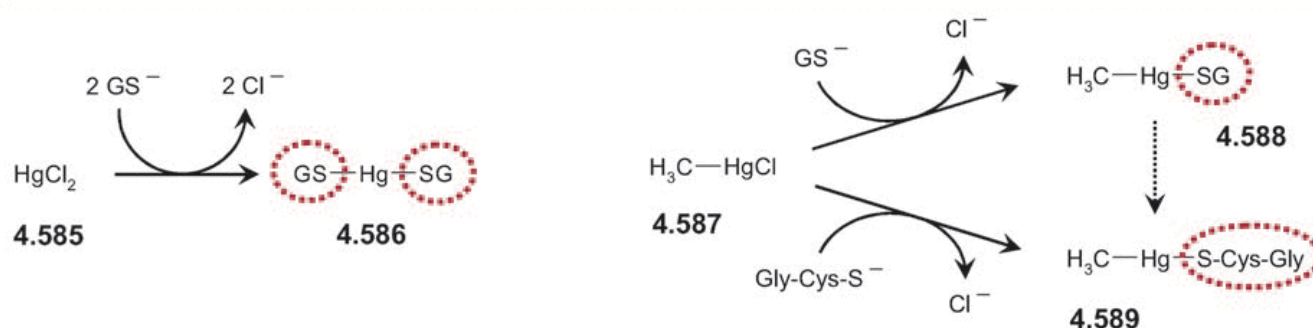
**E. Transacylation.** May contribute to the formation of glutathione conjugates from other primary metabolites. e.g.s Glucuronides, CoA thioesters



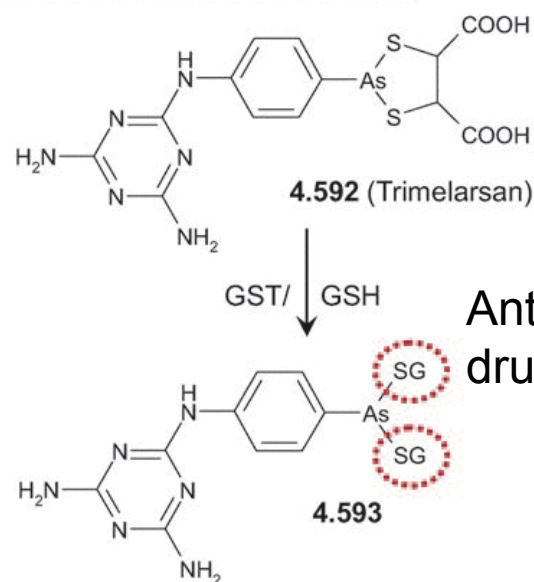
e.g.s zomepirac,  
diclofenac, clofibrate

## II. GSH Reactions: Substitution on Organometallics

### GSH Conjugation of mercury, arsenic, and platinum compounds

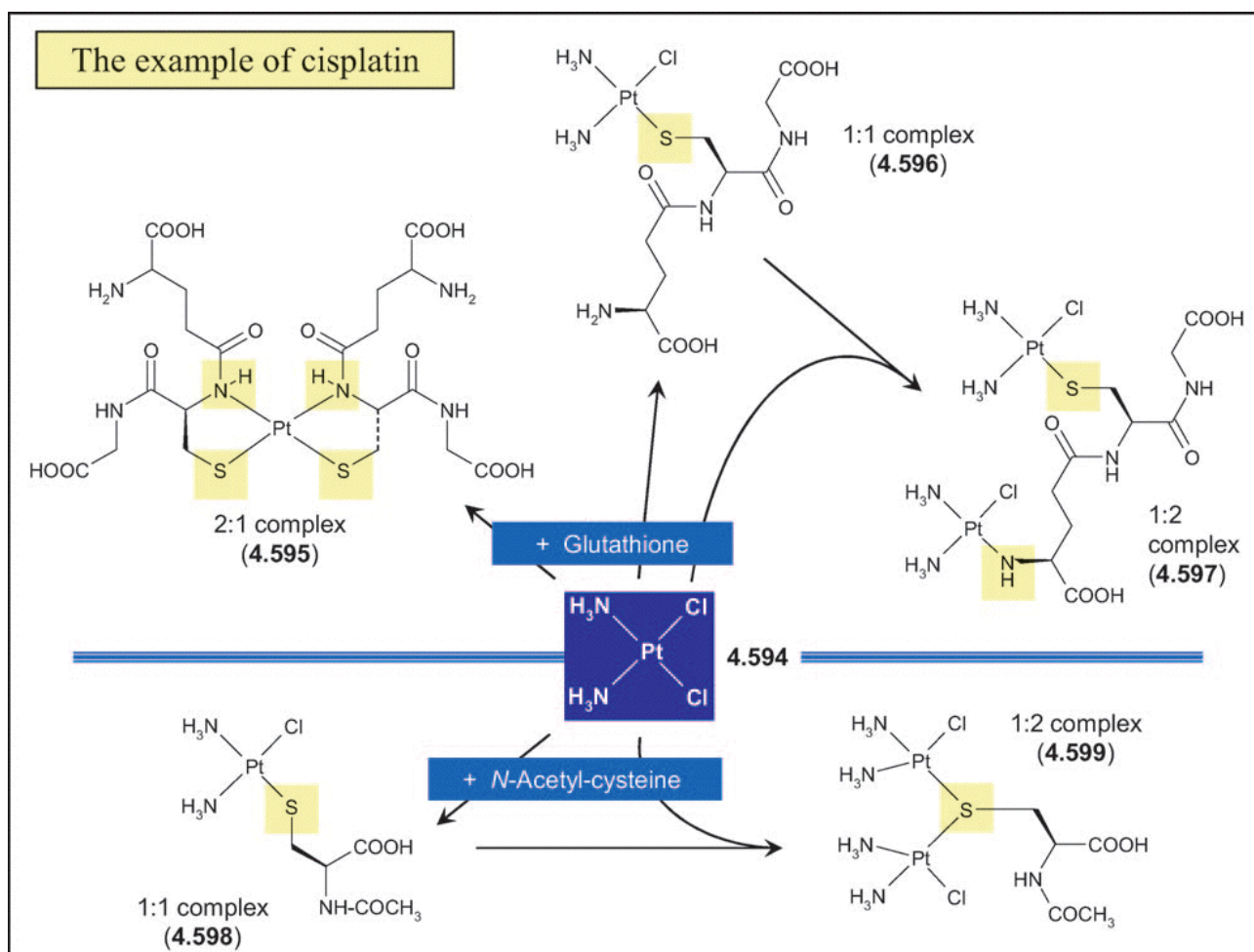


GST/GSH



Anti-trypanosomal drugs

## II. GSH Reactions: Cisplatin



# III. Glutathione S-Transferases

Two structurally unrelated families of GST:

**MAPEGs:** membrane-associated proteins involved in eicosanoid and glutathione metabolism.

- Family of 13 proteins is based on sequence and structural properties and functions. Six human members: 5-lipoxygenase-activating protein (FLAP), leukotriene C4 (LTC4) synthetase, Microsomal MGST1, MGST2, MGST3, and MGST-like 1 (mGSTL1).
- MAPEG GSTs are not closely related to the cytosolic GSTs. Relatively small proteins, ~17 kD.
- MGST1 appears to contribute to drug metabolism. MGST2 and 3 may contribute to drug metabolism, but also likely contribute to lipid peroxide metabolism, and hence protection from oxidative stress.

**Cytosolic GSTs** have multiple functions, including detoxification, drug metabolism, antioxidative stress.

Human GSTs involved in xenobiotic metabolism

The GST gene superfamilies	Human enzymes
<b>Microsomal GST superfamily</b> (homotrimers)	
<i>MGST1, MGST2, MGST3</i>	MGST1, MGST2, MGST3
<b>Cytoplasmic GST superfamily</b> ( $\mu$ : homodimers, and a few heterodimers)	
<i>GSTA1, GSTA2, GSTA3, GSTA4, GSTA5</i>	<b>Alpha class:</b> GST A1-1, A1-2, A2-2, A3-3, A4-4, A5-5
<i>GSTK1</i>	<b>Kappa class:</b> GST K1-1 (mitochondrial and peroxisomal)
<i>GSTM1, GSTM2, GSTM3, GSTM4, GSTM5</i>	<b>Mu class:</b> GST M1a-1a, M1a-1b, M1b-1b, M1b-2, M2-2, M2-3, M3-3, M4-4, M5-5
<i>GSTO1, GSTO2</i>	<b>Omega class:</b> GST O1-1, O2
<i>GSTP1</i>	<b>Pi class:</b> GST P1-1
<i>GSTT1</i>	<b>Sigma class:</b> GST S1
<i>GSTT1</i>	<b>Theta class:</b> GST T1-1, T2
<i>GSTZ1</i>	<b>Zeta class:</b> GST Z1-1

# III. MAPEG Structure and Function

MAPEGs are functional trimers

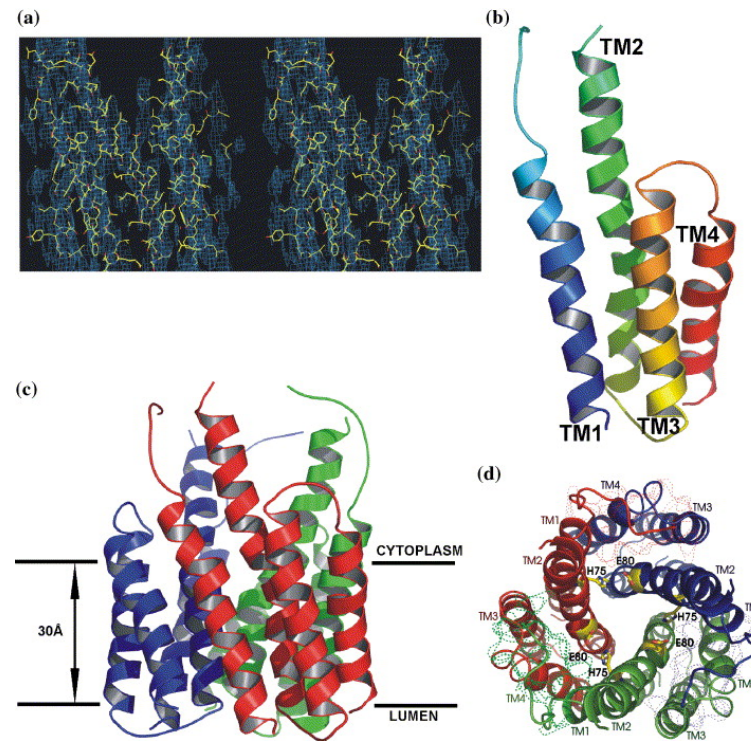
ER membrane proteins

4-transmembrane helices/monomer

# III. MAPEG Structure and Function

Structure available for MGST1 indicates a four helix bundle, with 4 transmembrane helices/subunit.

Trimers form along helical axes.



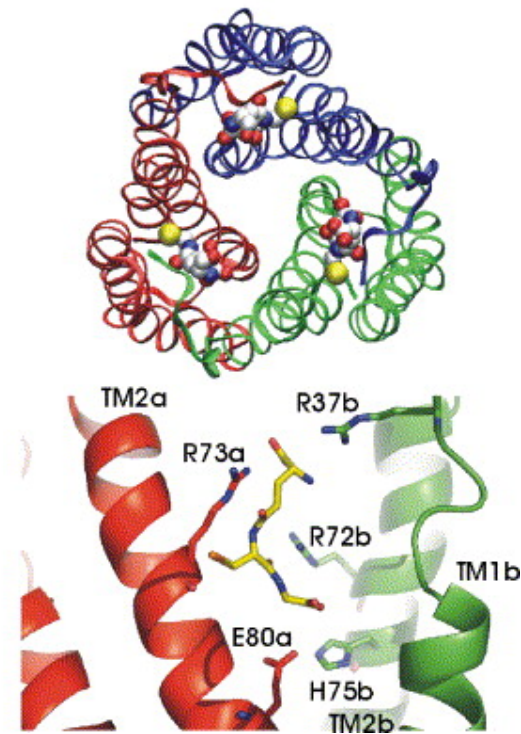


### III. MAPEGs: GSH Binding Site

GSH binding site includes intersubunit interactions.

Contacts to each GSH include two subunits within the trimer.

Binding sites near the top of the trimer at the cytosolic surface - no 'latency' as with UGTs.



# III. Cytosolic Glutathione S-transferases:

## Nomenclature

Isoforms designated on the basis of sequence homology, but unlike CYPs, SULTs, and UGTs the abbreviations include letter (family) and two numbers designating subunit composition.

In humans, A- (alpha-), P- (pi-), M- (mu-), O- (omega) and T- (theta), sigma (S)-classes are cytosolic, xenobiotic metabolizing enzymes.

The **cytosolic GSTs are dimeric enzymes**, usually homodimers, although intra-class heterodimers are documented. No inter-class heterodimers have been reported. The **nomenclature attempts to define gene class and subunit composition**: A1-1, M3-3 etc, where A1-2 represents a heterodimer of alpha-class subunits 1 and 2.

**Formally - not 'S-transferases'**

### III. Glutathione S-transferases: Tissue distribution

#### Human Isoform

#### Tissues

A1-1

Liver > kidney, adrenal > pancreas

A2-2

Liver, pancreas > kidney

A3-3

Ovary, testis, adrenal, placenta

A4-4

low but everywhere

M1-1

Liver > testis > brain

M2-2

Brain > testis > heart

M3-3

Testis > brain

M4-4

M5-5

P1-1

Brain > lung, placenta > kidney, pancreas

T1-1

Kidney, liver > small intestine > brain

T2-2

liver ?

O1-1

liver, heart

O2-2

testis

K1-1

many - mitochondrial

# III. Glutathione S-transferases

## Polymorphisms

GSTM1-1, high frequency of Caucasians are homozygous for a GSTM1-1 deletion. Possible increased risk of bladder and lung cancer.

M3-3, lower frequency in Caucasians may be associated with increased risk of skin cell carcinomas.

A1-1, small percentage (?) have deletion in the promoter region that leads to null phenotype. Possible increased risk of colon cancer.

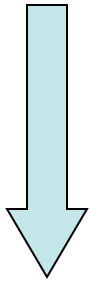
P1-1, several allelic 'SNP' variants produce GSTP's with altered function: Ile104→Val, Ala113→Val, double mutant. Have altered catalytic function/ substrate selectivity.

T1-1, gene deletion in 12-60% depending on ethnic origin. May be associated with increased risk of tumors in several tissues, but this is not well studied.

# III. Glutathione S-transferases

## Functional Overview

Sequestration/  
Passive binding



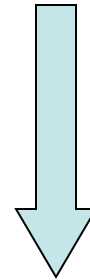
'ligandin' function,  
Cellular uptake/ transport:  
Hemes, steroids

Catalysis



GSH conjugation,  
endogenous/exogenous  
substrates

Signal Transduction



Regulation of  
stress kinases,  
JunK/ASK1

# III. Glutathione S-transferases

## Over-expression of GSTs in Cancer Cells

- A-, P-, M-, and T-class cytosolic enzymes have each been shown to be induced 3 -50-fold in various transformed model cell lines. The increased levels of GSTs may contribute to multidrug resistance of tumor cells, but the causal relationship between GST levels and drug resistance is still debated.
- GSTP1-1 may have a 'unique' function related to cancer cell response – GST inhibits c-Jun kinase. That is, GSTP1-1 also regulates signal transduction pathways involved in apoptotic/proliferative responses.

# III. Glutathione S-transferases

## Substrate Selectivity:

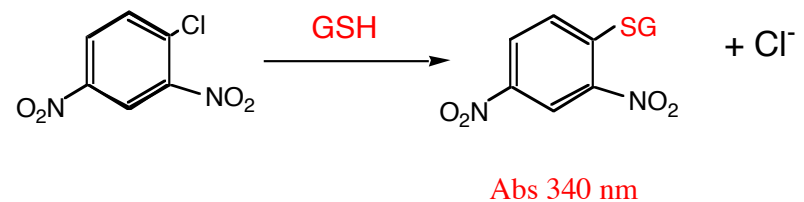
Class and isoform-dependent substrate selectivity is broad and overlapping, as with the other detoxification enzymes.

A few generalizations are:

M-class GSTs have high activity toward planar aromatic hydrocarbon epoxides.

T-class have high affinity for aryl sulfates, catalyze desulfation.

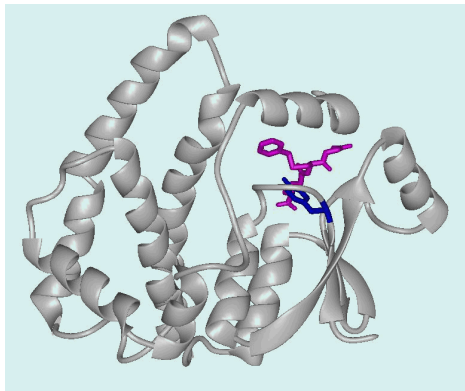
A1-1, A2-2 have relatively high activity toward organic peroxides. GSTA4-4 selective for lipid hydroxy-enals (4-Hydroxy-Nonenal, HNE). A3-3 great with androstenedione.



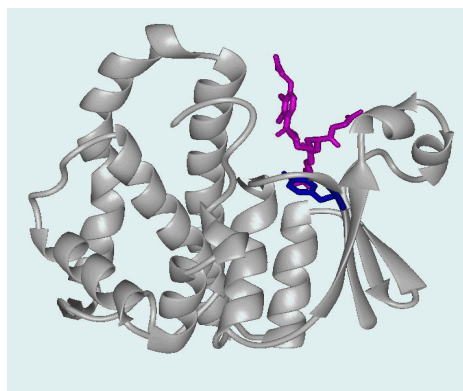
Universal substrate:  
CDNB

# III. Glutathione S-transferases

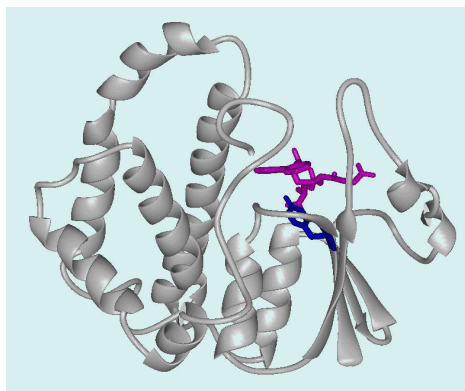
## Structures: Overall fold



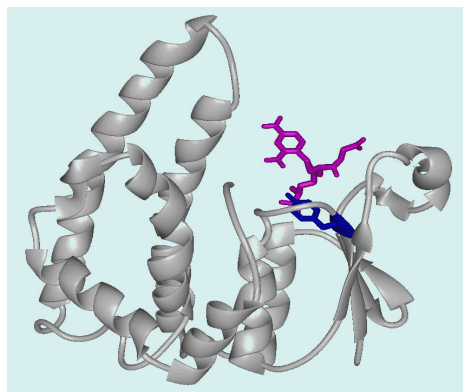
GST A1-1



GST P1-1



GST M3-3



Sigma GST

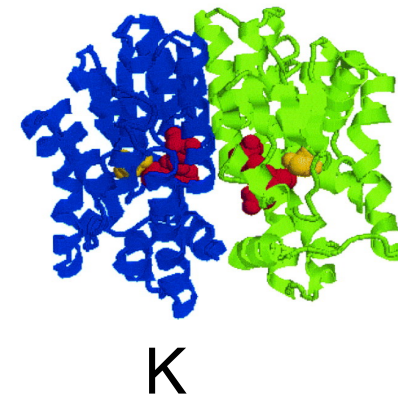
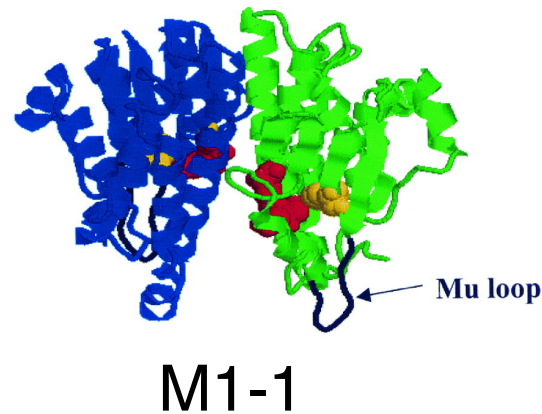
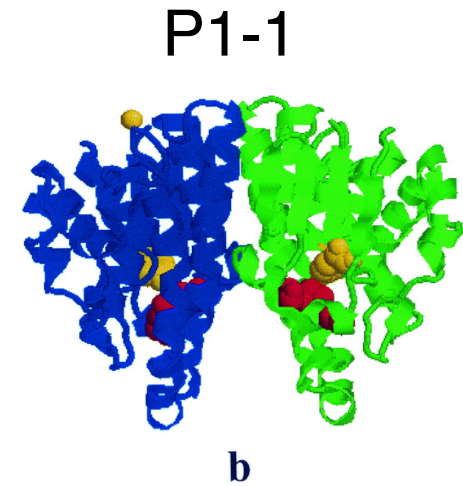
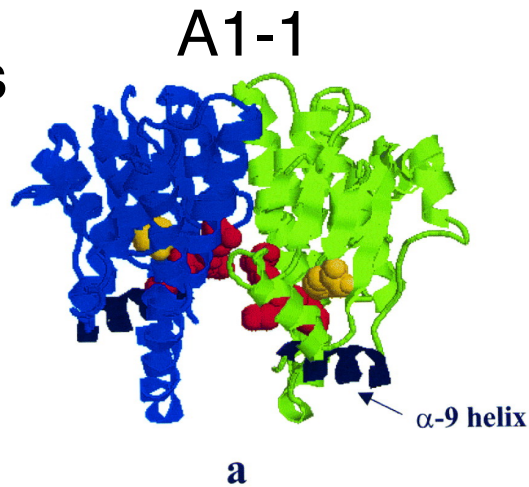
Cytosolic GSTs are dimers. Overall subunit structure is very similar, with two distinct domains. The N-terminal Domain binds glutathione (G-site) and consists of 3 or 4 helices and a 4-stranded  $\beta$ -sheet. The much larger C-terminal domain is loops and helices, and contributes to the xenobiotic binding site (H-site) which lies between the domains.



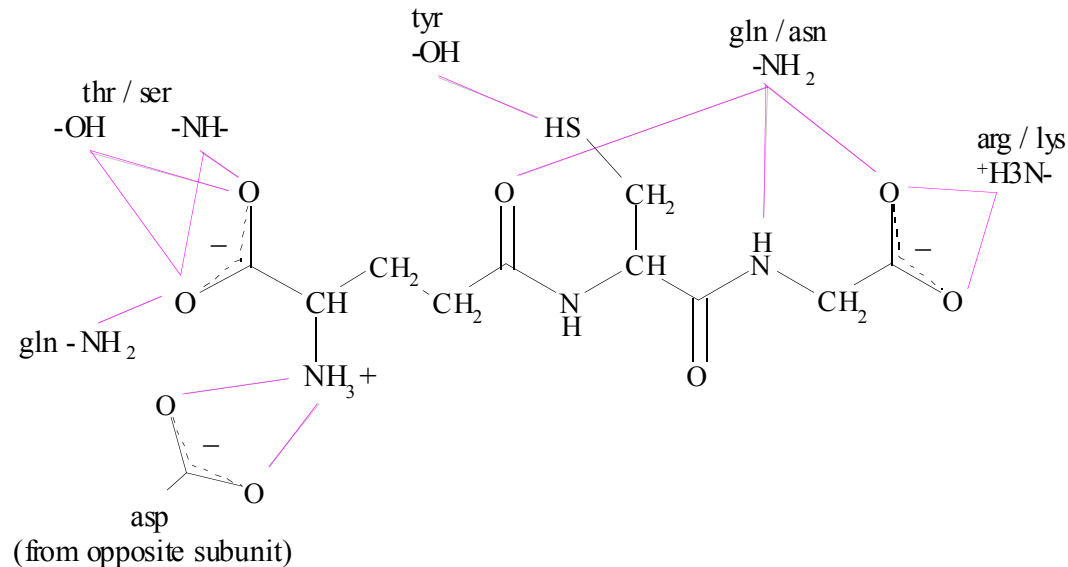
### III. GSTs

GSTs are **dimers**,  
interclass subunit-  
subunit interactions  
are incompatible.

Large intersubunit  
cleft of varying size  
and character.



# III. GSTs: G-site



Spectroscopic experiments indicate that GSH bound at the active site of GST has a pK<sub>a</sub> of ~ 6.5 (M)- (P) -7.4 (A). Thus, the **nucleophilic GS<sup>-</sup> is bound**, rather than GSH. This is due to a hydrogen bond to a conserved tyrosine (M, P, A), Cys (O) or ser (T). For some A class isoforms, the tyrosine has 'unusual' properties as well.

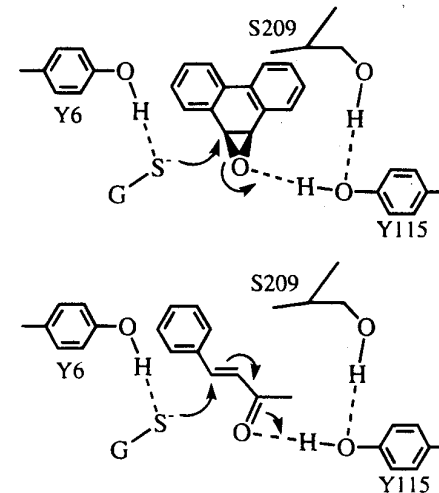
structures from each class indicate that salt bridges and hydrogen bonds between active site and GSH peptide are functionally conserved, but structurally distinct.

# III. GSTs: H-site, M-class

For each of the GST isoforms, the H-site is lined with hydrophobic residues. In some cases the H-site includes an appropriately placed residue that aids in catalysis of specific substrates i.e. substrate specificity.

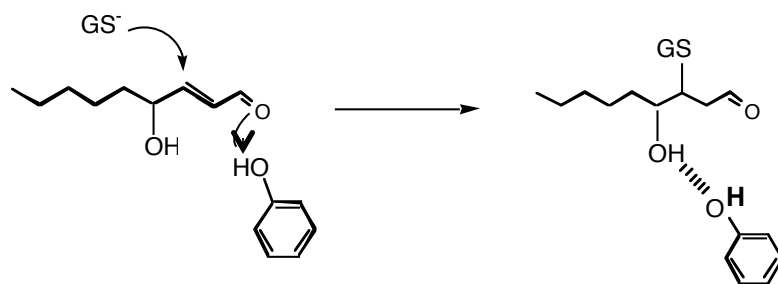
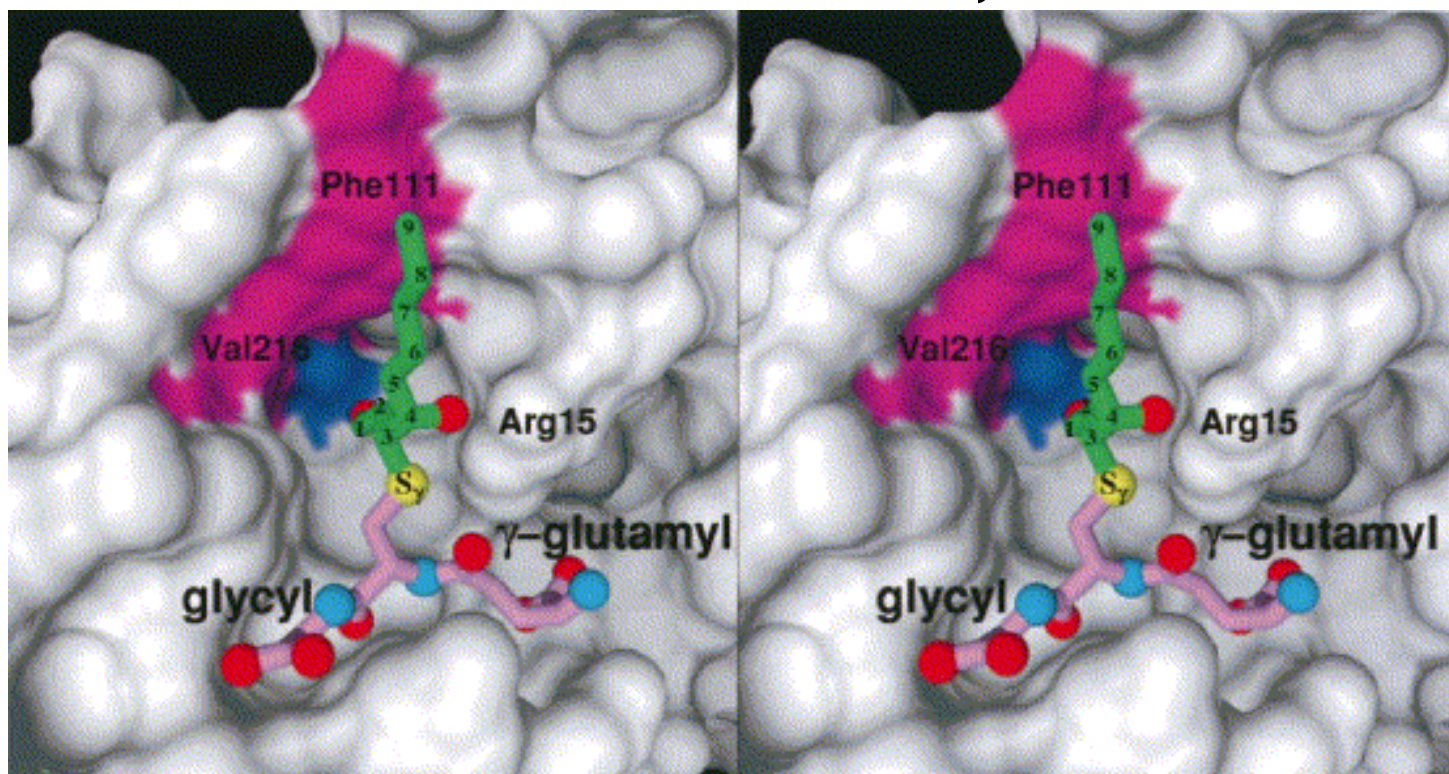
e.g. In GSTM an H-site Tyr provides a general base to the 'leaving' oxygen of epoxide substrates.

## M-class



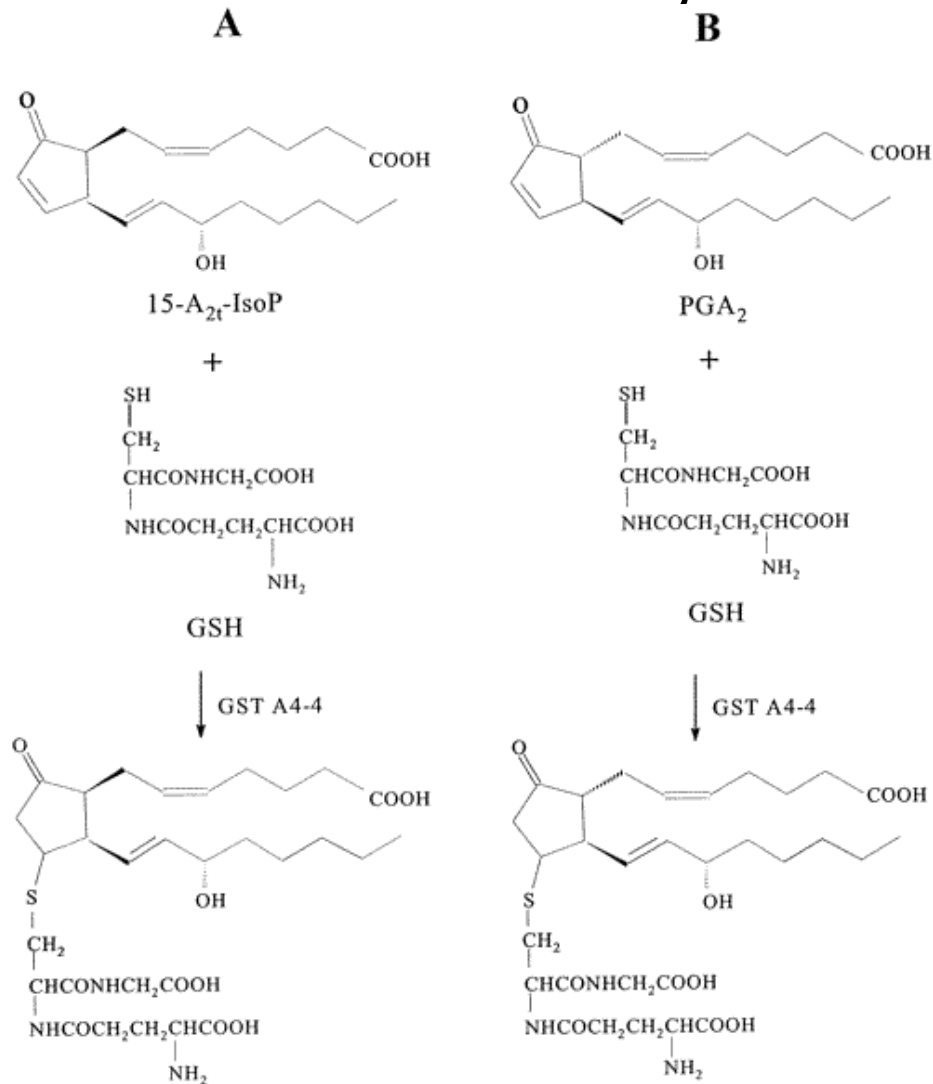
**Figure 9.** Proposed role of the hydroxyl group of Y115 as an electrophilic participant in catalysis of Michael additions and oxirane ring openings. Taken from ref 26.

### III. GSTs: H-sites, A4-4



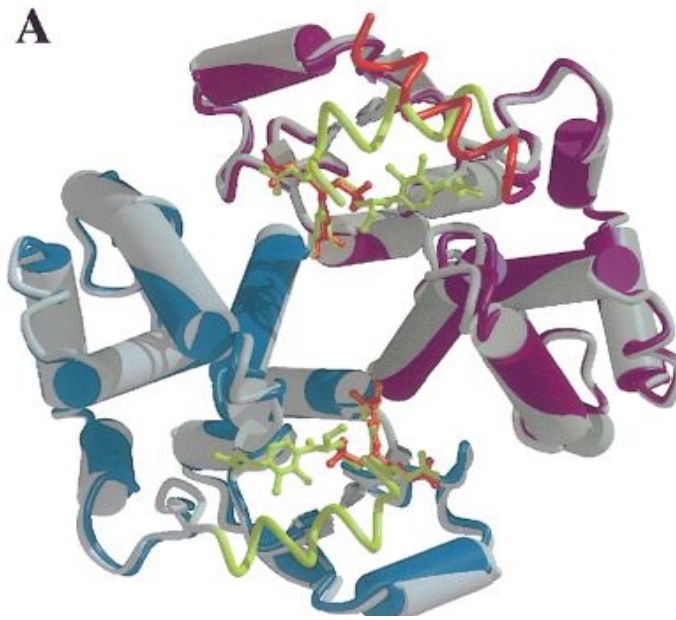
A4-4 has Tyr-212 which acts as general acid for 4-HNE conjugation

# III. GSTs, H-sites A4-4



A4-4 also likes isoprostane oxidation products of prostaglandins A(2) and J(2), possible toxic metabolites of oxidative stress.

### III. GSTs: Dynamics in GSTA1-1

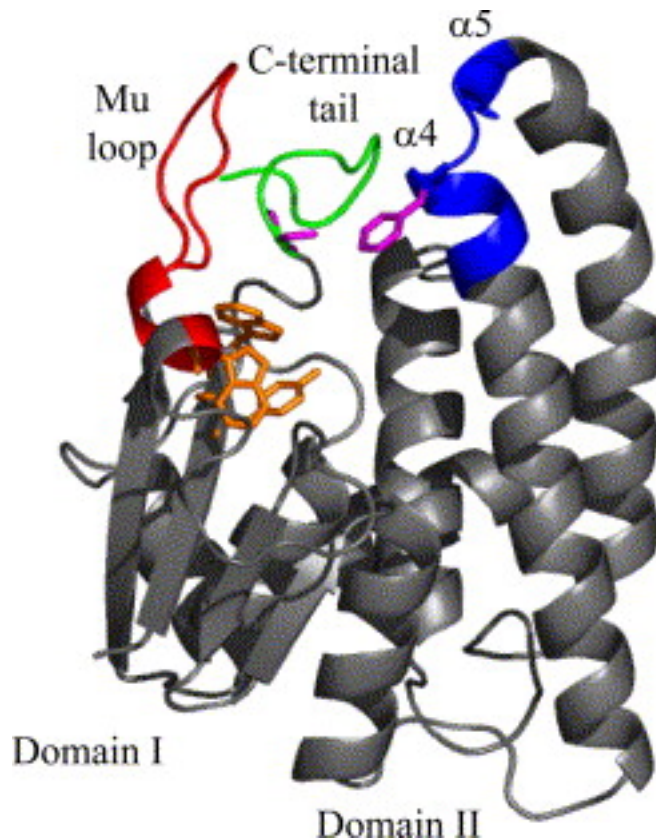


C-terminal helix  
of A1-1  
(208-222) is  
dynamic.

Location  
depends on  
which ligands  
are bound: helix  
closes over the  
active site with  
GS-conjugates



### III. Dynamics in GSTM3-3



'Mu-loop' (red) and C-terminus (green) are mobile and restrict egress of products from the active site.

Blue segment is also dynamic and includes the catalytic Tyr-115

# IV. Reactions of Glutathione Conjugates

Degradation by peptidases to Cysteine Conjugates occurs in the kidneys. Normally, the cysteine conjugate is eliminated in urine. Conversion of GSH conjugate to Cys conjugate is the 'Mercapturic Acid Pathway'.

## Mercapturic acid pathway

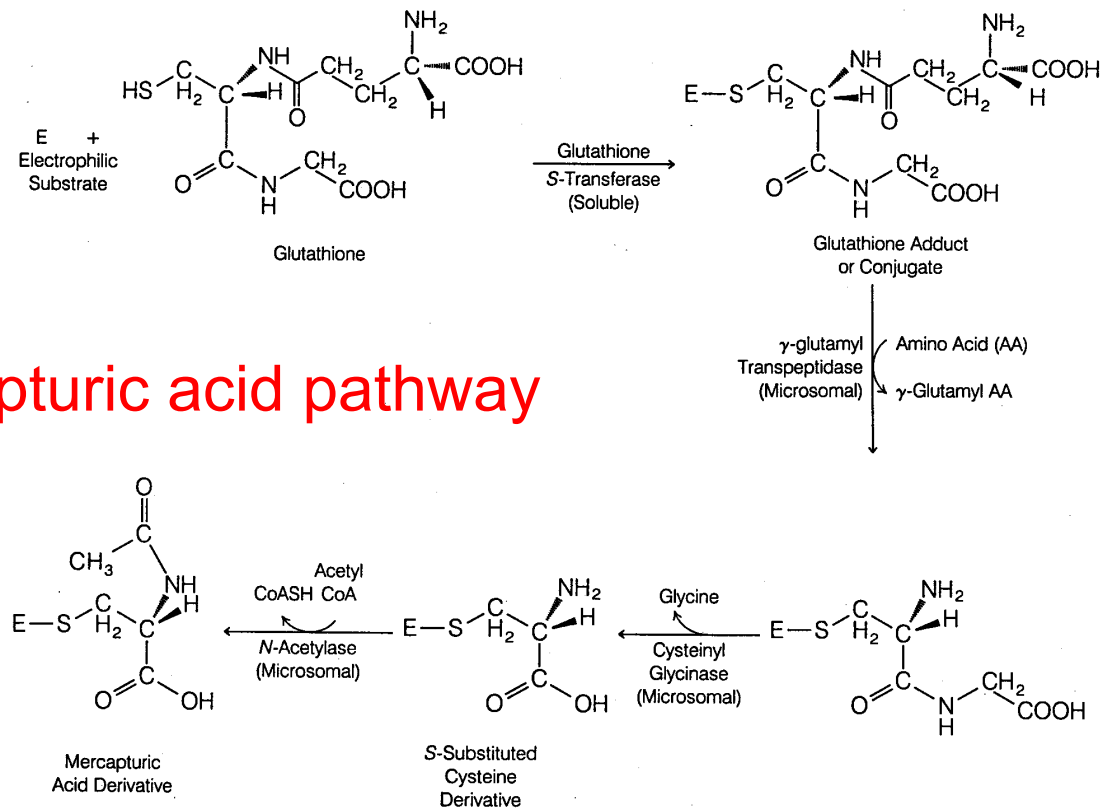


FIG. 3-15. Formation of glutathione conjugates of electrophilic xenobiotics or metabolites (E) and their conversion to mercapturic acids.



# IV. Reactions of Glutathione Conjugates: Cysteine Conjugate Metabolism

The enzyme Cysteine S-Conjugate  $\beta$ -Lyase further degrades Cys conjugates. The product may be toxic.

e.g.'s

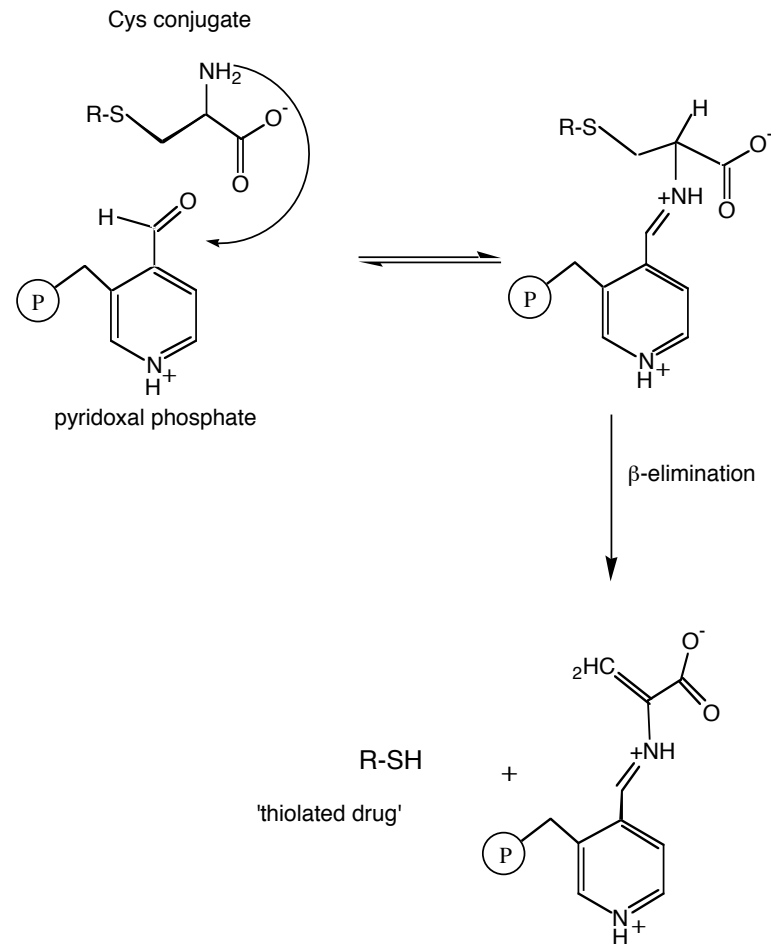
Cisplatin

Halogenated anesthetics

TCE and halogenated toxins

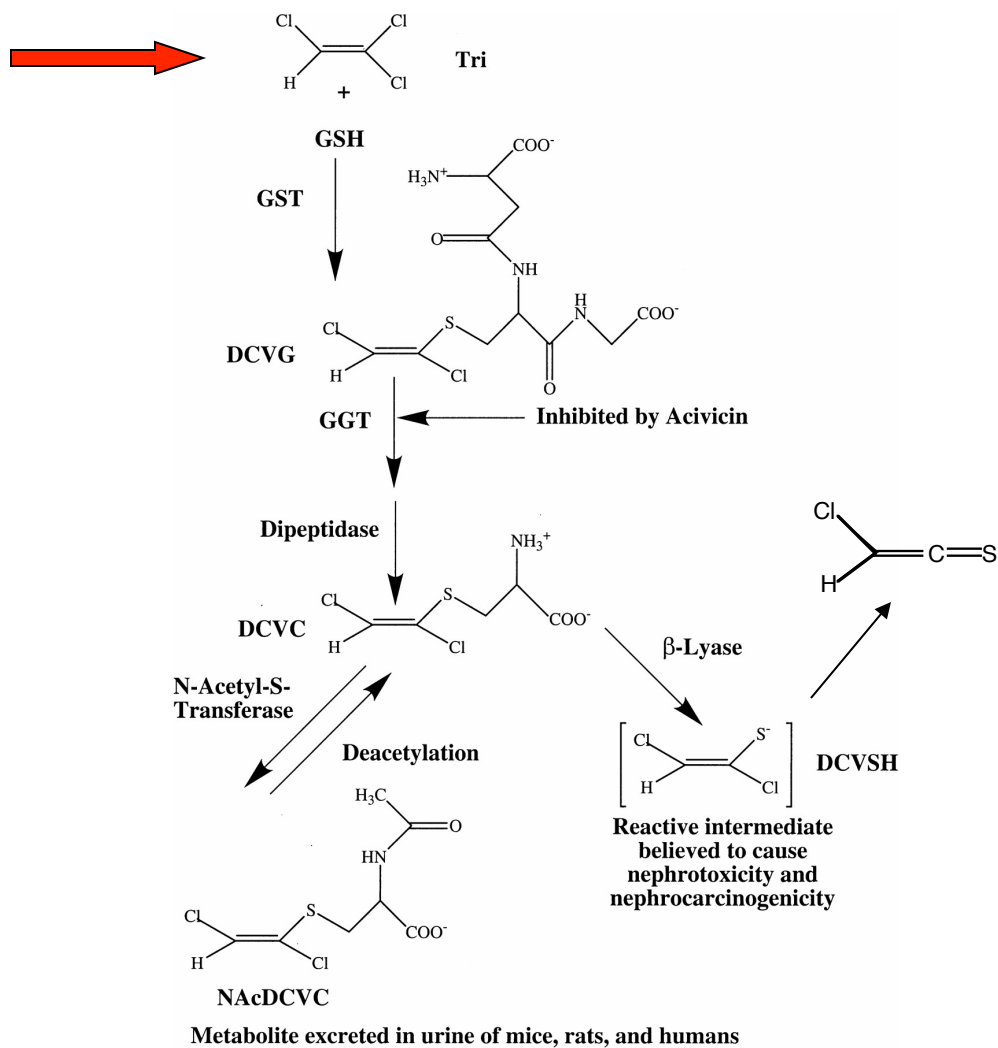
Catechol neurotransmitters

**Cysteine  $\beta$ -lyase pathway**

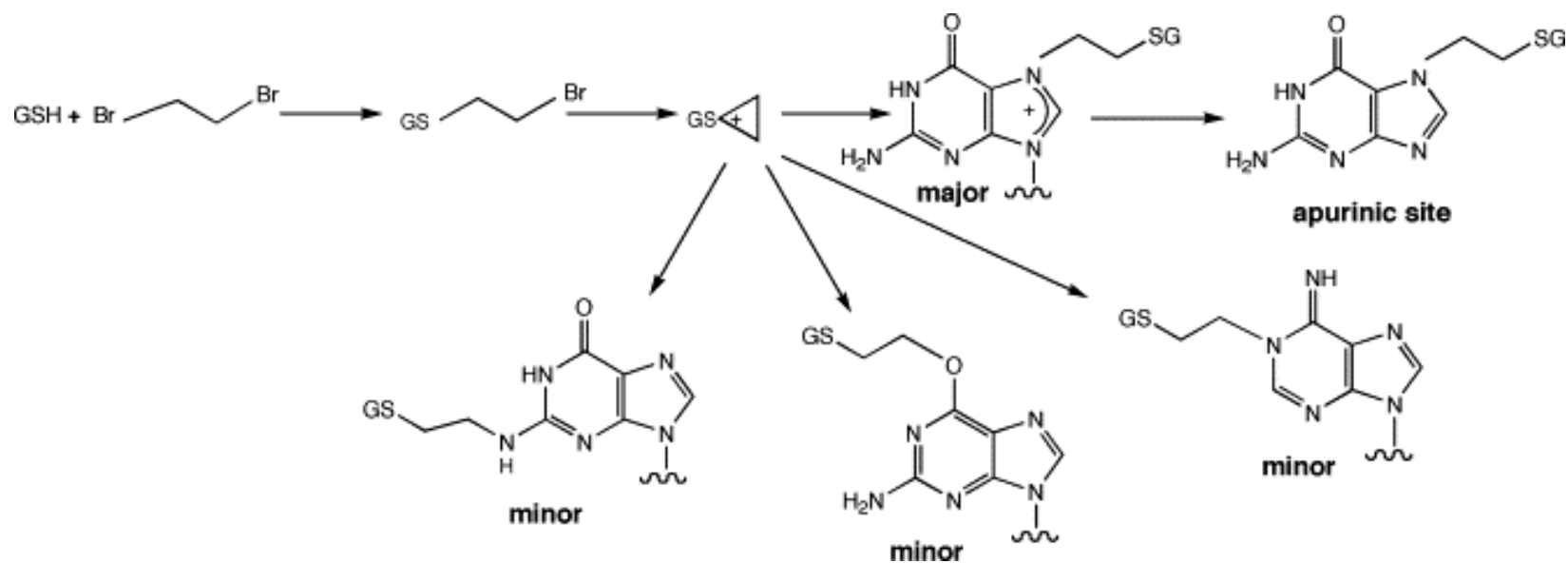


# IV. Reactions of Glutathione Conjugates

TCE, industrial solvent

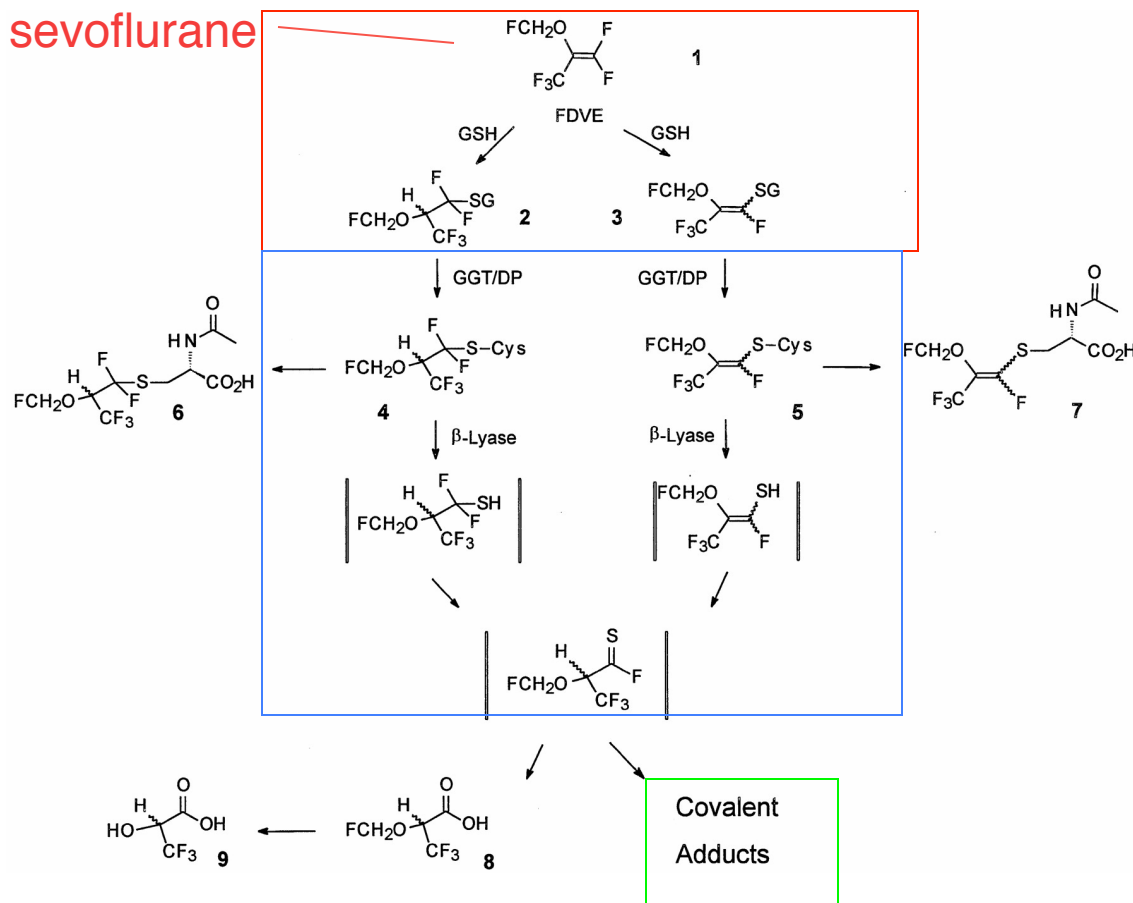


# IV. Reactions of Glutathione Conjugates: Ethylene Dibromide

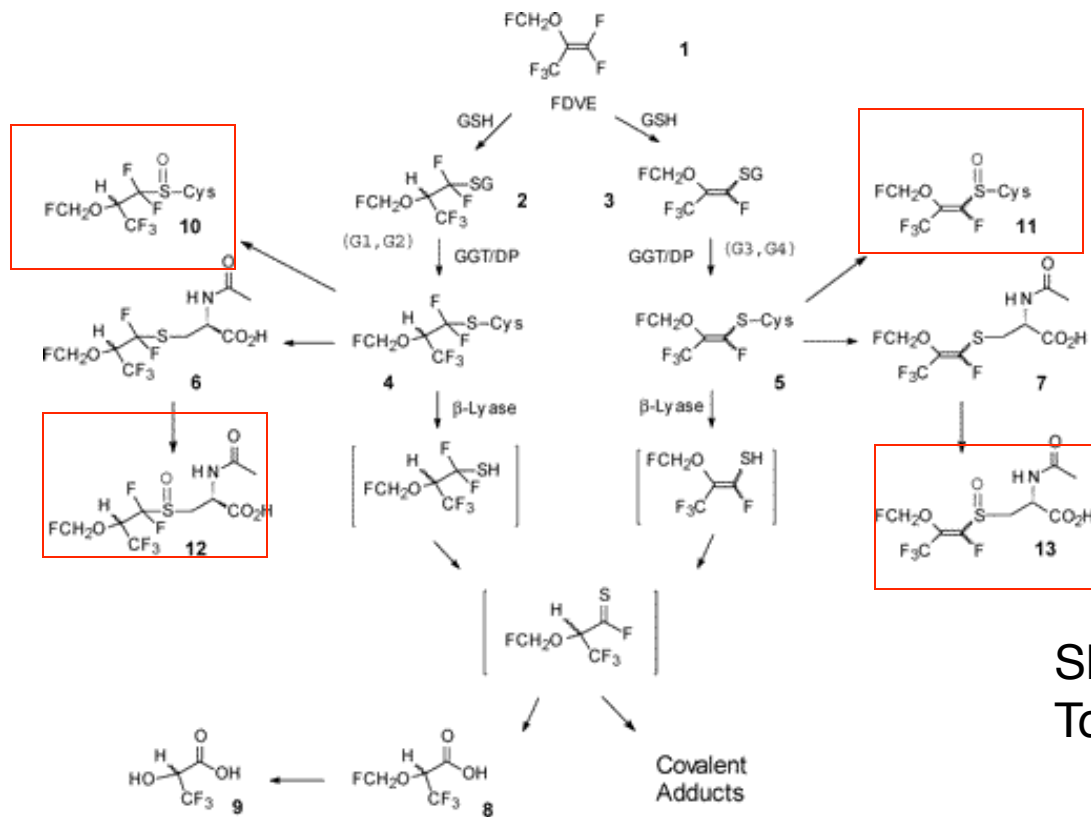


# IV. Reactions of GSH/CYS Conjugates- Sevoflurane

Contaminant in sevoflurane



## IV. Reactions of GSH/CYS Conjugates- Sevoflurane



More recently,  
CYP-dependent  
sulfoxidation of the  
Cys-conjugates  
has been  
demonstrated.

Sheffels et al Chem Res  
Toxicol (2004) 17:1177.

Other eg.'s: Cys conjugate of hexachlorobutadiene- Werner et al. Chem Res Toxicol (1995) 8: 917.

## IV. Reactions of GSH Conjugates: Transport

GSH conjugates are effluxed by several ATP-dependent transporters, MRP1, MRP2, RLIP.

In addition, GSH and GSH conjugates stimulate the transport by MRP's of other drugs and conjugates including glucuronides.

Mechanism? - free thiol not required, allosteric site?

# IV. Reduction of GS-Conjugates

## 4.7.4. Reductions Following Glutathione Conjugation

