

# **UNIVERSITI PUTRA MALAYSIA**

# PARTIAL PURIFICATION AND CHARACTERIZATION OF GLUTATHIONE S-TRANSFERASES FROM KEDAH-KELANTAN CATTLE (BOS INDICUS) AND WATER BUFFALO (BUBALUS BUBALIS) LIVER

LAILATUL JUMAIYAH BINTI SALEH HUDDIN.

FBSB 2006 5



## PARTIAL PURIFICATION AND CHARACTERIZATION OF GLUTATHIONE S-TRANSFERASES FROM KEDAH-KELANTAN CATTLE (*BOS INDICUS*) AND WATER BUFFALO (*BUBALUS BUBALIS*) LIVER

By

## LAILATUL JUMAIYAH BINTI SALEH HUDDIN

Thesis submitted to the School of Graduate Sudies, Universiti Putra Malaysia, in Fulfilment of the Requirement for the Master of Science

2006



For dearest family and friends

"...man will occasionally stumble over the truth, but usually manages to pick himself up, walk over or around it, and carry on."

Churchill, Winston S.





Abstract of thesis presented to the Senate of Universiti Putra Malaysia in fulfillment of the requirement for the degree of Master of Science

### PURIFICATION AND CHARACTERIZATION OF GLUTATHIONE S-TRANSFERASES FROM KEDAH-KELANTAN CATTLE (Bos indicus) AND WATER BUFFALO (Bubalus bubalis) LIVERS

By

### LAILATUL JUMAIYAH BINTI SALEH HUDDIN

### April 2006

#### Chairman: Professor Nor Aripin Shamaan, PhD

### Faculty: Biotechnology and Biomolecular Sciences

Biotransformation and detoxification process in living organisms consists of two phases, phase I and phase II. Phase I involves in the introduction of functional group into molecule while the phase II involves the conjugation of phase I metabolites. In phase II, glutathione S-transferases (GSTs; EC 2.5.1.18) has aroused much interest because of its involvement in the biotransformation and detoxification of wide spectrum of xenobiotics which can be from pesticides, herbicides and insecticides. The present study was undertaken to purify and characterized cytosolic GSTs from livers of Kedah-Kelantan cattle (*Bos indicus*) and Malaysia water buffalo (*Bubalus bubalis*). The glutathione S-transferases were isolated from two important livestock livers, Kedah-Kelantan cattle (*Bos indicus*) and Malaysian water buffalo (*Bubalus bubalis*) by glutathione affinity chromatography. The affinity-glutathione chromatography successfully purifies the GSTs isoenzymes with 14.73% yield (62.77 purification fold) and 19.71% yield (20.44 purification fold) for KK cattle and water buffalo livers respectively. Initial methods of purification included centrifugation and ultracentrifugation. The affinity elution with



highest activity towards CDNB was estimated for the pI values using isoelectric focusing method via LKB-8100 ampholyte type (LKB Bromna) apparatus. pI values for affinity purified KK cattle liver are 5.7 (C-34), 6.9 (C-38) and 8.8 (C-42). While for the water buffalo liver, the pI values for glutathione affinity purified isoenzymes are 6.85 (B-23) and 7.2 (B-24). The isoenzymes were then tested using SDS-PAGE method for purity and also to estimate the molecular weight estimation. It has been estimated that molecular weight for water buffalo isoenzymes of B-23 was 29.3  $\pm$  0.05 kDa and B-24 was 30.74  $\pm$  0.16 kDa. The KK cattle liver isoenzymes molecular weight was estimated with C-34 was 29.9  $\pm$  0.14; C-38 was 28.3  $\pm$  0.09 and 27.7  $\pm$ 0.03 for C-42. The study showed that KK cattle liver GSTs exist as isoenzymes (pI 8.8, 6.9 and 5.7), and have high activity towards CDNB, low towards DCNB and no activity towards the ethacrynic acid for the substrate specificities. On the other hand, the water buffalo liver GSTs exist as isoenzymes are specificities, the isoenzymes are isoenzymes are isoenzymes are isoenzymes are isoenzymes are isoenzymes are other hand, the water buffalo liver GSTs exist as isoenzymes are buffalo liver GSTs exist as isoenzymes with pI 6.85 and 7.2. For the substrate specificities, the isoenzymes also have high activity for CDNB, but low for DCNB and could not be detected for the ethacrynic acid.



Abstrak tesis yang dikemukakan kepada Senat Universiti Putra Malaysia sebagai memenuhi keperluan untuk ijazah Master Sains

### PENULENAN DAN PENCIRIAN GLUTATHIONE S-TRANSFERASES SEPARA DARI HATI LEMBU KEDAH-KELANTAN (Bos indicus) DAN KERBAU (Bubalus bubalis)

Oleh

### LAILATUL JUMAIYAH BINTI SALEH HUDDIN

April 2006

#### Pengerusi: Professor Nor Aripin Shamaan, PhD

### Fakulti: Bioteknologi dan Sains Biomolekul

Proses biotransformasi dan detoksifikasi di dalam organisma hidup merangkumi dua fasa; fasa I dan fasa II. Fasa I melibatkan penambahan kumpulan berfungsi kepada molekul asing manakala fasa II melibatkan konjugasi metabolit fasa I. Dalam fasa II, glutathione S-transferases telah (GSTs;EC 2.5.1.18) telah menarik minat saintis dengan kaitannya dalam biotransformasi dan detoksifikasi bagi sebahagian besar xenobiotik yang biasanya boleh didapati dari racun perosak. Kajian ini dijalankan untuk menulen dan mencirikan GST sitosolik dari hati lembu lembu Kedah-Kelantan (KK) (*Bos indicus*) dan kerbau Malaysia (*Bubalus bubalis*). Glutathion S-transferase telah ditulenkan dari hati dua ternakan penting, lembu Kedah-Kelantan (KK) (*Bos indicus*) dan kerbau Malaysia (*Bubalus bubalis*), dengan menggunakan teknik kromatografi affiniti-glutathione. Kromatografi affiniti-glutathion ini telah berjaya menulenkan isoenzim glutathion S-transferase dengan hasil penulenan sebanyak 14.73% dan 62.7 kali tahap penulenan telah berjaya didapati daripada hati kerbau. Langkah



awal penulenan adalah termasuk teknik pengemparan dan ultrapengemparan. Elusi affiniti yang mempunyai aktiviti enzim yang tertinggi terhadap substrat CDNB telah dianggar bagi nilai pI dengan menggunakan kaedah 'isoelectric focusing' dengan menggunakan alat LKB-8100 jenis 'ampholyte' (LKB Bromna). Nilai pI bagi lembu KK vang ditulenkan adalah 5.7 (C-34), 6.9 (C-38) dan 8.8 (C-42). Manakala bagi hati kerbau, nilai pI bagi isoenzim yang ditulenkan adalah 6.85 (B-23) dan 7.2 (B-24). Isoenzim yang didapati telah diuji dengan menggunakan kaedah SDS-PAGE bagi menganggarkan ketulenan dan berat molekul. Telah dianggarkan bahawa berat molekul bagi isoenzim dari hati kerbau adalah B-23 adalah 29.3 ± 0.05 kDa dan B-24 adalah 30.74 ± 0.16 kDa. Bagi isoenzim hati lembu KK dianggarkan berat molekul; C-34 (29.9  $\pm$  0.14 kDa), C-38 (28.3  $\pm$  0.09 kDa) dan 27.7  $\pm$  0.03 kDa bagi isoezim C-42). Mengikut pemerhatian yang dilakukan GST dari hati lembu KK wujud dalam bentuk isoenzim dengan nilai pI 8.8, 6.9 dan 5.17 dan mempunyai aktiviti enzim yang tinggi terhadap substrat CDNB dan rendah terhadap DCNB dan tiada aktiviti terhadap substrat asid 'ethacrynic' bagi ujian substrat spesifik. Manakala bagi GST yang ditulenkan daripada hati kerbau wujud dalam bentuk isoenzim dengan pI 6.85 (B-23) dan 7.2 (B-24). Bagi penentuan kadar substrat spesifik, isoenzim B-23 dan B-24 tidak menunjukkan sebarang aktiviti terhadap asid 'ethacrynic', rendah terhadap DCNB dan mempunyai aktiviti yang tinggi terhadap substrat CDNB.



### ACKNOWLEDGEMENTS

First and foremost, I would like to express my sincere appreciation to my supervisor Professor Nor Aripin Shamaan, for his invaluable advices and wide ranging discussions on the project and things in general. My sincere gratitude is extended to my cosupervisors, Dr. Mohd. Yunus Abd. Shukor and En. Ismail Omar, for the advices, guidance and encouragement given to me in this project.

I would also like to thank all the supporting staff in the Department of Biotechnology and Molecular Sciences for the help given in acquiring chemicals, reagents and equipments.

To my family, thank you so much for the moral supports, patience and also money support (especially for the registrations fees!! Thanks MAK). Thank you so much for being there for me during my worst days. I owe all of you more than I can ever say...

To my labmates and friends, thanks for all the ideas, advices, and cheers and tears that we share along the way. This appreciation especially goes to Tony, Farah, Nina, Che' Wan, Jewe, Fara, Eddie and Vani. My appreciation do also goes to Dr. Noor Rain Abdullah and staffs in IMR for the morals supports and time spare.

You guys are always in my heart.

Lailatul Jumaiyah Saleh Huddin Bandar Baru Bangi, September 2006

vii



I certify that an Examination Committee has met on 7 April 2006 to conduct the final examination of Lailatul Jumaiyah Binti Saleh Huddin on her Master of Science thesis entitled "Partial Purification and Characterization of Glutathione S-Transferases from Kedah-Kelantan Cattle (*Bos indicus*) and Water Buffalo (*Bubalus bubalis*) Livers" in accordance with Universiti Pertanian Malaysia (Higher Degree) Act 1980 and Universiti Pertanian Malaysia (Higher Degree) Regulations 1981. The Committee recommends that the candidate be awarded the relevant degree. Members of the Examination Committee are as follows:

### Johari Ramli, PhD

Associate Professor Faculty of Biotechnology and Biomolecular Sciences Universiti Putra Malaysia (Chairman)

### Mohd. Arif Syed, PhD

Professor Faculty of Biotechnology and Biomolecular Sciences Universiti Putra Malaysia (Internal Examiner)

### Juzu Hayati Arshad, PhD

Associate Professor Faculty of Biotechnology and Biomolecular Sciences Universiti Putra Malaysia (Internal Examiner)

Musalmah Mazlan, PhD Professor Faculty of Medicine Universiti Kebangsaan Malaysia (External Examiner)

HASANAH MATHD. GHAZALI, PhD Professor/Deputy Dean

School of Graduate Studies Universiti Putra Malaysia

Date: 11 JUL 2006



viii

This thesis submitted to the Senate of Universiti Putra Malaysia and has been accepted as fulfillment nof the requirement for the degree of Master of Science. The members of the Supervisory Committee are as follows:

# Nor Aripin Shamaan, PhD

Professor Faculty of Biotechnology and Molecular Sciences Universiti Putra Malaysia (Chairman)

## Mohd. Yunus Abdul Shukor, PhD

Lecturer Faculty of Biotechnology and Molecular Sciences Universiti Putra Malaysia (Member)

eij

AINI IDERIS, PhD Professor/Dean School of Graduate Studies Universiti Putra Malaysia

Date : 10 AUG 2006



### **DECLARATION**

I hereby declare that the thesis is based on my original work except for quotations and citations which have been duly acknowledged. I also declare that it has not been previously or concurrently submitted for any other degree at UPM or other institutions.

Lailas

# LAILATUL JUMAIYAH BINTI SALEH HUDDIN

Date: 30 June 2006



# **TABLE OF CONTENTS**

DEDICATION	ii
ABSTRACT	iii
ABSTRAK	v
ACKNOWLEDGEMNT	vii
APPROVAL	viii
LIST OF TABLES	xiii
LIST OF FIGURES	xiv
LIST OF ABBREVIATIONS	xv

## CHAPTER

Ι	INTRODUCTION	1
	Objectives of study	4
II	LITERATURE REVIEW	5
	Glutathione	5 8
	Glutathione S-transferases	
	Molecular properties of glutathione S-transferases	9
	Substrates specification	12
	Classes of glutathione S-transferases	15
	Alpha class	15
	Mu class	16
	Pi class	16
	Theta class	16
	Sigma class	17
	Kappa class	17
	Zeta class	17
	Distribution of glutathione S-transferases	18
	Glutathione S-transferases purification	19
	Kedah-Kelantan cattle (Bos indicus) and water buffalo (Bubalus	23
	bubalis)	
III	METHODOLOGY	25
	Enzyme source	25
	Chemicals and Equipments	25
	Purification of the Kedah-Kelantan cattle (Bos indicus) and water	26
	buffalo (Bubalus bubalis) liver glutathione S-transferases	
	Cytosol purification by glutathione-agarose affinity	26
	chromatography	•
	Measurement of protein concentration	28
	Dialysis	29
	Enzyme activity assay and substrate specificity	29



	1-chloro-2,4-dinitrobenzene (CDNB)	29
	Enzyme activity assay and substrate specificity	29
	Sodium Dodecyl Sulphate – Polyacrylamide Gel Electrophoresis	32
	(SDS-PAGE)	
	Sample preparation	
	Isoelectric focusing	35
	Summary of of purification of glutathione S-transferases	35
	isoenzymes from the livers of Kedah-Kelantan cattle (Bos indicus) and water buffalo (Bubalus bubalis)	
IV	RESULTS AND DISCUSSIONS	38
	Purification of cytosol by agarose-glutathione affinity chromatography	38
	Characterization of glutathione S-transferases	45
	Isoelectric focusing	45
	SDS-PAGE	52
	Substrates specificities	60
v	Conclusions	61
REFERE	NCES	64
APPEND	ICES	72
BIODAT	A OF THE AUTHOR	77



# LIST OF TABLES

Tables		Page
1	Classification of rat Glutathione S-transferases	10
2	Classification and nomenclature of the human glutathione S- transferases.	13
3	Methodology used to purify the glutathione S-transferases from various sources	21
4	Purification step for Glutathione-affinity chromatography	26
5	Solution preparation for SDS-PAGE method	31
6	The solution preparation for heavy dense gradient, electrode and less dense gradient	35
7	Purification table of GSTs from liver of Kedah-Kelantan cattle ( <i>Bos indicus</i> )	41
8	Purification table of GSTs from liver of water buffalo (Bubalus bubalis)	41
9	Kedah-Kelantan cattle ( <i>Bos indicus</i> ) and water buffalo ( <i>Bubalus bubalis</i> ) sample fraction and the molecular weight and the isoelectric focusing points (pI)	54
10	Kedah-Kelantan cattle ( <i>Bos indicus</i> ) and water buffalo ( <i>Bubalus bubalis</i> ) glutathione-affinity chromatography purified isoenzymes fractions for the substrate specificity	60

xiii



## **LIST OF FIGURES**

Figures 1	Structure of glutathione	Page 6
2	Mercapturic acid biosynthesis pathway	7
3	Glutathione-affinity chromatography purification elution profile for Kedah-Kelantan cattle ( <i>Bos indicus</i> ) liver	39
4	Glutathione-affinity chromatography purification elution profile for water buffalo ( <i>Bubalus bubalis</i> ) liver	40
5	Elution graph for the isoelectric focusing method using LKB-8100 for Kedah-Kelantan cattle ( <i>Bos indicus</i> ) affinity purified.	47
6	Elution graph for the isoelectric focusing method using LKB-8100 for Kedah-Kelantan cattle ( <i>Bos indicus</i> ) cytosolic fraction	48
7	Elution graph for the isoelectric focusing method using LKB-8100 for water buffalo ( <i>Bubalus bubalis</i> ) affinity purified.	49
8	Elution graph for the isoelectric focusing method using LKB-8100 for water buffalo ( <i>Bubalus bubalis</i> ) cytosolic fraction.	50
9	SDS-PAGE of Kedah-Kelantan cattle ( <i>Bos indicus</i> ) liver (glutathione-affinity purified)	55
10	SDS-PAGE of water buffalo ( <i>Bubalus bubalis</i> ) liver (glutathione-affinity purified)	56
11	SDS-PAGE of water buffalo ( <i>Bubalus bubalis</i> ) liver cytosolic isoenzymes for fractions collected from isoelectric focusing	57
12	SDS-PAGE of Kedah-Kelantan cattle ( <i>Bos indicus</i> ) liver cytosolic isoenzymes for fractions collected from isoelectric focusing	58
13	SDS-PAGE of Kedah-Kelantan cattle ( <i>Bos indicus</i> ) and water buffalo ( <i>Bubalus bubalis</i> ) liver GSTs on 12% gel	59



## LIST OF ABBREVIATIONS

%	percent
°C	degree Celsius
μg	microgram
μΙ	microliter
CDNB	1-chloro-2,4-nitrobenzene
DCNB	1,2-dichloro-4-nitrobenzene
EA	ethacrynic acid
EDTA	ethylenediaminetetra acetic acid
g	gram
HCl	hydrochloric acid
KCl	Potassium Chloride
kDa	kiloDalton
NaCl	Sodium chloride
L	liter
Μ	molar
mA	miliAmpere
mg	milligram
min	minute
mL	milliliter
mM	milimolar
PBS	phosphate buffered saline
pН	- log concentration of $H^+$ ion ( <i>Puissance hydrogen</i> )



pI	Isoelectric point
TEMED	N,N,N',N'-Tetramethyl-ethylenediamine
U	units
V	Volts
v/v	volume/volume
W	Watts
w/v	weight/volume
x	times

xvi



#### **CHAPTER I**

#### **INTRODUCTION**

Living organisms are exposed to an increasing number of toxic compounds in the environment, as well as the increasing variety of drugs. The toxic compound, also referred to as xenobiotics, include chemicals in the water, air, food additive or drugs. To get rid of these xenobiotics, the body uses the process of detoxification; a complex series of reaction, to get rid of molecules (toxins) whose prolonged presence may have damaging effects on tissues or lead to undesirable effect.

The detoxification process which is the conversion of non-polar (lipophilic) toxins to polar (hydrophilic) and non-toxic metabolites, occurs in two steps, namely Phase I and Phase II. Most cells are equipped with both of these biotransforming enzymes. Phase I metabolism introduces a functional group into molecule, while phase II metabolism involves conjugation of the phase I metabolites with endogenous substrate such as sulfate, glutathione, glucuronic acid and amino acids (Ionnides *et al.*, 1984). The induction of enzymes involved in detoxification may be caused by substances that selectively unregulated a Phase I enzyme without co-induction of the corresponding Phase II enzyme.

Phase I reactions are catalyzed by a multitude of enzyme activities; the most significant one is the cytochrome P450 (CYP450) supergene family of isoenzymes which has a very broad specificity. The reaction of CYP 450 will generate reactive molecules which often maybe more toxic than the parent compound. The intermediate metabolite is further metabolized by phase II enzymes or otherwise it



may react and cause damage to protein, RNA and DNA within the cell. While in the Phase II metabolism, the biotransformed molecules generated in the Phase I are conjugated by the addition of a water-soluble group to the reactive site; this increase their solubility and thus facilitates excretion in the urine or bile (Grant, 1991). The main types of enzymes catalyzing the Phase II reaction are such as glucuronyl transferase, glutathione S-transferases, amino acid transferases and epoxide hydrolase. Yet, not all xenobiotics go to the same path of metabolism; from Phase I to Phase II route. Instead of going through the Phase I step, they initially undergo the detoxification directly to Phase II.

In the Phase II, glutathione S-transferases (GSTs; EC 2.5.1.18) a phase II enzyme, ubiquitous, inducible, dimeric protein and also the most abundant protein in the cytosolic fraction of the liver (Booth et al., 1961; Wilce and Parker, 1994; Perez-Lopez et al., 1998). The GST have aroused much interest because of its involvement in the biotransformation and detoxification of a wide spectrum of endogenous and xenobiotics compounds. These functions really suit its major role as the phase II detoxification enzyme that conjugates the cellular nucleophile glutathione with a wide range of endogenous or xenobiotic hydrophobic molecules (Mannervik and Danielson 1988; Hayes and Pulford 1995; Armstrong 1997).

The Kedah-Kelantan (KK) cattle (*Bos indicus*) and water buffalo (*Bubalus bubalis*) are two livestocks that are very important to the small holders. The KK cattle and the water buffalo normally are free-range. They are free to roam and feed on vegetation in surrounding areas in rural areas. Thus, they are exposed to the agrochemicals and pesticides which are applied by the villagers and farmers. It is very important to



gather information about biochemical functions especially the patterns of glutathione S-transferases of both animals as it might be a useful tool in environment monitoring.

,





### **Objectives of the study**

The present study is mainly concerned about the partial purification and characterization of cytosolic glutathione S-transferase (GST) from the livers of Kedah-Kelantan (KK) cattle (*Bos indicus*) and water buffalo (*Bubalus bubalis*). Both species are the most economic importance for the meat market in Malaysia.

Objectives of the study are:

- 1. To partially purify cytosolic GST from both species using the agaroseglutathione affinity chromatography gel.
- 2. To partially characterize the partially purified GST by preparative isoelectric focusing, and SDS-PAGE. Characterization of the partially purified and isolated isoforms are carried out using the different substrates; 1-chloro-2,4-dinitrobenzene (CDNB) (broad specificity), 1,2-dichloro-4-nitrobenzene (DCNB) (relatively selective for rat Mu class GST) and ethacrynic acid (EA) (selective for rat Pi class GST) (Habig and Jakoby 1981).

This study hopes to establish the patterns of cytosolic glutathione S-transferase in the KK cattle and water buffalo. This might be useful to further achieve an understanding towards this enzyme in view of using it as a tool in environmental monitoring.



#### CHAPTER II

#### LITERATURE REVIEW

#### Glutathione

Glutathione (GSH) is widely found in all forms of life and plays an essential role in the health of organisms. It is a submajor constituent of all cells and is almost always the major non protein thiol compound present in cells. Glutathione (GSH) is a tripeptide with the sequence of  $\gamma$ -glutamyl-cysteinyl-glycine and with molecular weight of 307.33 daltons. The disulfide derived from GSH by oxidation of the thiol group of the cysteine residue is usually denoted as glutathione disulfide (GSSG).

Glutathione concentration ranges between 0.1 and 10mM in mammalian cells and its sulfhydryl group comprises 10-20% of the non-protein sulfhydryl groups in the cell (Manoharan *et al.*, 1992) representing the major intracellular low molecular weight sulfhydryl compound in animals, plants and in most microorganisms (Sies, 1998). The liver acts as the principal site of glutathione synthesis, the most important chemically active group present in the GSH molecule with respect to its biological and biochemical activity is the thiol group. In healthy tissues, more than 90 percent of the GSH pooled is in the reduced form and less than 10 percent exist in the disulfide form. The enzyme glutathione reductase is the principal enzyme that maintains the GSH in reduced form.

5

The GSH molecule has two peptide bonds, two carboxylic acid groups, one amino group and one thiol (Figure 1). The high number of hydrophilic functional groups combined with a low molecular weight leads to a high water solubility for GSH.

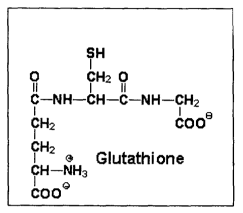


Figure 1: Structure of glutathione (γ-glu-cys-gly).

There are also a number of GSH-dependent enzymes that are part of the cellular protection against endogenous and xenobiotic toxic substances. Glutathione reductase (GR) catalyzes the reduction of GSSG using NADPH as a reductant (Krohne-Ehrich *et al.* 1977). GR is important to keep the high cellular reductive potential. Selenium dependent glutathione peroxidase (GPxs) is another GSH-linked enzyme that catalyzes the reduction of peroxides using GSH as the reducing agent (Krohne-Ehrich *et al.* 1977). Finally, last but not the least, glutathione S-transferases (GSTs) is also a GSH dependent enzymes with many properties among which catalyzing the conjugation of GSH to various electrophilic compounds is one of the most investigated function.

The cystenyl residue of glutathione provides a nucleophilic thiol important for the detoxification of electrophilic metabolites and metabolically produced oxidizing



agents. Its net negative charge and overall hydrophilicity greatly increases the aqueous solubility of the lipophilic moieties with which it becomes conjugated. Its molecular weight ensure that its adducts are preferentially secreted via the biliary system which selects molecules of molecular weight greater than 300 to 500 according to the species (Ketterer *et al.* 1983).

In mammals, GSH conjugates are often further metabolized by hydrolysis and Nacetylcystenyl conjugates known as mercapturic acids, which are excreted in the urine. This is presumed that GSH provides a means whereby the pool of cysteine for detoxification is kept separate from the pool of cysteine for protein synthesis (Ketterer *et al.* 1983).

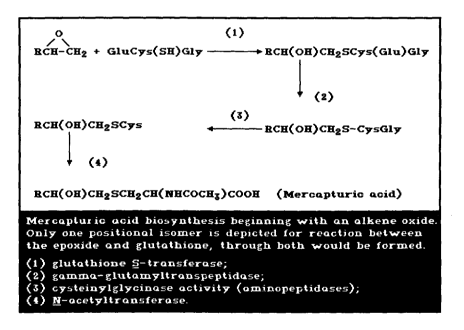


Figure 2: Mercapturic acid biosynthesis pathway.

Taken from www.inchem.org/documents/ ehc/ehc/ehc57.htm





GSH also plays roles in catalysis, metabolism, and signal transduction, gene expression and apoptosis. The most important is that glutathione is a cofactor for glutathione S-transferases (GST), enzymes which are involved in the detoxification of xenobiotics (Meyer *et al.*, 1985).

Though the GSH is undoubtedly a potent (Ioannides *et al.* 1984) antioxidant, indication for its use as supplement are not yet been well established. There is preliminary evidence that it might eventually prove to be useful in management of some cancers, atherosclerosis, and diabetes and also to help prevent or improve various toxicities.

#### **Glutathione S-Transferases**

The glutathione S- transferases (also known as GSTs; EC 2.5.1.18) are a family of multifunctional enzymes, found to play an important role in the detoxification of wide variety of xenobiotics (Hunaiti and Owais, 1985; (Habig *et al.* 1974; Prohaska 1980; Hunaiti and Owais 1985; Meyer *et al.* 1985; Ketterer 1986). First found as an enzyme by Booth and co-workers in 1961, GSTs are also proposed to act as carrier protein and were named ligandin (Litwack *et al.*, 1971).Today, despite of research for 40 years, the 'picture' of exact function of the superfamily, is more complex than ever.

GSTs which known as multifunction enzymes are capable of catalyzing a seemingly protean spectrum of reactions; widely distributed and are present at high concentrations in cytosol (Jakoby *et al.* 1984). GSTs can be found mostly in liver,

