

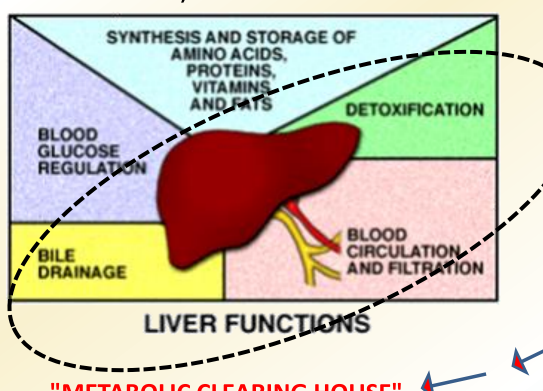


has multiple functions (>5000) → can be categorized into:

1. Regulation, synthesis & secretion. → utilization of glucose, lipids & proteins + bile for digesting fats.

2. Storage. → Glucose (as glycogen), fat soluble vitamins (A, D, E & K) & minerals

3. Purification, transformation & clearance → of endogenous (steroid hormones, cholestrol, FA, & proteins...) & exogenous (drugs, toxins, herbs...etc) chemicals.



Human body
identifies almost all
drugs as foreign
substances i.e.
XENOBIOTIC

Has to get rid of them

HEPATOTOXIC DRUGS

Subjects drugs to chemical transformation (METABOLISM) → to become inactive & easily excreted. Since most drugs are lipophilic they are changed into hydrophilic water soluble products → suitable for elimination through the bile or urine

Such metabolic transformation usually occur in 2 PHASES:

Phase 1 reactions
Oxidation, Reduction,
Hydrolysis, Hydration
Catalyzed by CYT P-450

Yields intermediates →
polar, transient, usually highly reactive →
far more toxic than parent substrates →
may result in liver injury

Drug Induced Liver Injury (DILI)

Phase 2 reactions
Conjugation with a moiety
(acetate, a.a., glutathione,
glucuronic a., sulfate)

Yields products of increased solubility

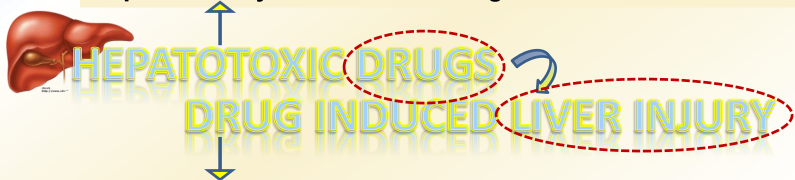
If of high molecular weight →

excreted in bile

If of low molecular weight → to blood →

excreted in urine

Hepatotoxicity → Is the Leading cause of ADRs



Injury / damage of the liver →

Caused by exposure to a drug → Inflict varying impairment in liver functions →

Manifests clinically a long range → hepatitis ⇒failure

Inflammation ⇒**Apoptosis** ⇒ **Necrosis**



Why the liver is the major site of ADRs?

- **♣It** is the first organ to come in contact with the drug after absorption from the GIT.
- **Learning House of the body** → it expresses the highest levels of drug metabolizing enzymes that converts some drugs(PROTOXINS) into intermediate (TOXINS) before being conjugated for elimination

Drug (Pro-toxin) \rightarrow Toxin \rightarrow Injury

Paracetamol $\rightarrow CYT P450 \rightarrow$ NABQI centrilobular

(NAPBQI): N-acetyl-p-benzoquinone imine



Can any drug cause liver-related ADRs?

Not all drugs do so.

Drugs that can cause ADRs in the liver (hepatotoxicity) → are called → HEPATOTOXIN

TOXICITY POTENTIAL OF THE DRUG

- **4Chemical composition of the drug itself**
- **Nature of its reactive metabolite**
- Conjugation reactions linked to it & their availability
- Mitochondrial effects of the drug
- Drug formulation

...etc

HEPATOTOXIC DRUGS

1. Nature of a Hepatotoxin 2. Types of drug-induced hepatotoxic ADRs it inflicts?

If the toxicity of HEPATOTOXIN is inflicted by:

SUPERTHERAPEUTIC or CUMULATIVE DOSE of the drug

→INTRINSIC HEPATOTOXIN

The hepatotoxicity it inflicts is → <u>DIRECT HEPATOTOXICITY</u> → belong to TYPE A ADRs: <u>PREDICTABLE</u> / <u>DIRECT</u>

If the toxicity is inflicted by NORMAL DOSE of the drug → IDIOSYNCRATIC HEPATOTOXIN

The hepatotoxicity it inflicts is → INDIRECT HEPATOTOXICITY

→belong to TYPE B ADRs: UNPREDICTABLE / BIZZAR /

IDIOSYNCRATIC

Types of drug-induced hepatotoxic ADRs?

1. DIRECT HEPATOTOXICITY caused by INTRINSIC HEPATOTOXIN

Type A

Dose-dependent hepatotoxicity

Direct increased dose dependent hepatotoxicity

- Acetaminophen
- Salicylates
- Statins
- **Amiodarone**
- Methotrexate
- Alcohol

Increased Dose

Increased Dose

Increased Dose

Cumulative Dose/effect

Increased & Cumulative

Oral contraceptives Cumulative Dose/effect

Increased & Cumulative Doses/effect

Direct cumulative hepatotoxicity

2. INDIRECT HEPATOTOXICITY caused by IDIOSYNCRATIC HEPATOTOXIN

Type B Dose-independent hepatotoxicity → divided into:

- Hypersensitivity or immunoallergic reactions
- Metabolic-idiosyncratic reactions

2.a. Immunoallergic Idiosyncratic Hepatotoxicity

Type B

A drug or its metabolite binds to hepatic membranes or proteins

→act as hapten to induce a variety of immune reactions

Inflammatory cholestasis	Viral hepatitis-like pattern
Chlorpromazine.Chlorpropamide.Erythromycin.	↓ Isoniazid.↓ Phenytoin.↓ Methyldopa.

2. INDIRECT HEPATOTOXICITY caused by IDIOSYNCRATIC HEPATOTOXIN

2.b. Metabolic Idiosyncratic Hepatotoxicity

TypeB

The metabolite of the offending drug interferes with hepatic metabolism as that of bilirubin or protein synthesis....etc

Interfere with bilirubin metabolism

- Erythromycin
- **A** Rifampicin

Interfere with protein synthesis

- **Les Corticosteroids**
- **Tetracycline**

N.B. Not all drugs fall neatly into one of these categories, and overlapping mechanisms may occur with some drugs

HOW CAN A DRUG INDUCE HEPATOTOXICITY?

Drug or its reactive metabolites can form <u>covalent bonds</u> with target molecules or alter the target molecule by <u>non-covalent interactions</u> or both

COVALENT INTERACTIONS

It is <u>adduct</u> formation between the metabolite of the drug & cellular macromolecules

If covalent binding to protein → immunogenic reaction

If binding to DNA → carcinogenesis

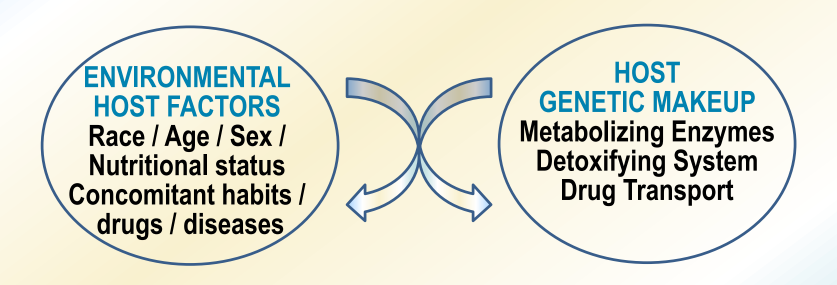
NON-COVALENT INTERACTIONS

- Lipid peroxidation → generation of cytotoxic oxygen radicals
- Impairment of mitochondrial respiration
- Depletion of GSH reactions → 'oxidative stress'
- Modification of sulfhydryl groups → impair Ca²⁺homostasis
- Protein synthesis inhibition

....etc

Do hepatotoxins cause liver disease in all persons?

Most hepatotoxins cause liver disease only in certain persons depending on:





DRUG-INDUCED LIVER INJURY

DRUG-INDUCED HEPATIC INJURY

Is DIHI common?

INCIDENCE of DILI

Drugs produce about 10% of all cases of hepatitis in young adults and 40% of cases in patients older than 50 years.

Are certain persons or population more susceptible?

Upon exposure to hepatotoxins people are categorized as;

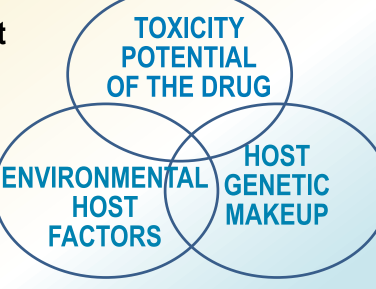
4Tolerators → No injury

♣Adaptors **→** Mild transient injury but adapt

Language Susceptibles → Develop overt symptoms

depending on existing predisposing factors

Lin Threat; DILI accelerates beyond initial targets due to **→** loss of synthetic & clearance function of hepatocyte with recruitment of inflammatory cells provoke apoptotic & necrotic signals



What are the presenting manifestations?

Individual drugs tend to have → CHARACTERISTIC SIGNATURE → composed of: A particular latency period

A clinical pattern

A particular pathological finding

LATENCY PERIOD → short (hrs/dys), intermediate (1-8ws), long (1-12ms)

- **Latency period** → SHORT as it occurs after a threshold of toxicity is reached
 - **→** acetaminophen (toxic dose)
- **Latency Latency Latency Period INTERMEDIATE Latency Direct Latency Direct INTERMEDIATE Latency Direct Direct Latency Direct Latency Latency Direct Latency Latency Direct Latency Direct Latency Direct Latency Latency Latency Latency Latency Latency Latency Latency La**

CLINICAL PATTERNS

The clinical presentation could be of variable intensity, ranging from asymptomatic ♠ of liver enzymes → → fulminant hepatic failure

Some drugs just induce → ASYMPTOMATIC

▲ IN AMINOTRANSFERASES

Statins

Sulfonamides

♣ Sulfonylureas

Other drugs induce →

SYMPTOMATIC MANIFESTATIONS

- ♣If injury targets hepatocytes → apoptosis or necrosis → HEPATITIS
 (cytotoxic) develops → rapid onset of malaise, severe anorexia and
 jaundice + ♠ in alanine aminotransferases (ALT)
- ↓If injury targets biliary system (canalicular or ductal) → CHOLESTASIS develop → jaundice + severe pruritus predominate → ♠ in alkaline phosphatase (ALP) + hyperbilirubinaemia
- <u>If injury targets both</u> hepatocytes & biliary system → MIXED TYPE

Some PATTERNS of SYMPTOMATIC drug-induced liver disease				
Hepatic injury	Hepatocellular	Cholestatic	Mixed	
	Flu-like, malaise, m. aches weakness, loss of appetite, GIT symptoms, diarrhea, jaundice, urine discolored,	Yellowish discoloration of skin, dark urine, rash, <u>pruritus</u> , stool may be light		
<u>ALT</u>	≥ 3 fold rise	Normal or slight	≥ 3 fold rise	
ALP	Normal	≥ 2 fold rise	≥ 2 fold rise	
Examples	Acetaminophen NSAIDs Isoniazid Amiodarone	Chlorpropamide Erythromycin Rifamycin Oral contraceptives	Phenytoin Carbamazepine Sulfonamides ACE Inhibitors	

→ A long standing rheumatoid arthritic patient developed tuberculosis 2 month ago. Today she was received in E.R complaining of yellowish discoloration of skin, severe anorexia, vomiting and flue like manifestations since two days. She is very weak and looks toxic. Her drug history reveals that she has been 4 month ago on cyclosporine to control the arthiritic exacerbations. A month ago, she was put on isoniazid when she developed T.B. and multivitamins because she is weak. Currently she is given domperidone for the emesis. Lab results reveals severe elevation in ALT but no elevation in ALP.

Hepatocellular

- **Which one of the following drugs is the likely cause of her symptoms?**
- a. Cyclosporine b. Multivitamines
- c. Isoniazid d. Domperidone
- **Which type of hepatotoxin is it considered?**
- **What is the likely hepatotoxic pattern inflicted by the drug?**

Treatment????

A hypercholestrolemic patient was received in E.R complaining of yellowish discoloration of skin, change in color of urine & stools, and severe itching. He has been receiving statins fro the long time for the hypercholestrolemia. Three month ago he was diagnosed as being diabetic and hypertensive and since then he is receiving chlorpropamide for the diabetes and nadolol for the hypertension. The last couple of days he had a flue; for which he was given acetaminophen for muscle aches and nasal drops for his nasal stuffiness. Lab investigations shows severe elevation in ALP and no significant elevation in ALT

Cholestatic

- **Which one of the following drug is the likely cause of his symptoms?**
- a. Nadolol b. Chlorpropamide c. Acetominophen d. Statins
- **Which type of hepatotoxin it is considered?**
- **What is the hepatotoxic pattern inflicted by the drug?**

Treatment????

HISTOPATHOLOGICAL PATTERNS

No universal histo-pathological pattern of DIHI exist.

The commonest are; Hepatocellular necrosis

Cholestasis

Steatosis

More than one type of injury may occur in the same patient
Any one agent may produce different types of injury in different patients

Ballooning & degeneration of hepatocyte



What are the lines of treatment?

Immediate withdrawal → of any suspected drug

No specific treatment → largely symptomatic & supportive Symptomatic:

If a <u>severe allergic reaction</u> is observed → Corticosteroids

If <u>pruritus</u> → enhance bile acid excretion → Cholestyramine

If <u>cholestatic liver injury</u> → <u>Ursodeoxycholic acid</u> (Ursodiol)

If coagulopathy or encephalopathy develop → treat accordingly

Supportive:

High carbohydrate, moderate protein diet adequate in calories

Specific antidotes

N-acetylcysteine → acetaminophen toxicity
L-carnitine → valproate toxicity

Emergency liver transplantation → for drug induced fulminant hepatic failure

