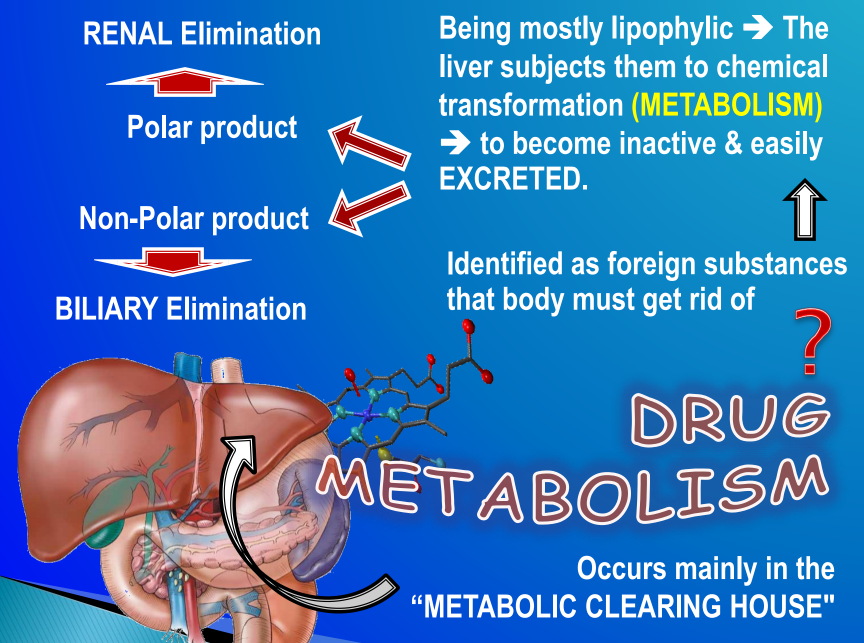


CYTOCHROME SYSTEM & DRUG METABOLISM

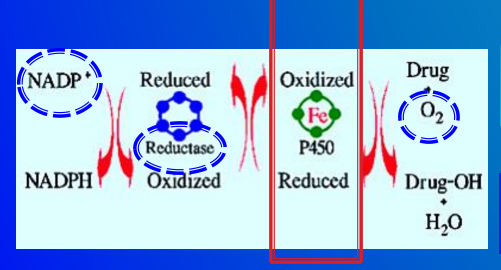
- Revise the aim & phases of drug metabolism
- > Define the role of cytochrome system in relation to drug metabolism
- Expand on the nature, location, nomenclature, structure, distribution & function of CYT P450
- Focus on its regulation; directly & indirectly, its induction & inhibition its relevance to drug interactions
- ►Interpret the molecular mechanism of interactions by CYT P450
- Classify its different isoforms, their substrates, inducers
 & inhibitors
- > Delineate some of its genetic variations.

Where do drug biotransformations occur?



" Cytochrome P450" " CYT 450"

Superfamily is the terminal rate limiting oxidase of this system



Its enzymes are part of a cascade

transfers electrons from molecular oxygen to oxidize the drugs

Inactive product

Active metabolite;

Similar to parent

More active than parent

* A product with different effect

*** Toxic metabolite**

Phase II Conjugation

Create a conjugation site

Phase I

OXIDATION / Reduction/Hydrolysis

CYTOCHROME SYSTEM



Cytochrome P450 cycle in drug oxidations

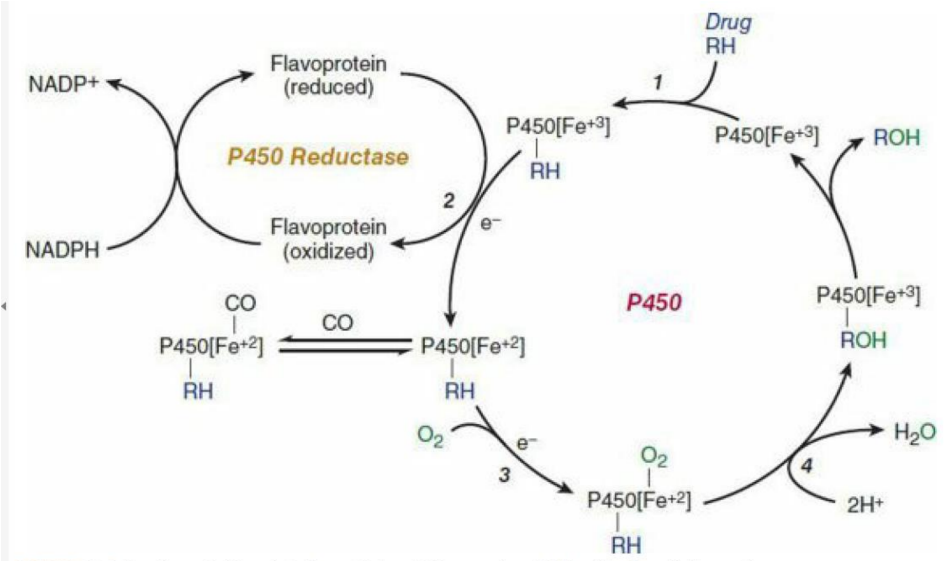
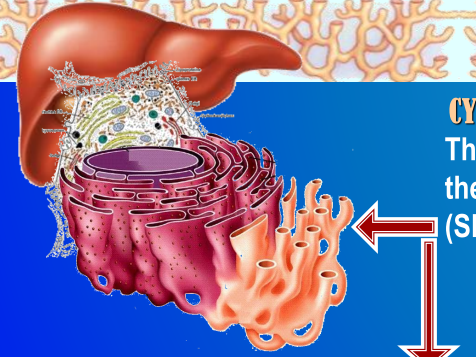


FIGURE 4-3 Cytochrome P450 cycle in drug oxidations. RH, parent drug; ROH, oxidized metabolite; e-, electron.

- ➤ Microsomal drug oxidations require: P450, P450 reductase, NADPH, & molecular O₂
- ➤Oxidized (Fe³⁺) P450 combines with drug substrate to form a binary complex (step 1)
- ➤NADPH donates an electron to the flavoprotein P450 reductase, which in turn reduces the oxidized-P450-drug complex (step 2)

- \triangleright A 2nd electron is introduced from NADPH via the same P450 reductase to form an activated O₂-P450 substrate complex (step 3)
- This complex in turn transfers activated O_2 to the drug substrate to form the oxidized product (step 4).



CYTOCHROME P450 FAMILY OF ENZYMES

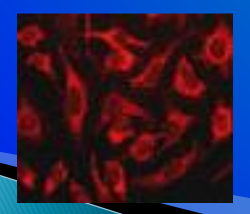
They are located mainly attached to the smooth endoplasmic reticulum (SER) of hepatocytes.

"Cytochrome" = colored cells
They color the liver cells dark red
as they contain iron

"P450" absorbs a very characteristic wavelength (450 nm) of UV light when it is exposed to carbon monoxide.

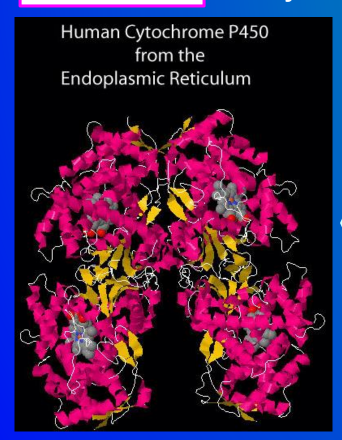
They are isolated in the subcellular fraction termed the MICROSOMES

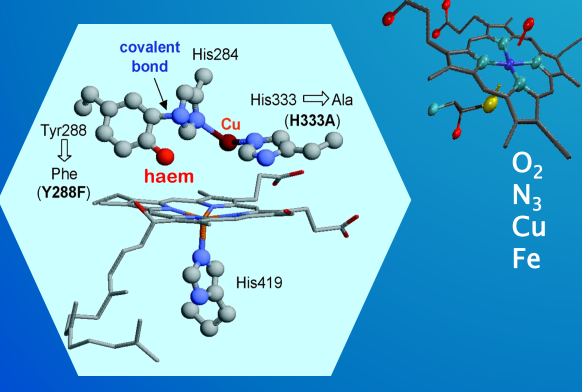
→ Liver microsomal enzymes



STRUCTURE

They are heme-containing isoenzymes





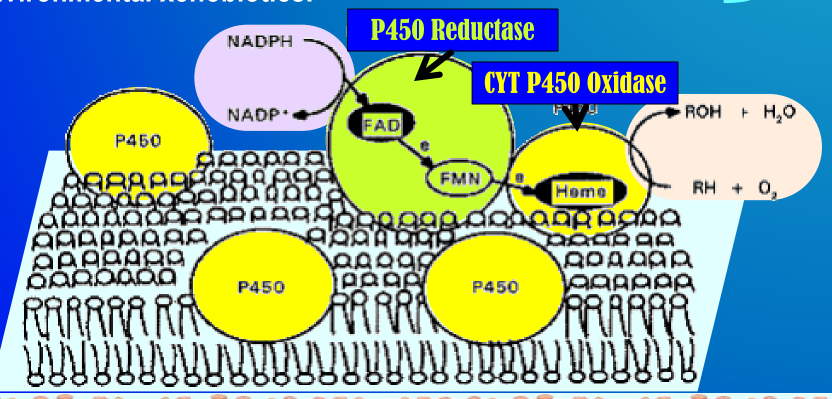
DISTRIBUTION

- Highly concentrated in hepatocytes
- Enterocytes of the small intestine present their principal extra-hepatic source
- **▶Very small quantities in kidneys, lungs, & brain.**

Function

Responsible for most of the **OXIDATIVE METABOLISM** of:

- Endogenous substances: steroid hormones, prostaglandins, lipids, & fatty acids
 Substrates
- Exogenous compounds: diet (food & beverages) / Drugs/ environmental xenobiotics.

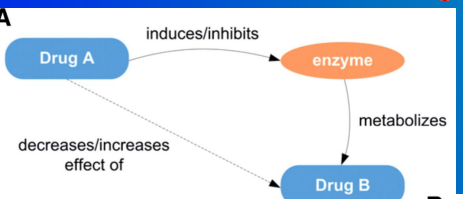


Regulation

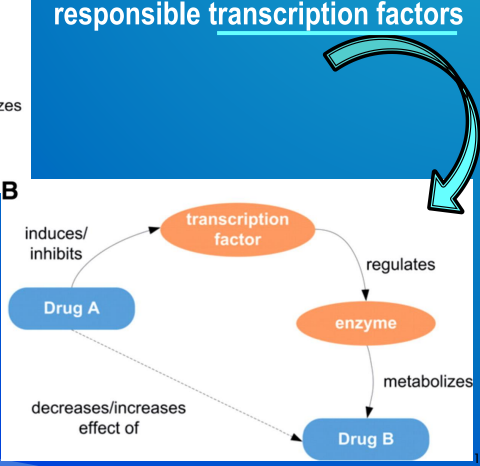
Activation or Inactivation of the CYT P450 can be achieved either

A: Directly

B: Indirectly by expression or repression of its relevant genes by



Activation or Inactivation can be processed by any food, intrinsic products or extrinsic xenobiotics as drugs (usually the lipophylic) that have to be metabolized.



activation or inhibition of the

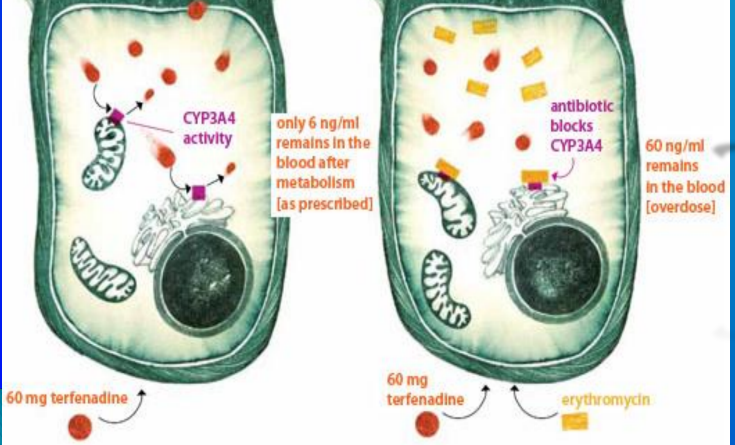
Regulation

When drugs play a role in regulation of the CYT P450 → they are termed

Enzyme Inducers if Activate the enzyme

Enzyme Inhibitors if Inactivate the enzyme

PHARMACOKINETIC DRUG-DRUG INTERACTION





"A 50 years old, patient was treated for the last 3 years by the hypocholestrolemic agent; atorvastatin. Yesterday he began to complain of severe muscle pains, weakness & reddish discoloration of urine

He receives daily <u>multivitamins</u> & his lab results last week, proved that he has become diabetic, for which he was prescribed <u>metformin</u>. He was also started on a course of fluconazole for a concomitant fungal infection.

From drug history, the diagnosis of his current state was likely rhabdomyositis (severe muscloskeletal toxicity) & was verified by the lab finding of severe elevation in creatinine phosphokinase. "

Which one of the following drug-drug interaction on CYT 3A4 is the likely cause of his current state?

Metformin + **Atrovastatin**

Atrovastatin + Fluconazole

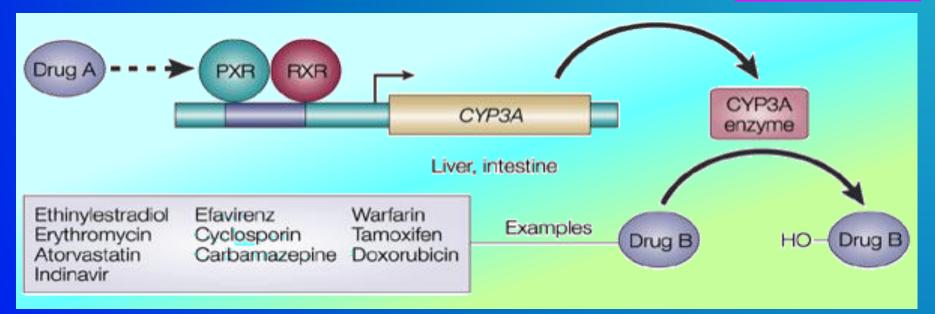
Metformin + Fluconazole

Fluconazole+ Multivitamins



Molecular Basis Of Drug-drug Interaction

Regulation



The orphan nuclear receptor PXR is a TRANSCRIPTION FACTOR that regulates the expression of the CYP P450 genes. If Drug A is INDUCER → it binds & activates PXR → which translocates in nucleus → dimerize with RXR → the heterodiamer PXR / RXR will induce EXPRESSION of CYT P450 isoenzymes to → ↑ metabolism of Drug B If Drug A is an INHIBITOR, its binding will prevent activation → **REPRESSION** of CYT P450 isoenzymes to →

metabolism of Drug B

Outcome Of Drug-drug Interactions Mediated By CYT P450

Regulation

IN RELATION TO ENZ INDUCERS

- ★ metabolism of the inducer + ★ its pharmacological action.
 Tolerance or complete nullification
- **↑** → metabolism of co-administered drugs



IN RELATION TO ENZ INHIBITORS

- →/ Retard metabolism & excretion of inhibitor & co-administered drugs
- ♠ / prolong action of the inhibitor & co-administered drugs.



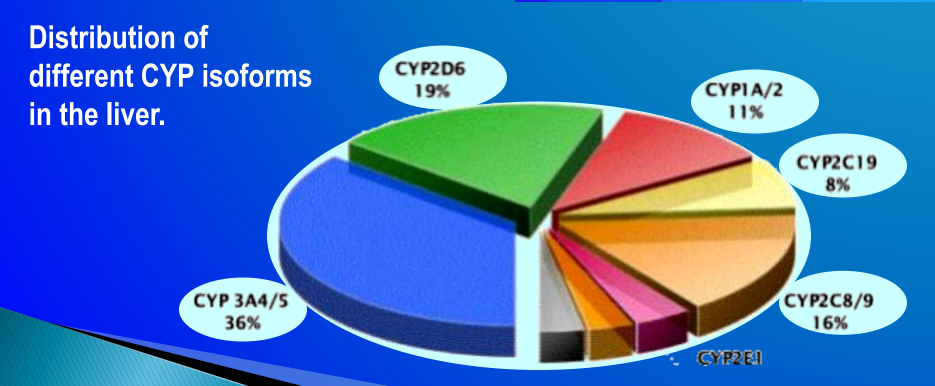
Classification

CYT P450 has been classified into

- **Property** Families designated by Numbers
- Sub families designated by Letters

Cytochrome P450 Isoforms

- CYP1A2
- CYP3A
- CYP2C9
- CYP2C19
- CYP2D6

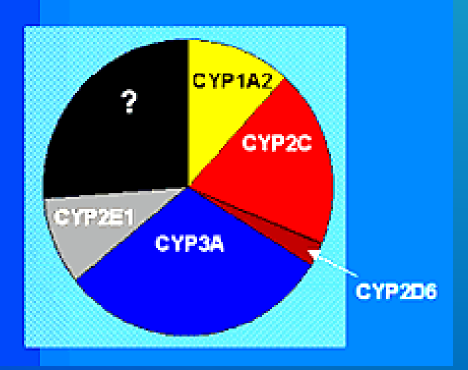


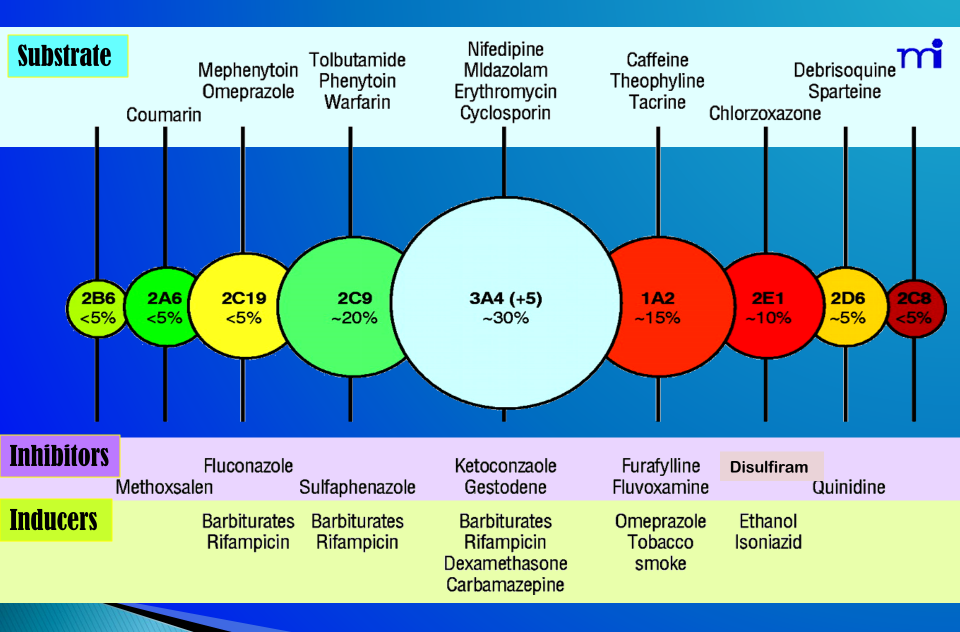
CYP450 → Major Contributor to Phase I Metabolism

Relative Importance of P450s in Drug Metabolism CYP2E1 CYP1A2



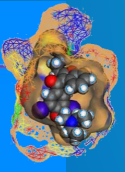
Relative Quantities of P450s in Liver

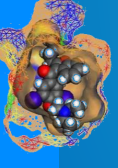




Cytochrome P450 3A

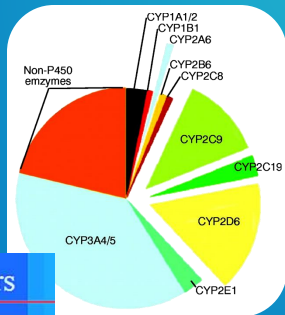
- Responsible for metabolism of:
 - Most calcium channel blockers
 - Most benzodiazepines
 - Most HIV protease inhibitors
 - Most HMG-CoA-reductase inhibitors
 - Cyclosporine
 - Most non-sedating antihistamines
 - Cisapride
- Present in GI tract and liver





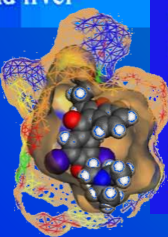
CYP3A Inhibitors

- Ketoconazole
- Itraconazole
- Fluconazole
- Cimetidine
- Clarithromycin
- Erythromycin
- Troleandomycin
- Grapefruit juice
- Ritonavir



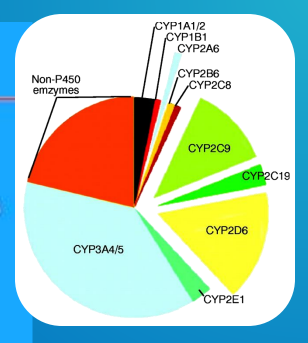
CYP3A Inducers

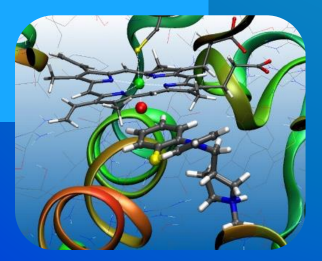
- Carbamazepine
- Rifampin
- Rifabutin
- **Barbiturates**



Cytochrome P450 2D6

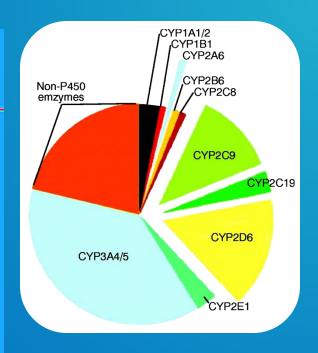
- Absent in 7% of Caucasians,
 1–2% non-Caucasians
- Hyperactive in up to 30% of East Africans
- Catalyzes primary metabolism of:
 - Codeine
 - Many β-blockers
 - Many tricyclic antidepressants
- Inhibited by:
 - -Fluoxetine
 - Haloperidol
 - Paroxetine
 - Quinidine
- Induced by Rifampicin

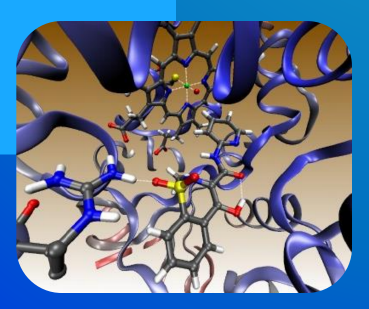




Cytochrome P450 2C9

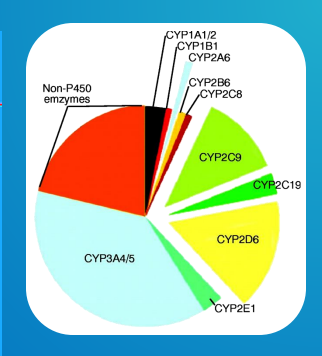
- Absent in 1% Caucasians and African-Americans
- Primary metabolism of:
 - Most NSAIDs (including COX-2)
 - S-warfarin (the active form)
 - Phenytoin
- Inhibited by:
 - Fluconazole
- Induced by
- Barbiturates
- Rifampicin

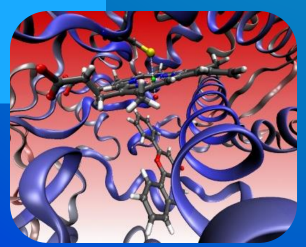




Cytochrome P450 1A2

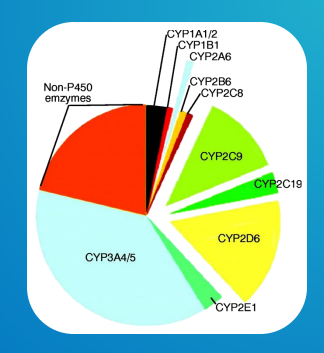
- Induced by smoking tobacco
- Catalyzes primary metabolism of:
 - Theophylline
 - Imipramine
 - Propranolol
 - Clozapine
- Inhibited by:
 - Many fluoroquinolone antibiotics
 - Fluvoxamine
 - Cimetidine

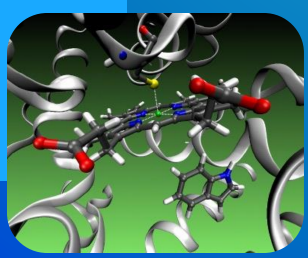




Cytochrome P450 2C19

- Absent in 20–30% of Asians,
 3–5% Caucasians
- Primary metabolism of:
 - Diazepam
 - Phenytoin
 - Omeprazole
- Inhibited by:
 - Omeprazole
 - Isoniazid
 - Ketoconazole
- Induced by
- Barbiturates
- Rifampicin





CYT P450 3A4

Substrates In	hibitors	Inducers
 Statins (Atorvastatin) Cancer Chemotherapy (Cyclophosphamide, Tamoxifen) Non-Sedating Antihistamines (Astamizole) Benzodiazipines (Midazolam, (Rito Cimeting Chlora Nefaza 	amphenicol	Phenytoin Carbamazepine Barbiturates Rifampicin Dexamethazone Progestins

"A 50 years old, patient was treated for the last 3 years by the hypocholestrolemic agent; atorvastatin. Yesterday he began to complain of severe muscle pains, weakness & reddish discoloration of urine

He receives daily <u>multivitamins</u> & his lab results last week, proved that he has become diabetic, for which he was prescribed <u>metformin</u>. He was also started on a course of <u>fluconazole</u> for a concomitant fungal infection.

From drug history, the diagnosis of his current state was likely rhabdomyositis (severe muscloskeletal toxicity) & was verified by the lab finding of severe elevation in creatinine phosphokinase. "

Which one of the following drug-drug interaction on CYT 3A4 is the likely cause of his current state?

Metformin + Atrovastatin

Atrovastatin + Fluconazole

Metformin + Fluconazole

Fluconazole+ Multivitamins

Genetic Variation

Genetic polymorphisms in CYT P450 isoenzymes have been observed & are reasons behind the **ALTERED RESPONSE** to drug therapy

CYP2D6

This isoenzyme has the most frequent polymorphisms in all CYT P450 When polymorphism occurs → → metabolizing capacity of CYP2D6 i.e those who exhibit the polymorphism become poor metabolizers:

- 1. Metabolism of some neuroleptics, tricyclic antidepressants, antianginals agent (perihexiline), antiarrhythmics (propafenone & metoprolol) is suppressed → so side effects & toxicity develop. i.e.
 - Neuropathy after therapeutic doses of perihexiline
 - Bradycardias & arrhythmias on therapeutic dose of propafenone or metaprolol
- 2. The pro-drugs cannot be converted to their therapeutically active metabolite; e.g poor analgesia with codeine & tramadol because they are not transformed into active forms

Genetic Variation

CYP2C9.

Warfarin, phenytoin, & tolbutamide are examples of drugs with narrow therapeutic index that are metabolized by CYP2C9.

Clearance of these drugs is impaired in genetic variation of the enzyme

CYP2C19

Polymorphism in CYP2C19 shows increased & prolonged action of its substrates as omeprazole

This has been an advantage as in those variants → ↑ cure rates in peptic ulcer patient with Helicobacter pylori.

Benefit

28

