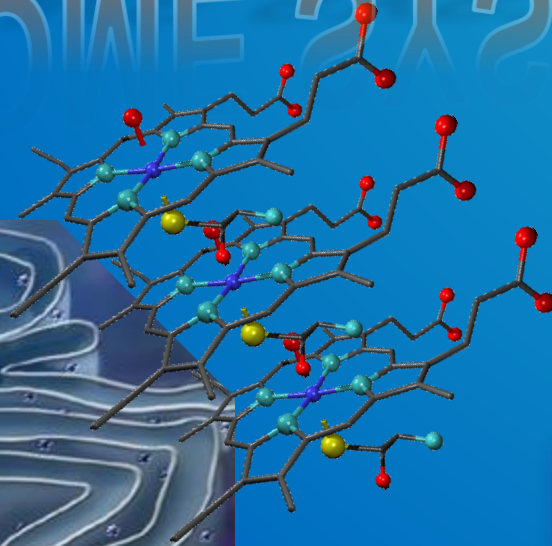
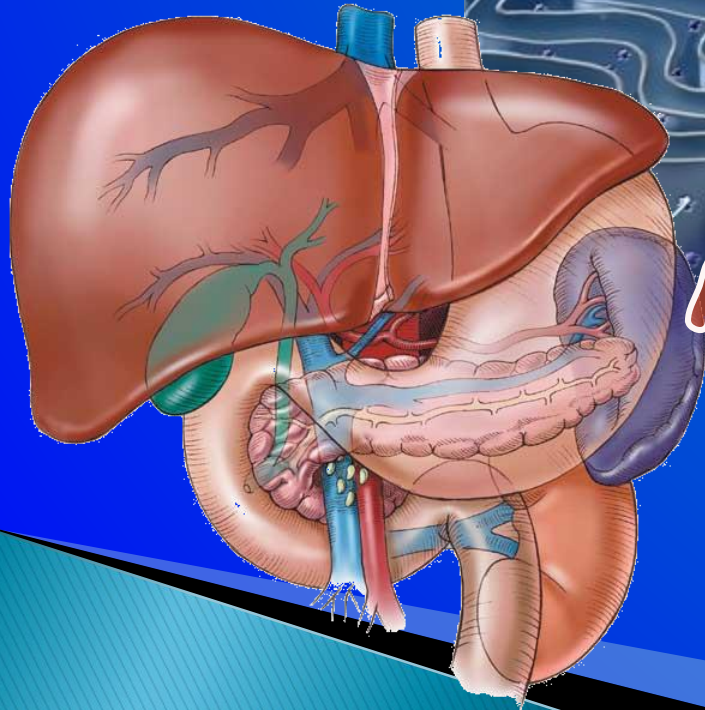


# CYTOCHROME SYSTEM

&

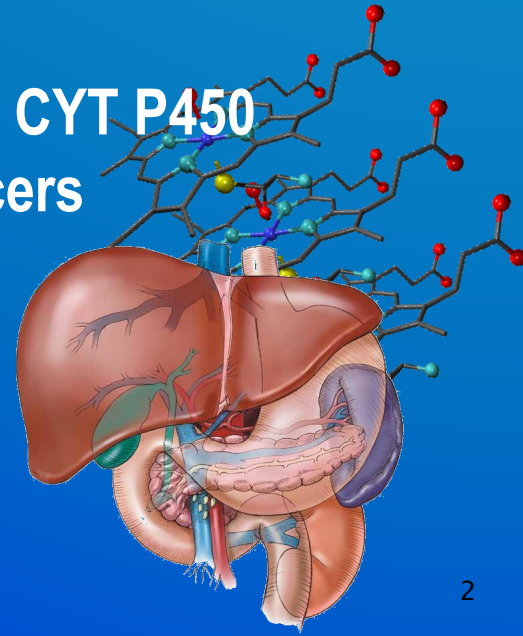
DRUG  
METABOLISM



Dec 18

# 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 ?

RENAL Elimination



Polar product

Non-Polar product



BILIARY Elimination

Being mostly lipophylic → The liver subjects them to chemical transformation (**METABOLISM**) → to become inactive & easily EXCRETED.

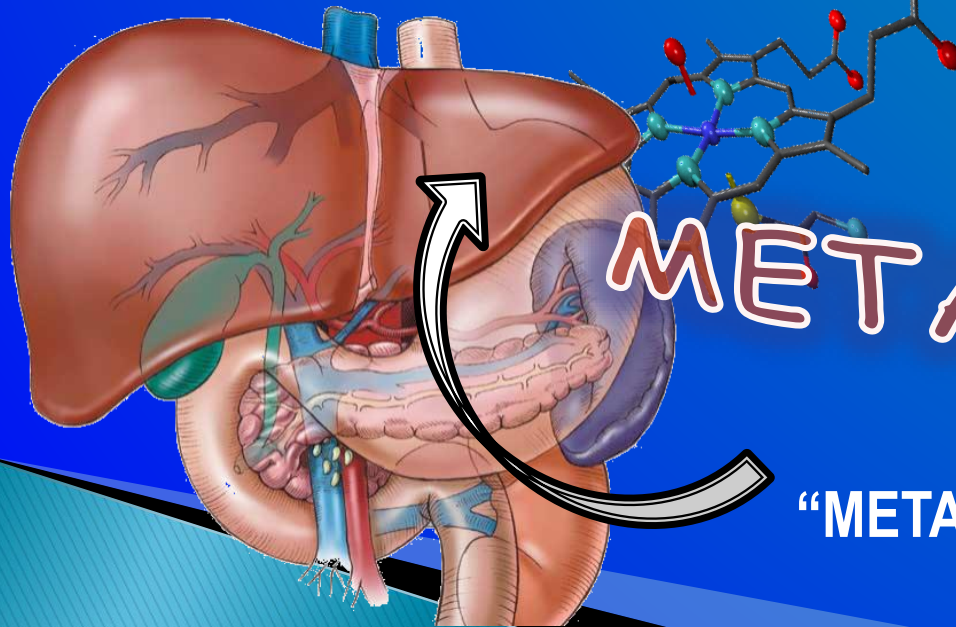


Identified as foreign substances that body must get rid of



DRUG  
METABOLISM

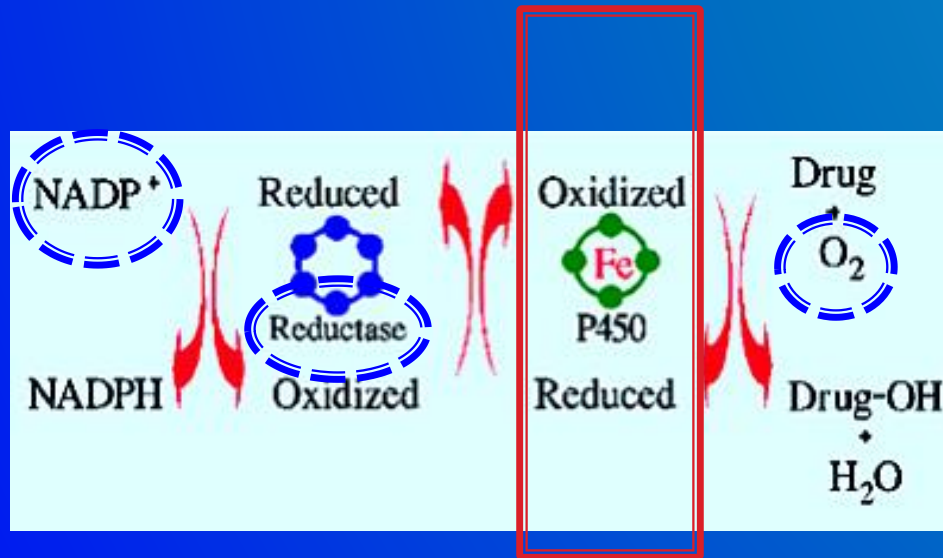
Occurs mainly in the  
“METABOLIC CLEARING HOUSE”



# “ Cytochrome P450“ “ CYT 450”

Superfamily is the terminal rate limiting oxidase of this system

- \* Inactive product
- \* Active metabolite;
- \* Similar to parent
- \* More active than parent
- \* A product with different effect
- \* Toxic metabolite



Its enzymes are part of a cascade  
→ transfers electrons from molecular oxygen to oxidize the drugs

## Phase II Conjugation

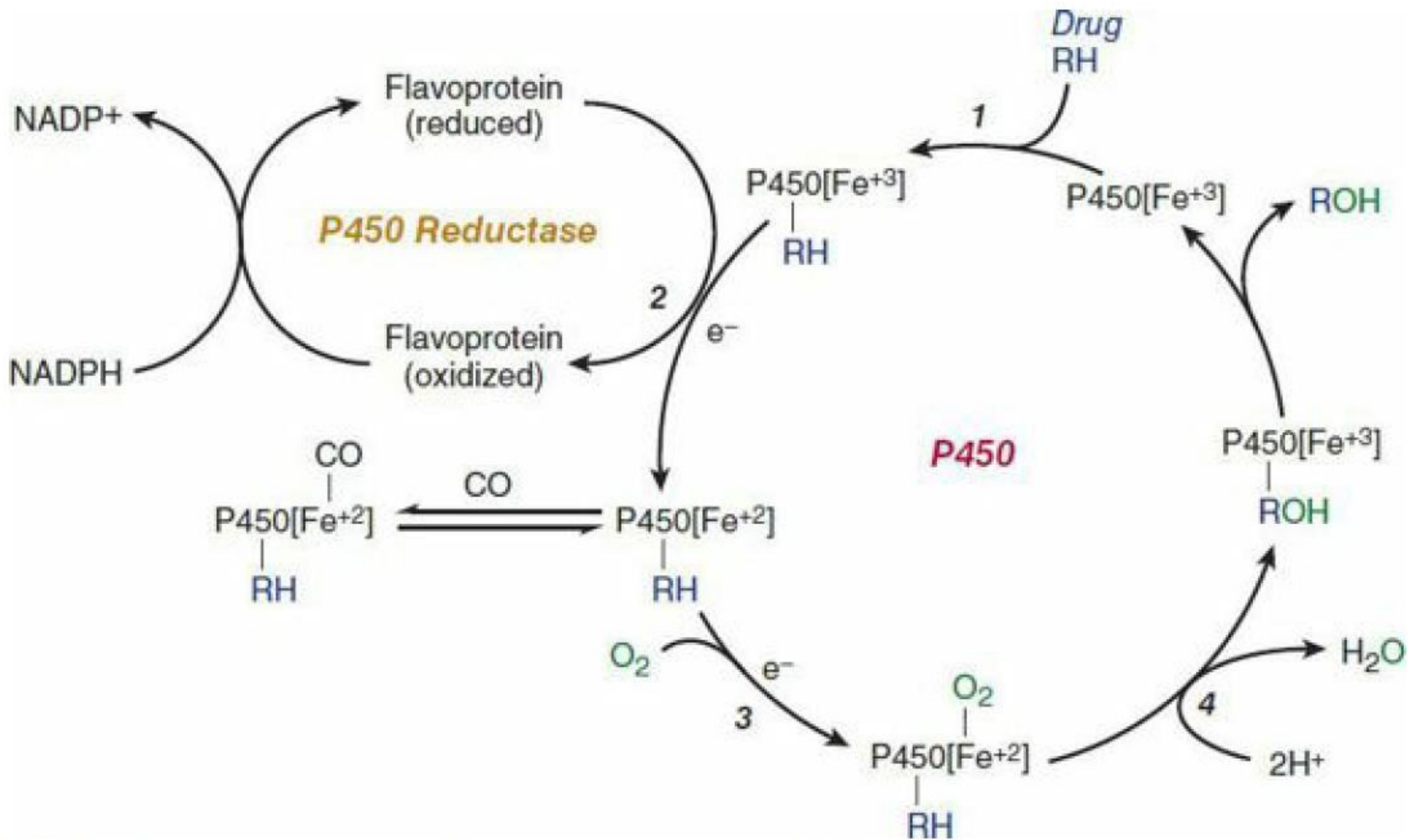
Create a conjugation site

## Phase I OXIDATION /Reduction/Hydrolysis

## CYTOCHROME SYSTEM

# DRUG METABOLISM

# Cytochrome P450 cycle in drug oxidations

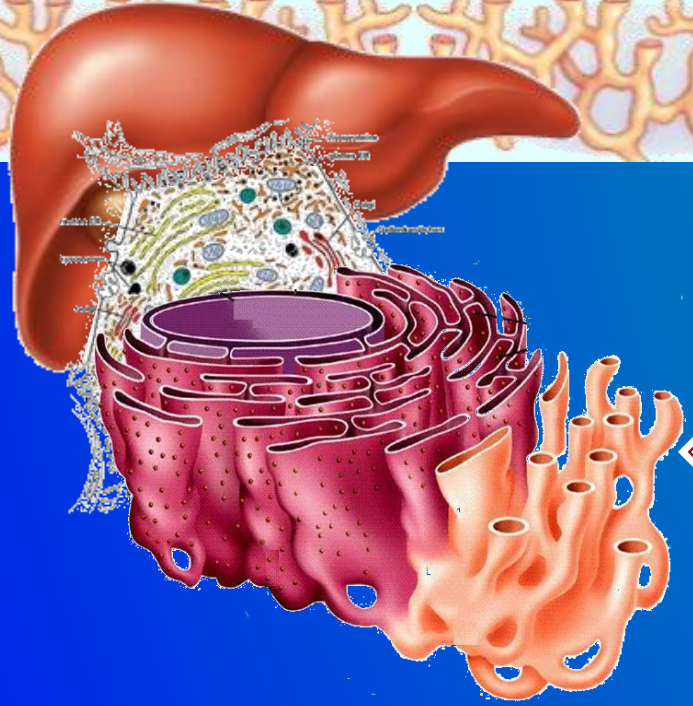


**FIGURE 4–3** Cytochrome P450 cycle in drug oxidations. RH, parent drug; ROH, oxidized metabolite; e<sup>-</sup>, electron.

- Microsomal drug oxidations require: P450, P450 reductase, NADPH, & molecular O<sub>2</sub>
- Oxidized (Fe<sup>3+</sup>) 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**)

- A 2<sup>nd</sup> electron is introduced from NADPH via the same P450 reductase to form an activated O<sub>2</sub>-P450 substrate complex (**step 3**)
- This complex in turn transfers activated O<sub>2</sub> to the drug substrate to form the oxidized product (**step 4**).



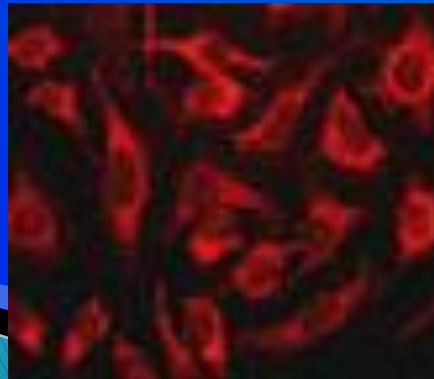


## CYTOCHROME P450 FAMILY OF ENZYMES

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



They are isolated in the subcellular fraction termed the **MICROSOMES**  
→ **Liver microsomal enzymes**



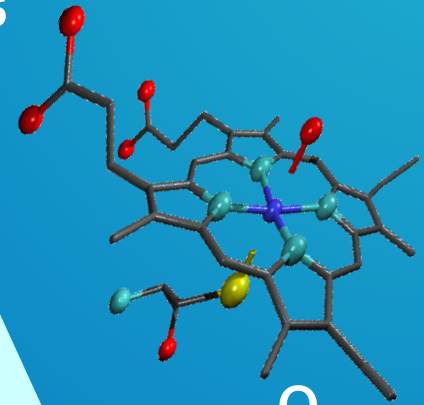
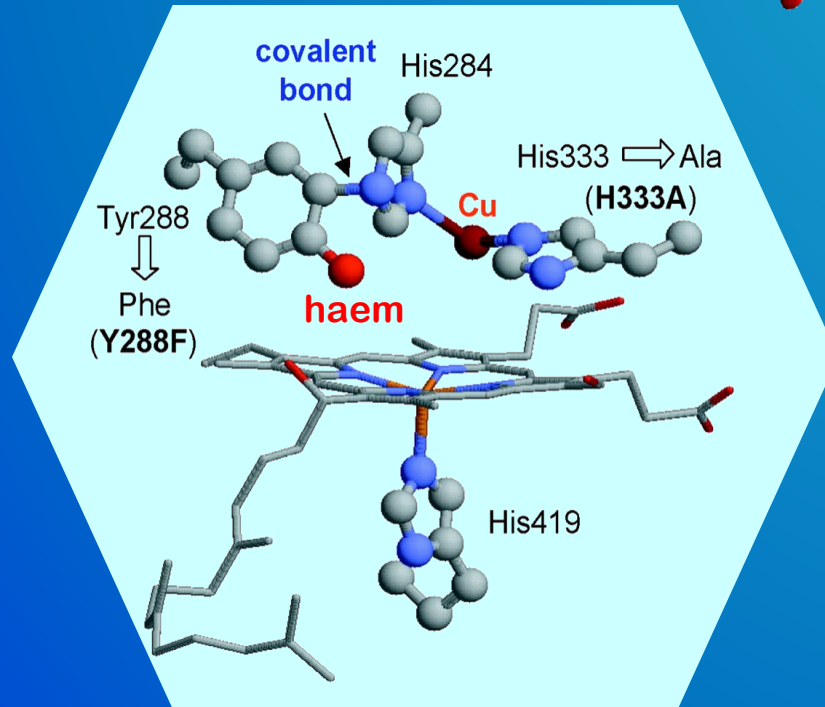
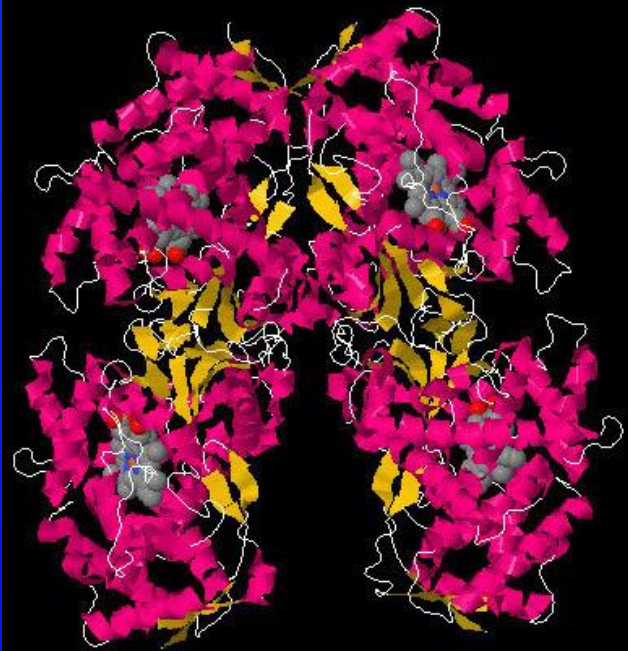
**"Cytochrome"** = colored cells  
They color the liver cells dark red as they contain iron

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

# STRUCTURE

They are heme-containing isoenzymes

Human Cytochrome P450  
from the  
Endoplasmic Reticulum



O<sub>2</sub>  
N<sub>3</sub>  
Cu  
Fe

# DISTRIBUTION

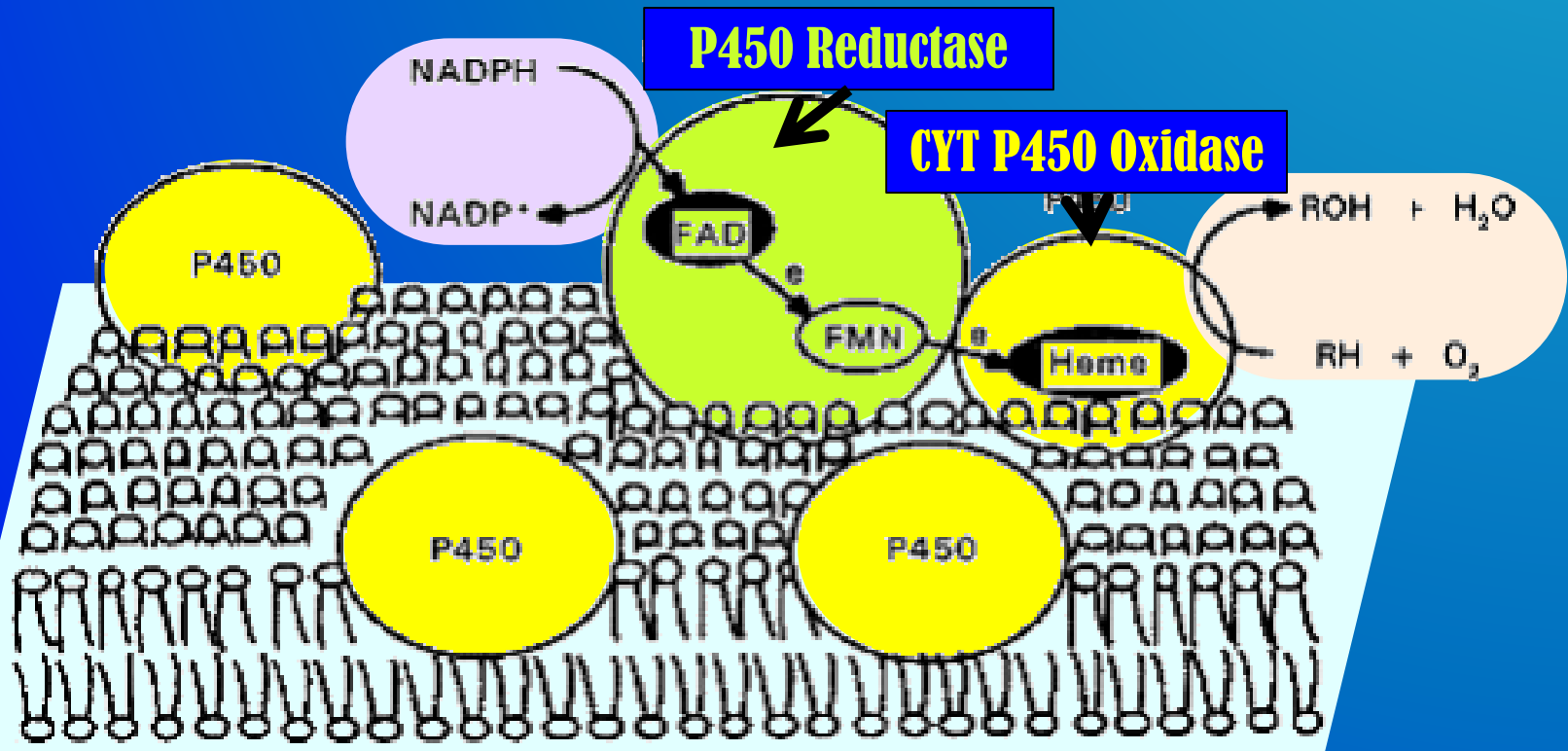
- Highly concentrated in hepatocytes
- Enterocytes of the small intestine present their principal extra-hepatic source
- Very small quantities in kidneys, lungs, & brain.

## Function

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

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

Substrates

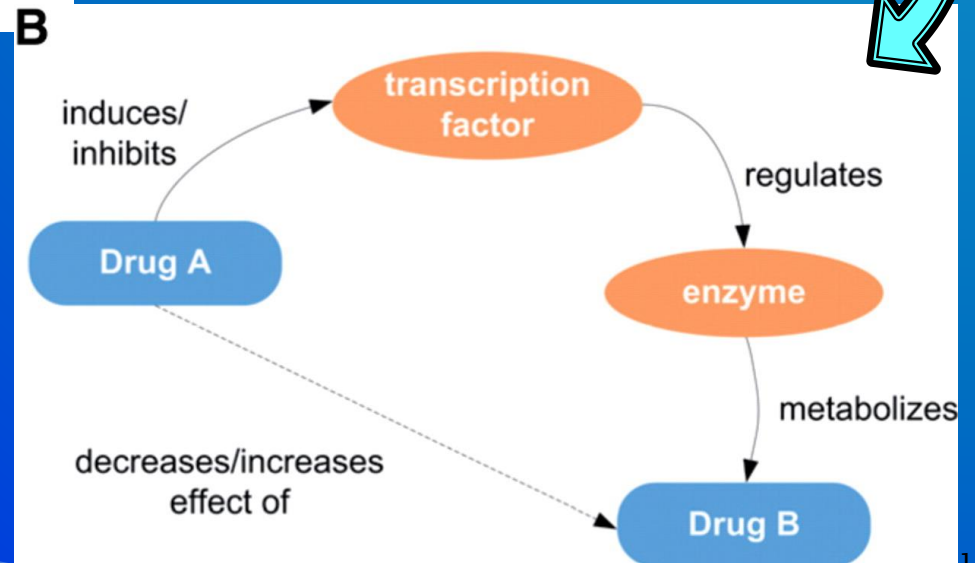
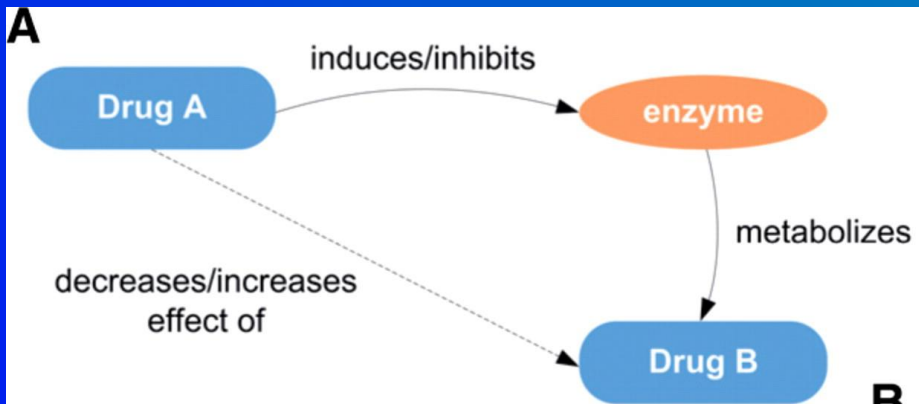


# Regulation

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 inhibition of the responsible transcription factors**



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

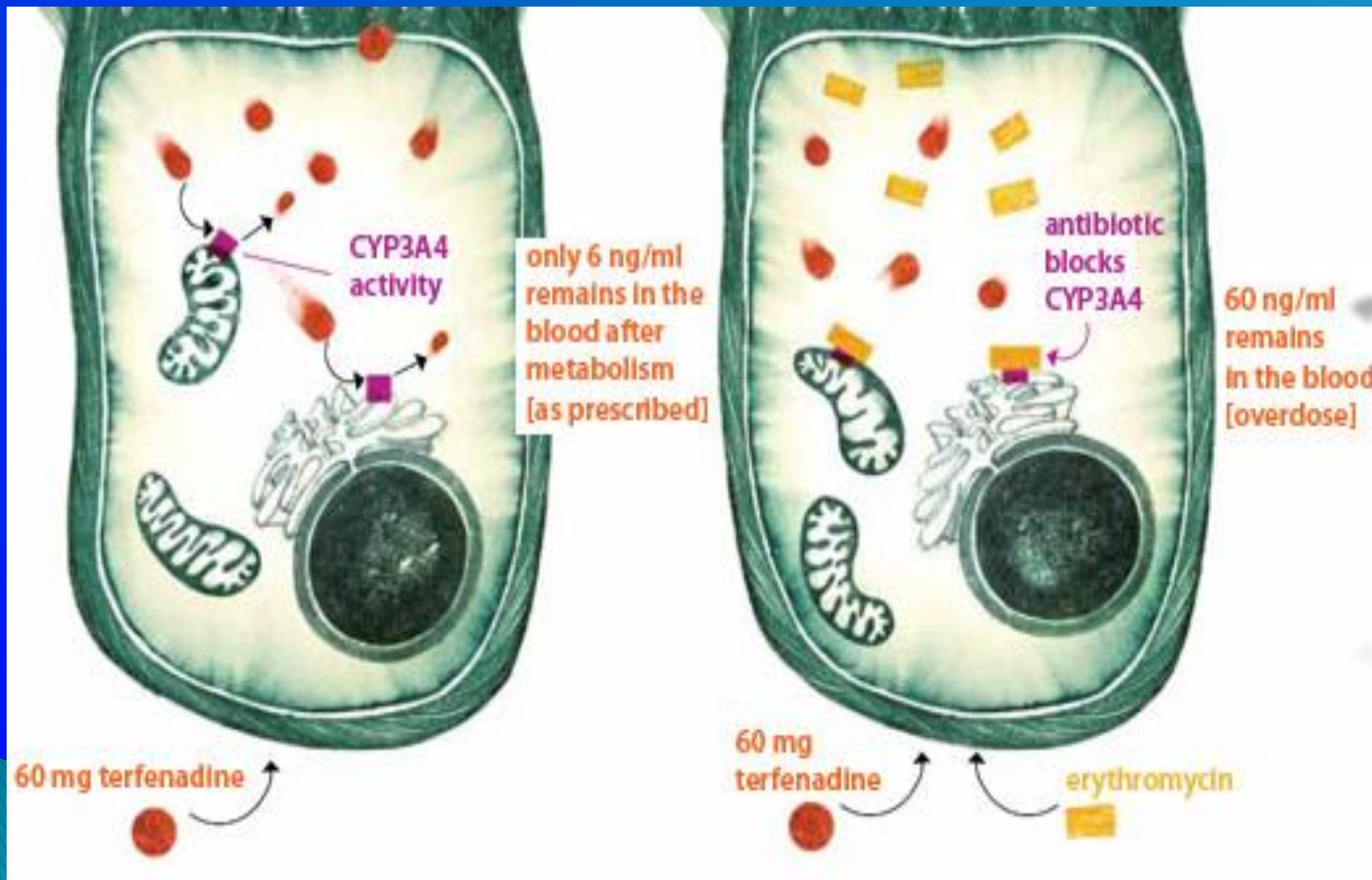
# Regulation

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 multivitamins & his lab results last week, proved that he has become diabetic, for which he was prescribed metformin. 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 musculoskeletal toxicity) & was verified by the lab finding of severe elevation in creatinine phosphokinase. “***

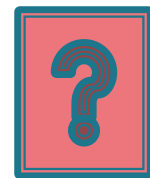
**Which one of the following drug-drug interaction on CYT 3A4 is the likely cause of his current state?**

Metformin + Atrovastatin

Atrovastatin + Fluconazole

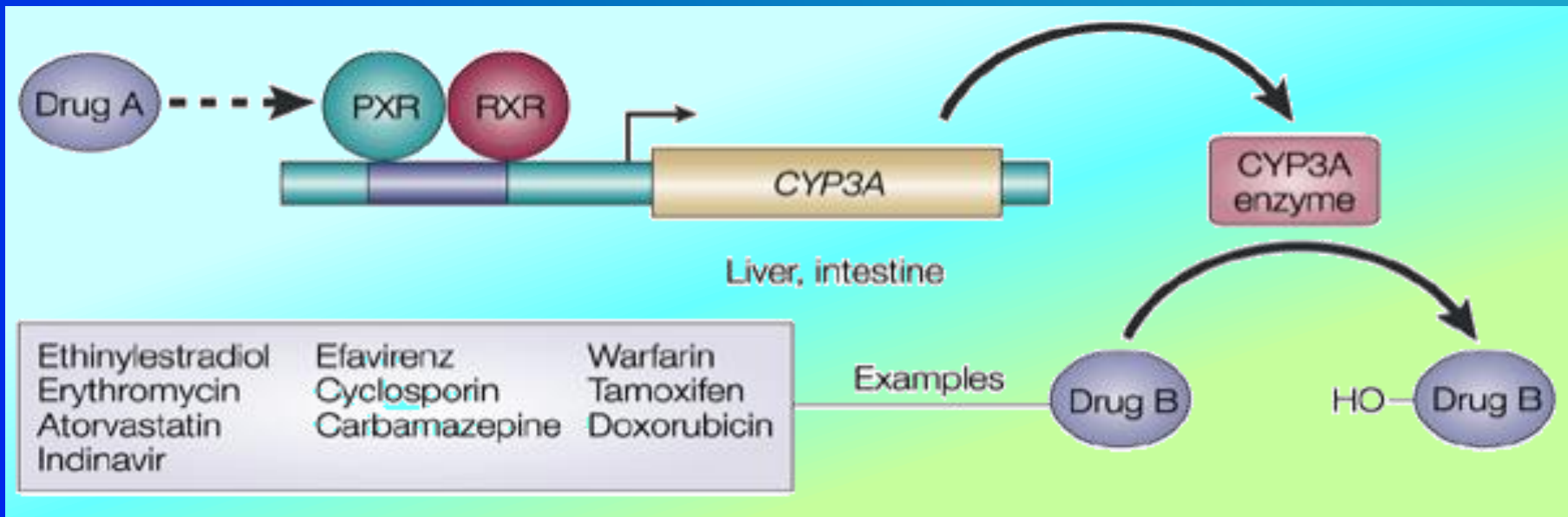
Metformin + Fluconazole

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 heterodimer **PXR / RXR** will induce **EXPRESSION** of *CYP P450* isoenzymes to → ↑ metabolism of Drug B

If Drug A is an **INHIBITOR**, its binding will prevent activation → **REPRESSION** of *CYP P450* isoenzymes to → ↓ metabolism of Drug B

*PXR, pregnane X receptor*  
*RXR, retinoid X receptor.* 15

## 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

↓ EFFICACY

### IN RELATION TO ENZ INHIBITORS

↓ / Retard metabolism & excretion of inhibitor & co-administered drugs

↑ / prolong action of the inhibitor & co-administered drugs.

↑ TOXICITY



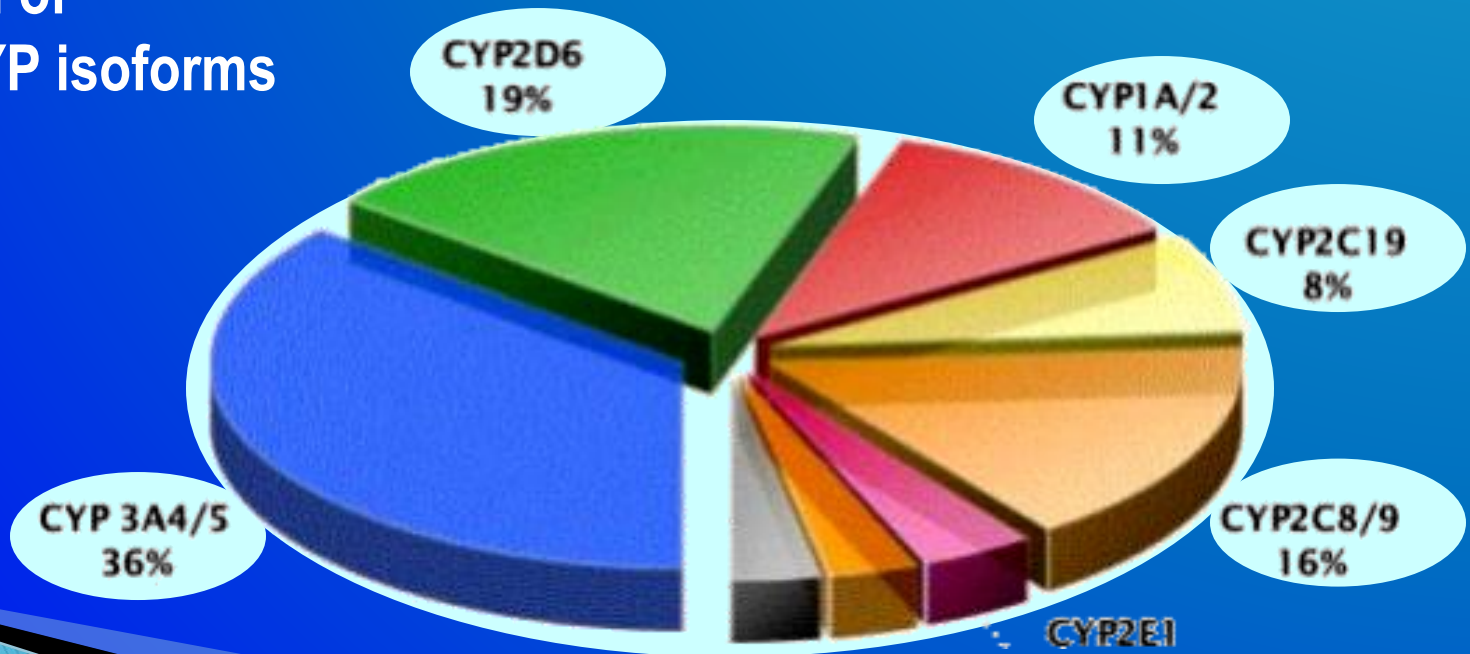
# Classification

- CYT P450 has been classified into
- Families designated by Numbers
  - Sub families designated by Letters

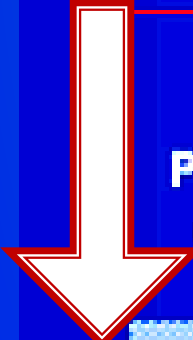
## Cytochrome P450 Isoforms

- CYP1A2
- CYP3A
- CYP2C9
- CYP2C19
- CYP2D6

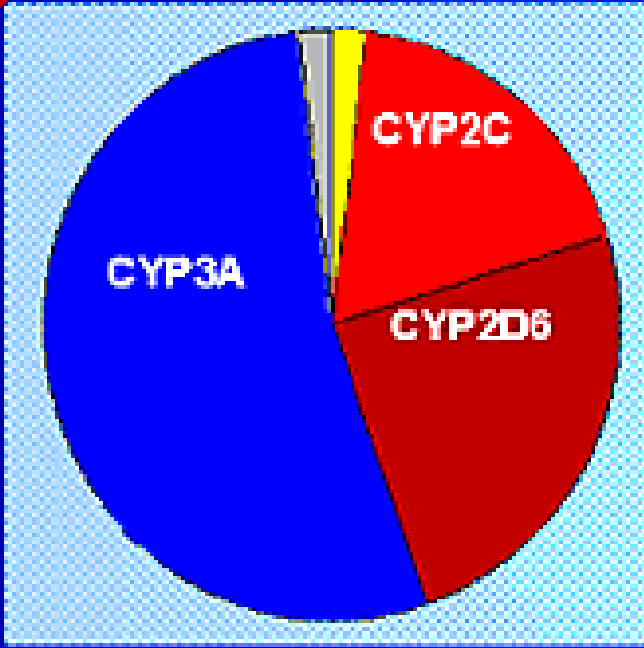
Distribution of different CYP isoforms in the liver.



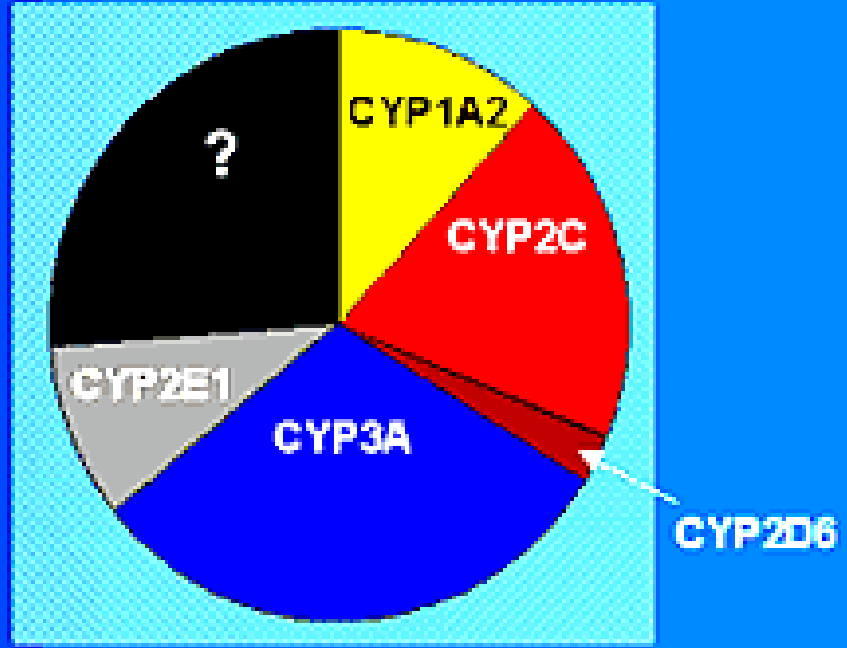
# CYP450 → Major Contributor to Phase I Metabolism



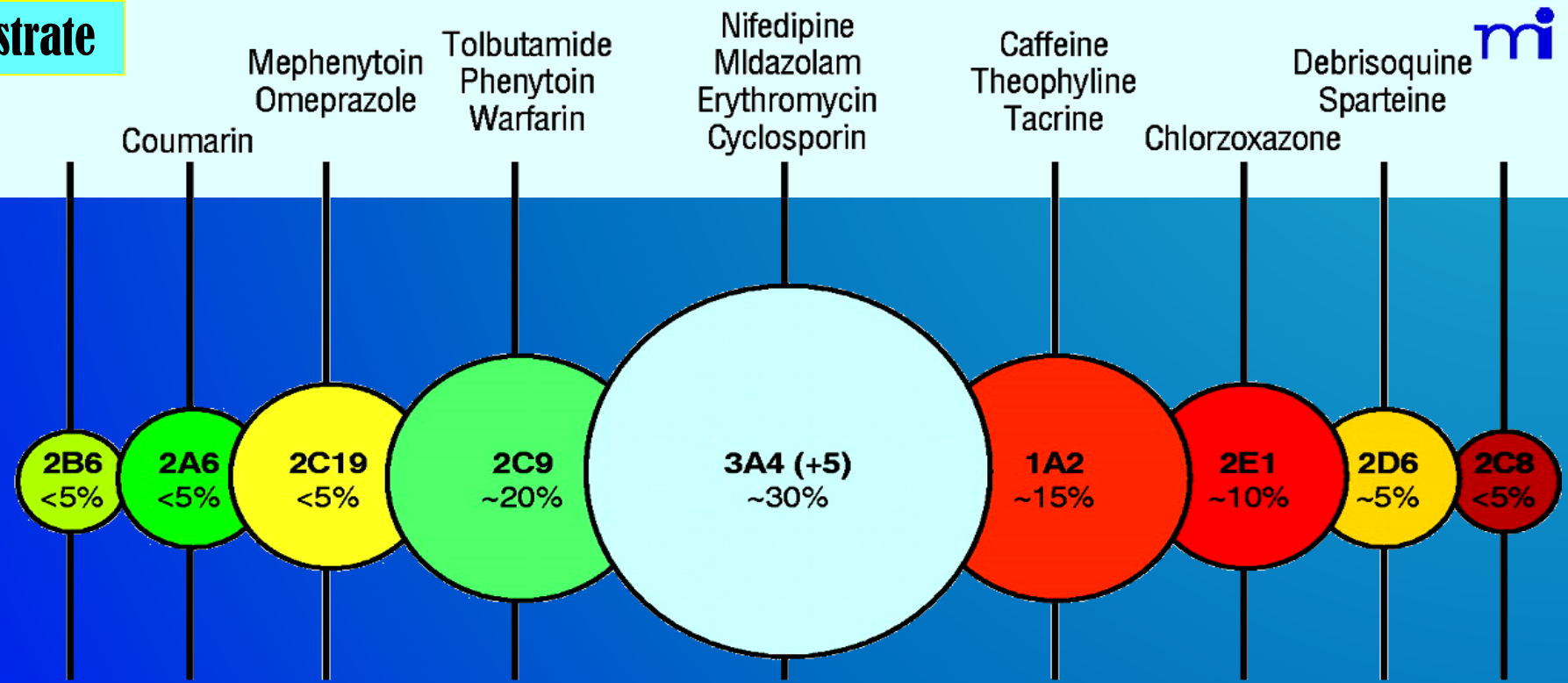
Relative Importance of P450s in Drug Metabolism



Relative Quantities of P450s in Liver



# Substrate



# Inhibitors

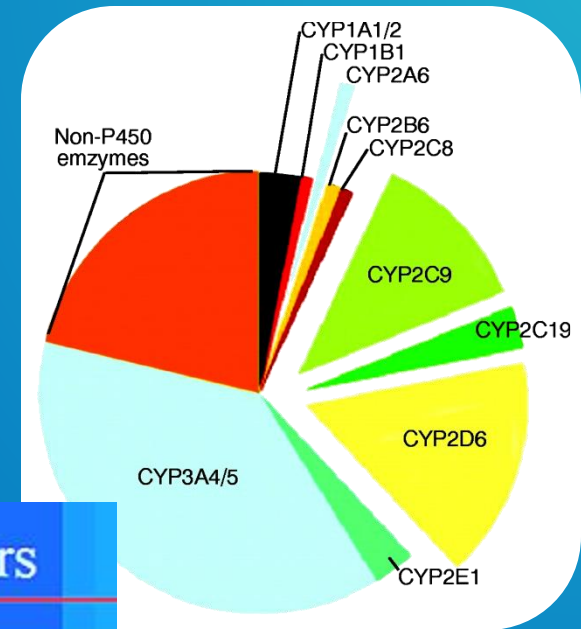
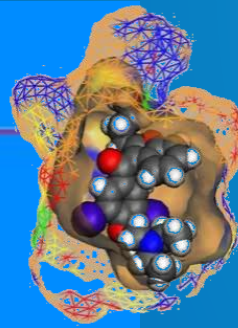
Fluconazole	Ketoconazole	Furafylline	Disulfiram
Methoxsalen	Gestodene	Fluvoxamine	Quinidine
Sulfaphenazole			

# Inducers

Barbiturates	Barbiturates	Omeprazole	Ethanol
Rifampicin	Rifampicin	Tobacco smoke	Isoniazid
Dexamethasone	Carbamazepine		

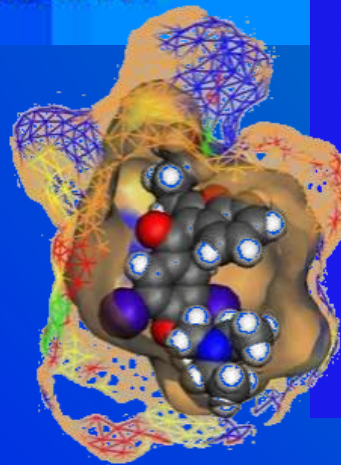
# Cytochrome P450 3A

- 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

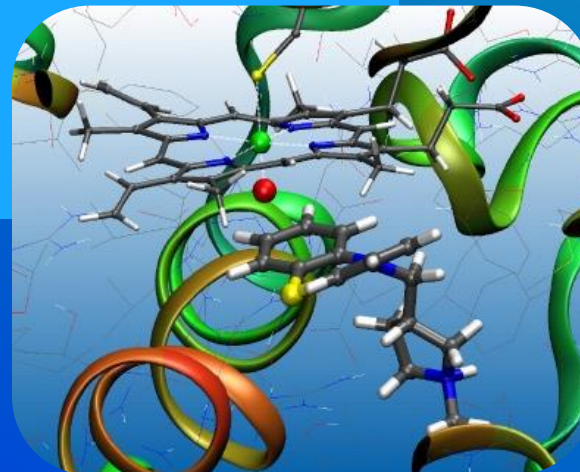
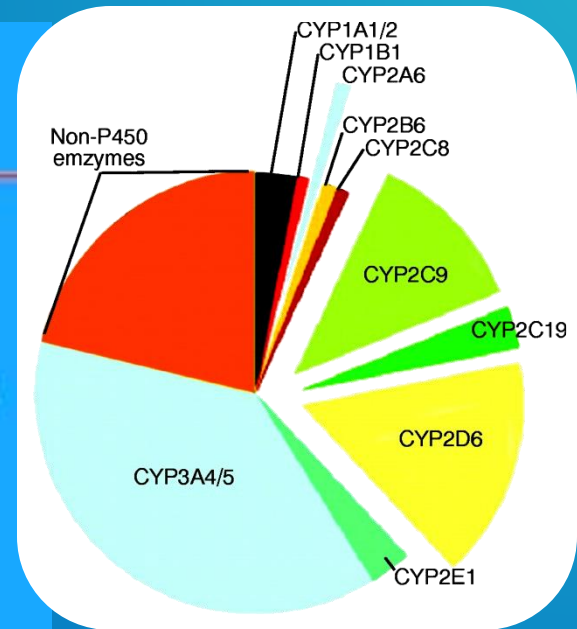


## CYP3A Inducers

- Carbamazepine
- Rifampin
- Rifabutin
- Barbiturates

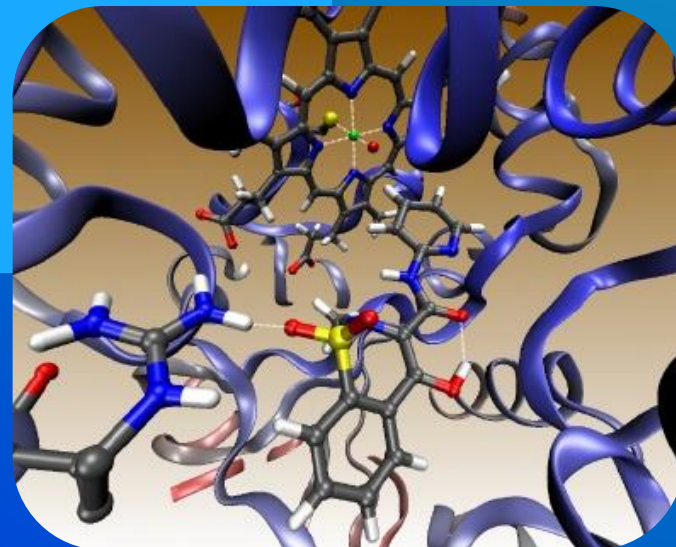
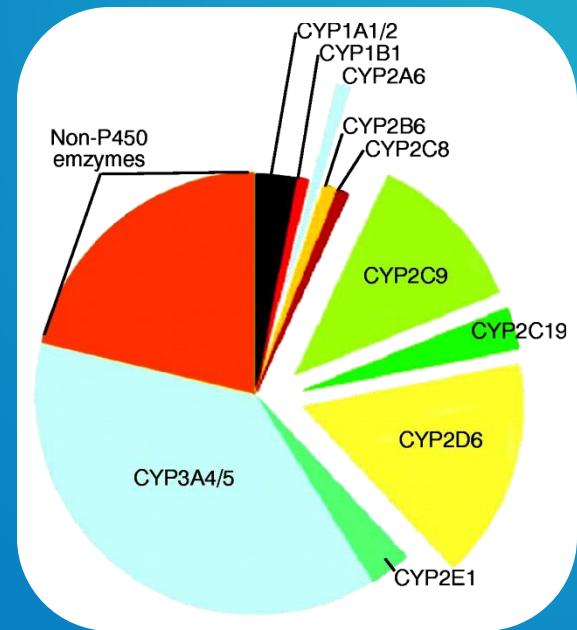
# Cytochrome P450 2D6

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



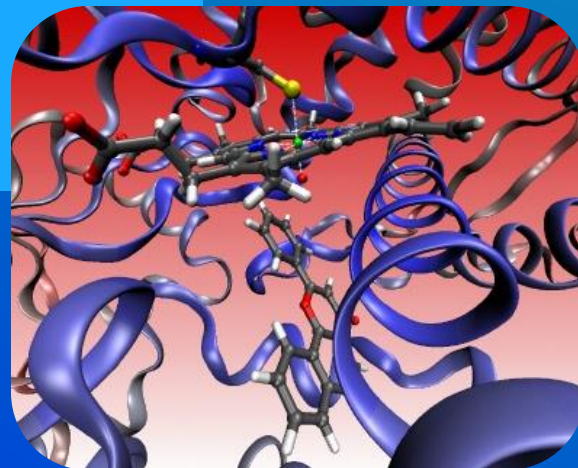
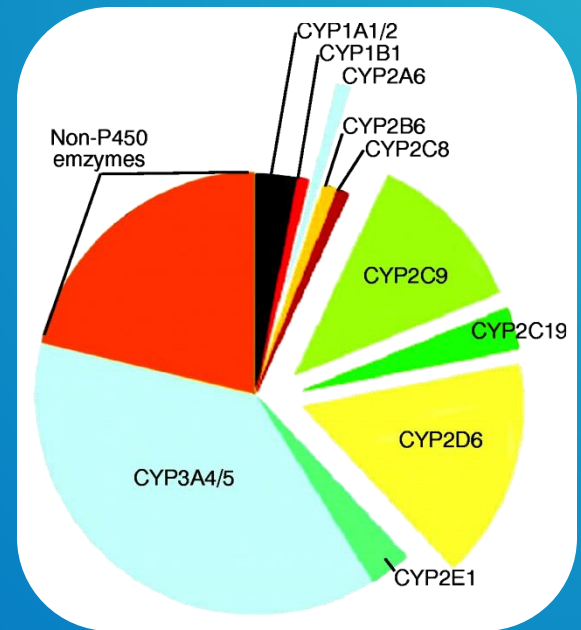
# Cytochrome P450 2C9

- 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



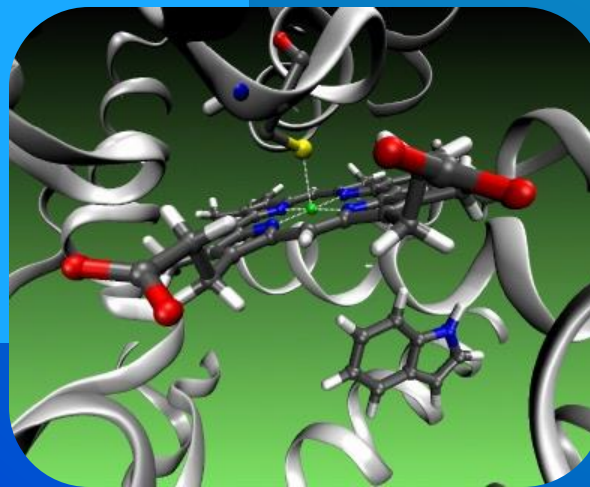
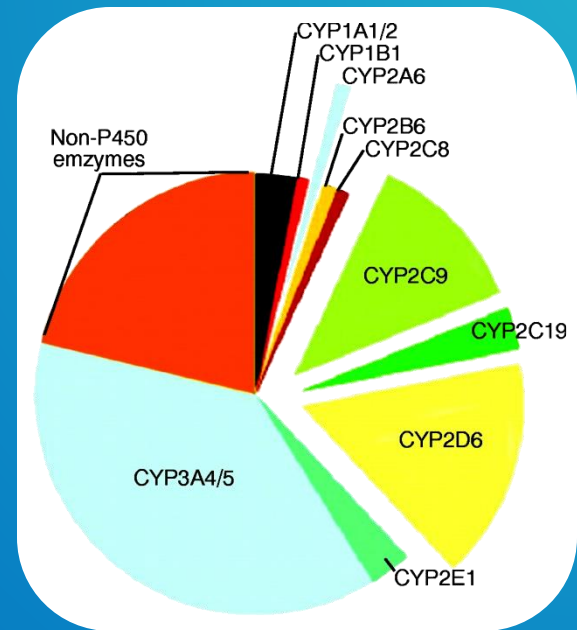
# Cytochrome P450 1A2

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



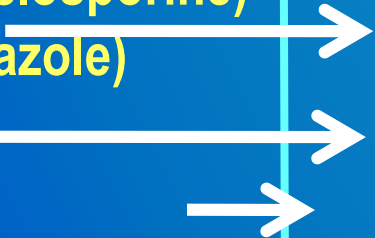
# Cytochrome P450 2C19

- 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	Inhibitors	Inducers
<ul style="list-style-type: none"><li>• Immunosuppressants (<b>Cyclosporine</b>)</li><li>• Azole Antifungals (<b>Fluconazole</b>)</li><li>• Antibiotics (<b>Erythromycin, Clarithromycin</b>)</li><li>• Ca channel blockers (<b>Amlodipine, Verapamil</b>)</li><li>• Statins (<b>Atorvastatin</b>)</li><li>• Cancer Chemotherapy (<b>Cyclophosphamide, Tamoxifen</b>)</li><li>• Non-Sedating Antihistamines (<b>Astemizole</b>)</li><li>• Benzodiazepines (<b>Midazolam, Clonazepam</b>).</li></ul>	 <ul style="list-style-type: none"><li>Protease inhibitors (<b>Ritonavir</b>)</li><li><b>Cimetidine</b></li><li><b>Chloramphenicol</b></li><li><b>Nefazadone</b></li><li><b>Grape Fruits</b></li></ul>	<ul style="list-style-type: none"><li><b>Phenytoin</b></li><li><b>Carbamazepine</b></li><li><b>Barbiturates</b></li><li><b>Rifampicin</b></li><li><b>Dexamethazone</b></li><li><b>Progestins</b></li></ul>

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

*He receives daily multivitamins & his lab results last week, proved that he has become diabetic, for which he was prescribed metformin. 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 musculoskeletal 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

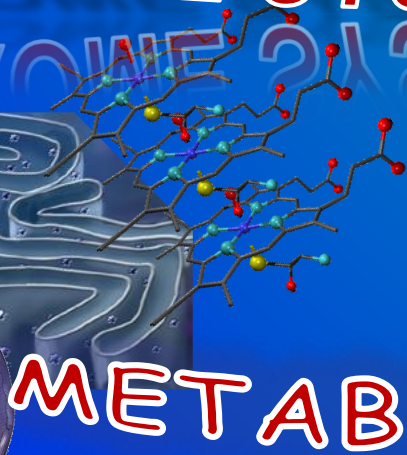
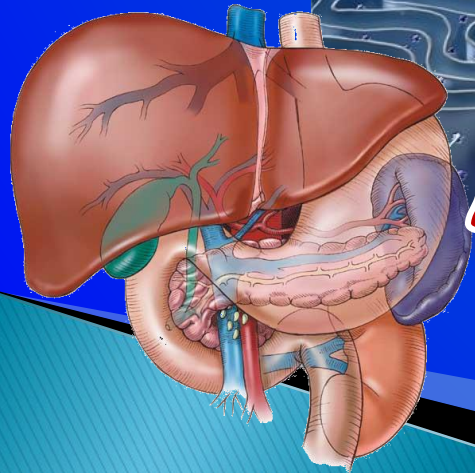
## CYP2C19

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

# CYTOCHROME SYSTEM



&

# DRUG METABOLISM

G

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