

Drugs for hyperlipidemia



ILOs

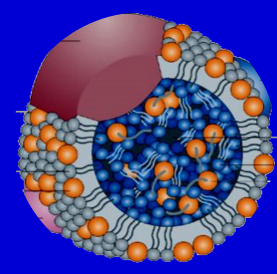
By the end of those 2 lectures the student will be able to:

- Define hyperlipidemia vs normal lipid levels
- Discuss the non-pharmacological treatment of hyperlipidemia
- Classify lipid lowering agents targeting exogenous & endogenous pathways
- Expand on the pharmacology of drugs related to each group
- Hint on adjuvant drugs that can help in lipid lowering

Hyperlipidemia

- Hyperlipidemia is a major cause of atherosclerosis which may lead to CAD and ischemic cerebrovascular disease
- Denotes abnormally

 Lipoproteins [LP] in blood
- **▶ Lipids** originate from two sources:
 - endogenous lipids, synthesized in the liver
 - exogenous lipids, ingested and processed in the intestine
- The principle lipids in the blood are:
 - Cholesterol (C) Triglycerides (TG)
 - Phospholipids (PL)- Cholesterol esters (CE)
 - Non-estrified fatty acids (NEFA)



Familial Hyperlipoproteinemia

LProteinemia	↑LP	↑ Lipids	Risk
Туре I	CM	TGs	-
Туре Па	LDL	С	***
Type IIb	VLDL & LDL	TG & C	^
Type III	IDL	TGs & C	↑
Type IV	VLDL	TGs	↑
Type V	VLDL & CM	TGs & C	_

Therapeutic strategies for treatment of hyperlipidemia

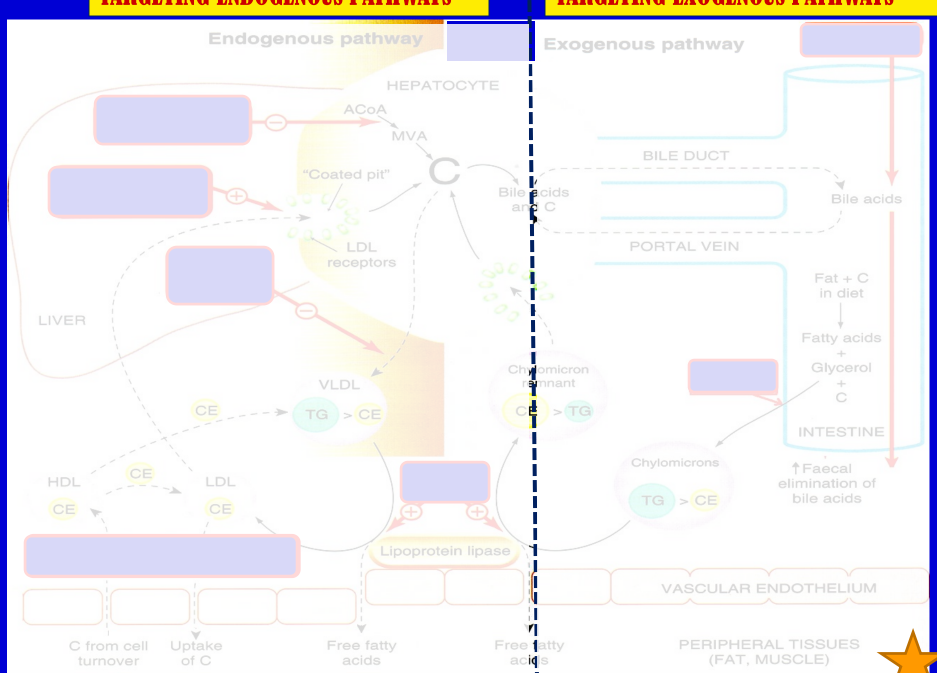
Therapeutic lifestyle changes

- 1. Healthy diet; optimal Quantitative & Qualitative faticontentic agents
- Diet has <30% of calories as fat, <7% as saturated fat and <200mg cholesterol/day
- Avoid trans-fatty acids & acute increase in C intake
- Use vegetable oils rich in unsaturated fatty acids: oleic acid,
- linoleic acid & linolenic acids. Diet should also contain plant
- stanols (interfere with the formation of micellar cholesterol)
- & soluble fibers
- Eat food high in antioxidants vitamins
- 2. Regular exercise
- 3. Cessation of hazardous habits; smoking, alcohol,
- 4. Losing weight
- Can achieve a fall in LDL-C of 8-15% ... but long-term compliance is a problem



TARGETING ENDOGENOUS PATHWAYS

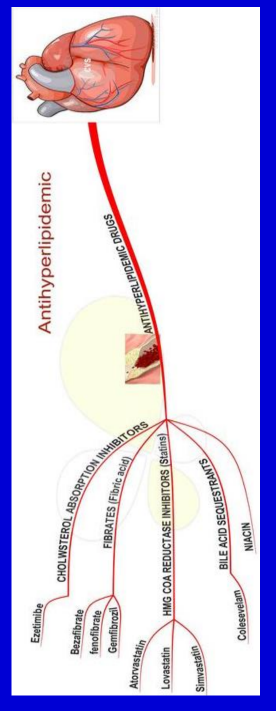
TARGETING EXOGENOUS PATHWAYS



Antihyperlipidemic agents

- A-According to the mechanism of action:
- 1- Inhibits cholesterol absorption in the intestine Ezetimibe
- 2-Sequester bile acids in the intestine Exchange resins
- 3-Inhibits synthesis of cholesterol
 Inhibitors of hydroxymethylglutaryl coenzyme
 A reductase (Statins)
- 4-Alter relative levels & patterns of different plasma LPs

Fibrates, Nicotinic acids



B-According to site of action

I-Agents targeting exogenous cholesterol

- Ezetimibe
- Colestipol & cholestyramine

II-Agents targeting endogenous cholesterol

- Statins
- Fibrates
- Nicotinic acid

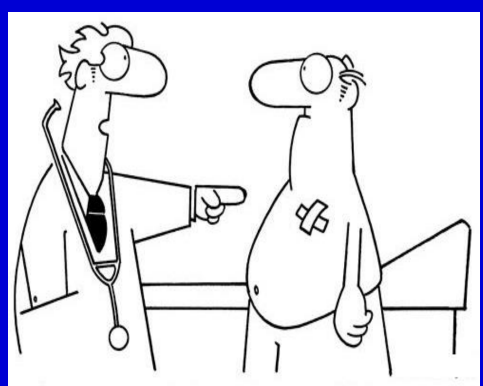
III-Adjuvant agents

Omega-3-Fatty Acids, Stanols



Cholesterol Absorption Inhibitors

Ezetimibe



"Whenever your cholesterol gets too high, a sensor will send out a signal that automatically locks the kitchen door and turns on your treadmill."

Mechanism of action of Ezetimibe

Blocks C transporter located on brush border of small intestine → → pool of C available to the liver → upregulate LDL receptor, trapping more LDL particles from blood.

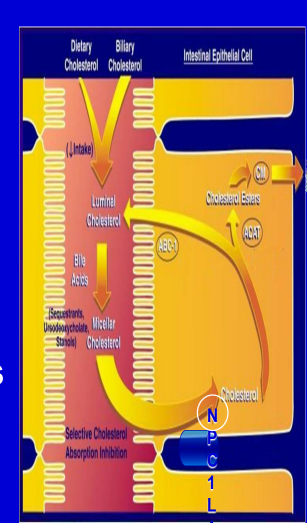
Pharmacological action

→LDL 20% → TG 8%, **→ HDL 1-4%** No effect on steroids, lipid-soluble vitamins,

bile acids.

Pharmacokinetics

- Absorbed & conjugated in intestine to
- active glucuronide
- Reaches peak blood level in 12–14 hours
- Undergoes enterohepatic circulation
- Its half-life is 22 hours
- Most of the drug is excreted in feces



Indications

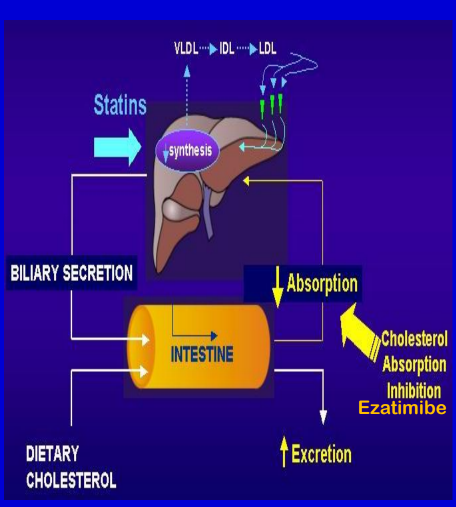
As Monotherapy;
Primary prevention of low risk of CHD which needs modest ↓ LDL

As Combination Therapy; safe

- -With statins; synergistic in moderate/severe ↑ LDL
- -Or with other lipid lowering drugs; as fibrates

ADRs

Not common GIT disturbance, headache, fatigue, artheralgia & myalgia



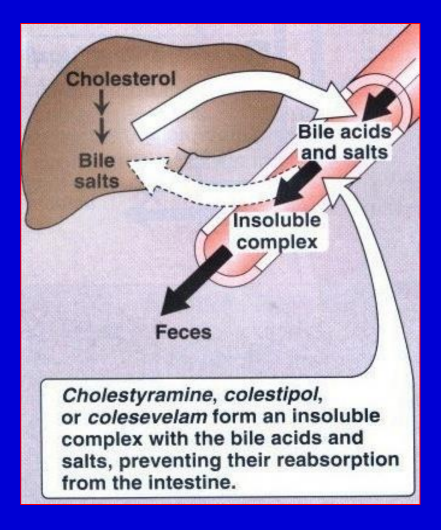
Exchange resins Bile acid sequestrants

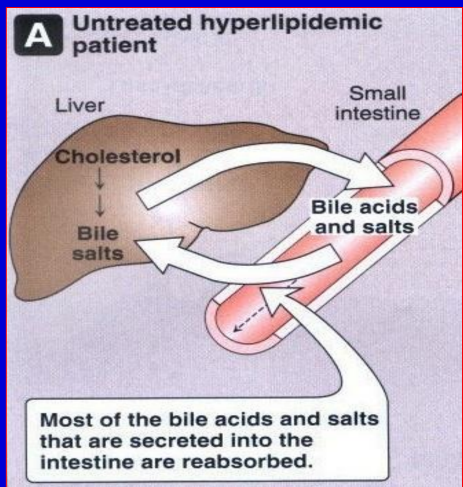
Cholestyramine & Colestipol Colesevelam



"If I'm digging my grave with a fork and spoon, wouldn't that burn a lot of calories?"

Resins: Mechanism of Action





Bile Acid-Binding Resins

- Moderately effective with excellent safety record
- Large MW polymers which bind to bile acids and the acid-resin complex is excreted so their fecal excretion
 10 folds
 - prevents enterohepatic cycling of bile acids
 - obligates the liver to synthesize replacement bile acids from cholesterol
- The liver increases the number of LDL receptors to obtain more cholesterol
- The levels of LDL-C in the serum are reduced as more cholesterol is delivered to the liver
- Excellent choice for people that cannot tolerate other types of drugs

Resins: Adverse Effects

- Resins are clinically safe as they are not systemically absorbed
- GIT upset: abdominal discomfort, bloating, constipation
- Decreased absorption of: fat soluble vitamins (Vitamin A, D, K)
- The concentration of HDL-C is unchanged

Resins: Drugs interactions Interfere with the absorption of:

- Statins, Ezetimibe
- Chlorothiazides, Digoxin, Warfarin
- N.B. wait 1 hour before or 4 hrs after administration of resins
- Colesevelam has not been shown to interfere with the absorption of co-administered medications and is a better choice for patients on multiple drug regimens

Contraindications of resins

- 1- Complete biliary obstruction (because bile is not secreted into the intestine)
- 2- Chronic constipation
- 3-Severe hypertriglyceridemia (TG >400 mg/dL) ??

HMG-Co A Reductase Inhibitors

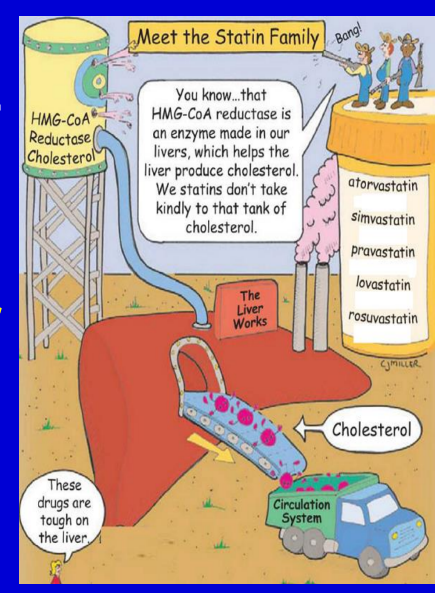
Statins



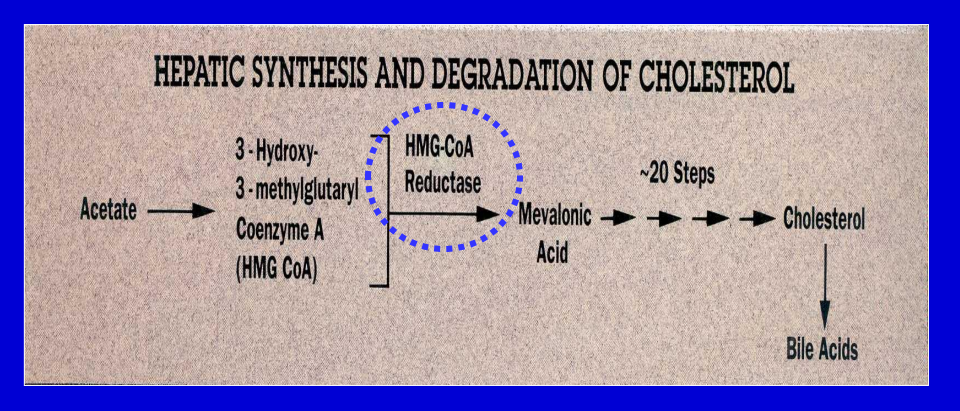
"Listen, when the side effects of this medication kick in, you'll forget what was wrong in the first place!"

HMG-Co A Reductase Inhibitors

- Hydroxy MethylGlutaryl-Coenzyme A reductase inhibitors or statins are the most effective and best-tolerated agents for treating hyperlipidemia
- Statins are considered as first-line drugs when LDLlowering drugs are indicated



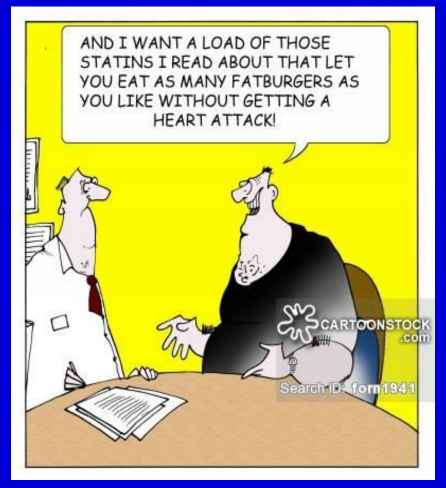
Statins: Mechanism of Action



Statins are potent competitive inhibitors of 3-hydroxy-3-methyl glutaryl coenzyme A (HMG-CoA) reductase, which catalyzes an early, ratelimiting step in do-novo hepatic C synthesis. Thus, HMG-Co A is not converted to mevalonic acid

Statins: Preparations

- Rosuvastatin
- Atorvastatin
- Simvastatin
- Pravastatin
- Lovastatin



 Used alone or with other anti-hyperlipidemic drugs (ezetimibe) for treatment of drugresistant dyslipidaemia

PLEIOTROPIC ANTIATHEROGENIC effects [> in Vessels]

- Improve endothelial function
- ◆ ▼ vascular inflammation
- Stabilization of atherosclerotic
- Plaque
- ◆ ◆ platelet aggregability
- Antithrombotic actions
- Enhanced fibrinolysis ...etc



"Your good cholesterol is fine, but your bad cholesterol is plotting to hack into your computer, empty your bank account and steal your wife."

Statins: Pharmacokinetics

- Most statins have a high first-pass clearance by the liver
- Greater than 95% of most of these drugs are bound to plasma proteins with short half-life
- Drug-drug interactions involve specific interactions with the cytochrome P-450 drug metabolizing system, especially CYP3A4
- All statins are taken orally at bedtime because of hepatic C synthesis is maximal between midnight and 2:00 a.m., except atorvastatin taken at any time because of its long half-life (14 hours)

Indications

As monotherapy;

2nd ry Prevention; In all ischemic insults [stroke, AMI,etc.] So given from1st day of ischemic attack

Pry Prevention;

- 1. Patients with hyperlipidemia and with other risks for ischemic insults.
- 2. Type Ila Hyperlipoprotinemia.

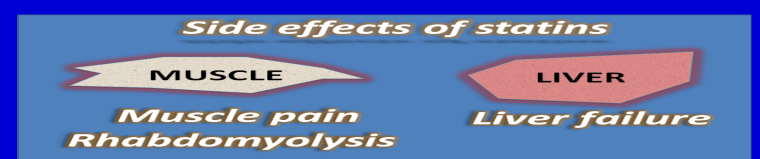
If no control → combine (sequestrants / ezatimibe, niacin,...) to → C

As Combination therapy:

- 1. Mixed dyslipidaemias; added to fibrates or niacin if necessary
- 2. In diabetics and patients with insulin resistance [metabolic syndrome] because these patients will possess small dense LDL (severely atherogenic) + evident endothelial dysfunction + increased thrombotic profile

Statins: Adverse Effects

- Common side effects: Headache, myalgia, fatigue, Gl intolerance, and flu-like symptoms
- Hepatotoxicity, raised concentrations of liver enzymes (serum aminotransferases)
- Myopathy (increased creatine kinase [CK] released from muscles)
- Teratogenicity, statins should be avoided during pregnancy



Statins: Drug Interactions

- Statins potentiate the action of oral anticoagulant and anti-diabetic drugs (by displacement from plasma protein binding sites)
- Drugs that increase the risk of statin-induced myopathy include:
- Other antihyperlipidemics (fibrates)
- Drugs metabolized by 3A4 isoform of cytochrome P450: erythromycin, verapamil, cyclosporin, ketoconazole
- Pravastatin and fluvastatin are the statins of choice in patients taking other drugs metabolized by cytochrome 3A4 system.

Statin-induced myopathy

- Muscle aches, soreness, or weakness associated with an elevation of creatine kinase (CK), are the best indicator of statin-induced myopathy.
- Failure to recognize myopathy and to discontinue drug therapy can lead to rhabdomyolysis, myoglobinuria, and acute renal necrosis.
- ♣ serum transaminase → can progress to evident hepatotoxicity
 So lab investigations recommended every 6 month → if levels ↑ up
 to 3 folds at any time, statin must be stopped then dose adjusted.
- ↑ creatine kinase activity (index of muscle injury) →
 Measured only if myalgia or myositis develops → if ↑ 3-5 folds → we
 ↓ statin doses / omit combination with fibrates.....

Niacin (Nicotinic Acid)



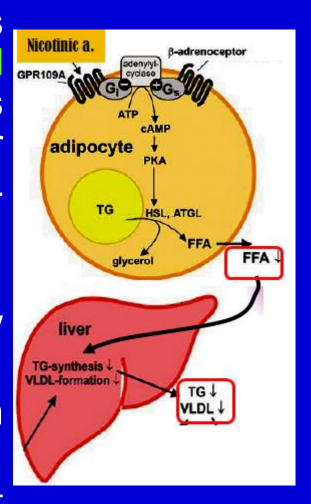
Niacin (Nicotinic Acid)

Water soluble B-complex vitamin with multiple actions

- Niacin is the most effective medication for increasing HDL cholesterol levels and it has positive effects on the complete lipid profile
- It is useful for patients with mixed dyslipidemias
- Niacin exerts greatest beneficial effects on wide range of lipoprotein abnormalities

Mechanism of action:

- 1. In adipose tissue: it binds to adipocytes nicotinic acid receptors, this will lead to decrease in free fatty acids mobilization from adipocytes to the liver resulting in ↓ TG and thus VLDL synthesis
- 2. In liver: niacin inhibits hepatocyte
 2-diacylglycerol acyltransferase, a key
 enzyme for TG synthesis
- Thus, it decreases VLDL production (decreased TG synthesis and estrification)
- 3. In plasma: it increases LPL activity that increases clearance of VLDL & chylomicron



Pharmacological actions

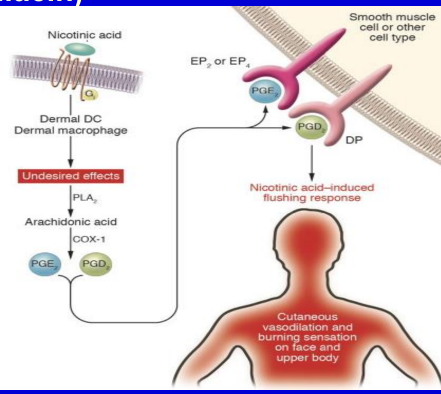
- Effect on VLDL:

 VLDL by:
- 1) **\(\sqrt** synthesis in liver
- 2) increased clearance in plasma
- Effect on LDL:
 ↓ LDL due to reduction in its precursor (VLDL)
- Effect on HDL: Induces remarkable increase in HDL-C (The catabolism of HDL can be inhibited by nicotinic acid through a mechanism that is largely unknown)
- Niacin also promotes hepatic apoA-I production and slows hepatic clearance of apoA-I and HDL

Niacin: Adverse Effects

 The most common side effect is cutaneous flushing, (which is prostaglandin -mediated, can be avoided by low dose aspirin ½ h before niacin)

- GIT disturbances: Dyspepsia reactivation of peptic ulcer (ca after meal)
- High doses:
- Reversible ↑ liver enzymes →
- Impairment of glucose tolerar
- uric acid



Indications

Monotherapy or in combination with fibrate, resin or statin

Type IIa hypercholestrolemia
Type IIa, IIb hypercholesterolemia & any combined hyperlipidemia

Patient with hypertriglyceridemia & low HDL-C

Contraindications

- Gout
- Peptic ulcer Hepatotoxicity Diabetes mellitus



Fibric acid Derivatives (Fibrates)

Fibrates: Mechanism of Action

- Fibrates are agonists of peroxisome proliferator activated receptors (PPARα) which are a class of intracellular receptors that modulate fat metabolism
- They increase genes transcription for lipoprotein lipase (LPL) leading to increased catabolism of TG in VLDL and chylomicrons
- Examples: Clofibrate & Gemfibrozil & Fenofibrate

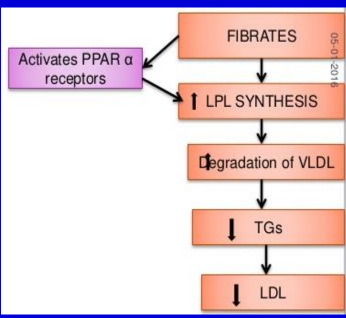
Fibrates: pharmacological effects

 — ► LPL activity, which increases clearance of VLDL &

chylomicron in plasma

 A marked reduction in TG (due to stimulation of catabolism of VLDL)

- FFA uptake by the liver
- + LDL-C uptake by the liver
- ★in HDL-C (by increasing the
- production of the apoprotein components of HDL)
- reparting the excretion of hepatic C in bile, thus endogenous hepatic C synthesis may be decreased



Fibrates: Adverse Effects

- GIT (indigestion, abdominal pain, diarrhea)
- Myositis: can occur resulting in weakness and tenderness of muscles, use of fibrates with statins is generally inadvisable
- Gallstones: Clofibrate increases C content of bile, predisposes to gallstones, and its use is therefore limited to patients who have cholecystectomy

Indication of Fibrates

1st-line defense for:

- *mixed dyslipidemia (i.e. raised serum TG and C)
- * Patients with low HDL and high risk of atheromatous disease (often type 2 diabetic patients)
- * Patients with severe treatment- resistant dyslipidemia (combination with other lipid-lowering drugs).

ADRs

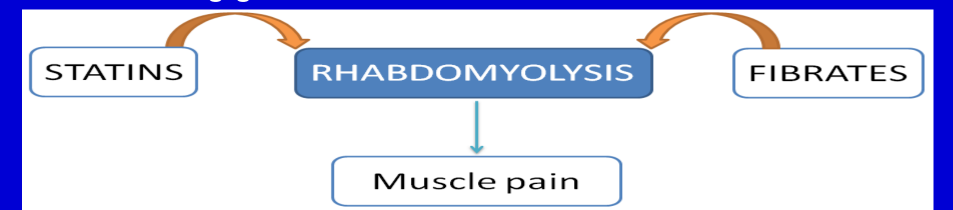
- 1. G.I.T upset, headache, fatigue, weight gain
- 2. Rash, urticaria, hair loss
- 3. Myalagia, Myositis, Rhabdomyolysis → Acute renal failure → Occurs >
 - -In alcoholics,
- -If combined with statins (each -ve metabolism of other)
 - -Or In impaired renal function
- 4. fibrates should be used with caution in patients with biliary tract disease, as they increase the risk of cholesterol gallstones as a result of an increase in the cholesterol content of bile.

Drug interactions

- Increased risk of myopathy when combined with statins.
- Displace drugs from plasma proteins (e.g.oral anticoagulants and oral hypoglycemic drugs)

Contraindications

- Patients with impaired renal functions
- Pregnant or nursing women
- Preexisting gall bladder disease



Adjuvants in hyperlipidemia

Omega -3-FA

β-Sitosterol



Omega -3-FA found in fish oils containing highly unsaturated FA

Mechanism

- enzymes involved in TG synthesis
- beta-oxidation of FFA
- ◆ platelet function
- Prolongation of bleeding time
- Anti-inflammatory effects

Pharmacological Effects

→ TGs

Some vascular protection

Indications Approved as adjunctive for treatment of very high TGs

β-Sitosterol

found in plants with structure similar to C

Mechanism & Pharmacological Effects

Compete with dietary & biliary C absorption → ↓ levels LDL levels ±10%

Indications Given as food supplement before meal in hypercholestrolemia

Eat Butter.

Scientists labeled fat the enemy. Why they were wrong

A meta-analysis of prospective epidemiologic studies showed that there is no significant evidence for concluding that dietary saturated fat is associated with an increased risk of CHD or CVD.