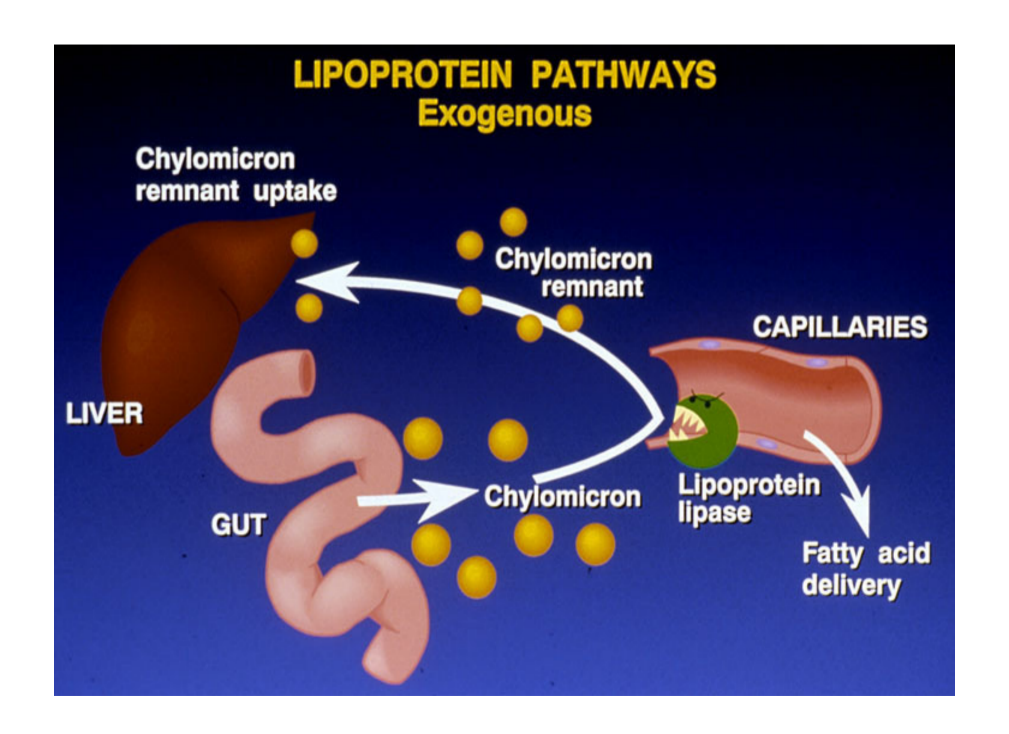
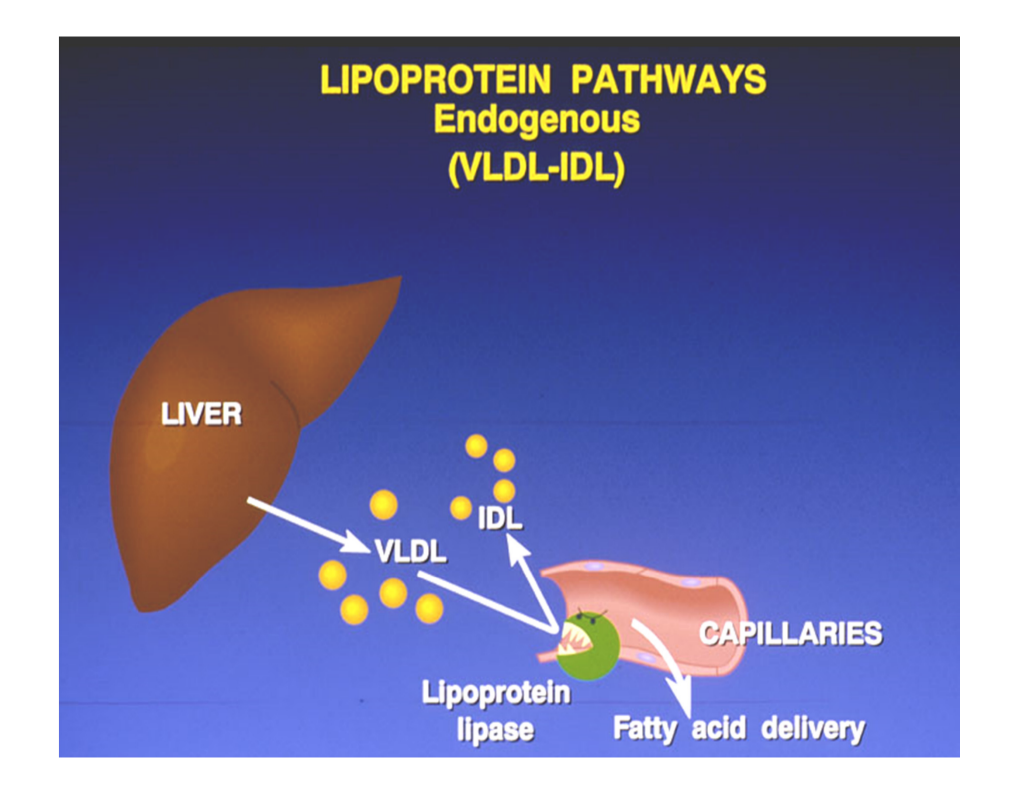
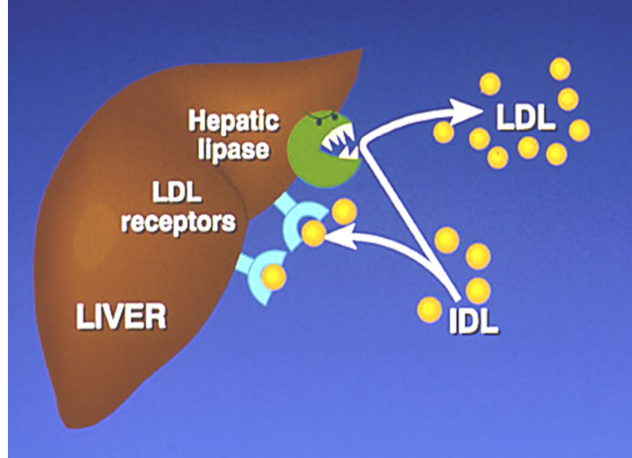
Dyslipidemia (Med-341)

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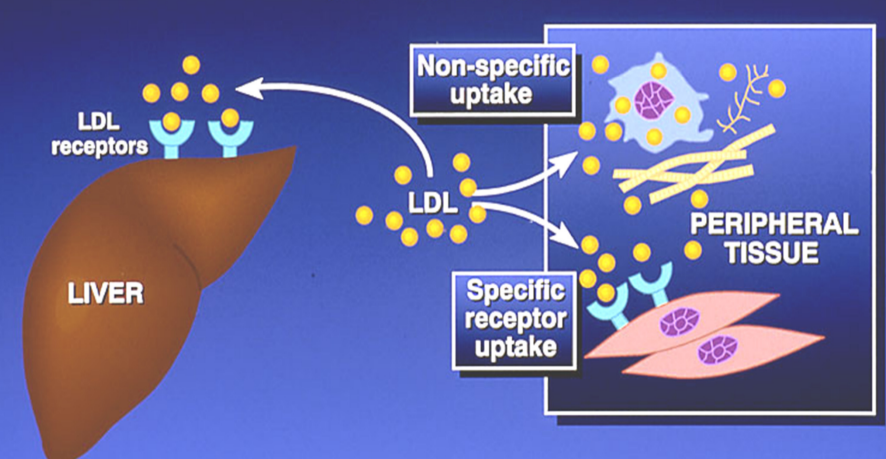


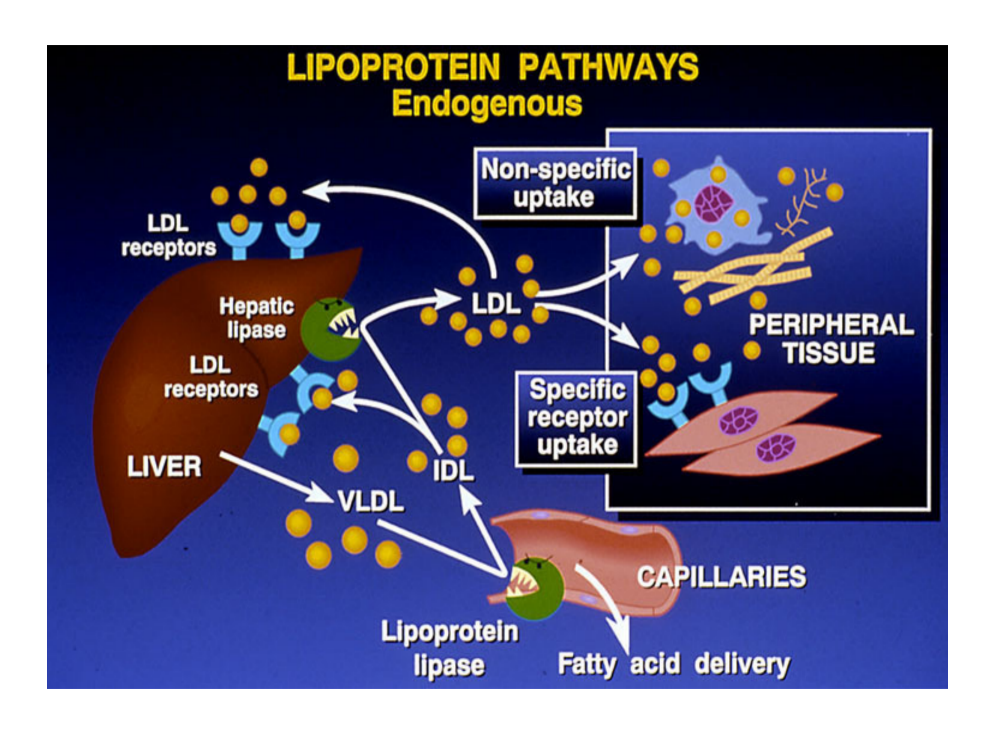


LIPOPROTEIN PATHWAYS Endogenous (IDL-LDL)









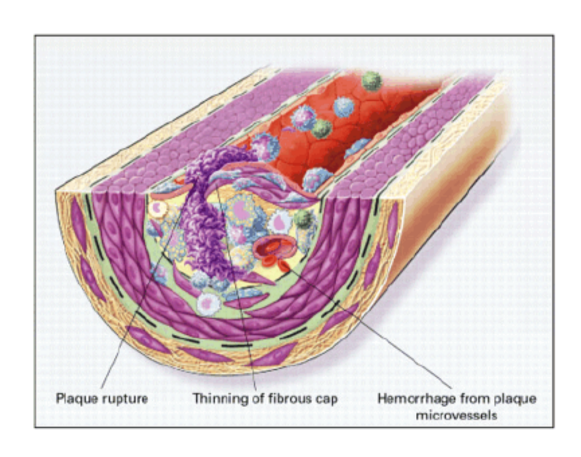
The story of lipids

- ☐ Chylomicrons transport fats from the intestinal mucosa to the liver
- ☐ In the liver, the chylomicrons release triglycerides and some cholesterol and become low-density lipoproteins (LDL).
- □LDL then carries fat and cholesterol to the body's cells. LDL receptors in Liver take the LDL to Liver.
- ☐ High-density lipoproteins (HDL) carry fat and cholesterol back to the liver for excretion.

The story of lipids (cont.)

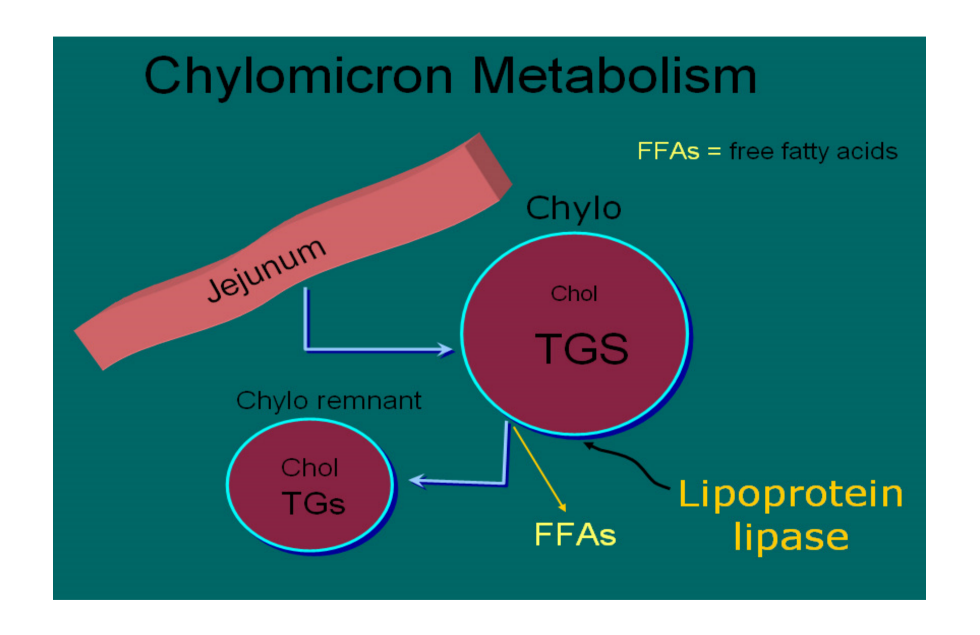
- ☐ When oxidized LDL cholesterol gets high, atheroma formation in the walls of arteries occurs, which causes atherosclerosis.
- ☐HDL cholesterol is able to go and remove cholesterol from the atheroma.
- □ Atherogenic cholesterol → LDL, VLDL, IDL

Atherosclerosis



Lipid Transport Blood Peripheral tissues Intestine ABCA1 • Free cholesteral Chylomicrons Fatty acids Fatty acids I-Aoqs ABCG1 Adipose Muscle tissue LPL/Apo C2 Cholesteryl Nascent LCAT Nucleus HDL Chylomicron remnants ::0 Hepatic lipase ABCA1 Liver Nascent Endothelial lipase HDL Mature HDL ABCA1 Macrophage PLTP CETP Triglyceride Cholesteryl ester LDLR VLDL and LDL

Rader DJ, Daugherty, A Nature 2008; 451:904-913

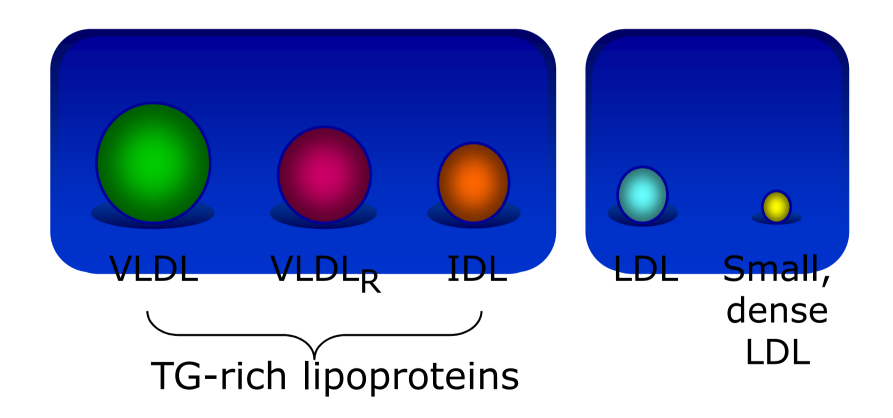


VLDL Metabolism

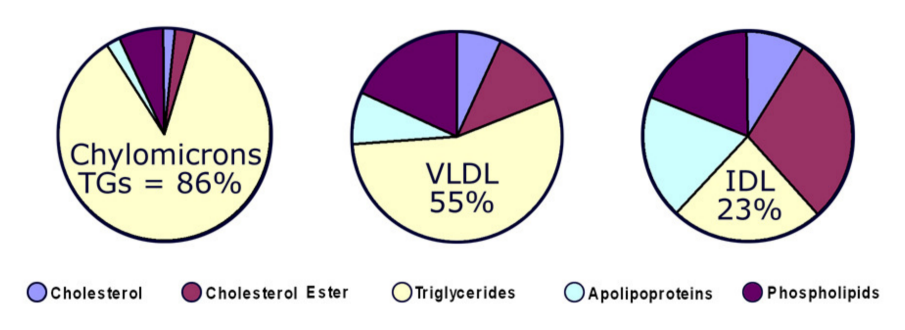
FFAs = free fatty acids **VLDL** Chol TGs LDL IDL Chol Chol TGs Lipoprotein TGs lipase Hepatic **FFAs** lipase **FFAs**

Atherogenic Particles

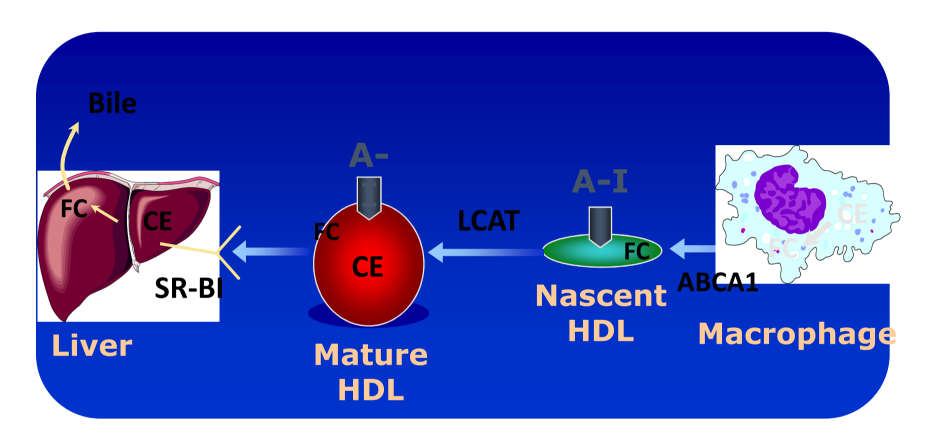
MEASUREMENTS:



Composition of Triglyceride-Rich Lipoproteins (% dry mass)



HDL and Reverse Cholesterol Transport



Plasma lipoproteins

Туре	Source	Major lipid	Apoproteins	ELFO	Athero- genicity
Chylomicrons	Gut	Dietary TGs	A-I, B-48, C-I, C-III, E	no mobility	– (pancreatiti s)
VLDL	Liver	Endogenous TGs	B-100, E, C- II, C-III,	Pre-β	+
IDL	VLDL remnant	Ch esters, TGs	B-100, C-III, E	Slow pre- β	+
LDL	VLDL, IDL	Ch esters	B-100	β	+++
HDL	Gut, liver	Ch esters, PLs	A-I, A-II, C-II, C-III, D, E	α	anti- atherogenic

Hereditary Causes of Hyperlipidemia

Physical findings





Fredrickson classification of hyperlipidemias

Phenotype	Lipoprotein(s) elevated	Plasma cholesterol	Plasma TGs	Athero- genicity	Rel. freq.	Treatment
Ι	Chylomicrons	Norm. to ↑	个个个个	– pancreatiti s	<1%	Diet control
IIa	LDL	$\uparrow \uparrow$	Norm.	+++	10%	Bile acid sequestrants, statins, niacin
IIb	LDL and VLDL	$\uparrow \uparrow$	$\uparrow \uparrow$	+++	40%	Statins, niacin, fibrates
III	IDL	$\uparrow \uparrow$	$\uparrow \uparrow \uparrow$	+++	<1%	Fibrates
IV	VLDL	Norm. to ↑	$\uparrow \uparrow$	+	45%	Niacin, fibrates
V	VLDL and chylomicrons	↑ to ↑↑	ተተተተ	+ pancreatiti s	5%	Niacin, fibrates

Primary hypercholesterolemias

Disorder	Genetic defect	Inheritance	Prevalence	Clinical features
Familial hyper- cholesterolemia	LDL receptor	dominant	heteroz.:1/500 5% of MIs <60 yr homoz.: 1/1 million	premature CAD (ages 30–50) TC: 7-13 mM CAD before age 18 TC > 13 mM
Familial defective apo B-100	apo B-100	dominant	1/700	premature CAD TC: 7-13 mM
Polygenic hypercholestero lemia	multiple defects and mechanisms	variable	common 10% of MIs <60 yr	premature CAD TC: 6.5-9 mM
Familial hyper- alphalipoprotein emia	unknown	variable	rare	less CHD, longer life elevated HDL

Primary hypertriglyceridemias

Disorder	Genetic defect	Inheritance	Prevalence	Clinical features
LPL deficiency	endothelial LPL	recessive	rare 1/1 million	hepatosplenomegaly abd. cramps, pancreatitis TG: > 8.5 mM
Apo C-II deficiency	Apo C-II	recessive	rare 1/1 million	abd. cramps, pancreatitis TG: > 8.5 mM
Familial hyper- triglyceridemia	unknown enhanced hepatic TG- production	dominant	1/100	abd. cramps, pancreatitis TG: 2.3-6 mM

Primary mixed hyperlipidemias

Disorder	Genetic defect	Inheritance	Prevalence	Clinical features
Familial dysbeta- lipoproteinemia	Apo E high VLDL, chylo.	recessive rarely dominant	1/5000	premature CAD TC: 6.5 -13 mM TG: 2.8 - 5.6 mM
Familial combined	unknown high Apo B-100	dominant	1/50 - 1/100 15% of MIs <60 yr	premature CAD TC: 6.5 -13 mM TG: 2.8 - 8.5 mM

Dietary sources of Cholesterol

Type of Fat	Main Source	Effect on Cholesterol levels
Monounsaturated	Olives, olive oil, canola oil, peanut oil, cashews, almonds, peanuts and most other nuts; avocados	Lowers LDL, Raises HDL
Polyunsaturated	Corn, soybean, safflower and cottonseed oil; fish	Lowers LDL, Raises HDL
Saturated	Whole milk, butter, cheese, and ice cream; red meat; chocolate; coconuts, coconut milk, coconut oil, egg yolks, chicken skin	Raises both LDL and HDL
Trans	Most margarines; vegetable shortening; partially hydrogenated vegetable oil; deepfried chips; many fast foods; most commercial baked goods	Raises LDL

Causes of Hyperlipidemia

- Diet
- Hypothyroidism
- Nephrotic syndrome
- Anorexia nervosa
- Obstructive liver disease
- Obesity
- Diabetes mellitus
- Pregnancy

- Obstructive liver disease
- Acute heaptitis
- Systemic lupus erythematousus
- AIDS (protease inhibitors)

Secondary hyperlipidemias

Disorder	VLDL	LDL	HDL	Mechanism
Diabetes mellitus	↑ ↑ ↑	1	Ţ	VLDL production ↑, LPL ↓, altered LDL
Hypothyroidism	1	$\uparrow\uparrow\uparrow$	J	LDL-rec.↓, LPL ↓
Obesity	11	1	Ţ	VLDL production ↑
Anorexia	-	1 1	-	bile secretion ↓, LDL catab. ↓
Nephrotic sy	11	1 1 1	Ţ	Apo B-100 ↑ LPL ↓ LDL- rec. ↓
Uremia, dialysis	1 1 1	-	\	LPL ↓, HTGL ↓ (inhibitors ↑)
Pregnancy	1 1	1 1	1	oestrogen ↑ VLDL production ↑, LPL ↓
Biliary obstruction PBC	-	-	J	Lp-X ↑ ↑ no CAD; xanthomas
Alcohol	↑↑ chylomicr. ↑	-	↑	dep. on dose, diet, genetics

When to check lipid panel

- Different Recommendations
 - Adult Treatment Panel (ATP III) of the National Cholesterol Education Program (NCEP)
 - Beginning at age 20: obtain a fasting (9 to 12 hour) serum lipid profile consisting of total cholesterol, LDL, HDL and triglycerides
 - Repeat testing every 5 years for acceptable values

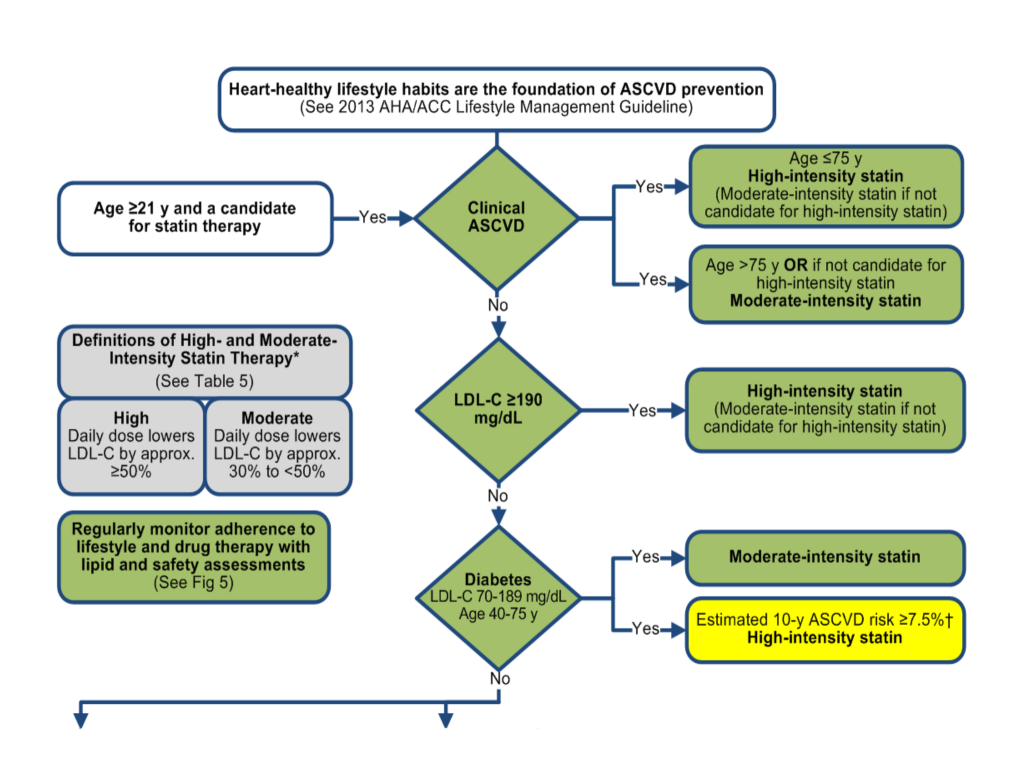
United States Preventative Services Task Force

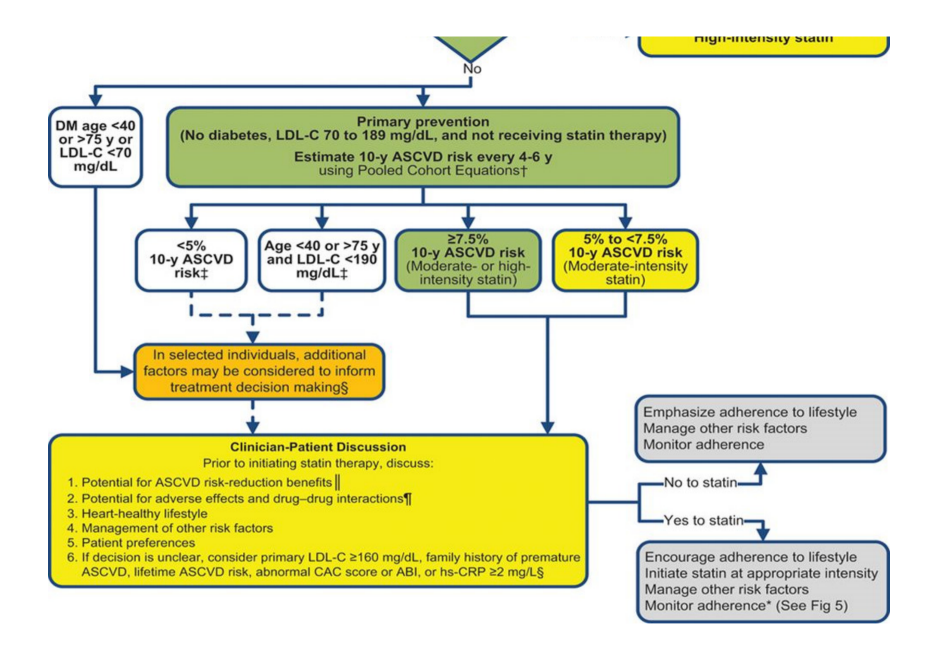
- ☐ Women aged 45 years and older, and men ages 35 years and older undergo screening with a total and HDL cholesterol every 5 years.
- □ If total cholesterol > 200 or HDL < 40, then a fasting panel should be obtained
- □ Cholesterol screening should begin at 20 years in patients with a history of multiple cardiovascular risk factors, diabetes, or family history of either elevated cholesteral levels or premature cardiovascular disease.

Treatment

Targets

- □LDL: To prevent coronary heart disease outcomes (myocardial infarction and coronary death)
- □Non LDL(TC/HDL): To prevent coronary heart disease outcomes (myocardial infarction and coronary death)
- ☐ Triglyceride: To prevent **pancreatitis** and may be coronary heart disease outcomes (myocardial infarction and coronary death)





Stone N J et al. Circulation. 2014;129:S1-S45



Guideline of therapy

Age	Risk Factors	Statin Intensity*
>29 Age	ASCVD	High
>29 years	LDL >190 mg/dl (4.9 mmol/l)	High
NO DAA	estimate 10-year risk for ASCVD <5%	No
NO DM LDL <190	estimate 10-year risk for ASCVD 5-7.5%	Moderate
	estimate 10-year risk for ASCVD >7.5%	High

Estimate 10-year risk for ASCVD

http://tools.acc.org/ASCVD-Risk-Estimator-Plus/#!/calculate/estimate/

AGE

SBP/DBP

T cholesterol

HDL

LDL

DM

Smoking

On Anti HTN

On statin

On asprin

Estimate 10-year risk for ASCVD

AMERICAN COLLEGE of CARDIOLOGY ASCV	D Risk Estimato	r Plus Estimate	Risk Ø 1	Therapy Impact	⊘ Advice
		••••			
Current Age 🛭 *	Sex *		Race *		
	Male	Female	White	African American	Other
Systolic Blood Pressure (mm Hg) * Value must be between 90-200	Value	stolic Blood Pressure (mm Hg) o			
Total Cholesterol (mg/dL) *	HDI	. Cholesterol (mg/dL) *	L	DL Cholesterol (mg/dL) 🐧 ^O	
Value must be between 130 - 320	Value	must be between 20 - 100	V	alue must be between 30-300	
History of Diabetes? *	Smo	oker: 🛛 *			
Yes	No	Yes	Forme		No
On Hypertension Treatment? * Yes	On No	a Statin? 🛭 ° Yes	No	On Aspirin Therapy? 🛭 ° Yes	No

Recommendations in DM

Age	Risk Factors	Statin Intensity*
	None	None
<40 years	ASCVD risk factor(s)	Moderate or high
	ASCVD	High
	None	Moderate
40-75	ASCVD risk factors	High
years	ACS & LDL ≥50 or in patients with history of ASCVD who can't tolerate high dose statin	Moderate + ezetimibe
	None	Moderate
>75 years	ASCVD risk factors	Moderate or high
	ASCVD	High
	ACS & LDL ≥50 or in patients with history of ASCVD who can't tolerate high dose statin	Moderate + ezetimibe

Statin Treatment

High-Intensity Statin Therapy	Moderate-Intensity Statin Therapy	Low-Intensity Statin Therapy
Daily dose lowers LDL-C, on average, by approximately ≥50%	Daily dose lowers LDL- C, on average, by approximately 30% to <50%	Daily dose lowers LDL- C, on average, by <30% Simvastatin 10 mg
Atorvastatin (40†)-80 mg Rosuvastatin 20 (40) mg	Atorvastatin 10 (20) mg Rosuvastatin (5) 10 mg Simvastatin 20-40 mg‡ Pravastatin 40 (80) mg Lovastatin 40 mg Fluvastatin XL 80 mg Fluvastatin 40 mg BID Pitavastatin 2-4 mg	Pravastatin 10-20 mg Lovastatin 20 mg Fluvastatin 20-40 mg Pitavastatin 1 mg

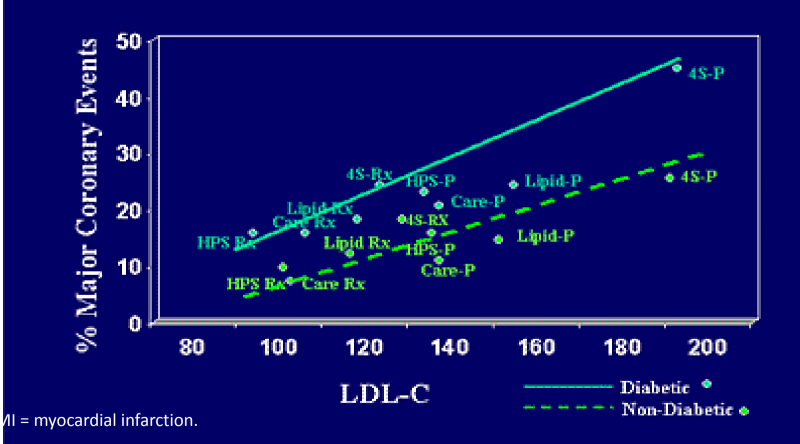
Treatment of Hyperlipidemia

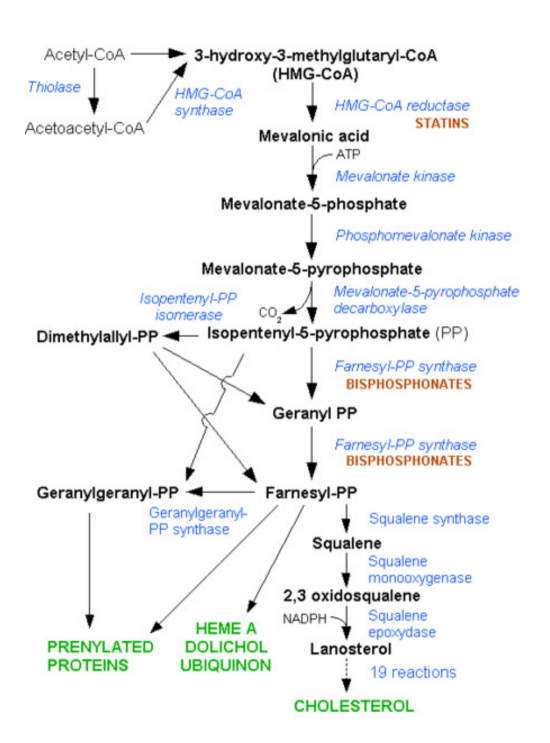
- ☐ Lifestyle modification
 - □Low-cholesterol diet
 - **□**Exercise
 - **□**Smoking
 - **□**Alcohol

Medications for Hyperlipidemia

Drug Class	<u>Agents</u>	Effects (% change)	Side Effects
HMG CoA reductase inhibitors	Statins	↓LDL (18-55),↑ HDL (5-15) ↓ Triglycerides (7-30)	Myopathy, increased liver enzymes
Cholesterol absorption inhibitor	Ezetimibe	↓ LDL(14-18), ↑ HDL (1-3) ↓Triglyceride (2)	Headache, GI distress
Nicotinic Acid		↓LDL (15-30), ↑ HDL (15-35) ↓ Triglyceride (20-50)	Flushing, Hyperglycemia, Hyperuricemia, GI distress, hepatotoxicity
Fibric Acids	Gemfibrozil Fenofibrate	↓LDL (5-20), ↑HDL (10-20) ↓Triglyceride (20-50)	Dyspepsia, gallstones, myopathy
Bile Acid sequestrants	Cholestyramine	↓ LDL↑ HDL No change in triglycerides	GI distress, constipation, decreased absorption of other drugs
PCSK9	Evolocumab Alirocumab	↓ LDL (50-60%)	injection-site reactions, muscle pain, neurocognitive adverse events. These included memory impairment and confusion

Statin Risk Reduction in Diabetic Patients and Non-Diabetic Patients





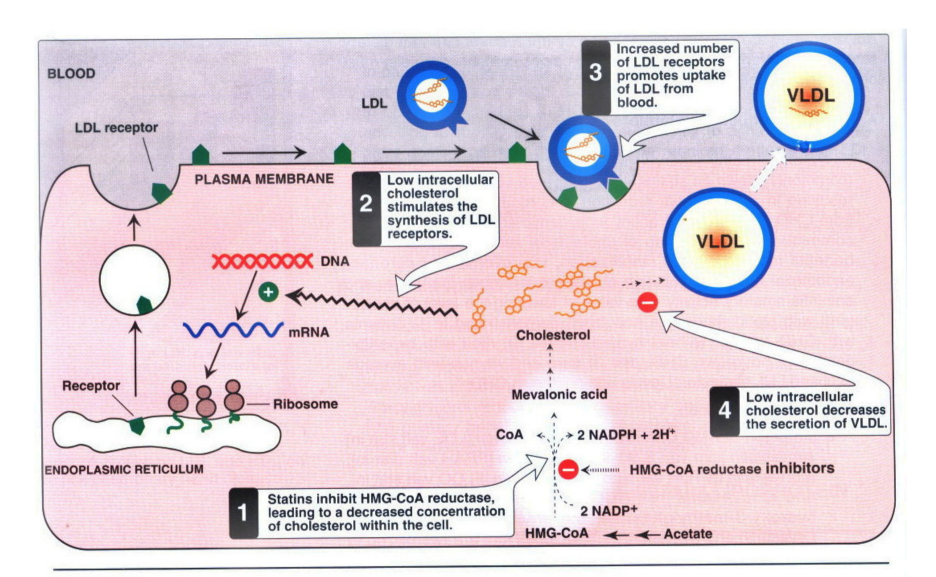


Figure 21.5 Inhibition of HMG-CoA reductase by the statin drugs.

Table 1: Assessment and action strategies for elevated plasma triglyceride concentrations [TG]

[TG], mmol/L	Step Action and comments	Retest interval, mo*
< 2	Continue current managementReassess lipid profile regularly, to ensure that [LDL-C] is at target	6-12
≥ 2, < 5	 1. Therapeutic lifestyle measures Weight control Reduce dietary fat, simple sugars Reduce alcohol intake Increase physical activity Reassess lipid profile regularly, to ensure th [LDL-C] is at target 	3-6 at
	 2. Manage other secondary factors Control glycemia, if diabetic Reassess medications; consider lipid-neutral alternatives 	al
	 3. Consider pharmacologic treatment Intensify LDL-lowering (e.g., statin therapy Fish oil (omega-3 fatty acid) Niacin (e.g., extended release) 	y)

Table 1: Assessment and action strategies for elevated plasma triglyceride concentrations [TG]

≥ 5, < 10	 4. Intensify steps 1-3, above [LDL-C] cannot be estimated when [triglycerides] > 5 mmol/L Apolipoprotein B determination might be helpful 	2-3
	 5. Consider fibrate therapy, e.g., Bezafibrate (Bezalip) 400 mg/d Fenofibrate Lipidil micro 200 mg/d Lipidil supra 160 mg/d Lipidil EZ 145 mg/d Gemfibrozil (Lopid) 600-1200 mg/d 	
≥ 10	 6. Further intensify steps 1-3 With acute pancreatitis: • Very-low-fat diet (10%-15% of energy intake) • Cessation of alcohol • Insulin, if indicated for glycemic control • Admit patient to hospital — Nothing by mouth: IV fluid replacement — Plasma exchange is unhelpful 	1-2
	7. Initiate fibrate therapyMonitor serum [creatinine]	

George Yuan, Khalid Z. Al-Shali, Robert A. Hegele

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THANK YOU

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See you in 5th year MED-441 Course