Preventive Cardiology Beyond Statins for

Cardiovascular Risk Reduction

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Introduction

Objectives

- 1. Identify clinical scenarios in which statins and/or non-statin lipid lowering treatments are indicated
- 2. Describe the mechanism of action of and indications for PCSK9 inhibitors, SGLT2 inhibitors, and high dose <u>omega-3-polyunsaturated fatty</u> <u>acids</u>
- No competing interests /financial relationships to disclose
- I will discuss what is currently <u>off-label use of</u> <u>icosapent ethyl</u> (Vascepa®)
- Branded Rx/OTC products shown: not an endorsement









Treatment	algorithm (20	: NLA Par 15)	t 1 / Part 2
0-1	Major ASCVD ⊨ HTN Age (M≥45, F≥ Low HDL EmHx early Cl Smoking	risk factors (<u>HA</u> :55 y) HD (M<55, F<6	<u>LFS)</u> 5)
≥3			
	Risk Category	Non-HDL-C goal	LDL-C goal
	Low	< 130 mg/dl	< 100
	Moderate	< 130	< 100
	High	< 130	< 100
	Very High	< 100	< 70
	ASCVD, or [Diabe	etes mellitus + end	organ damage]
Jacobson TA, Ito MK, Maki KC, et a	al. J Clin Lipidology 2015;9:129-1	169.	



LDL-C vs	. No	n-HC	DL-C
	Lipoprotein	Cholesterol	TG
		(approx. % of lipid content)	(approx. % of lipid content)
	LDL	70	30
VLDL	VLDL or Chylomicron remnant	20	80
СМ	Chylomicron	5	95
Walker HK, Hall WD, Hurst JW, eds. Chapter 31, Chole	esterol, Triglycerides, a	and Associated Lipoprote	ins. Butterworths 1990.

LDL-	C vs.	Non-H	DL-C
• We live in a	n LDL-C parad	igm. Why?	С тс
Diagnosis → ↓ Lipids (mg/dl)	Normal	Familial Hyperchol.	Metabolic Syndrome / DM
Total-C	158	342	318
HDL-C	59	49	23
LDL-C	88	280	?
Triglycerides	53	67	1,621
Non-HDL-C	99	293	295
Depiction LDL	-		
VLDL / CM remn.			
Walker HK, Hall WD, Hurst JW	, eds. Chapter 31, Cholesterol, Tr	iglycerides, and Associated Lipo	proteins. Butterworths 1990.

LDL-C	Non- HDL-C	N (MACE)	N (Total)	HR (95% CI)	
≥ 100 mg/dl	≥ 130 mg/dl	1,877	10,419	1.21 (1.13-1.29)	-
≥ 100 mg/dl	< 130 mg/dl	467	2,873	1.02 (0.92-1.12)	
< 100 mg/dl	≥ 130 mg/dl	283	1,435	1.32 (1.17-1.50)	
< 100 mg/dl	< 130 mg/dl	2,760	23,426	1.00 (Reference)	1.0 1.5 HR (95% CI)

• HRs adjusted for sex, age, smoking, DM, SBP, and trial

Boekholdt SM, Arsenault BJ, Mora S, et al. JAMA. 2012;307:1302–1309

Cited in Jacobson TA, Ito MK, Maki KC, et al. J Clin Lipidology 2015;9:129-169.



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St	atins
Citrate ATP citrate lyase Acetyl-CoA	 Dosing + Effects: Potency varies by statin and dose High intensity: >50% ↓LDL- C
HMG-CoA STATINS HMG-CoA reductase	 May also decrease TRG and HDL PO formulations Once daily administration* Newer agents can be taken
Mevalonic acid	 at any time of day Pleiotropic effects Adverse Effects: Mvalgias, Gl upset
Rosuvastatin [package insert]/ AstraZeneca. Wilmington,	Drug interactions *exception: fluvastatin DE. 11/2018. Graphic original.

Stati	ins	– P	oter	ıcy	+ Li	рор	hilicity
Intensity:	Lova-	Prava-	Simva-	Fluva-	Pitava-	Atorva-	Rosuvastatin
Low	20mg	20mg	10mg	40mg	1mg		
Mod	40mg	40mg	20mg	80mg	2mg	10mg	5mg
WOO.	80mg	80mg	40mg		4mg	20mg	10mg
			(80mg)			40mg	20mg
High						80mg	40mg
		Lipop	ohilic	Atorvasta simvasta	atin, lovast tin	atin,	
		Hydro	philic	Pravasta rosuvast	tin, atin, fluvas	tatin	



























PCSK9 inhibitors					
	FOURIER	ODYSSEY OUTCOMES			
Primary endpoint (composite)	CV death, MI, stroke, hospitalization for unstable angina, or cor. revascularization	CHD death, non-fatal MI, fatal or non-fatal ischemic stroke, or UA requiring hospitalization			
Treatment vs. placebo	9.8% vs. 11.3%	9.5% vs. 11.1%			
Median follow up	2.2 years	2.8 years			
HR	0.85	0.85			
NNT	67	64			



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ω-3-poly-unsaturated fatty acids

Does treatment change outcomes?

Trial	Endpoints / Mean Follow-up	Daily dose	Outcome
Meta-analysis of 10 trials Aung et al. <i>JAMA <u>Cardiol</u></i> 2018 (n=77,917) prior CHD, CVA, or high ASCVD risk	Any CHD (fatal/nonfatal) or major vascular events 4.4 years	Generally 1 g EPA/DHA	No effect
GISSI- <u>Prevenzione</u> investigators <i>Lancet</i> 1999; (n=11,324) with recent MI (2x2 design also with vit. E)	Death, non-fatal MI, CVA 3.5 years	1 g EPA/DHA vs. Placebo	Benefit Composite RRR 10% Death RRR 14%
JELIS Yokogama et al. <i>Lancet</i> 2007 (n=18,645) unselected hypercholesterolemic (Total-C > 252 mg/dl) Japanese patients	Any CHD event (CHD death, SCD, fatal/nonfatal MI, UA, PCI, CABG) 4.6 years	Statin + [1.8 g EPA- only or placebo]	Benefit Composite RRR 19% No difference in LDL





ω-3-	poly-	unsat	turat	ted
	fatty	/ acid	S	

Antiarrhythmic or not?				
	Trial	Dose	Outcomes	
	REDUCE-IT 2019	4 g/d EPA only	↑47% excess atrial fib/flutter	
	Cochrane Review 2018 79 RCTs, (n=112,059)	Varies (0.5 to >5 g/d)	Marine: No difference arrhythmia Plant-based (ALA): ↓21% arrhythmias	
	GISSI-HF (n=6,975) with HF	1 g/d mixed	No difference in atrial fibrillation ↓9% mortality; ↓8% HF admissions	

 Animal studies suggest DHA may have antiarrhythmic properties in AF

Bhatt et al. New Engl J Med 2019;380(1):11-22. Abdelhamid et al. Cochrane Database Syst Rev 2018;11:CD003177. Aleksova et al. Eur J Heart Fail 2013;15(11):1289-95. Tavazzi et al. Lancet 2008;372(9645):1223-30. Ninio et al. J Cardiovasc Electrophysiol 2005 16:1189-1194.

•	ω-3-poly-unsaturated fatty acids Current Rx products and labeling							
	Agent	Trade Name Composition	Dose	Labeled Indication				
	Icosapent ethyl	<u>Vascepa</u> ®	2 g bid with food	 Significant hypertriglyceridemia (>500 mg/dl) as adjunct to diet and exercise 				
	ω-3 acid ethyl esters	Lovaza® 55% EPA / 45% DHA	4 g gd or 2 g bid +/- food	 Significant hypertriglyceridemia (>500 mg/dl) as adjunct to diet and exercise For use as adjunct to simvastatin for hyper-TG 				
	ω-3 carboxylic acids	Epanova® Mostly EPA	2-4 g gd +/- food	 Significant hypertriglyceridemia (>500 mg/dl) as adjunct to diet and exercise 				
	Source: Drug Mono	ographs. Gold Stand	dard. Accessed	120 Jan 2019.				





"Nutraceuticals" and lifestyle changes

ntervention	Mechanism of action	Dose	Expected Δ LDL-C (relative)
ncreased physical activity	Multifactorial	200-300 min/week	↓ ~ 5%
Loss of body weight	Multifactorial	↓5% body weight	↓ 3-5%
Diet low in saturated and trans fats	↓LDL-C production		↓ 5-10%
Viscous fiber	Bile acid sequestration, ↑satiety	5-10 g/day	↓ 5-20%
Plant sterols/ <u>stanols</u>	Competitive inhibition of cholesterol absorption	2 g/day	↓ ~10%
	cholesterol absorption		

Agent	Mechanism of action	Dose	∆ LDL-C (absolut <u>e)</u>
Berberine	Has PCSK9 inhibitory properties, increases LDLR expression and decreases intestinal <u>chol</u> . absorption	300 mg/day	↓25 mg/dl
Artichoke	Luteolin interacts with HMG-CoA reductase, SREBPs, ACAT	500-2,700 mg/d	↓15 mg/dl
Garlic	Inhibition of HMG-CoA reductase	5-6 g/d	↓9 mg/d
Green tea	Inhibition of inducible NO synthase, inhibition of HMG-CoA reductase	170-1,200 mg/d	↓7 mg/dl



Take Home Points

- Recent history of and important concepts in clinical lipidology
 - Please consider non-HDL-C as well as LDL-C lowering, especially in hypertriglyceridemics
- New ACC/AHA Blood Cholesterol guidelines
 - Goal atherogenic cholesterol levels are both motivating and evidence based
- PCSK9 inhibition: when and how?
 - FH or ASCVD and LDL-C > 70 or non-HDL-C > 100 mg/dl

