State-of-the-Art Review

Same evidence, varying viewpoints: Three questions illustrating important differences between United States and European cholesterol guideline recommendations

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ABSTRACT
In 2018, the AHA/ACC Multisociety Guideline on the Management of Blood Cholesterol was released. Less than one year later, the 2019 ESC/EAS Dyslipidemia Guideline was published. While both provide important recommendations for managing atherosclerotic cardiovascular disease (ASCVD) risk through lipid management, differences exist. Prior to the publication of both guidelines, important randomized clinical trial data emerged on non-statin lipid lowering therapy and ASCVD risk reduction. To illustrate important differences in guideline recommendations, we use this data to help answer three key questions: 1) Are ASCVD event rates similar in high-risk primary and stable secondary prevention? 2) Does imaging evidence of subclinical atherosclerosis justify aggressive use of statin and non-statin therapy (if needed) to reduce LDL-C levels below 55 mg/dL as recommended in the European Guideline? 3) Do LDL-C levels below 70 mg/dL achieve a large absolute risk reduction in secondary ASCVD prevention? The US guideline prioritizes both the added efficacy and cost implications of non-statin therapy, which limits intensive therapy to individuals with the highest risk of ASCVD. The European approach broadens the eligibility criteria by incorporating goals of therapy in both primary and secondary prevention. The current cost and access constraints of healthcare worldwide, especially amidst a COVID-19 pandemic, makes the European recommendations more challenging to implement. By restricting non-statin therapy to a subgroup of high- and, in particular, very high-risk individuals, the US guideline provides primary and secondary ASCVD prevention recommendations that are more affordable and attainable. Ultimately, finding a common ground for both guidelines rests on our ability to design trials that assess cost-effectiveness in addition to efficacy and safety.

1. Introduction

In 2018, the American Heart Association (AHA)/American College of Cardiology (ACC) Multisociety Guideline on the Management of Blood Cholesterol was released [1]. Less than one year later, the 2019 European Society of Cardiology (ESC)/European Atherosclerosis Society (EAS) Dyslipidemia Guideline was published [2]. Both guidelines highlight approaches for atherosclerotic cardiovascular disease (ASCVD) risk reduction through lipid management. While agreement exists that low-density lipoprotein-cholesterol (LDL-C) is the primary target for risk reduction and statin therapy is the first-line therapy to reduce LDL-C levels and ASCVD risk, they diverge at some important clinical decision points in the management of ASCVD risk for both primary and secondary prevention.

This narrative review examines both guidelines and focuses on three key questions:
1. Are ASCVD event rates similar in high-risk primary and stable secondary prevention?

2. Does imaging evidence of subclinical atherosclerosis justify aggressive use of statin and non-statin therapy (if needed) to reduce LDL-C levels below 55 mg/dL as recommended in the ESC/EAS Guideline?

3. Do LDL-C levels below 70 mg/dL achieve a large absolute risk reduction (ARR) in secondary ASCVD prevention?

1.1. Examining the AHA/ACC Multisociety and ESC/EAS guidelines

The AHA/ACC Multisociety Guideline utilizes established criteria in primary prevention to help determine when statin therapy should be initiated or intensified (Table 1). It suggests that the decision-making should be based on a clinician–patient risk discussion, risk-enhancing factors (Fig. 1) and a coronary artery calcium (CAC) score may be used to further guide decision-making. Incorporation of these other factors into the clinician–patient risk discussion promotes personalized treatment decisions for reducing risk in the primary prevention of ASCVD [3].

For secondary prevention and high-risk primary prevention patients, maximally tolerated statin therapy is recommended to reduce LDL-C by at least 50%. For the highest risk secondary prevention patients, where LDL-C remains at or above the treatment threshold of 70 mg/dL despite maximally tolerated statin therapy, non-statin therapy is recommended in preference to ezetimibe as the first line therapy, based on its ease of accessibility and low cost. If the LDL-C level remains ≥70 mg/dL (or non-high density lipoprotein cholesterol (non-HDL-C) remains ≥100 mg/dL) on maximally tolerated statin therapy and ezetimibe, then the addition of a proprotein convertase subtilisin/kexin type 9 inhibitor (PCSK9i) is reasonable, though this is more likely to be cost-effective if the LDL-C is >100 mg/dL.

The ESC/EAS Dyslipidemia Guideline has prioritized treatment goals, including percent LDL-C reduction and fixed targets, for guiding lipid-lowering therapy (Table 1). If a maximally tolerated statin is unable to achieve an LDL-C goal based on an individual’s estimated risk or comorbidities, non-statin therapy should be considered for both primary and secondary prevention to achieve that desired target. While the ESC/EAS Guideline recommends consideration of risk modifiers (Fig. 1) and a CAC score for additional risk stratification, it also includes carotid and femoral plaque imaging, which does not carry the same negative predictive value as a CAC score of zero [4].

While the ESC/EAS Guideline includes the option to add on non-statin therapy for all subgroups to achieve specific LDL-C goals based on data that ‘lower is better’, the AHA/ACC Multisociety Guideline defines a smaller, more specific subgroup where use of non-statin therapy is most beneficial. Ultimately, constructing a framework to optimize the use of lipid-lowering therapy requires careful consideration of an individual’s absolute risk over 10 years, LDL-C levels on optimized treatment regimen, and the cost of different forms of non-statin therapy. A ‘highest risk–highest benefit’ matrix has been proposed as one approach [5]. It should be noted that both guidelines are largely in agreement on the general principle of tailoring the intensity of interventions to the level of risk. However, there are some differences regarding which aspects should be prioritized when determining their recommendations. In an attempt to further inform the comparative reviews of both guidelines [6,7], important differences between the two guidelines can be clarified by answering three specific questions (Fig. 2).

Question

1. Are ASCVD event rates similar in high-risk primary and stable secondary prevention?

Both guidelines leverage scoring systems derived from population-based studies to estimate an individual’s risk for total and fatal ASCVD events – the Pooled Cohort Equations in the United States (US) and the Systematic Coronary Risk Estimation in Europe, respectively. While there are imprecisions with risk estimation at the individual level [8], both scoring systems are based on the principle that the intensity of prevention efforts should match the absolute ASCVD risk of the individual [9].

Currently, there is no randomized clinical trial (RCT) data to support the use of non-statin therapy as an add-on for primary prevention other than in individuals with a baseline LDL-C ≥190 mg/dL or heterozygous familial hypercholesterolemia. The AHA/ACC Multisociety Guideline adheres closely to the trial evidence related to this approach, whereas the ESC/EAS Guideline extends use of non-statin therapy in primary prevention based on the belief that event rates are similar in high-risk primary and stable secondary prevention.

Data from two key trials can be helpful in clarifying this (Fig. 3). In the Reduction of Cardiovascular Events with Icosapent Ethyl Intervention Trial (REDUCE-IT) [10], which enrolled patients on statin therapy with established ASCVD or diabetes and additional cardiovascular risk factors and followed them over an ~5-year period, the percentage of patients with an adverse cardiovascular event was almost double in the placebo arm (25.5%) of the secondary prevention cohort (patients with established ASCVD) versus the placebo arm (13.6%) of the high-risk primary prevention cohort, which included patients with diabetes and other cardiovascular risk factors, but without ASCVD.

In the Dapagliflozin and Cardiovascular Outcomes in Type 2 Diabetes Trial (DECLARE-TIMI 58), which enrolled patients with type 2 diabetes on statin therapy or ezetimibe and followed them over an ~4 year period, the percentage of patients with a major adverse cardiovascular events (MACE) was almost three times greater in the placebo arm of the secondary (15.3%) versus primary (5.2%) prevention cohort [11]. While both groups included patients with diabetes, this trial still allows for comparison of event rates among high-risk primary and stable secondary prevention patients.

First, both RCTs include a unique population with significant lipid and inflammatory risk, which may limit the generalizability to all primary and secondary cohorts. Collectively though, without adjusting for baseline differences between both subgroups, these studies indicate that event rates in secondary prevention are considerably higher than in high-risk primary prevention. In addition, despite similar risk factor burden in these patients, development of an ASCVD event in one identifies inherent host susceptibility to those risk factors or yet unknown factors that may make a host more susceptible to another event.

As a result, the AHA/ACC Multisociety Guideline largely limits drug treatment to statin therapy for primary prevention, with the exception of ezetimibe and/or a PCSK9i in patients with severe hypercholesterolemia and an LDL-C ≥100 mg/dL despite adherence to high-intensity statin therapy. Non-statin therapy is more likely to be used in secondary prevention but is informed by the level of ASCVD risk. The more conservative use of non-statin therapy is based on a desire to add value by considering both net clinical benefit and cost, with a number needed to treat (NNT) ≤50 being reasonably indicative of this [12].

This concept when applied to ezetimibe and PCSK9i was previously validated and clarifies the thresholds to initiate non-statin therapy utilized in the AHA/ACC Multisociety Guideline [13]. Use of ezetimibe (with an approximate 24% decrease in LDL-C) has been suggested to be reasonable at a NNT ≤50 for very high-risk patients with a LDL-C ≥130 mg/dL or high-risk patients with a LDL-C ≥190 mg/dL. The addition of a PCSK9i (with a 50–65% reduction in LDL-C) is felt to be reasonable at a NNT ≤50 for very high-risk patients with a LDL-C ≥70 mg/dL or high-risk patients with a LDL-C ≥100 mg/dL.

In contrast, the ESC/EAS Guideline maintains that adding generic ezetimibe is reasonable to achieve lower LDL-C levels in primary and secondary prevention, which is often separated by a fine line with just an abrupt and adverse event being the only transition. They cite the heterogeneity of these patients, whom upon further stratification reveals a group with a wide range of risk based on varying levels of subclinical atherosclerosis. For example, based on retrospective cohort data, the ACC/AHA previously described individuals with a CAC >400 as highest
| Statin therapy only; >30% reduction in LDL-C | AHA/ACC Multisociety |
|----------------------------------------------|---------------------|
| **Primary Prevention**                       |                     |
| In adults 40-75 years with LDL-C 70-189 mg/dl|                     |
| • 10-year ASCVD risk ≥ 7.5% to 19% (Consider when 10-year ASCVD risk ≥ 5 to 7.5% + REF); Diabetes without multiple diabetes risk factors |                     |

| Statin therapy only in most cases; >50% reduction in LDL-C | AHA/ACC Multisociety |
|-----------------------------------------------------------|---------------------|
| **Primary Prevention**                                    |                     |
| • 10-year ASCVD risk ≥ 20%; LDL-C ≥190 mg/dl; Diabetes + multiple risk factors |                     |
| **Secondary Prevention**                                  |                     |
| • Clinical ASCVD                                           |                     |
|   • ACS; History of MI; Stable or unstable angina; Coronary or other arterial revascularization; Stroke/ TIA; PAD (including aortic aneurysm of atherosclerotic origin) |                     |

| Statin + add-on therapy; >50% reduction in LDL-C | AHA/ACC Multisociety |
|--------------------------------------------------|---------------------|
| **Primary Prevention**                           |                     |
| • HeFH or LDL-C ≥190 mg/dL (1st ezetimibe; 2nd PCSK9i) |                     |
| **Very High Risk**                               |                     |
| **Secondary Prevention**                         |                     |
| • Multiple major ASCVD events                    |                     |
|   • ACS within the past 12 months; A history of MI other than the ACS event; Ischemic stroke; Symptomatic PAD with claudication and an ABI <0.85 or prior peripheral revascularization/amputation |                     |
| • 1 major ASCVD event + multiple high-risk conditions |                     |
|   • Age ≥65 years; HeFH; Prior coronary artery revascularization; Diabetes mellitus; Hypertension; CKD; Current smoking; History of heart failure |                     |
| • A persistently elevated LDL-C level (≥100 mg/dL on maximal statin therapy and ezetimibe) |                     |

| **ESC/EAS**                                         |                     |
| **LDL-C Goal <116 mg/dL**                          |                     |

| **Low Risk**                                       |                     |
| **Primary Prevention**                             |                     |
| **SCORE <1%**                                      |                     |

| **Moderate Risk**                                  |                     |
| **Primary Prevention**                             |                     |
| • SCORE 1-5%; Type 1 Diabetes ≥35 years; Type 2 Diabetes ≥50 years |                     |

| **High Risk**                                      |                     |
| **Primary Prevention**                             |                     |
| • SCORE 5-10%; Total cholesterol ≥310 mg/dL or LDL-C ≥190 mg/dL; BP >180/110; HeFH without other major risk factors; Moderate CKD; Type 2 Diabetes >10 years or + risk factors (no organ damage) |                     |

| **Very High Risk**                                 |                     |
| **Primary Prevention**                             |                     |
| • SCORE >10%; HeFH with another major risk factor; Severe CKD; Type 2 Diabetes + target organ damage |                     |
| **Secondary Prevention**                           |                     |
| • Clinical ASCVD                                    |                     |
|   • Previous ACS; Stable Angina; Coronary revascularization; Stroke/TIA; PAD |                     |
| • Imaging ASCVD                                     |                     |
|   • Significant plaque on coronary angiography; Multivessel CAD with 2 major epicardial arteries >50% stenosis on CCTA or carotid ultrasound |                     |
| • HeFH with ASCVD                                   |                     |

*For patients with ASCVD who experience a second vascular event within 2 years (not necessarily of the same type as the first event) while taking maximally tolerated statin-based therapy, an LDL-C goal of <40 mg/dL may be considered.

Abbreviations: AHA – American Heart Association; ACC – American College of Cardiology; ESC – European Society of Cardiology; EAS – European Atherosclerosis Society; ASCVD – Atherosclerotic Cardiovascular Disease; LDL-C – Low-Density Lipoprotein Cholesterol; HeFH – Heterozygous Familial Hypercholesterolemia; SCORE – Systematic Coronary Risk Estimation; PAD – peripheral arterial disease; BP – Blood Pressure; CKD – Chronic Kidney Disease; ABI – Ankle-brachial index; MI – myocardial infarction; ACS – acute coronary syndrome; REF – risk enhancing factors
**Fig. 1.** List of AHA/ACC risk enhancing factors and ESC/EAS risk Modifiers

**Abbreviations:** AHA – American heart association; ACC – American college of cardiology; ESC – european society of cardiology; EAS – european atherosclerosis society; ASCVD – atherosclerotic cardiovascular disease; LDL-C – low-density lipoprotein cholesterol; CVD – cardiovascular disease; BMI – body mass index.

### AHA/ACC Risk Enhancing Factors
- Family History of Premature ASCVD
- Primary Hypercholesterolemia
  - LDL-C 160-189 mg/dL (4.1-4.8 mmol/L);
  - non-HDL-C 190-219 mg/dL (4.9-5.6 mmol/L)
- Metabolic Syndrome
- Chronic Kidney Disease
- Chronic Inflammatory Conditions
- History of Premature Menopause or Preeclampsia
- High-Risk Ethnicities
- Persistently Elevated Primary Hypertriglyceridemia >175 mg/dL

*If measured, consider these additional factors:*
- High-Sensitivity C-Reactive Protein ≥2 mg/L
- Lipoprotein (a) ≥50 mg/dL or ≥125 nmol/L
- Apolipoprotein B ≥130 mg/dL
- Ankle Brachial Index <0.9

### ESC/EAS Risk Modifiers
- Social Deprivation
- Obesity (Measured by BMI)
- Central Obesity (Measured by Waist Circumference)
- Physical Inactivity
- Psychosocial Stress
- Family History of Premature CVD
- Chronic Immune-Mediated Inflammatory Disorder
- Major Psychiatric Disorders
- Treatment for Human Immunodeficiency Virus Infection
- Atrial Fibrillation
- Left Ventricular Hypertrophy
- Chronic Kidney Disease
- Obstructive Sleep Apnea
- Non-Alcoholic Fatty Liver Disease

**Fig. 2.** A Comparison of the Evidence Favoring the AHA/ACC and ESC/EAS Approach to the Three Key Questions

**Abbreviations:** AHA – American Heart Association; ACC – American College of Cardiology; ESC – European Society of Cardiology; EAS – European Atherosclerosis Society; ASCVD – Atherosclerotic Cardiovascular Disease; LDL-C – Low-Density Lipoprotein Cholesterol; REDUCE-IT – Reduction of Cardiovascular Events with Icosapent Ethyl-Intervention Trial; DECLARE-TIMI 58 – Dapaglifozin and Cardiovascular Outcomes in Type 2 Diabetes Trial; 1 Primary; 2 Secondary; IMPROVE-IT – Improved Reduction of Outcomes: Vytorin Efficacy International Trial; TIMI – Thrombolysis In Myocardial Infarction; CABG – Coronary Artery Bypass Graft; PCSK9i – Proprotein Convertase Subtilisin/Kexin Type 9 Inhibitor; Rx – Prescriptions; ARR – Absolute Risk Reduction; FOURIER – Further Cardiovascular Outcomes Research With PCSK9 Inhibition in Subjects with Elevated Risk; ODYSSEY OUTCOMES – Evaluation of Cardiovascular Outcomes after an Acute Coronary Syndrome During Treatment with Alirocumab; CAC – Coronary Artery Calcium; SCCT – Society of Cardiovascular Computed Tomography; MACE – Major Adverse Cardiovascular Events.
vascular death rates (0.80%/year) were very similar to those of a stable very-high risk primary prevention population where annualized cardio-event rates in secondary prevention patients compared to those with secondary prevention cohort. These factors could lower the observed use and intensity of statin therapy, which was much higher in the stable primary prevention group. However, it does not account, however, for variation in the follow-up duration or the practice patterns, specifically appropriate statin therapy allocation and optimal blood pressure control, which was much higher in the stable secondary prevention cohort. These factors could lower the observed event rates in secondary prevention patients compared to those with significant subclinical atherosclerosis.

With improvement in early initiation, adherence and intensification of risk factor modification, including diet, exercise and lipid lowering, hypoglycemic and anti-hypertensive medications, high-risk primary prevention individuals with significant subclinical atherosclerosis can further reduce their ASCVD risk [18–20]. Therefore, if more current practice patterns, specifically appropriate statin therapy allocation and optimal blood pressure control, were implemented equitably among the high-risk primary and stable secondary prevention patients, the event rates may more accurately reflect modern RCT data.

Based on the REDUCE-IT and DECLARE-TIMI 58 trial findings, as well as cohort data from high-risk primary prevention patients with significant subclinical atherosclerosis, a spectrum of risk in primary and secondary prevention exists. Varying interpretation of the event rates in high-risk primary and stable secondary prevention patients has led to differences in recommendations for the treatment of ASCVD risk in the US and Europe.

According to the AHA/ACC Multisociety Guideline, ezetimibe, based on its ease of accessibility, cost, and ability to reduce LDL-C levels by 15–24% on top of statin therapy, is currently first-line non-statin add-on therapy only in secondary prevention or select primary prevention groups. The EVAPORATE trial showed that icosapent ethyl can reduce low attenuation plaque volume providing mechanistic insight into the benefits of icosapent ethyl in REDUCE-IT [21]. However, further confirmatory RCT evidence of the benefit of icosapent ethyl in other subgroups such as high-risk primary prevention patients without diabetes is needed. As access to icosapent ethyl improves and additional RCT data emerges, it will be important to consider this therapy given its significant effect on hard ASCVD events in patients with a triglycerides level ≥150 mg/dL. Based on initial data, the effect of icosapent ethyl is greater for secondary than primary prevention (NNT 17 vs. 61) as well [10]. Additional analyses have also demonstrated icosapent ethyl’s benefit in not only reducing initial but also subsequent events in a population at high risk for ischemic events with an annualized placebo primary end point rate of 5.7% [22]. While these effects are more pronounced than those noted with ezetimibe in the Improved Reduction of Outcomes: Vytorin Efficacy International Trial (IMPROVE-IT), the current cost of icosapent ethyl is likely to limit broader use.

Question 2 Does imaging evidence of subclinical atherosclerosis justify aggressive use of statin and non-statin therapy (if needed) to reduce LDL-C levels below 55 mg/dL as recommended in the ESC/EAS Guideline?

In the AHA/ACC Multisociety Guideline, imaging to detect subclinical atherosclerosis is limited to a CAC score when the decision to initiate statin therapy in primary prevention is uncertain. If CAC is present, statin therapy is recommended, especially if CAC is ≥ 100. However, once statin therapy is initiated, CAC is no longer indicated for guiding treatment decisions, including adding non-statin therapy.

The ESC/EAS Guideline recommends that a CAC score be used to help guide decision-making in individuals with low to moderate risk for ASCVD. However, by having a combined treatment goal of both a ≥50% reduction in LDL-C and a LDL-C level <55 mg/dL, non-statin therapy is likely to be needed frequently.

While extremely high levels of calcified coronary plaque (a CAC score ≥1000), which is strongly correlated with total plaque, can pose substantial risk in primary prevention [16], the US guideline maintains that patients with advanced subclinical atherosclerosis on appropriate statin therapy are unlikely to derive significant enough clinical benefit to justify routine non-statin therapy as part of aggressive LDL-C lowering.

For example, in IMPROVE-IT [23], which enrolled a very high-risk secondary prevention population with an acute coronary syndrome (ACS) within the preceding 10 days, the addition of ezetimibe to statin therapy (32.7%) resulted in a ~2% ARR compared to statin therapy alone (34.7%) for the primary outcome (CV mortality, major CV event, or nonfatal stroke) over a 7-year period. Even among this very high-risk
population, the ARR of adding ezetimibe to statin therapy was most pronounced in patients with recent ACS and additional high-risk features, including those with a high Thrombolysis In Myocardial Infarction (TIMI) Risk Score, diabetes, and age ≥ 75 years, as well as prior stroke, or prior coronary artery bypass graft surgery [24].

Therefore, the approach in the ESC/EAS Guideline to target LDL-C levels below 55 mg/dL based on the presence of subclinical atherosclerosis is unlikely to be cost-effective if ezetimibe and/or PCSK9i are needed to get the LDL-C this low. In most cases, the presence of moderate or advanced subclinical atherosclerosis in those on optimal medical therapy does not usually raise ASCVD risk to the level of a patient with clinical ASCVD especially after moderate-to high-intensity statin therapy is used. Therefore, the use of non-statín therapy (especially with a PCSK9i) in this population is likely to have a NNT considerably >50.

Question 3 Do LDL-C levels below 70 mg/dl achieve a large ARR in secondary ASCVD prevention?

The Cholesterol Treatment Trials’ Collaboration demonstrated an ~22% relative risk reduction in major ASCVD events per mmol/L reduction in LDL-C (~39 mg/dL) [25]. This analysis was pivotal in validating the benefits of LDL-C lowering (with statin therapy) to reduce ASCVD risk. As can be seen, multiplying relative risk reduction (~22%/mmol) from LDL-C lowering therapies by the patient’s absolute risk determines the absolute risk reduction achieved. Additionally, when LDL-C is higher, the greater the LDL-C reduction from such therapies leads to greater absolute risk reduction and therefore increased benefit.

Based on this, the AHA/ACC Multisociety Guideline defines a secondary prevention population that is most likely to benefit from addition of non-statin therapy and further reduction in LDL-C levels. This includes individuals with a history of multiple major ASCVD events or 1 major ASCVD event plus multiple high-risk conditions. According to the AHA/ACC Multisociety Guideline, for these very high-risk individuals, it is reasonable to consider adding ezetimibe to maximally tolerated statin therapy when the LDL-C level remains ≥70 mg/dL; it is also reasonable to consider adding a PCSK9i to maximally tolerated statin therapy and ezetimibe when the LDL-C level remains ≥70 mg/dL or non-HDL-C level remains ≥100 mg/dL.

The ESC/EAS Guideline broadly extends the addition of non-statin therapy to a much larger group – primary prevention patients with moderate atherosclerosis on imaging and patients with stable ASCVD. The most relevant data documenting the benefit of intensifying LDL-C lowering therapy beyond statin therapy comes from the IMPROVE-IT, Further Cardiovascular Outcomes Research With PCSK9 Inhibition in Subjects with Elevated Risk (FOURIER), and Evaluation of Cardiovascular Outcomes after an Acute Coronary Syndrome During Treatment with Alirocumab (ODYSSEY OUTCOMES) trials – studies which specifically enrolled patients with clinical ASCVD and prior events. Acknowledging that these studies provide support that ‘lower is better’, the degree to which LDL-C should be lowered is of key importance.

In IMPROVE-IT, which included a high-risk secondary prevention population with a recent ACS, addition of ezetimibe (compared to placebo) was associated with a LDL-C level difference of 24% (LDL-C 53.2 mg/dL vs. 69.9 mg/dL) at 1 year and an ARR of 2.0% (32.7% vs. 34.7%) at 7 years. The between group difference in LDL-C levels (12.8 mg/dL) and proportional reduction in rate of major vascular events (7.2%) was also consistent with the reduction that occurs with statin therapy.

If this population was further risk stratified, those with three or more high-risk features had a much higher recurrent event rate (40% vs. 14% for 0–1 high-risk features) and greater risk reduction with the addition of ezetimibe (6.3% vs. 2.2% ARR in patients with two high-risk features) [21]. This data suggest a clear benefit of ezetimibe in a very high-risk secondary prevention population, where the reduction in LDL-C is directly proportional to the relative risk reduction in ASCVD events [26]. However, once LDL-C levels are below 70 mg/dL, the ARR from LDL-C lowering is less significant. In fact, the addition of ezetimibe did not result in a statistically significant graded decrease in the hazard ratio for the primary outcome with lower LDL-C strata (<30 mg/dl or 30–49 mg/dl) once below 70 mg/dl (50–69 mg/dl) [27].

In the FOURIER trial, patients with established ASCVD on statin therapy with a LDL-C ≥70 mg/dL were randomized to evolocumab or placebo [17]. The primary outcome of incident cardiovascular death, MI, stroke, hospitalization for unstable angina or coronary revascularization occurred in 12.6% in the evolocumab versus 14.6% in the placebo group over a mean of 2.2 years. While LDL-C levels were reduced on average by 56 mg/dl in those on evolocumab, with an on-treatment mean LDL-C level of 30 mg/dL (down from a baseline median LDL-C level of 92 mg/dL), the magnitude of ARR was more pronounced among individuals with an MI within 2 years (3.4%), ≥2 MIs (3.7%), or residual multivessel coronary artery disease (3.6%) [28].

While a prespecified secondary analysis of the FOURIER trial also demonstrated a linear association with the rate of primary and secondary endpoints and LDL-C levels to very low values, once LDL-C levels reached below 70 mg/dL, the ARR significantly diminished [29]. In IMPROVE-IT vs. FOURIER, while the ARR of non-statin add-on therapy extends to all secondary prevention patients, there is a significant increase in the ARR when LDL-C levels are >70 mg/dL as the baseline risk of the individual increases. This highlights that the benefit of LDL-C lowering diminishes with lower LDL-C levels – a 50% relative reduction in LDL-C at lower baseline LDL-C levels of 60 mg/dL and 80 mg/dL leads to 30 and 40 mg/dL absolute reductions from baseline [30]. Therefore, adding non-statin therapy at lower LDL-C levels results in less LDL-C reduction and lower relative risk reduction for ASCVD events [31].

In the ODYSSEY OUTCOMES trial, patients with an ACS within the preceding 1–12 months on statin therapy and an LDL-C ≥70 mg/dL were randomized to alirocumab or placebo [32]. The primary endpoint, which was a composite of death from coronary heart disease, nonfatal MI, fatal or nonfatal ischemic stroke, or unstable angina requiring hospitalization, occurred in 9.5% in the alirocumab versus 11.1% in the placebo group over a median of 2.8 years.

In a secondary analysis, the population was classified as very high-risk and non-very high-risk according to the AHA/ACC Multisociety Guideline. Very high-risk participants, despite an incidence rate of MACE per 100 patient-years more than double that of non-very high-risk individuals, benefited most from alirocumab with an ARR of 2.1% [33].

Consistent with the AHA/ACC Multisociety Guideline, these very high-risk individuals represent those most likely to benefit from addition of non-statin therapy with a PCSK9i to achieve an LDL-C level <70 mg/dL. Acknowledging that adherence to an appropriate intensity of statin therapy is the first step in lipid lowering, statin therapy alone is often insufficient to lower LDL-C levels and ASCVD risk enough in those at very high-risk.

In a simulated analysis of the SWEDHEART registry, it was estimated that half of MI patients would require PCSK9i therapy after maximizing statin intensity and adding ezetimibe to achieve a LDL-C level <55 mg/dL [34]. In the Pravastatin or Atorvastatin Evaluation and Infection Therapy-Thrombolysis in Myocardial Infarction (PROVE IT-TIMI 22) study, ~55% and ~75% of all individuals LDL-C levels would be above the AHA/ACC (≥70 mg/dL) and ESC/EAS (≥55 mg/dL) very high-risk LDL-C thresholds, respectively, with high-intensity statin therapy alone [35].

In a more modern data set from the Veterans Affairs Health care system, 1,038,903 individuals with ASCVD were identified, of which 43% met criteria for very high-risk ASCVD [36]. Despite use of statin therapy in 82% of these very high-risk individuals, 67% of them had an LDL-C level >70 mg/dL. Even after titrating to a high-intensity statin, ~37% of the very high-risk individuals were found to have an LDL-C level ≥70 mg/dL. After the addition of ezetimibe to high-intensity statin, 24% of these individuals still had a LDL-C level ≥70 mg/dL, and therefore were candidates for a PCSK9i.

Data from IMPROVE-IT, FOURIER and ODYSSEY OUTCOMES support an additional ARR when individuals LDL-C levels fall below 70 mg/
Enrichment of this population with those at highest risk, though, associated with PCSK9i therapy, all high-risk individuals should first be titrated to a high-intensity statin with strong emphasis on adherence [38, 39], which in doing so will also reduce clinical inertia [40]. Then individuals with the highest risk of recurrent ASCVD events among those with clinical ASCVD should be considered for additional non-statin therapies if LDL-C levels are >70 mg/dL, as recommended in the AHA/ACC Multisociety Guideline. This approach identifies a subgroup likely to derive the greatest benefit and reduces the NNT and economic challenges of achieving the ESC/EAS guideline recommendations [41].

2. Conclusions

Both guidelines incorporate advances from recent RCTs. The 2018 AHA/ACC Multisociety Guideline adheres closely to trial evidence and strongly considers both the added efficacy and cost implications of broadly reducing LDL-C levels in all high-risk individuals. This provides the basis for its more conservative recommendations pertaining to the highest risk subgroup where addition of non-statin therapy should be considered. In contrast, the 2019 ESC/EAS Guideline focuses primarily on trial data demonstrating that lower LDL-C levels resulted in lower recurrent ASCVD event rates, without strongly weighing the additive benefit and financial cost. Given current challenges, where issues with cost and access to some non-statin therapies exist, the ESC/EAS guideline provides recommendations that may be difficult to attain both in the US (if implemented) and in Europe, particularly in a context of limited healthcare resources. This provides support for the US guideline that selectively recommends non-statin therapy use in high- and, in particular, very high-risk individuals optimized on statin therapy. Future studies that incorporate cost-effectiveness in addition to efficacy and safety may help answer questions that could bring the guidelines closer together.

Declaration of competing interest

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