Combination therapy of statin and ezetimibe for the treatment of familial hypercholesterolemia

Ian Hamilton-Craig 1  
Karam Kostner 2  
David Colquhoun 2  
Stan Woodhouse 2  
1Griffith University School of Medicine, Southport, Queensland, Australia; 2University of Queensland, Brisbane, Queensland, Australia

Abstract: High-dose potent statin therapy in combination with ezetimibe is now standard practice for the treatment of adult patients with heterozygous familial hypercholesterolemia (heFH), as the result of numerous studies in patients with primary hypercholesterolemia or heFH. These studies have shown the combination to be both effective and safe in the short to medium term. Recently, short-term ezetimibe therapy has also been shown to be effective and safe in combination with statin therapy for children and adolescents with heFH. Effective statin–ezetimibe combination therapy is capable of achieving near-normal lipid profiles in heFH patients, with expected improvement in risk for cardiovascular disease (CVD) and improved life expectancy resulting predominantly from reduction in levels of low-density lipoprotein cholesterol. There are few data to support a pleiotropic action of ezetimibe with regard to CVD benefit, unlike therapy with statins. No serious and unexpected clinical adverse effects of combination statin–ezetimibe therapy have emerged till date, although data are limited in children and adolescents, for whom longer-term studies are required. Recent data suggesting possible proatherogenic effects of ezetimibe require confirmation. One large long-term randomized controlled clinical outcomes trial is in progress in non-FH patients to determine the efficacy and safety of ezetimibe therapy; it is unlikely that such a trial will ever be performed in patients with FH.

Keywords: familial hypercholesterolemia, ezetimibe, statin, combination therapy, low-density lipoprotein cholesterol

Introduction

Ezetimibe, the first specific inhibitor of the intestinal cholesterol uptake transporter Niemann–Pick C1 Like 1 (NPC1 L1) protein, was developed as an agent to lower plasma levels of low-density lipoprotein cholesterol (LDL-C).1

Statins are first-line drugs for treatment of elevated LDL-C levels, whereas ezetimibe remains one of the available second-line drugs for use in patients whose LDL-C levels remain above target in spite of maximally tolerated doses of statins, or for those who are unable to tolerate statins.2 Alternative therapy to lower LDL-C levels include fibrates, niacin (nicotinic acid [NA]), resins (bile acid sequesterants), and plant stanols and sterols.3,4

Ezetimibe has an additive and at times synergistic effect on the reduction of LDL-C and total cholesterol (TC) concentrations when combined with statin therapy. Thus, although doubling the dose of statin therapy and switching to an alternative statin generally lead to a further reduction in baseline LDL-C concentrations of approximately 6% and 8%, respectively, the addition of ezetimibe to statin therapy is likely to lead to...
greater incremental reductions in LDL-C concentrations of 15%–20% or more. The variation in response between individuals may in part be due to genetic variation. The intuitive hypothesis for improved efficacy of ezetimibe in individuals with high cholesterol absorption and low hepatic synthesis versus improved efficacy of statins in individuals with low absorption and high hepatic synthesis was not supported by the results of a recent study. Responsiveness to statin and ezetimibe were highly correlated; suggesting that factors downstream of the primary sites of action are major determinants of response.

In this article, we present an overview of the results of studies of ezetimibe in patients with primary hypercholesterolemia (HC) and make recommendations for the use of ezetimibe in patients with familial hypercholesterolemia (FH). This has also been the subject of two recent reviews.

**Familial hypercholesterolemia**

FH is an autosomal dominant genetic disorder due to a mutation in the gene coding for the LDL receptor (LDL-R). Heterozygous FH (heFH) results in functionally half of the normal number of hepatic LDL-Rs. As a consequence, decreased uptake of LDL-C from the blood occurs. This increases the activity of hepatic β-hydroxy-β-methylglutaryl-CoA (HMG-CoA) reductase, the rate-limiting enzyme, in the pathway of cholesterol synthesis. Increased production of cholesterol by hepatic cells, coupled with reduced uptake of LDL by its hepatic receptor result in elevated plasma LDL-C levels 2–3 times the normal in individuals with one abnormal allele (heFH). Those with two abnormal alleles, which may be either compound heFH or true homozygous FH (hoFH) with identical mutant alleles, have grossly elevated plasma LDL-C levels 4–6 times the normal. The elevation in plasma LDL-C levels leads to the development of early and aggressive atherosclerosis and premature atherothrombotic vascular disease. This includes coronary artery disease (CAD), cerebrovascular attacks or stroke, transient ischemic attacks, and peripheral arterial disease.

The incidence of heFH in the community is approximately 1 in 300 to 1 in 500 and that of hoFH approximately 1 per million. In the absence of genetic screening, only 10% of those affected are identified before the onset of symptomatic disease. One reason for this is that the majority of affected individuals may not have tendon or cutaneous xanthomas, the presence of which would normally bring them into early medical management.

The key to diagnosis of FH is measurement of serum TC levels, often routinely performed in automated biochemistry laboratories. A TC level >8 mmol/L with triglyceride (TG) level <2 mmol/L should alert the practitioner to the probable diagnosis of HeFH and the need for treatment and follow-up as a family. A good family history taken by the primary practitioner may help to identify FH through a history of premature CVD or sudden death on one side of the family, usually affecting males at a younger age than females. Linking this with follow-up of families of those presenting with early CVD should allow for cascade screening of cholesterol levels in near relatives and, where available, genetic screening with DNA analysis.

A consequence of the potential for early disease and death has provoked an attempt at early childhood diagnosis and early treatment with statins. Several community-wide and often nationwide programs have been instituted, including MEDPED (Make Early Diagnosis, Prevent early Death), begun in Utah by the late Professor Roger Williams. Extensive screening programs are now being conducted in the Netherlands and Scandinavia, and less ambitious programs have been initiated elsewhere. Family history taking and follow-up are often difficult to accomplish because multiple health practitioners are involved and the family group is constantly changing due to births, deaths, and possible intermarriage. A continuous process is required, which needs funding by a central government agency.

Clinical management of FH patients is primarily with the use of statins (HMG-CoA reductase inhibitors). In the limited studies which have been carried out, affected heFH persons, even though they may have reached presently accepted target levels of LDL-C, may have an additional residual risk. This may be related to cumulative LDL exposure before beginning treatment, elevated lipoprotein(a) levels, a need to reduce target levels further, as yet unknown factors, or the so-called legacy effect caused by delay in treatment. It may also be due to a longer exposure to elevated LDL-C levels from birth, in contrast to hypercholesterolemic individuals without FH, whose raised LDL-C levels usually occur after puberty. The presence of other metabolic factors such as obesity, diabetes, hypertension, and cigarette smoking will also impact on residual risk.

The management of FH also focuses on the need to achieve general lifestyle changes involving diet modification, abstinence from tobacco, weight reduction, and the undertaking of regular exercise. Identification of those at special risk includes patients with renal disease, albuminuria, depression and schizophrenia and particular ethnic groups including indigenous people, South Sea Islanders, and South
Ezetimibe with statins in familial hypercholesterolemia

Table 1 Primary hypercholesterolemia: change from baseline in calculated plasma LDL-C for ezetimibe alone and combined with statins or placebo²,

<table>
<thead>
<tr>
<th></th>
<th>AV</th>
<th>SV</th>
<th>PV</th>
<th>LV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Placebo</td>
<td>0.20 (+4%)</td>
<td>−0.08 (−1%)</td>
<td>−0.03 (−1%)</td>
<td>0.00 (0%)</td>
</tr>
<tr>
<td>E</td>
<td>−0.92 (−20%)</td>
<td>−0.92 (−19%)</td>
<td>−0.91 (−20%)</td>
<td>−0.86 (−19%)</td>
</tr>
<tr>
<td>10 mg S</td>
<td>−0.16 (−37%)</td>
<td>−1.25 (−27%)</td>
<td>−0.96 (−21%)</td>
<td>−0.94 (−20%)</td>
</tr>
<tr>
<td>E + 10 mg S</td>
<td>−2.46 (−35%)</td>
<td>−2.10 (−46%)</td>
<td>−1.55 (−34%)</td>
<td>−1.56 (−34%)</td>
</tr>
<tr>
<td>20 mg S</td>
<td>−1.91 (−42%)</td>
<td>−1.74 (−36%)</td>
<td>−1.10 (−23%)</td>
<td>−1.18 (−26%)</td>
</tr>
<tr>
<td>E + 20 mg S</td>
<td>−2.59 (−54%)</td>
<td>−2.16 (−46%)</td>
<td>−1.82 (−40%)</td>
<td>−1.87 (−41%)</td>
</tr>
<tr>
<td>40 mg S</td>
<td>−2.09 (−45%)</td>
<td>−1.75 (−38%)</td>
<td>−1.43 (−31%)</td>
<td>−1.44 (−30%)</td>
</tr>
<tr>
<td>E + 40 mg S</td>
<td>−2.69 (−56%)</td>
<td>−2.55 (−56%)</td>
<td>−1.97 (−42%)</td>
<td>−2.15 (−46%)</td>
</tr>
<tr>
<td>80 mg S</td>
<td>−2.57 (−54%)</td>
<td>−2.11 (−45%)</td>
<td>−1.21 (−25%)</td>
<td>−1.86 (−40%)</td>
</tr>
<tr>
<td>E + 80 mg S</td>
<td>−2.93 (−61%)</td>
<td>−2.64 (−58%)</td>
<td>−1.76 (−36%)</td>
<td>−2.19 (−54%)</td>
</tr>
<tr>
<td>Pooled data: All S</td>
<td>−2.08 (−44%)</td>
<td>−1.71 (−36%)</td>
<td>−1.16 (−25%)</td>
<td>−1.19 (−25%)</td>
</tr>
<tr>
<td>Pooled data: All E + S</td>
<td>−2.67 (−56%)</td>
<td>−2.36 (−51%)</td>
<td>−1.78 (−39%)</td>
<td>−1.86 (−40%)</td>
</tr>
</tbody>
</table>

Note: Values represent mean absolute change (in mmol/L) from baseline, and values in parenthesis represent mean percent change from baseline.

Abbreviations: AV, atorvastatin; SV, simvastatin; PV, pravastatin; LV, lovastatin; E, ezetimibe; S, statin.
compared with statin–ezetimibe combination therapy at various doses of statins.

Table 2 shows the results for a pooled analysis of all ezetimibe plus statin doses for changes from baseline in TC, apolipoprotein B-100 (apoB), TG, and high-density lipoprotein cholesterol (HDL-C) levels.1

Table 3 shows the response to addition of ezetimibe to on-going statin therapy in patients with primary HC. Percentages of patients receiving each statin are as follows: 40% atorvastatin, 31% simvastatin, and 29% others (pravastatin, fluvastatin, cerivastatin, and lovastatin).1

In summary, the above studies in patients with primary HC (predominantly non-FH) showed that ezetimibe monotherapy or ezetimibe combined with a statin significantly reduces TC, LDL-C, apoB, and TG and increases HDL-C levels compared with placebo. Reduction in LDL-C is consistent across age, sex, race, and baseline LDL-C. In addition, ezetimibe has no effect on the plasma concentrations of the fat-soluble vitamins A, D, and E and on the prothrombin time, and it does not impair adrenocortical steroid hormone production.1

The UK National Institute for Health and Clinical Excellence (NICE) guidance recently reviewed the results of several randomized controlled clinical trials with ezetimibe in patients with primary HC, with and without prior CVD.7 Average baseline LDL-C concentrations ranged from 3.4 to 6.5 mmol/L. Thirteen trials met the criteria of their review, and all were considered to be well designed and conducted. They varied in duration from 12 to 48 weeks. No studies reported health-related quality of life or clinical end-points such as CVD morbidity and mortality. Levels of TC, LDL-C, HDL-C, and TG were used as indicators of outcomes. No information was available on pretrial treatment history.

To represent the population of people with HC that is not appropriately controlled with statin therapy, six 12-week, fixed-dose randomized controlled trials (RCTs, n = 3,610) were identified that compared ezetimibe plus statin therapy with statin therapy alone.7 The NICE Assessment Group carried out a meta-analysis on the RCTs. Ezetimibe plus statin therapy was associated with an additional mean reduction in TC and LDL-C levels of

![Figure 1](image-url)
Panel (NCeP-ATP iii) goals1 cardiovascular risk factors on statin monotherapy who had not achieved National Cholesterol education Program Adult Treatment therapy is considered inappropriate or is not tolerated.7 All concentrations by 23.2% more than statin therapy alone. The results showed that addition of ezetimibe to statin therapy reduced LDL-C from baseline.

Table 2 Pooled analysis of absolute and percent change from baseline in total cholesterol (TC), apolipoprotein B (apoB), triglyceride (TG), and high-density lipoprotein cholesterol (HDL-C) for ezetimibe therapy in combination with various statin doses.1 Data from four multicenter double-blind randomized controlled trials of 12-week duration in patients with hyperlipidemia1

<table>
<thead>
<tr>
<th>Treatment</th>
<th>TC</th>
<th>ApoB</th>
<th>TG</th>
<th>HDL-C</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td>E + AV</td>
<td>−2.86 (−41%)</td>
<td>−0.78 (−45%)</td>
<td>−0.55 (−33%)</td>
<td>0.09 (+7%)</td>
<td>255</td>
</tr>
<tr>
<td>AV alone</td>
<td>−2.24 (−32%)</td>
<td>−0.61 (−36%)</td>
<td>−0.40 (−24%)</td>
<td>0.05 (+4%)</td>
<td>248</td>
</tr>
<tr>
<td>E + SV</td>
<td>−2.49 (−37%)</td>
<td>−0.69 (−41%)</td>
<td>−0.53 (−29%)</td>
<td>0.11 (+9%)</td>
<td>274</td>
</tr>
<tr>
<td>SV alone</td>
<td>−1.78 (−26%)</td>
<td>−0.51 (−30%)</td>
<td>−0.32 (−20%)</td>
<td>0.09 (+7%)</td>
<td>263</td>
</tr>
<tr>
<td>E + PV</td>
<td>−1.86 (−27%)</td>
<td>−0.51 (−30%)</td>
<td>−0.36 (−21%)</td>
<td>0.10 (+8%)</td>
<td>204</td>
</tr>
<tr>
<td>PV alone</td>
<td>−1.17 (−17%)</td>
<td>−0.35 (−20%)</td>
<td>−0.26 (−14%)</td>
<td>0.08 (+7%)</td>
<td>205</td>
</tr>
<tr>
<td>E + LV</td>
<td>−1.96 (−29%)</td>
<td>−0.57 (−33%)</td>
<td>−0.44 (−25%)</td>
<td>0.10 (+9%)</td>
<td>192</td>
</tr>
<tr>
<td>LV alone</td>
<td>−1.25 (−18%)</td>
<td>−0.36 (−21%)</td>
<td>0.21 (−12%)</td>
<td>0.04 (+4%)</td>
<td>220</td>
</tr>
</tbody>
</table>

Note: Values represent mean absolute change from baseline, mmol/L for lipid levels and mg/dL for apoB levels, and values in parenthesis represent mean percent change from baseline.

Abbreviations: E, ezetimibe; AV, atorvastatin; SV, simvastatin; PV, pravastatin; LV, lovastatin.

10.4% (95% confidence interval [CI]: 11.1–9.6) and 13.9% (95% CI: 14.9–13.0), respectively, for prestatin treatment concentrations compared with statin therapy alone. This equated to a 22.4% reduction achieved by the combination of ezetimibe plus statin compared with an on-statin baseline LDL-C level.

Four extension studies (n = 1,800) compared ezetimibe plus statin therapy with a titrated statin dose.7 One study included an heFH subgroup; in the ezetimibe plus statin arm, 17% reached the LDL-C target (2.6 mmol/L or less) compared with 4% in the statin monotherapy arm.

The NICE Assessment Group carried out an additional meta-analysis of shorter-term studies (less than 12 weeks in duration) comparing ezetimibe coadministered with statin therapy vs statin therapy alone.7 The results showed that the addition of ezetimibe to statin therapy reduced LDL-C concentrations by 23.2% more than statin therapy alone.

Seven RCTs (n = 2,577) comparing ezetimibe monotherapy with placebo represented the population in which statin therapy is considered inappropriate or is not tolerated.7 All were 12-week studies and were included in a meta-analysis performed by the NICE Assessment Group. Ezetimibe monotherapy was associated with a statistically significant mean reduction in TC concentrations (13.4%; 95% CI: 14.2–12.6) and LDL-C concentrations (18.6%; 95% CI: 19.7–17.4) compared with placebo.

Four studies demonstrated LDL-C-lowering effects of ezetimibe treatment across subgroups, including different ethnic groups and people with or without conditions such as CVD, diabetes, and heFH.7 None of the subgroup comparisons showed statistically significant differences between subgroups. All other trials reported that the effect of ezetimibe therapy on LDL-C levels was generally consistent across all subgroups. There was no evidence to suggest a difference in the effectiveness of ezetimibe in any subgroup, including people with heFH or diabetes, or people with or without a history of CVD.7

The NICE guidance meta-analysis showed that ezetimibe plus statin therapy reduces LDL-C levels by an additional 13.9% compared with statin therapy alone.7 This absolute change was approximately 22% when calculated as a proportion of the poststatin LDL-C levels.

The clinical effectiveness of ezetimibe, based on its mode of action, is unlikely to differ markedly between different ethnic groups; therefore, separate recommendations for different ethnic groups were not made in the NICE guidelines.7

Table 3 Response to addition of ezetimibe to on-going statin therapy in patients with primary hypercholesterolemia (HC): absolute and percent changes from baseline.1 Data from 8-week trials of patients with primary HC, known coronary heart disease of multiple cardiovascular risk factors on statin monotherapy who had not achieved National Cholesterol Education Program Adult Treatment Panel (NCEP-ATP III) goals1

<table>
<thead>
<tr>
<th>Treatment</th>
<th>TC</th>
<th>LDL-C</th>
<th>ApoB</th>
<th>TG</th>
<th>HDL-C</th>
</tr>
</thead>
<tbody>
<tr>
<td>S + PV</td>
<td>−0.16 (−2%)</td>
<td>−0.16 (−4%)</td>
<td>0.05 (−3%)</td>
<td>0.05 (−3%)</td>
<td>+0.00 (+1%)</td>
</tr>
<tr>
<td>S + E</td>
<td>−0.99 (−17%)</td>
<td>−0.92 (−25%)</td>
<td>−0.27 (−19%)</td>
<td>−0.19 (−14%)</td>
<td>+0.03 (+3%)</td>
</tr>
<tr>
<td>E − PV</td>
<td>−0.83 (−15%)</td>
<td>−0.76 (−21%)</td>
<td>−0.22 (−16%)</td>
<td>−0.14 (−11%)</td>
<td>+0.03 (+2%)</td>
</tr>
</tbody>
</table>

Note: Values represent mean absolute change (in mmol/L) from baseline, and values in parenthesis represent mean percent change from baseline.

Abbreviations: TC, total cholesterol; LDL-C, low-density lipoprotein cholesterol; ApoB, apolipoprotein B; TG, triglyceride; HDL-C, high-density lipoprotein cholesterol; S, statin; PV, pravastatin; E, ezetimibe.
Ezetimibe in FH

The initial, multicenter, double-blind, 14-week study of 621 hypercholesterolemia patients included approximately 60% of patients with heFH. Those receiving atorvastatin 10 mg daily with an LDL-C >3.36 mmol/L were randomized to receive atorvastatin 20 mg or ezetimibe 10 mg added to atorvastatin 10 mg therapy. The atorvastatin dose could be titrated up to 80 mg in the atorvastatin arm and up to 40 mg in the ezetimibe plus atorvastatin coadministration arm, based on patients not attaining LDL-C goal (<2.59 mmol/L). The mean baseline LDL-C was 4.84 mmol/L. At study end, there was a significant difference in attainment of LDL-C goal between patients in the ezetimibe coadministration arm (22%) and patients on atorvastatin monotherapy (7%). At week 4, there was a significant difference in LDL-C reductions between coadministration patients (24%; ezetimibe plus atorvastatin 10 mg) and monotherapy patients (9%; atorvastatin 20 mg). In the subgroup of patients with heFH, similar results for LDL-C goal attainment and LDL-C reductions were achieved.

Several subsequent studies confirmed these results in heFH patients. The NICE Assessment Group carried out an additional subgroup analysis of the effect of ezetimibe therapy in people with or without heFH. The greater reductions in LDL and TC concentrations in the heFH group were not found to be statistically significant.7

Ezetimibe in hoFH

The initial study was a double-blind, randomized, 12-week study of 50 patients, with a clinical and/or genotypic diagnosis of hoFH, with or without concomitant LDL apheresis, who were already receiving atorvastatin or simvastatin (40 mg). Patients were randomized to 1 of 3 treatment groups, atorvastatin or simvastatin (80 mg), ezetimibe 10 mg administered with atorvastatin or simvastatin (40 mg), or ezetimibe 10 mg administered with atorvastatin or simvastatin (80 mg). Results are shown in Table 4.1 Ezetimibe administered with atorvastatin (40 or 80 mg) or simvastatin (40 or 80 mg) significantly reduced LDL-C compared with increasing the dose of simvastatin or atorvastatin monotherapy from 40 to 80 mg.

Several subsequent studies have confirmed the efficacy of ezetimibe in hoFH.45–52

Ezetimibe in children with FH

A small number of short-term studies have investigated the efficacy and tolerability of ezetimibe in children with FH.53–56

In general, ezetimibe was well tolerated with similar results to those observed in adult subjects with FH. No long-term data are available.

Ezetimibe in addition to other therapies in FH

Isolated reports have been published on the use of ezetimibe in addition to LDL-apheresis in patients with resistant FH. Further LDL-C reductions of 11%–25% have been observed with no evident adverse effects.

Table 4 Mean LDL-C response to ezetimibe in patients with homozygous familial hypercholesterolemia1

<table>
<thead>
<tr>
<th>Treatment (daily dose)</th>
<th>N</th>
<th>LDL-C (mmol/L)</th>
<th>Changea</th>
<th>% Changeb</th>
</tr>
</thead>
<tbody>
<tr>
<td>AV or SV (80 mg)</td>
<td>17</td>
<td>−0.51</td>
<td>−7%</td>
<td></td>
</tr>
<tr>
<td>E + AV or SV (40, 80 mg)</td>
<td>33</td>
<td>−1.76</td>
<td>−21%</td>
<td></td>
</tr>
<tr>
<td>E + AV or SV (80 mg)</td>
<td>17</td>
<td>−2.00</td>
<td>−27%</td>
<td></td>
</tr>
</tbody>
</table>

Notes: aMean absolute change from baseline (mmol/L). bMean percent change from baseline.

Abbreviations: AV, atorvastatin; SV, simvastatin; E, ezetimibe.

Ezetimibe in human immunodeficiency virus infection

Many of the protease inhibitors used in the treatment of human immunodeficiency virus (HIV) infection will induce mixed forms of dyslipidemias, and in FH, patients have the potential for further deterioration. Studies in children, however, show the changes were of the same degree as non-FH patients.58

Antiviral drugs and HIV protease inhibitors have an increased risk of myopathy when used with statins, and so they should be used with care. Pravastatin is least likely to be a problem, as it is not substantially metabolized by cytochrome P450 in the liver. Simvastatin should be avoided.59 Rosuvastatin has some promise, as it is excreted 90% intact in the feces and so it would not have a great effect on the P450 enzymes to induce dangerous levels of the drug. However, small trials have shown increases in serum rosuvastatin varying from 61% to 76%, depending on the antiviral drug used.60 Although no clinical problems were encountered in these small short-term trials, there is clearly need for care. When any statin is used patients must be alerted to watch for muscle symptoms such as muscle pain, stiffness, weakness, or cramps.

Ezetimibe therapy may play an important role in reducing the dose of statin required, thus reducing adverse side effects. Hence, it may be considered for use when target levels of

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LDL-C have not been reached and side effects or risk of side effects are present. The studies that have been conducted using ezetimibe in conjunction with pravastatin in the presence of the protease inhibitors are of limited usefulness because of the small number of patients. These have shown only modest decrease in LDL-C levels (22%) and small increase in HDL-C levels but with no new or enhanced side effects noted. Ezetimibe, therefore, seems to be a safe drug when used in this context. A Canadian study is presently underway in which rosuvastatin 10 mg plus ezetimibe 10 mg is compared with rosuvastatin 20 mg in HIV patients being treated with protease inhibitors.

**Antiatherosclerotic and pleiotropic effects of ezetimibe**

Animal experiments have shown antiatherosclerotic effects with ezetimibe therapy, in part possibly mediated by nonlipid (pleiotropic) mechanisms, the significance of which is controversial. In apoe knockout (ko) mice, aortic lesion formation was significantly reduced by ezetimibe therapy. Ezetimibe treatment resulted in a significant reduction in plaque size and macrophage and fibronectin extra domain-B immunoreactivity in brachiocephalic lesions, indicating plaque regression. Similar results were also shown with high-intensity (7 T) magnetic resonance imaging.

In rabbits, femoral atherosclerosis was induced by a combination of endothelial desiccation and atherogenic diet. Ezetimibe treatment reduced the intima/media ratio by 13%, simvastatin therapy by 27%, and ezetimibe plus simvastatin therapy by 28% compared with control rabbits. Ezetimibe decreased macrophage content and monocyte chemoattractant protein 1 (MCP-1) expression in atherosclerotic lesions and reduced the increased activity of nuclear factor-κB in peripheral blood leucocytes and plasma CRP levels. In THP-1 cells, ezetimibe decreased MCP-1-induced monocyte migration. The combination of ezetimibe with simvastatin was associated with a more significant reduction in plaque monocyte/macrophage content and some proinflammatory markers than observed with either drug alone.

In another study, apoe ko and apoe/endothelial nitric oxide synthase (eNOS) double ko (dko) mice received a high-fat diet with or without 0.05% ezetimibe. Ezetimibe therapy significantly reduced plasma cholesterol concentrations and atherogenic lipoproteins in both genotypes to a similar extent. Moreover, the drug reduced vascular inflammation, as it significantly reduced vascular cell adhesion molecule 1 expression and vascular CD14 expression, a marker for mononuclear cell infiltration, in both genotypes. Neither NOS protein expression nor vascular reactivity of aortic rings was changed in apoE ko mice following ezetimibe treatment. Significant lesion reduction was seen in ezetimibe-treated male and female apoe ko and apoE/eNOS dko animals (P ≤ 0.05). The drug-mediated additional atheroprotection in male apoe ko mice compared with male eNOS dko mice suggests that lipid lowering does provide additional eNOS-dependent atheroprotection in this experimental group.

Yorkshire pigs treated with streptozotocin to induce diabetes mellitus (DM) were treated with either atorvastatin or ezetimibe and evaluated for the number of bone marrow and circulating endothelial progenitor cells (EPCs) and for femoral artery endothelial function. There was no effect of either medication on cholesterol level. One month after induction of DM prior to administration of drugs, the number of bone marrow and circulating EPCs significantly decreased (P < 0.0001) compared with baseline. Three months after DM induction, the mean proportion of circulating EPCs significantly increased in the atorvastatin group, but not in the control or ezetimibe groups. The control group showed progressive reduction in percentage of flow-mediated vasodilatation (no dilatation at 3 months), whereas the atorvastatin group and ezetimibe group exhibited 6% and 4% vasodilatation, respectively.

One study investigated endothelial function in 20 patients with heart failure treated with either simvastatin or simvastatin plus ezetimibe. Simvastatin and ezetimibe treatment reduced LDL-C to a similar extent (15.6% vs 15.4%; P = not significant [NS]), whereas changes in mevalonate, the product of HMG-CoA reductase, differed between groups (∆0.62 vs 1.04 ± 0.62 vs ∆mevalonate–simvastatin, 1.79 ± 0.94 ng/mL; P < 0.05 between groups). Flow-mediated dilatation (FMD) was markedly improved after simvastatin (10.5 ± 0.6% vs 5.1 ± 0.7%; P < 0.01) but not after ezetimibe treatment (5.6 ± 0.5% vs 5.8 ± 0.6%; P = NS). The FMD before and after intra-arterial infusion of vitamin C to determine the portion of FMD inhibited by radicals was substantially reduced after simvastatin but not after ezetimibe treatment. Extracellular superoxide dismutase activity was increased by >100% (P < 0.05) after simvastatin but not ezetimibe treatment. Simvastatin treatment increased the number of functionally active EPCs, whereas ezetimibe had no effect.

Similar results were observed in a study of forearm blood flow (FBF) responses to acetylcholine (ACH) and sodium nitroprusside, measured by venous occlusion plethysmography, in four prospectively defined groups of patients with stable CAD before and after 4 weeks of
lipid-lowering therapy. Ezetimibe 10 mg/d monotherapy (N = 15) was compared with long-term administration of simvastatin 20 mg/d plus add-on ezetimibe (N = 15). After 4 weeks of therapy, LDL-C levels were significantly reduced in both groups. Neither ezetimibe monotherapy nor ezetimibe combined with 20 mg simvastatin was associated with an increase in ACH-mediated FBF responses after 4 weeks. It was concluded that both statins and ezetimibe effectively lower LDL levels within 4 weeks of therapy, but only statin therapy is associated with improved endothelial vasodilator function.

Another study investigated synthesis of isoprenoids, which are important for mediating signalling through the Rho-associated coiled-coil containing protein kinase (ROCK) pathway. Increased ROCK activity has been implicated in endothelial dysfunction and vascular inflammation, and statins reduce isoprenoid synthesis and ROCK activity. Dyslipidemic subjects (N = 60) without cardiovascular disease (CVD) were randomized to treatment with simvastatin 40 mg/d, simvastatin/ezetimibe 10/10 mg/d, or placebo tablets for 28 days (n = 20 in each arm). Compared with the placebo group, both treatment regimens decreased LDL-C by 38% and CRP by 38%–40% after 28 days (P < 0.01 for both compared with placebo). Although the LDL-C and CRP reductions were comparable with either lipid-lowering regimen, only simvastatin 40 mg reduced ROCK activity and improved FMD (P < 0.01 for both compared with baseline). Reduction in ROCK activity with simvastatin 40 mg remained significant even after controlling for changes in LDL-C (P = 0.01) and correlated with improvement in FMD (R² = -0.78, P < 0.01). No correlation was found between changes in FMD and changes in LDL-C or CRP.

The effects of ezetimibe on hs-CRP were reported in a meta-analysis of 13 randomized placebo-controlled trials with ezetimibe and statin therapy. Six were monotherapy trials of 12-week duration (N = 1,372), and 7 involved 6–8 weeks of add-on ezetimibe to stable statin therapy (N = 3,899). A 6% additional reduction was observed comparing ezetimibe with placebo (P = 0.094). A 10.4% reduction was observed for ezetimibe and statin therapy, which was significantly greater than placebo (P < 0.001). Weak significant correlation was observed between baseline hs-CRP and LDL-C levels only in the ezetimibe add-on groups. It was concluded that the lowering of hs-CRP with statin therapy was enhanced by ezetimibe to a small degree, the clinical significance of which is uncertain.

A review of the published literature characterizing the impact of ezetimibe-containing lipid-lowering regimens on endothelial function and other markers of cardiovascular risk, and the potential relevance of these effects on the clinical benefit of ezetimibe, concluded that ezetimibe, either as monotherapy or in combination with a statin, exerts minimal beneficial effects on endothelial function and other ancillary measures of CVD risk beyond those conferred by its cholesterol-lowering effects. Recent studies in patients with CAD, heart failure, and hypercholesterolemia demonstrated that treatment with ezetimibe for 4–12 weeks elicits no improvement of endothelial function or other measures of CVD risk. In contrast, other studies have reported that ezetimibe improves endothelial function in certain patient populations, including those with rheumatoid arthritis, CAD with type 2 diabetes, and metabolic syndrome. However, the statin monotherapy comparator groups in these studies that yielded equivalent reductions in cholesterol were superior, or at least equivalent to, ezetimibe-containing regimens in the improvement of these ancillary end-points. It was suggested that studies with larger sample sizes and follow-up beyond 12 weeks were necessary to further define the impact of ezetimibe on the processes integral to the pathogenesis and progression of CVD.

**Outcomes of ezetimibe therapy**

**LDL-C level as a surrogate outcome**

The NICE guidance on ezetimibe for the treatment of primary HC considered the published evidence on the correlation between changes in lipid concentrations to reductions in CVD events, in which lowering LDL-C levels is associated with CVD outcome benefits independent of the treatment used. It was concluded there is sufficient evidence to link reductions in LDL-C levels from ezetimibe therapy with future reductions in CVD events. This has been the approach of regulatory bodies around the world, which have not required demonstration of benefit in clinical CVD outcomes in order for ezetimibe to be registered, although the validity of this approach is now being questioned.

**Carotid intima-media thickness as a surrogate outcome**

Limited studies of surrogate imaging outcomes for CVD have been performed with ezetimibe. The three trials (SANDS, ENHANCE, and ARBITER 6-HALTS) have investigated the effects of ezetimibe on carotid intima-media thickness (CIMT), a validated surrogate marker for CVD.
SANDS
The Stop Atherosclerosis in Native Diabetics Study (SANDS) investigated the effects of standard vs aggressive management of risk factors in 499 North American Indian men and women aged >40 with type 2 diabetes, hypertension, and dyslipidemia.75,76 Multiple risk factors (blood pressure [BP], glucose, and lipids) were targeted with stepped treatment algorithms. Aggressive therapy lowered LDL-C and systolic BP (SBP) levels to 1.7 mmol/L and 117 mm Hg compared with 2.7 mmol/L and 129 mm Hg, respectively, in the standard group. Ezetimibe was used more often in the aggressive treatment group in which regression of CIMT and greater reduction in left ventricular mass index were observed (see Table 5). However, ezetimibe use was not randomized. Baseline levels of CIMT in SANDS trial were increased (mean, 0.9 mm).

ENHANCE
The Effect of Combination Ezetimibe and High-Dose Simvastatin vs Simvastatin Alone on the Atherosclerotic Process in Patients with Heterozygous Familial Hypercholesterolemia (ENHANCE) trial investigated the effects of therapy on CIMT in patients with heFH.40,77–86

ENHANCE was an imaging study designed to examine change in CIMT in a select population of FH patients, with a mean LDL-C baseline 8.2 mmol/L (317 mg/dL) after a 6-week washout, treated with either maximum dose statin (simvastatin 80 mg/d) compared with ezetimibe 10 mg/d plus simvastatin 80 mg/d.40 There was no measurable CIMT impact by the addition of ezetimibe to simvastatin 80 mg/d.

Patients were followed over a 24-month period with measures of CIMT taken at 6-month intervals. The study found similar rates of CIMT progression in the simvastatin and the simvastatin–ezetimibe groups. Baseline mean CIMT levels were within the normal range (<0.7 mm).

Although no significant CIMT regression was seen in the selected population (N = 720), the majority (over 80%) had previously been treated for FH with statin therapy, and no adverse outcomes were seen. Results for the remaining 20% of statin-naïve patients were not published. Importantly, there was no control group treated with ezetimibe, which could be compared with the statin–ezetimibe group in order to determine the effects of ezetimibe therapy alone on CIMT progression.

The ENHANCE study results provided little insight to the benefit or nonbenefit of statin and ezetimibe combinations in this patient group because of problematic study design, especially with regard to patient selection; there were no prewashout criteria for either CIMT or LDL-C levels. This was a high-risk population that did not attain treatment targets, and furthermore, the prewashout lipid values and prestudy lipid-lowering treatments of subjects were not included in the data collected for the ENHANCE study.

Although this trial showed no benefit in reducing CIMT by the combination of simvastatin and ezetimibe, it did show the expected lowering of LDL-C and apoB levels. The ENHANCE trial demonstrated a 16% greater reduction in LDL-C in the combination group compared with the simvastatin-alone group over the 24-month period, as well as 18% greater reduction in hs-CRP.

The earlier Effects of Atorvastatin and Simvastatin on Atherosclerosis Progression (ASAP) study compared the effects of statin therapy on CIMT in patients with FH.39 There was significantly greater CIMT regression with atorvastatin 80 mg/d compared with simvastatin 80 mg/d. Inclusion criteria for ASAP included LDL-C > 4.5 mmol/L (>173 mg/dL) and CIMT > 0.7 mm. In ENHANCE, patients were required to have LDL-C > 5.4 mmol/L (210 mg/dL) after washout of prior therapy, without minimal requirement for CIMT levels. Baseline CIMT levels of ASAP and

Table 5 Results of The Stop Atherosclerosis in Native Diabetics Study (SANDS)39

<table>
<thead>
<tr>
<th>End-point</th>
<th>Aggressive therapy</th>
<th>Standard therapy</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline CIMT (mm)</td>
<td>0.81 [0.78, 0.83]</td>
<td>0.80 [0.78, 0.82]</td>
<td>NS</td>
</tr>
<tr>
<td>Mean CIMT change at 36 mo (mm)</td>
<td>−0.012 [-0.03, 0.003]</td>
<td>+0.038 [0.02, 0.06]</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Left ventricular mass index (g/mm²)</td>
<td>−2.4 [-3.2, −1.6]</td>
<td>−1.2 [−1.9, −0.4]</td>
<td>0.03</td>
</tr>
<tr>
<td>Carotid artery area (mm²)</td>
<td>−0.02 [-0.33, +0.30]</td>
<td>+1.05 [0.73, 1.38]</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Adverse events (%)c</td>
<td>38.5 [32, 45]</td>
<td>26.7 [21, 32]</td>
<td>0.005</td>
</tr>
<tr>
<td>Serious adverse eventsda</td>
<td>29.4 [24, 35]</td>
<td>22.3 [17, 28]</td>
<td>NS</td>
</tr>
</tbody>
</table>

Notes: aAggressive therapy was to achieve primary targets of the following: low-density lipoprotein cholesterol (LDL-C) ≤ 70 mg/dL (1.8 mmol/L), nonhigh-density lipoprotein cholesterol (non-HDL-C) < 100 mg/dL (2.6 mmol/L), and systolic blood pressure (SBP) ≤ 115 mm Hg. Figures in brackets refer to 95% confidence intervals. bStandard therapy was to achieve primary targets of the following: LDL-C ≤ 100 mg/dL (2.6 mmol/L), non-HDL-C ≤ 130 mg/dL (3.4 mmol/L), and SBP ≤ 130 mm Hg. cThe nature of these events was not described in the original publication other than excluding cardiovascular events. dNo serious adverse events were related to lipid drugs; four events were related to BP drugs in the aggressive group and 1 in the standard group (P = 0.18).

Abbreviations: CIMT, carotid intima-media thickness; NS, not significant.
ENHANCE were –0.9 and –0.7 mm, respectively. This may explain to a large extent the different outcomes of ASAP and ENHANCE because changes in response to therapy are correlated with baseline values of LDL-C and CIMT (greater changes in CIMT occur with higher baseline values). The failure in ENHANCE to reduce CIMT in FH patients with the same mean age and same dose of the same drug as ASAP may therefore be due to higher baseline CIMT in ASAP. In both ASAP and ENHANCE studies the biggest predictor of change in CIMT in response to statin therapy was baseline CIMT ($r = 0.53$, ENHANCE; $r = 0.41$, ASAP).

Since the 1990s, the authors of these two studies have identified most FH patients in the Netherlands; the percentage of patients treated with statins increased from 39% to 91% in 2001. In the same year, the ASAP publication stated that adult FH patients should be treated with statins. Recently, ezetimibe has been shown to increase the proportions of small, dense LDL-particles (sdLDL) in normal subjects after 2 weeks of therapy. Ezetimibe significantly increased the sdLDL subfractions LDL-IVA and LDL-IVB (+14.2% and +16.7%, respectively), whereas simvastatin significantly decreased the LDL-IVB subfraction (−16.7%). With simvastatin–ezetimibe combination therapy, the LDL-IVB subfraction was increased (+14.3%, NS). Each of the three treatments decreased the large LDL-I subfraction, especially ezetimibe (ezetimibe −13.9%, $P < 0.0001$; combination therapy −7.3%, $P = 0.0743$; simvastatin −4.6%, $P < 0.0001$). The significance of this finding remains unclear for several reasons. Firstly, other studies have shown contradictory results, and LDL functional assays are required to assess whether or not ezetimibe-induced sdLDL particles behave differently from normal LDL particles with regard to propensity for oxidation and uptake by macrophages, among other potential effects.

In the past, almost all studies of CIMT have indicated a close correlation between CIMT, atherosclerosis severity, and CVD risk, leading to the concept of CIMT as a valid surrogate marker. Furthermore, ezetimibe therapy has shown to result in atherosclerosis regression in experimental models. Further prospective studies are, therefore, required to provide more information on this potential adverse effect of ezetimibe therapy in relation to CIMT, atherosclerosis progression and CVD events. It is now appropriate to discuss long-term clinical outcomes studies with ezetimibe.

**Clinical outcomes – non-FH patients**

The possible adverse effects of ezetimibe were explained on the basis of mild inhibition of acyl coenzyme A: cholesterol acyltransferase, which may worsen atherosclerosis. Ezetimibe has also been shown to inhibit SR-BI, the hepatic HDL receptor, a mechanism that may inhibit reverse cholesterol transport and promote atherogenesis. Ezetimibe was shown to increase the proportion of small, dense LDL-particles (sdLDL) in normal subjects after 2 weeks of therapy. Ezetimibe significantly increased the sdLDL subfractions LDL-IVA and LDL-IVB (+14.2% and +16.7%, respectively), whereas simvastatin significantly decreased the LDL-IVB subfraction (−16.7%). With simvastatin–ezetimibe combination therapy, the LDL-IVB subfraction was increased (+14.3%, NS). Each of the three treatments decreased the large LDL-I subfraction, especially ezetimibe (ezetimibe −13.9%, $P < 0.0001$; combination therapy −7.3%, $P = 0.0743$; simvastatin −4.6%, $P < 0.0001$). The significance of this finding remains unclear for several reasons. Firstly, other studies have shown contradictory results, and LDL functional assays are required to assess whether or not ezetimibe-induced sdLDL particles behave differently from normal LDL particles with regard to propensity for oxidation and uptake by macrophages, among other potential effects.

**Clinical outcomes – non-FH patients**

To date, no long-term studies of CVD outcomes have been published with the exception of the Simvastatin and Ezetimibe in Aortic Stenosis (SEAS) trial. The SEAS trial was designed to investigate the efficacy of ezetimibe and simvastatin on the progression of aortic valve disease. Compared with placebo, LDL-C was reduced by 61% (2.0 mmol/L). There was no difference in the primary end-point (a combination of aortic valve replacement (AVR), CV death, nonfatal myocardial infarction, congestive heart failure from aortic stenosis progression, coronary revascularization, hospitalized unstable angina, and nonhemorrhagic stroke). Compared with placebo, ischemic
CVD events were reduced by 4.4% from 20.1% to 15.7% in the simvastatin/ezetimibe group (P = 0.02). The event reduction was largely driven by reduction in coronary artery bypass graft (CABG) surgery procedures that were performed at the same time as AVR.91 The results of SEAS suggest either a more favourable symptomatic outcome (less angina requiring CABG) or a reduction in coronary atherosclerosis severity in the ezetimibe and simvastatin group, for which less frequent CABG surgery was required.92 No data on these end-points have been published, however.

**IMPROVE-IT**

The Improved Reduction of Outcomes: Vytorin Efficacy International Trial (IMPROVE-IT), a large-scale clinical end-point trial is currently being conducted. This trial compares the effects of simvastatin vs simvastatin plus ezetimibe (Vytorin) on CVD end-points in approximately 18,000 patients with high cardiovascular risk.93 IMPROVE-IT is expected to be completed in June 2013.

**Clinical outcomes – FH patients**

To the authors’ knowledge, no long-term CVD outcome trials in patients with genetic HC treated with ezetimibe are being conducted or are planned. This may reflect the number of patients required for such a trial because it would require a comparator study in which both arms were treated with LDL-lowering therapy, as it would be unethical to use a placebo arm for FH patients.

At the time of writing, several short-term trials are either currently recruiting FH patients for ezetimibe therapy or have been completed. They include studies of chylomicron metabolism and effects of therapy in children, adolescents, Japanese, and Filipinos.94

**Safety and adverse effects**

There are limited long-term data on adverse events relative to the use of statins, but no significant adverse events of ezetimibe have emerged outside of the trials other than isolated reports of ezetimibe-associated musculoskeletal symptoms that are clinically similar to those with statin therapy, the mechanisms of which are speculative.95–97 In addition, there has been no evidence of any existing increase in all-cause mortality, or of specific mortality, as a result of treatment with ezetimibe.1

An increased incidence of cancer was reported in the SEAS trial in the group receiving ezetimibe compared with placebo.93 These results prompted interim analysis of two other long-term trials being conducted at the time – the IMPROVE-IT and Study of Heart and Renal Protection (SHARP) trials.98 No increase in cancer incidence was observed in these trials, and it was concluded that the adverse results of SEAS trial were due to chance.99 Similar conclusions were reached from postmarketing analysis.99

**Cost-effectiveness**

No data have been published for cost-effectiveness of ezetimibe in FH patients. In the NICE guidance, several other clinical scenarios were discussed, and models were analyzed.7 It was suggested that ezetimibe coadministered with a statin should not be recommended as an alternative to dose titration of the initiated statin where dose titration is possible and not prevented by the emergence of adverse effects.

The NICE Committee agreed that in non-FH patients, adding ezetimibe to initial statin therapy as a treatment option is a cost-effective use of National Health Service resources when compared with switching to an alternative statin.7 Ezetimibe therapy in FH subjects is likely to be more cost-effective because of the higher CVD risk of FH compared with non-FH populations.

**Summary and recommendations**

Physicians using ezetimibe should be familiar with the product information and refer to it for specific details of drug interaction, tolerability, and other details.1 Ezetimibe, a cholesterol absorption inhibitor, is indicated for the treatment of HC due to elevation of LDL-C. The rationale for its use in the treatment of HC is that it provides 10%–20% further reduction in LDL-C levels compared with that achieved with a given dose of statin or any other LDL-C-lowering therapy. Slightly higher percentage reductions are achieved with ezetimibe monotherapy compared with combination statin–ezetimibe therapy (see Figure 1). On the basis of many LDL-C-lowering trials, epidemiological studies, and animal experiments, there is a strong positive correlation between LDL-C levels, atherosclerosis severity, and incidence of atherothrombotic CVD. A reduction of LDL-C of 20% equates to a similar reduction in CVD events over a 5-year period. This has yet to be confirmed in a long-term randomized controlled intervention trial with ezetimibe, although such a trial is expected to be completed in 2013. Until then, most authorities and lipidologists recommend the use of ezetimibe to further lower LDL-C in patients whose LDL-C is not at target despite other LDL-C-lowering therapy, or in patients intolerant of such therapies.

FH is a monogenic, autosomal dominant disorder caused by a mutation in the gene coding for the LDL-R. In heFH,
the presence of one abnormal LDL-R allele is associated with approximately 50% loss of LDL-R activity and two fold increase in plasma levels of LDL-C. This leads to accelerated atherosclerosis and premature CVD in the aged (50% of men with heFH have CVD by the age of 50 years and 50% of women by the age of 60–65 years). Young men with heFH have about 80 times higher standardized mortality rate than the general population. In hoFH patients, the presence of two abnormal LDL-R alleles is associated with approximately 100% loss of LDL-R activity and several-fold increases in plasma levels of LDL-C. This leads to markedly accelerated atherosclerosis and premature CVD often in the teenage years (some children suffer from CVD before the age of 10 years).

The prevalence of hoFH is about 1 per million of the population (but slightly higher in populations with increased LDL-R mutation gene frequency), and often results from consanguinity. The prevalence of heFH is about 1 per 500 of the population (up to 1:60–1:80 in populations with increased LDL-R mutation gene frequency such as French Canadians, Southern Afrikaners, and Lebanese). A survival value for the presence of LDL-R mutations has yet to be ascertained.

FH is recognized clinically by the combination of high LDL-C level, positive family history of premature CVD on one side of the family (males and females being equally affected, although clinical CVD usually occurs 10–15 years later in females), and clinical signs of LDL-C deposition. These include premature arcus senilis, xanthelasmas, and tendon and cutaneous xanthomas.

FH can be diagnosed at virtually any age, although sensitivity and specificity of LDL-C are lower in young children and older adults, when genetic testing may be more appropriate. Universal screening of children and adolescents (9–16 years) may be more cost-effective than cascade family screening, and it has been proposed that such screening takes place at the time of immunization. Use of the cutoff levels for median of the means of either TC or LDL-C levels may be more sensitive and specific than either lipid level alone. Universal screening of children also has the potential for diagnosis of the affected parent and for improved efficiency of detection.

The generally accepted paradigm for treatment of FH is as follows: the earlier the treatment and lower the LDL-C the better. Ideally, diagnosis is made in childhood, and statin therapy begun early. In adulthood, maximum doses of potent statins (atorvastatin 80 mg/d or rosuvastatin 40 mg/d) are used in combination with ezetimibe 10 mg/d. If necessary, additional measures to lower LDL-C are also used (NA, bile acid sequesterants, and/or fenofibrate therapy). At all ages, lifestyle measures are important, particularly avoidance of cigarette smoking. Dietary compliance can improve LDL-C control, as can weight control.

Some patients with ‘resistant’ heFH or hoFH require additional invasive measures to control LDL-C levels. If available, LDL-apheresis is highly effective but costly. An alternative is plasmapheresis, both requiring twice-weekly treatment sessions. For hoFH, apheresis may be bridging therapy for liver transplantation, the only proven effective long-term therapy for this condition. Previous treatment with attempted transfer of normal liver cells containing normal LDL-R alleles was unsuccessful. ApoB RNA-silencing therapy is currently under trial with the novel drug, mipomersin, which in Phase II studies lowered LDL-C levels by a somewhat disappointing ∼20% and increased liver transaminases.

Regression of atherosclerosis and reduction in CVD events have been observed in short-term studies of FH patients with LDL-C-lowering therapy. However, no long-term RCTs have been performed in which aggressive LDL-C lowering with high-dose statin therapy plus ezetimibe or NA has been compared with other therapy in patients with FH.

The efficacy of treating patients with heFH in the “real world” was recently determined in the Netherlands, a country renowned for the quality of its cascade family screening program for FH, as well as for the high quality of its medical care. European and Dutch guidelines currently recommend treatment for lowering LDL-C in heFH patients to plasma levels <2.5 mmol/L. A cross-sectional study of 5 outpatient lipid clinics included 1,249 patients with heFH; 96% of patients were on statin treatment. The LDL-C goal <2.5 mmol/L was achieved in only 21% of patients. Of those not reaching LDL-C goals, 27% were on maximum statin dose and ezetimibe, and in 32%, acceptance of a higher target LDL-C level by the treating physician was the main reason for goal achievement failure. An alternative treatment goal of >50% reduction in LDL-C levels, as recommended in the NICE guidelines, was achieved in 47% of patients who had LDL-C levels ≥2.5 mmol/L and were not using maximum therapy. These data suggest the necessity for greater education of physicians on the need to achieve LDL-C goals in heFH patients.

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References


