Bempedoic acid and its role in contemporary management of hyperlipidemia in atherosclerosis

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1. Introduction
Cardiovascular disease (CVD) is the leading cause of death in the United States of America (USA) [1]. Hyperlipidaemia is a major modifiable risk factor for the development of atherosclerotic cardiovascular disease (ASCVD) and represents an important target for therapeutic interventions. LDL cholesterol is a well-known risk factor for ASCVD and has been the primary target for LDL lowering agents and lifestyle modifications. Statins have been the mainstay of medical therapy in hyperlipidaemia management for almost three decades but have adverse effects on medication compliance [2]. Furthermore, maximally tolerated doses of statins often fail to achieve the target LDL goal. While observational studies have indicated that about 10–15% of patients have statin intolerance, almost 1.5–5% of patients in randomized controlled trials have been noted to have statin intolerance [3]. The most recent American Heart Association (AHA)/American College of Cardiology (ACC)/Multi-societies 2018 cholesterol management guidelines recommended target LDL-C goal of less than 70 mg/dL in high-risk ASCVD patients [4]. The 2019 European Society of Cardiology (ESC)/European Atherosclerosis (EAS) guidelines suggest a lower target goal LDL-C less than 55 mg/dL for patients with very high-risk of ASCVD for primary and secondary prevention [5]. High intensity or maximally tolerated statin therapy often fails to achieve this lipid goal, thus necessitating the use of other non-statin hypolipidemic agents. Ezetimibe and proprotein convertase subtilin-kexin type 9 (PCSK9) inhibitors are recommended by the ACC/AHA/Multi-societies 2018 and ESC/EAS 2019 lipid guidelines to achieve the target LDL-C levels [4]. The
use of ezetimibe along with statin therapy has been shown to cause an additional lowering of 13–20% in LDL-C levels [6]. There is an additional 43–64% of LDL-C reduction with the use of PCSK9 inhibitors along with statins [7–9]. PCSK9 inhibitors have not only been shown to reduce LDL-C level but also been shown to have significant mortality and cardiovascular morbidity benefits. However, there are major barriers in using PCSK9 inhibitors, including cost and subcutaneous administration [10].

Bempedoic acid is a new hypolipidemic drug with a novel mechanism of action that targets cholesterol synthesis through inhibition of the enzyme ATP citrate lyase. Bempedoic acid received USA Food and Drug Administration (FDA) approval in February 2020 for adults with heterozygous familial hypercholesterolaemia or established ASCVD [11]. The bempedoic acid and ezetimibe (Nexlizet) combination received USA FDA approval for the same indication [12]. We present a comprehensive review exploring the underlying mechanism, pre-clinical studies, and clinical trials of bempedoic acid with our opinion of the future therapeutic directions with this drug (Table 1).

2. Mechanism of action

Bempedoic acid (ETC-1002) (8-hydroxy-2,2,14,14-tetramethylpentadecanoic acid) inhibits the activity of the enzyme ATP citrate lyase (ACLY). The ACLY enzyme plays a prominent role in lipid synthesis by converting citrate to acetyl CoA [13] (Figure 1). It functions in the same biosynthetic pathway as the enzyme HMG CoA Reductase (the target enzyme for statins), but lies upstream along the path. Due to the critical position of ACLY in the cholesterol pathway, multiple attempts have been made in the past to develop an inhibitor [14].

Bempedoic acid is once a day prodrug agent, taken orally with liver activation of the enzyme acyl-CoA synthetase [15]. The inhibition of ACLY by bempedoic acid decreases cholesterol synthesis, leading to upregulation of LDL receptors. Increased LDL receptor expression is associated with increased liver LDL uptake and therefore a decrease in serum LDL-C levels [16] (Figure 1). Bempedoic acid has high bioavailability due to its small size and rapid intestinal absorption. Statins and bempedoic acid, both exert their mechanism of action through machinery in the liver. However, the receptors utilized by the two drugs to enter the liver are different. This property of the bempedoic acid ensures that it does not competitively interrupt the uptake of statins in the liver [17,18]. The biosynthetic property of bempedoic acid makes it unique compared to statin, and the liver specific nature of the mechanism of action may be the reason behind lack of muscle related adverse effects seen with this prodrug [15].

In a preclinical study by Pinkosky et al., the authors demonstrated that besides inhibiting ACLY enzyme, the drug also activates the hepatic AMP kinase pathway [19]. AMP kinase inhibits the rate limiting enzymes of fatty acid pathway (acyl CoA carboxylase) and cholesterol pathway (HMG-CoA reductase). Thus, bempedoic acid targets more than one pathway in lowering cholesterol levels [19].

Bempedoic acid affects its target enzymes only in the cells that can mediate its conversion from its inactive to active form. The enzyme that is responsible for this conversion is very long chain acyl CoAsynthetase-1 (ACSVL1) that is expressed in liver with minimal presence in the kidneys and skeletal muscles of humans. The lack of ACSVL1 in skeletal muscle has been hypothesised as the potential mechanism for the lower myotoxic effects of bempedoic acid as compared to statins [18]. The aggregate data from observational studies and randomized controlled trials suggest up to 29% of statin associated muscle side effects [20]. Pinkosky et al., demonstrated the mechanistic basis for ETC-1002 avoiding the myotoxic effects compared to statins due to the absence of ACSVL1 activity in skeletal muscles in both mice and humans [18]. Jadhav et al. recently presented findings of their study suggesting an equivalence in the LDL-C lowering capacity of statin and bempedoic acid combination, compared to high dose statin monotherapy, without the muscular side effects with the higher dose [21]. Additionally, bempedoic acid was not associated with prolongation of QT interval [22].

3. Preclinical studies

The function of bempedoic acid (ESP-55016 and then ETC-1002) in dyslipidemia was studied in multiple animal studies. It was initially recognised as ω-hydroxy-alkanedicarboxylic acid (ESP 55016) with lipid lowering effect in female Zucker (fa/fa) rats. ESP 55016 showed

| Table 1. Pharmacology of bempedoic acid [12–19]. |
| --- |
| 1. Bempedoic acid gets activated in liver to an active metabolite |
| 2. Activated drug inhibits the enzyme ATP Citrate lyase (ACLY) |
| 3. Oral dose, 180mg once daily |
| 4. Elimination half-life is 15–24 hours |
| 5. Initial trials showed LDL-C lowering efficacy of upto 27% as monotherapy |
| 6. Added LDL lowering of upto 48% with Bempedoic acid and ezetimibe combination |
| 7. Lower incidence of myopathy compared with statin |
Table 2. Comparing pharmacology of novel non statin hypolipidemic drugs.

| Drugs                  | Alirocumab (PCSK-9 inhibitor) | Evolocumab (PCSK-9 inhibitor) | Inclisiran | Bempedoic acid |
|------------------------|-------------------------------|-------------------------------|------------|----------------|
| Dose                   | 75–150mg Bi-weekly or 300mg monthly | 140mg Bi-weekly or 420mg monthly | 100–150mg Every 6 months | 180mg Once daily |
| Route                  | Subcutaneous                  | Subcutaneous                  | Subcutaneous | Oral          |
| Metabolism             | Binds to PCSK9 enzyme and proteolysis | Binds to PCSK9 enzyme and proteolysis | – | – |
| Elimination t1/2       | 11–20 days                    | 11–20 days                    | – | 15–24hrs      |
| Effectiveness (%LDL-C reduction) | 45–60% [28] | ~60% [29] | 35.5–52.6% [30] | 17–48% [31] |

**Figure 1.** Mechanism of action of bempedoic acid.

reduction in serum non-high-density lipoprotein (HDL) cholesterol, triglyceride and non-esterified fatty acids. There was a dose dependent increase in serum HDL cholesterol levels and decrease in serum glucose and insulin levels. In rat hepatocytes, this drug suppressed lipid synthesis [23]. This mechanism of action led to the discovery of ETC-1002, a novel agent developed in treatment of cardio-metabolic diseases and dyslipidemia. This molecule affects AMP-activated protein kinase in a Ca (2+)/calmodulin-dependent kinase β-independent and liver kinase β 1-dependent manner. It forms a CoA thioester in the liver leading to reduction of cholesterol synthesis by inhibiting ACLY. One of the ACLY products, oxaloacetate, serves as a substrate of gluconeogenesis, possibly explaining the role of the drug in reduction of gluconeogenesis [19]. In addition, ETC-1002 reduced glucose production through FOXO1, phosphoenolpyruvate carboxykinase (PEPCK), and glucose-6-phosphatase (G6Pase) protein levels [20]. This suggested that risk of diabetes was reduced with ETC-1002 compared to statins [24]. Ference et al. performed a study with genetically constructed scores mimicking bempedoic acid and did not find association of diabetes in the ACLY score [25]. Samsoondar et al. showed the effect of ETC-1002 on diet-induced metabolic dysregulation using LDLr−/− mice. These mice on high fat, high cholesterol diet was given bempedoic acid (3-30mg/kg/day for 12 weeks) which showed reduction in cholesterol by 50% and triglycerides by 64% [26]. Reduction in hepatic and aortic inflammation was also noted, as measure by decrease in markers for inflammatory macrophages in mice treated with bempedoic acid [26,27].
4. Bempedoic acid ameliorates inflammation

High sensitivity C-Reactive protein (hsCRP) level is an independent predictor of ASCVD risk [28]. Multiple studies have shown that hsCRP level < 2 mg/L with statin therapy or with LDL-C levels < 70 mg/dL provided significant ASCVD risk reduction, while an elevated hsCRP level were associated with residual risk of ASCVD. This inflammatory marker is considered a risk modulator in the latest 2018 ACC/AHA/Multi-societies lipid guidelines [31].

Unlike PCSK9 inhibitors, in addition to its cholesterol lowering potential, bempedoic acid has been shown to possess strong systemic anti-inflammatory effects in animal studies [26,32]. Multiple animal studies indicated the possible potential benefit of bempedoic acid in reducing insulin resistance and cardiac complications of metabolic syndrome through the activation of AMP-activated protein kinase which plays an important role in lipid and glucose metabolism. Another proposed mechanism is the downregulation of inflammatory signalling pathways through AMP kinase activation [33]. Additionally, inhibition of ACLY by bempedoic acid has been demonstrated to reduce prostaglandin production and contribute to anti-inflammatory action [34] and may have some role in NASH as well [35]. Lauf et al., in their CLEAR Serenity study, showed that bempedoic acid was associated with a 24.3% reduction in hsCRP levels from baseline, reducing inflammation and possibly ASCVD risk [36].

5. Clinical development of bempedoic acid

5.1. Phase I trials

The drug’s safety and tolerability were well established with phase 1 clinical studies evaluating the effect of varying doses of the drug as compared with placebo. In a phase 1a study involving 53 subjects with mild dyslipidemia (average LDL-C 126.7 mg/dL and TG 100-350 mg/dL), participants were treated with increasing doses of the drug up to 120 mg daily. A 17% reduction in LDL-C level was observed with an average dose of 100 mg daily. The phase 1b study enrolled 24 healthy subjects who received drug in dose increments (140, 180, and 220 mg daily). A reduction of 36% LDL-C levels was observed, compared to placebo (p < .0001). Finally, ETC-1002-011 study had 6 healthy male subjects to assess absorption, metabolism and excretion of single dose in urine and faeces [37]. No major adverse events were noted in phase I studies, despite renal impairment [38].

5.2. Phase II trials

Multiple phase 2 studies evaluated the lipid lowering efficacy of bempedoic acid as monotherapy or as an add on to statin therapy, ezetimibe or a combination of all three agents [24,39,40]. Doses of between 40 and 240 mg/day of bempedoic acid were used in these studies. The study population included hyperlipidemic patients with other cardiovascular risk factors including diabetes mellitus, hypertension and/or statin intolerance. The use of this drug as a single agent or in combination with other hypolipidemic agents was associated with significant improvements in lipid levels and inflammatory markers. The results of the studies are summarized in Table 3.

Another phase-II trial (NCT02659397) published by Lalwani et al. studied the effect of adding bempedoic acid to stable high intensity atorvastatin therapy in patients with hypercholesterolaemia. Patients received 80 mg atorvastatin for 4 weeks followed by randomization in ratio of 2:1 to receive bempedoic acid 180 mg daily or placebo for 4 weeks. The mean LDL level was lowered by 22% with bempedoic acid compared to baseline (p = .003). The addition of a stable dose of bempedoic acid to high intensity atorvastatin 80 mg reduced the non HDL cholesterol (-13%; p = .015), apolipoprotein B (-15%; p = .004), and high-sensitivity C-reactive protein (-44%; p = .002) significantly, with no significant alteration of the concentration of atorvastatin or its major metabolite ortho-hydroxy atorvastatin [39].

5.3. Phase III trials

The CLEAR trial series are all phase III randomised clinical trials focussed on evaluating the efficacy of bempedoic acid. The initial four trials (CLEAR Tranquillity, CLEAR Serenity, CLEAR Wisdom and CLEAR Harmony) have evaluated the LDL-C lowering efficacy of the drug while the CLEAR OUTCOME trial is an ongoing trial focussed on evaluating the cardiovascular outcomes with the drug.

The safety and efficacy of bempedoic acid 180 mg daily as an add on therapy to ezetimibe 10 mg daily in patients with a history of statin intolerance and those who had elevated LDL-C (LDL-C ≥ 100 mg/dL) on their current therapy was studied in CLEAR Tranquillity. The study subjects were randomized to oral bempedoic acid 180 mg or placebo once daily for 12 weeks. Bempedoic acid yielded an additional 28.5% LDL-C lowering compared with placebo (p < .001; −23.5% bempedoic acid, +5.0% placebo). A significant improvement was also seen in other lipid and
Table 3. Phase 2 and phase 3 clinical trials of bempedoic acid.

| Name of the trial                                                                 | Phase of study | Primary objective                                                                 | Status/Result                                                                 |
|-----------------------------------------------------------------------------------|----------------|-----------------------------------------------------------------------------------|-------------------------------------------------------------------------------|
| Bempedoic Acid + Ezetimibe fixed-dose combination (FDC) study in patients with type 2 diabetes and elevated LDL-C | Phase 2        | Percent change of LDL-C from baseline to week 12, by FDC vs. Placebo and FDC vs. Ezetimibe | Completed                                                                     |
| Evaluation of the efficacy and safety of bempedoic acid (ETC-1002) 180 mg, Ezetimibe 10 mg, and Atorvastatin 20 mg triplet therapy in patients with elevated LDL-C | Phase 2        | Percent change in low-density lipoprotein cholesterol (LDL-C) after 6 weeks       | Completed                                                                     |
| A study of pharmacokinetics, pharmacodynamics and safety of adding ETC-1002 to atorvastatin 80 mg | Phase 2        | Percent change in low-density lipoprotein cholesterol (LDL-C) from baseline to 4 weeks treatment. | Completed                                                                     |
| Evaluation of the efficacy and safety of bempedoic acid (ETC-1002) 180 mg, when added to PCSK9 inhibitor therapy | Phase 2        | Percent change in low-density lipoprotein cholesterol (LDL-C) after 12 weeks of therapy | Completed                                                                     |
| Evaluation of long-term efficacy of bempedoic acid (ETC-1002) in patients with hyperlipidaemia at high cardiovascular risk (CLEAR Wisdom) | Phase 3        | Percent change in low-density lipoprotein cholesterol (LDL-C) after 12 weeks of therapy | Completed                                                                     |
| Evaluation of the efficacy and safety of bempedoic acid (ETC-1002) in patients with hyperlipidaemia and statin intolerant (CLEAR serenity) | Phase 3        | Percent change in low-density lipoprotein cholesterol (LDL-C) after 12 weeks of therapy | Completed                                                                     |
| Evaluation of the efficacy and safety of bempedoic acid (ETC-1002) as add-on to Ezetimibe therapy in patients with elevated LDL-C (CLEAR tranquility) | Phase 3        | Percent change in low-density lipoprotein cholesterol (LDL-C) after 12 weeks of therapy | Completed                                                                     |
| A study evaluating the safety and efficacy of bempedoic acid plus ezetimibe fixed-dose combination compared to bempedoic acid, ezetimibe, and placebo in patients treated with maximally tolerated statin therapy | Phase 3        | Percent change in low-density lipoprotein cholesterol (LDL-C) after 12 weeks of therapy | Completed                                                                     |
| Evaluation of long-term safety and tolerability of ETC-1002 in high-risk patients with hyperlipidaemia and high CV risk (CLEAR HARMONY) | Phase 3        | Incidence of treatment related adverse effects                                  | Completed                                                                     |
| Assessment of the long-term safety and efficacy of bempedoic acid (CLEAR Harmony OLECl) | Phase 3        | Incidence of adverse events in patients with high cardiovascular risk and elevated LDL cholesterol | Active                                                                        |
| Evaluation of major cardiovascular events in patients with, or at high risk for, cardiovascular disease who are statin intolerant treated with bempedoic acid (ETC-1002) or Placebo (CLEAR OUTCOMES) | Phase 3        | Time to first occurrence of MACE                                            | Recruiting                                                                   |

lipoprotein parameters, including non-HDL-C, total cholesterol and apoB level. A median hsCRP reduction of 33% was noted at week 12 (p < .001). Besides showing lipid lowering efficacy, bempedoic acid was also well tolerated in the study arm. Adverse events of the drug were included increase in serum uric acid level (7.7% in bempedoic acid group compared to 2.3% in placebo group) and headache (4.4% in bempedoic acid and 3.4% in placebo group). Muscle-related adverse effects including muscular weakness, myalgias and rhabdomyolysis were similar in both the study and the placebo arms (3.3% in bempedoic acid and 3.4% placebo). The most important limitation of the trial was the short duration of 12 weeks over which outcomes were measured [41]. Subsequently, the CLEAR Harmony trial was a longer (24 weeks) study involving patients with either ASCVD, heterogeneous FH or both. In this trial, 2230 patients who were on guideline directed statin therapy were randomized into two groups (bempedoic acid versus placebo) and the baseline mean LDL-C was 103 ± 29.4 mg per decilitre. The incidence of serious adverse events was comparable in both the groups (14.5% in bempedoic acid group and 14.0% in the placebo group). Similar to the
results of the previous trials, myalgia and other muscle related serious effects were similar in both groups while gout occurred more often in the bempedoic acid group (1.2%) compared to the placebo group (0.3%). Bempedoic acid reduced LDL cholesterol level by 18.1% (95% CI: −20.0 to −16.1, \( p < .001 \)) at week 12 which persisted at week 24 (16.1%, 95% CI: −18.4 to −14, \( p < .001 \)) when compared to the placebo. There was a lower incidence of diabetes and worsening of the disease in bempedoic acid group (3.3% in bempedoic acid group compared to 5.4% in placebo group, \( p = .2 \)), but due to low event rate the study was under-powered to specifically study this effect [42].

While the above two trials evaluated the efficacy of bempedoic acid as an add-on therapy in patient on maximal statin therapy, the CLEAR Serenity trial studied 345 patients with hypercholesterolaemia who were intolerant to at least two statins (one of them at the lowest available dose). These patients were followed for over 24 weeks to the primary end point of percentage change in LDL-C level from baseline with bempedoic acid 180 mg. At week 12, bempedoic acid significantly lowered LDL-C level from baseline (95% CI: −25.1 to −17.7%, \( p < .001 \)). There was also a significant reduction in hsCRP compared to baseline in the bempedoic acid group. The adverse event rate was higher in bempedoic acid group 64.1% as compared to the 56.8% in placebo group. The common side effects included musculoskeletal and connective tissue disorders (22.2 versus 25.2% in bempedoic acid and placebo groups, respectively). This trial expanded the evidence regarding the use of bempedoic acid in statin intolerant population in long term period [36].

A reduction in LDL-C levels among ASCVD or heterogenous FH patients on maximum tolerated statin therapy using bempedoic acid was first investigated in the CLEAR Wisdom trial. It was a randomized controlled trial with 779 patients to the drug group (180 mg bempedoic acid) or placebo once daily. The baseline LDL-C was 120 mg/dL with a follow up of 52 weeks. The primary outcome was a significant change in the LDL-C levels from baseline to week 52 (−15.4 vs 2.4% in bempedoic acid group compared to placebo). The study was underpowered, making it difficult to interpret clinical outcomes [43]. Banach et al. performed a meta-analysis of these 4 trials and confirmed a statistically significant reduction of LDL-C with bempedoic acid, when compared to the 1.5% reduction in the placebo group at week 12 (−24.5%; 95% CI, −27.8% to −21.1%; \( p < .001 \)). The reduction in the levels of LDL-C levels persisted at week 52 in patients with ASCVD, HeFH or both receiving a maximally tolerated statin (−12.7% difference between bempedoic acid and placebo group) [44]. Similar results were confirmed in other meta-analysis, with no significant increase in adverse effects noted with bempedoic acid [45,46].

While all the trials have shown consistent reduction in LDL-C levels with bempedoic acid, two recently published systematic analyses have also shown a lower incidence of new onset and worsening of diabetes mellitus [47–50]. This has the additional potential in expanding the use of the drug in diabetes dyslipidemia and could be explained by the modest reduction in triglyceride levels mitigated by bempedoic acid. In a recently published meta-analysis of the four RCTs on bempedoic acid, it was shown to reduce the levels of hsCRP (−23.4% (CI 95% = 32.6 to −14.2)); \( p < .05; \chi^2 = 69\% \) [51,52]. The latest trial in the series (CLEAR Outcomes trial) is aimed at studying the cardiovascular outcomes with bempedoic acid 180 mg daily. It is the largest trial randomizing 14,014 high cardiovascular disease risk statin intolerant patients with a follow up period of 36–42 months [53].

The majority of the phase-3 trials compared bempedoic acid alone with either placebo or statins. One recently published phase-3 trial compared the safety and efficacy of fixed dose combination (FDC) of bempedoic acid 180 mg along with ezetimibe 10 mg in high-risk patients on maximum tolerated statin [54]. In this trial, 382 patients were randomized into 4 groups in ratio 2:2:2:1 to oral bempedoic acid 180 mg plus ezetimibe 10 mg daily combination, bempedoic acid 180 mg daily, ezetimibe 10 mg daily or placebo for 12 weeks. Significant reduction in percentage change in LDL level at end of week 12 was seen in FDC group (−36.2%) in comparison to bempedoic acid as single agent (−17.2%; \( p < .001 \)), ezetimibe only (−23.2%; \( p < .001 \)) and placebo (1.8%; \( p < .001 \)). The FDC bempedoic acid and ezetimibe also lowered the hs-CRP (35.1%) and other lipid levels including non-HDL-cholesterol, total cholesterol and apolipoprotein B. The
adverse events were frequently reported in FDC and bempedoic acid groups compared to ezetimibe alone or placebo groups. Among the FDC group, the most common side effects were constipation, muscle spasms, fatigue and blood uric acid increase. In a subgroup analysis of patients who were not receiving statin due to intolerance, FDC reduced LDL-C by 38.8% and subgroup analysis of patients on high-intensity statin, FDC reduced LDL-C by 38.9% [54]. These interesting findings support the use of bempedoic acid and ezetimibe combination in lowering LDL, even in population not on a medication from the statin class of drugs [54]. In a pooled analysis of 388 patients, Bhagavathula et al. also noted a significant reduction of LDL-C (−29.14%, 95% CI −39.52 to −18.76; p < .001) and hsCRP (−30.48%, 95% CI −44.69 to −16.28; p = .04) with the combination of bempedoic acid and ezetimibe in this patient population, as compared to the control group, after 12 weeks [55]. In a smaller trial enrolling 59 patients on background therapy with PCSK9 inhibitors, the addition of bempedoic acid showed a further reduction of LDL-C by 30.3% after 2 months of therapy, when compared with placebo [56].

6. Conclusion and future direction

For many decades, statins were the only hypolipidemic drugs available with a proven cardiovascular mortality benefit. However, many patients either could not achieve optimal lipid goals on maximally tolerated statin doses or experienced statin intolerance. The last 5 years have seen significant advances in the development and evidence basis for novel therapies for hyperlipidaemia and cardiovascular diseases, with many new molecules receiving approval in both North America and Europe. There have been multiple updated guidelines from the ACC/AHA and ESA/ESC to keep pace with this advancement. Parallel to this surge in development of new classes of drugs, recent clinical trials have shown improvement in cardiovascular mortality with additional reduction in the LDL-C levels. Thus, the target LDL-C goal for primary and secondary prevention for “very high-risk” and “high-risk” ASCVD patients has progressively become lower over the decades. Drugs including ezetimibe, PCSK9 inhibitors, bempedoic acid are now available for clinicians to target these LDL-C goals, and pick the most optimized combination with maximum cost effectiveness of hypolipidemic drugs for their patients [57]. Keeping focus on this patient-centric approach, the recently approved bempedoic acid-ezetimibe combination could be tried prior to attempting PCSK9 inhibitors, which have certain limitations including injectable preparation and higher costs. The fixed drug combination of bempedoic acid and ezetimibe has shown LDL-C reduction comparable to PCSK9 inhibitors in clinical trials. In February 2020, US FDA (Table 4) approved oral bempedoic acid, once a day medication, for use in heterozygous hypercholesterolaemia or established ASCVD patients, who need further LDL-C lowering in addition to dietary modifications and maximally tolerated statin. The once daily oral regimen and the favourable adverse effect profile makes bempedoic acid an appealing alternative. However, one major limitation of use of bempedoic acid includes lack of CVD outcome data, which will be addressed in future clinical trials. A recent study identified insurance approvals and cost as additional barriers which will also require to be addressed to promote in real world clinical practice [58].

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Author contributions

Authors RT and RK were involved in the conception and design; all authors were involved the drafting of the paper, revising it critically for intellectual content; and the final approval of the version to be published. All authors agree to be accountable for all aspects of the work.

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**Data availability statement**

The authors confirm that the data supporting the findings of this study are available within the article [and/or] its supplementary materials.

**References**

[1] Benjamin EJ, Muntner P, Alonso A, American Heart Association Council on Epidemiology and Prevention Statistics Committee and Stroke Statistics Subcommittee, et al. Heart disease and stroke statistics-2019 update: a report from the american heart association. Circulation. 2019;139(10):e56–e528.

[2] Jacobson TA, Ito MK, Maki KC, et al. National lipid association recommendations for patient-centered management of dyslipidemia: part 1-full report. J Clin Lipidol. 2015;9(2):1287–169.

[3] Ward NC, Watts GF, Eckel RH. Statin toxicity. Circ Res. 2019;124(2):328–350.

[4] Grundy SM, Stone NJ, Bailey AL, et al. 2018 AHA/ACC/AACVPR/ABC/ACPM/ADA/AGS/APhA/ASP/NLA/PCNA guideline on the management of blood cholesterol: executive summary: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines. J Am Coll Cardiol. 2019;73(24):3168–3209.

[5] Mach F, Baigent C, Catapano AL, ESC Scientific Document Group, et al. 2019 ESC/EAS guidelines for the management of dyslipidaemias: lipid modification to reduce cardiovascular risk. Eur Heart J. 2020;41(1):111–188.

[6] Cannon CP, Blazing MA, Giugliano RP, et al. Ezetimibe added to statin therapy after acute coronary syndromes. N Engl J Med. 2015;372(25):2387–2397.

[7] Sabatine MS, Giugliano RP, Wiviott SD, et al. Efficacy and safety of evolocumab in reducing lipids and cardiovascular events. N Engl J Med. 2015;372(16):1500–1509.

[8] Schwartz GG, Steg PG, Szarek M, et al. Alirocumab and cardiovascular outcomes after acute coronary syndrome. N Engl J Med. 2018;379(22):2097–2107.

[9] Kastelein JJP, Kereiakes DJ, Cannon CP, et al. Effect of alirocumab dose on LDL lowering and lipid goal attainment in patients with dyslipidemia. Coron Artery Dis. 2017;28(3):190–197.

[10] Chaudhary R, Garg J, Shah N, et al. PCSK9 inhibitors: a new era of lipid lowering therapy. World J Cardiol. 2017;9(2):76–91.

[11] Agha AM, Jones PH, Ballantyne CM, et al. Greater than expected reduction in low-density lipoprotein-cholesterol (LDL-C) with bempedoic acid in a patient with heterozygous familial hypercholesterolemia (HeFH). J Clin Lipidol. 2021;15(5):649–652.

[12] Bempedoic acid FDA approval [Internet]. Available from: [https://www.accessdata.fda.gov/drugsatfda_docs/label/2020/211616s000lbl.pdf](https://www.accessdata.fda.gov/drugsatfda_docs/label/2020/211616s000lbl.pdf).

[13] Daikuha Y, Sunemi T, Takeda Y. The role of ATP citrate lyase in the transfer of acetyl groups in rat liver. Biochim Biophys Acta. 1968;158(1):51–61.

[14] Ameer F, Scanduzzi L, Hasnain S, et al. De novo lipogenesis in health and disease. Metabolism. 2014;63(7):895–902.

[15] Cicero AFG, Fogacci F, Cinzone I. Evaluating pharmacokinetics of bempedoic acid in the treatment of hypercholesterolemia. Expert Opin Drug Metab Toxicol. 2021;17(9):1031–1038.

[16] Hamilton JG, Sullivan AC, Kritchevsky D. Hupolipidemic activity of (-)-hydroxycitrate. Lipids. 1977;12(1):1–9.

[17] Bilien O, Ballantyne CM. Bempedoic acid (ETC-1002): an investigational inhibitor of ATP citrate lyase. Curr Atheroscler Rep. 2016;18(10):61.

[18] Pinkosky SL, Newton RS, Day EA, et al. Liver-specific ATP-citrate lyase inhibition by bempedoic acid decreases LDL-C and attenuates atherosclerosis. Nat Commun. 2016;7:13457.

[19] Pinkosky SL, Filippov S, Srivastava RAK, et al. AMP-activated protein kinase and statin combinations: predictions from a dose–response model. Eur Heart J Cardiovasc Pharmacother. 2021::pvab064.

[20] Cohen JD, Brinton EA, Ito MK, et al. Understanding statin use in america and gaps in patient education (USAGE): an internet-based survey of 10,138 current and former statin users. J Clin Lipidol. 2012;6(3):208–215.

[21] Jadhav SB, Crass RL, Chapel S, et al. Pharmacodynamic effect of bempedoic acid and statin combinations: predictions from a dose–response model. Eur Heart J Cardiovasc Pharmacother. 2021::pvab064.

[22] Amore BM, Cramer CT, MacDougal DE, et al. Absence of effect of steady state bempedoic acid on cardiac repolarization: results of a thorough QT/QTc study in healthy volunteers. Clin Transl Sci. 2021;14(6):2487–2496.

[23] Cramer CT, Goetz B, Hopson KLM, et al. Effects of a novel dual lipid synthesis inhibitor and its potential utility in treating dyslipidemia and metabolic syndrome. J Lipid Res. 2004;45(7):1289–1301.

[24] Gutierrez MJ, Rosenberg NL, Macdougall DE, Hanselman JC, et al. Efficacy and safety of ETC-1002, a novel investigational low-density lipoprotein-cholesterol-lowering therapy for the treatment of patients with hypercholesterolemia and type 2 diabetes mellitus. ATVB. 2014;34(3):676–683.

[25] Ference BA, Ray KK, Catapano AL, et al. Mendelian randomization study of ACLY and cardiovascular disease. N Engl J Med. 2019;380(11):1033–1042.

[26] Samsoondar JP, Burke AC, Sutherland BG, et al. Prevention of Diet-Induced metabolic dysregulation, inflammation, and atherosclerosis in LDLr−/− mice by treatment with the ATP-Citrate lyase inhibitor
bempedoic acid. Arterioscler Thromb Vasc Biol. 2017;37(4):647–656.

[27] Galic S, Fullerton MD, Schertzer JD, et al. Hematopoietic AMPK β1 reduces mouse adipose tissue macrophage inflammation and insulin resistance in obesity. J Clin Invest. 2011;121(12):4903–4915.

[28] Kaptoge S, Di Angelantonio E, Lowe G, Emerging Risk Factors Collaboration, et al. C-reactive protein concentration and risk of coronary heart disease, stroke, and mortality: an individual participant meta-analysis. Lancet. 2010;375:132–140.

[29] Bohula EA, Giugliano RP, Cannon CP, et al. Achievement of dual low-density lipoprotein cholesterol and high-sensitivity C-reactive protein targets more frequent with the addition of ezetimibe to simvastatin and associated with better outcomes in IMPROVE-IT. Circulation. 2015;132(13):1224–1233.

[30] Ruscica M, Corsini A, Ferri N, et al. Clinical approach to the inflammatory etiology of cardiovascular diseases. Pharmacol Res. 2020;159:104916.

[31] Lloyd-Jones DM, Braun LT, Ndumele CE, et al. Use of risk assessment tools to guide decision-making in the primary prevention of atherosclerotic cardiovascular disease: a special report from the American Heart Association and American College of Cardiology. J Am Coll Cardiol. 2019;73(24):3153–3167.

[32] Ruscica M, Tokgözöglu L, Corsini A, et al. PCSK9 inhibition and inflammation: a narrative review. Atherosclerosis. 2019;288:146–155.

[33] Filippov S, Pinkosky SL, Lister RJ, et al. ETC-1002 regulates immune response, leukocyte homing, and adipose tissue inflammation via LKB1-dependent activation of macrophage AMPK. J Lipid Res. 2013;54(8):2095–2108.

[34] Verberk SGS, Kuiper KL, Lauterbach MA, et al. The multifaceted therapeutic value of targeting ATP-citrate lyase in atherosclerosis. Trends Mol Med. 2021;27(12):1095–1105.

[35] Sanjay KV, Vishwakarma S, Zope BR, et al. ATP citrate lyase inhibitor bempedoic acid alleviates long term HFD induced NASH through improvement in glycemic control, reduction of hepatic triglycerides & total cholesterol, modulation of inflammatory & fibrotic genes and improvement in NAS score. Curr Res Pharmacol Drug Discov. 2021;2:100051.

[36] Laufs U, Banach M, Mancini GBJ, et al. Efficacy and safety of bempedoic acid in patients with hypercholesterolemia and statin intolerance. J Am Heart Assoc. 2019;8:e011662.

[37] Nikolic D, Mikhailidis DP, Davidson MH, et al. ETC-1002: a future option for lipid disorders? Atherosclerosis. 2014;237(2):705–710.

[38] Amore BM, Sasiela WJ, Ries DK, et al. Pharmacokinetics of bempedoic acid in patients with renal impairment. Clin Transl Sci. 2021;cts.13202.

[39] Thompson PD, Rubino J, Janik MJ, et al. Use of ETC-1002 to treat hypercholesterolemia in patients with statin intolerance. J Clin Lipidol. 2015;9(3):295–304.

[40] Ballantyne CM, Mckenney JM, MacDougall DE, et al. Effect of ETC-1002 on serum Low-Density lipoprotein cholesterol in hypercholesterolemic patients receiving statin therapy. Am J Cardiol. 2016;117(12):1928–1933.

[41] Ballantyne CM, Banach M, Mancini GBJ, et al. Efficacy and safety of bempedoic acid added to ezetimibe in statin-intolerant patients with hypercholesterolemia: a randomized, placebo-controlled study. Atherosclerosis. 2018;277:195–203.

[42] Ray KK, Bays HE, Catapano AL, et al. Safety and efficacy of bempedoic acid to reduce LDL cholesterol. N Engl J Med. 2019;380(11):1022–1032.

[43] Goldberg AC, Leiter LA, Stroes ES, et al. Effect of bempedoic acid vs placebo added to maximally tolerated statins on Low-Density lipoprotein cholesterol in patients at high risk for cardiovascular disease: the CLEAR wisdom randomized clinical trial. JAMA. 2019;322(18):1780–1788.

[44] Banach M, Duell PB, Gotto AM, et al. Association of bempedoic acid administration with atherogenic lipid levels in phase 3 randomized clinical trials of patients with hypercholesterolemia. JAMA Cardiol. 2020;5(10):1124.

[45] Dai L, Zuo Y, You Q, et al. Efficacy and safety of bempedoic acid in patients with hypercholesterolemia: a systematic review and meta-analysis of randomized controlled trials. Eur J Prev Cardiol. 2021;28(8):825–833.

[46] Bays HE, Banach M, Catapano AL, et al. Bempedoic acid safety analysis: pooled data from four phase 3 clinical trials. J Clin Lipidol. 2020;14(5):649–659.e6.

[47] Leiter LA, Banach M, Catapano AL, et al. Bempedoic acid in patients with type 2 diabetes mellitus, prediabetes, and normoglycemia: a post hoc analysis of efficacy and glycaemic control using pooled data from phase 3 clinical trials. Diabetes Obes Metab. 2022;14645.

[48] Masson W, Lobo M, Lavalle-Cobo A, et al. Effect of bempedoic acid on new onset or worsening diabetes: a meta-analysis. Diabetes Res Clin Pract. 2020;168:108369.

[49] Leiter L, Banach M, Catapano A, et al. Bempedoic acid and glycemic control: a pooled analysis of 4 phase 3 clinical trials. J Clin Lipidol. 2020;14(5):577–579.

[50] Bays HE, Baum SJ, Brinton EA, et al. Effect of bempedoic acid plus ezetimibe fixed-dose combination vs ezetimibe or placebo on low-density lipoprotein cholesterol in patients with type 2 diabetes and hypercholesterolemia not treated with statins. Am J Prev Cardiol. 2021;8:100278.

[51] Masson W, Lobo M, Lavalle-Cobo A, et al. Effect of bempedoic acid on atherogenic lipids and inflammation: a meta-analysis. Clinica Rev Investig En Arterioscler. 2021;33(3):117–126.

[52] Ballantyne CM, Bays H, Catapano AL, et al. Role of bempedoic acid in clinical practice. Cardiovasc Drugs Ther. 2021;35(4):853–864.

[53] Nichols S, Lincoff AM, Bays HE, et al. Rationale and design of the CLEAR-outcomes trial: evaluating the effect of bempedoic acid on cardiovascular events in patients with statin intolerance. Am Heart J. 2021;235:104–112.

[54] Ballantyne CM, Laufs U, Ray KK, et al. Bempedoic acid plus ezetimibe fixed-dose combination in patients with hypercholesterolemia and high CVD risk treated.
with maximally tolerated statin therapy. Eur J Prev Cardiol. 2020;27(6):593–603.

[55] Bhagavathula AS, Al Matrooshi NO, Clark CCT, et al. Bempedoic acid and ezetimibe for the treatment of hypercholesterolemia: a systematic review and meta-analysis of randomized phase II/III trials. Clin Drug Investig. 2021;41(1):19–28.

[56] Rubino J, MacDougall DE, Sterling LR, Kelly SE, et al. Lipid lowering with bempedoic acid added to a pro-protein convertase subtilisin/kexin type 9 inhibitor therapy: a randomized, controlled trial. J Clin Lipidol. 2021;

[57] Blaum C, Brunner FJ, Gößling A, et al. Target populations and treatment cost for bempedoic acid and PCSK9 inhibitors: a simulation study in a contemporary CAD cohort. Clin Ther. 2021;43(9):1583–1600.

[58] Warden BA, Cardiology B-A, Purnell JQ, et al. Real-world utilization of bempedoic acid in an academic preventive cardiology practice. J Clin Lipidol. 2022;16(1):94–103.