Legacy effects of statins on cardiovascular and all-cause mortality: a meta-analysis

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Legacy effects of statins on cardiovascular and all-cause mortality: a meta-analysis

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ABSTRACT

Objectives To assess evidence for ‘legacy’ (post-trial) effects on cardiovascular disease (CVD) mortality and all-cause mortality among adult participants of placebo-controlled randomised controlled trials (RCTs) of statins.

Design Meta-analysis of aggregate data.

Setting/Participants Placebo-controlled statin RCTs for primary and secondary CVD prevention.

Methods Data sources: PubMed, Embase from inception and forward citations of Cholesterol Treatment Trials’ Collaborators RCTs to 16 June 2016. Study selection: Two independent reviewers identified all statin RCT follow-up reports including ≥1000 participants, and cardiovascular and all-cause mortality. Data extraction and synthesis: Two independent reviewers extracted data in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines. Main outcomes: Post-trial CVD and all-cause mortality.

Results We included eight trials, with mean post-trial follow-up ranging from 1.6 to 15.1 years, and including 13 781 post-trial deaths (6685 CVD). Direct effects of statins within trials were greater than legacy effects post-trials. The pooled data from all eight studies showed no evidence overall of legacy effects on CVD mortality, but some evidence of legacy effects on all-cause mortality (p=0.01). Exploratory subgroup analysis found possible differences in legacy effect for primary prevention trials compared with secondary prevention trials for both CVD mortality (p=0.15) and all-cause mortality (p=0.02). Pooled post-trial HR for the three primary prevention studies demonstrated possible post-trial legacy effects on CVD mortality (HR=0.87; 95% CI 0.79 to 0.95) and on all-cause mortality (HR=0.90; 95% CI 0.85 to 0.96).

Conclusions Possible post-trial statin legacy effects on all-cause mortality appear to be driven by the primary prevention studies. Although these relative benefits were smaller than those observed within the trial, the absolute benefits may be similar for the two time periods. Analysis of individual patient data from follow-up studies after placebo-controlled statin RCTs in lower-risk populations may provide more definitive evidence on whether early treatment of subclinical atherosclerosis is likely to be beneficial.

INTRODUCTION

‘Legacy effects’ are treatment effects that persist or emerge at some time after trial treatment ends. The existence, or not, of such legacy effects have important clinical implications, including the potential value of starting preventative treatment at a younger age, and the risks of treatment cessation. Although technically even short-term or transient improvement or worsening of symptoms and signs may be classified as a legacy effect, most people appear to use the term to indicate sustained clinical benefit or harm. 1-2 Legacy effects have been explained as the ‘memory of a treatment given in an early phase of a disease which produces benefits long after the cessation of intervention’. 3 They are an extension of the belief that we should intervene with treatment early on in the course of a chronic disease/condition; the legacy effect assumes that the duration of the condition predicts permanent pathologica changes which in turn are strong modifiers of treatment effectiveness.

Recently, there has been considerable interest in the possible legacy effects of statins, 4-5 sparked by reports on the long-term outcomes for participants of large placebo-controlled trials. In some of these
reports, there is still a persistent survival advantage to participants who were randomised to statin during the trial, even though there was no or minimal difference in the management of participants after the trial ended. Legacy effects could indicate that earlier treatment with statins slows atherosclerotic plaque build-up in arteries and so alters the natural disease progression during a person’s lifetime. As with the direct effects of statins, these legacy effects may be pleiotropic, and act through anti-inflammatory, anticoagulation and/or lipid lowering. This hypothesis has some support from the finding that statins have minimal effect on cardiovascular disease (CVD) prevention in patients with advanced kidney disease who require haemodialysis, and who have high short-term risk of CVD, but reduce CVD events in patients with earlier chronic kidney disease who are not yet requiring haemodialysis. There are also some data from a small imaging study of patients with angiogram to support the early treatment hypothesis, where similar reduction in lipid levels appeared to result in reduction in plaque volume only in participants younger than 65 years. Finally, differences in long-term response to statins are noted for primary prevention trials compared with secondary prevention trials. To this end, aggressive lipid-lowering therapy in much younger individuals with lower risk for CVD has been suggested as a possible means of primary prevention. Some have argued for universal screening of cholesterol levels in young people and offering early statin treatment to those with raised levels, whereas others have argued that statins be offered to all young people, regardless of cholesterol levels.

At least some of the survival benefit observed on long-term follow-up is attributable to the direct treatment effects on CVD outcomes observed during the within-trial period. To illustrate this point, we generated data to simulate the situation where there was a legacy effect (see online supplementary eFigure 1A and 1B), and where there was not a legacy effect (online supplementary eFigure 1C and 1D). We simulated two scenarios where an intervention has effects during the trial period, and either (1) has an additional effect after the trial (legacy effect) or (2) has no additional effect after the trial (no legacy effect). In the survival curves of both scenarios, the apparent legacy effect is exaggerated because the cumulative incidence includes the direct effects during the initial trial period (see online supplementary eFigure 1A and 1C). If hazard curves are constructed instead, the direct effects during the initial trial period are not included in the instantaneous hazard of the post-trial periods, allowing an unbiased estimation of the legacy effect (see online supplementary eFigure 1B and 1D; note that these are curves of the instantaneous hazard at each time point, and are not curves of hazard ratios (HRs). Details of the methods for the simulation are provided in the Appendix). Although survival curves like online supplementary eFigure 1A and 1C demonstrate that the direct effects of the intervention (observed during the trial period) are still apparent many years later, they do not provide evidence of legacy effects after the intervention has ceased. From the hazard curves in online supplementary eFigure 1B and 1D, it is clear that to estimate legacy effects, we should instead focus on outcomes observed during the post-trial period. To this end, we aimed to identify and combine estimates of the effect of trial treatment group allocation on post-trial all-cause and CVD mortality from published reports on the long-term follow-up after placebo-controlled trials of statins.

METHODS

Protocol and registration

The review protocol was not registered.

Selection

We performed a systematic search and meta-analysis of all reports on follow-up after randomised, placebo-controlled studies of adults (age >18 years) of statins with ≥1000 participants. As the legacy effect relates to the difference in treatment received within the trial period, we focused our analysis on follow-up reports of high-quality, large randomised controlled trials (RCTs). We chose to limit our studies to those with ≥1000 participants in the original trial for consistency with the Cholesterol Treatment Trialists’ Collaboration. These large trials were designed to assess effects on mortality within the trial period, and their follow-up reports are the most appropriate studies to address post-trial effects on mortality. We excluded studies that did not report mortality data during post-trial follow-up. The primary outcomes were death due to all-causes and death due to CVD.

Search strategy

We identified placebo-controlled RCTs of cholesterol-lowering treatment from the Cholesterol Treatment Trialists’ Collaboration19 and ran forward citation searches in Scopus; our search was limited to those citations which included one of the investigators from the RCT. We searched for additional reports in Medline and Embase with no earliest date restriction, though to 16 June 2016 using the terms listed in box 1, with no restrictions on year published, type of publication or language. We checked references of included studies to identify further relevant papers and contacted trialists to identify updated or additional reports.

Box 1 Search terms for Medline Search

1. Follow-Up Studies/
2. random$tw.
3. placebo.tw
4. Hydroxymethylglutaryl-CoA Reductase Inhibitors/
5. cholesterol/
6. lipids/
7. (#1) AND (#2) AND (#3) AND (#4 OR #5 OR #6).
## Table 1 Characteristics of the eight included studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Target population</th>
<th>Dates of recruitment (years)</th>
<th>Mean follow-up (years)</th>
<th>Statin</th>
<th>Number of participants</th>
<th>Proportion of women (%)</th>
<th>Mean age (range, years)</th>
<th>Diabetes (%)</th>
<th>History of CVD (%)</th>
<th>Duration of post-trial follow-up</th>
<th>Difference in proportion taking statins post-trial (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primary prevention/primary care population</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>ALLHAT-LLT</td>
<td>Treated for high BP with high cholesterol</td>
<td>1994–1998</td>
<td>4.8</td>
<td>Pravastatin 40mg</td>
<td>10355</td>
<td>49</td>
<td>66 (55-NA)</td>
<td>35</td>
<td>11</td>
<td>4</td>
<td>NA</td>
</tr>
<tr>
<td>ASCOT-LLA</td>
<td>High BP and no history of CHD, with 3+ other CVD risk factors</td>
<td>1998–2000</td>
<td>3.2</td>
<td>Atorvastatin 10mg</td>
<td>10305</td>
<td>19</td>
<td>63 (40–79)</td>
<td>25</td>
<td>14</td>
<td>8.3</td>
<td>4</td>
</tr>
<tr>
<td>WOS-COPS</td>
<td>Men with high cholesterol and no history of myocardial infarction</td>
<td>1989–1991</td>
<td>4.8</td>
<td>Pravastatin 40mg</td>
<td>6595</td>
<td>0</td>
<td>55 (45–64)</td>
<td>1</td>
<td>8</td>
<td>15.1</td>
<td>4</td>
</tr>
<tr>
<td><strong>Secondary prevention/clinical population</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>ALERT</td>
<td>Renal or combined renal and pancreas transplant recipients at high risk of CVD</td>
<td>1996–1997</td>
<td>5.1</td>
<td>Fluvastatin 40mg</td>
<td>2102</td>
<td>34</td>
<td>50 (30–75)</td>
<td>19</td>
<td>19</td>
<td>1.6</td>
<td>0</td>
</tr>
<tr>
<td>SSSS</td>
<td>History of angina or myocardial infarction</td>
<td>1988–1989</td>
<td>5.2</td>
<td>Simvastatin 20–40mg</td>
<td>4444</td>
<td>19</td>
<td>60+ (35-70)</td>
<td>5</td>
<td>100</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>PROSPER</td>
<td>≥70 years with history of CVD or at high risk of CVD</td>
<td>1997–1999</td>
<td>3.2</td>
<td>Pravastatin 40mg</td>
<td>5804</td>
<td>52</td>
<td>75 (70–82)</td>
<td>11</td>
<td>44</td>
<td>5.4</td>
<td>NA</td>
</tr>
<tr>
<td>HPS</td>
<td>Coronary disease, other occlusive arterial disease, diabetes or treated for high BP</td>
<td>1994–1997</td>
<td>5.0</td>
<td>Simvastatin 40mg</td>
<td>20536</td>
<td>25</td>
<td>64 (40–80)</td>
<td>29</td>
<td>85</td>
<td>5.7</td>
<td>0</td>
</tr>
<tr>
<td>LIPID</td>
<td>Myocardial infarction or hospitalisation for unstable angina</td>
<td>1990–1992</td>
<td>5.6</td>
<td>Pravastatin 40mg</td>
<td>9014</td>
<td>17</td>
<td>62 (31–75)</td>
<td>&gt;99</td>
<td>10</td>
<td>1</td>
<td></td>
</tr>
</tbody>
</table>

Order of trials within primary prevention and secondary prevention order is from shortest post-trial follow-up to longest. BP, blood pressure; CVD, cardiovascular disease; NA, not available.
Figure 1  Direct (within-trial) and legacy (post-trial) effects of statins on cardiovascular disease mortality for eight trials. Note: Within primary and secondary prevention subgroups, studies are ordered by duration of follow-up.

Figure 2  Direct (within-trial) and legacy (post-trial) effects of statins on all-cause mortality for eight trials. Note: Within primary and secondary prevention subgroups, studies are ordered by duration of follow-up.
Figure 3  Random effects meta-analysis of relative risks for legacy (post-trial) effects of statins on cardiovascular disease mortality for eight trials. Note: Within primary and secondary prevention subgroups, studies are ordered by duration of follow-up. RR, relative risk.

Figure 4  Random effects meta-analysis of relative risks for legacy (post-trial) effects of statins on all-cause mortality for eight trials. Note: Within primary and secondary prevention subgroups, studies are ordered by duration of follow-up. RR, relative risk.
Validity assessment
Two authors (AN and LZ) extracted data on the following characteristics which may bias the estimated legacy effect: mean follow-up (years) and difference in proportion taking statins. Data were extracted separately for the within trial and post trial periods for each study.

Study selection and data abstraction
Two authors (AN and KB) independently checked the titles and abstracts of all citations identified through the database searches and forward citation search. Full text was obtained if either author judged the article potentially relevant. The same two authors then independently checked all the full-text articles for eligibility, resolving disagreements through discussion.

Two authors independently extracted clinical data (AN and LZ) using standardised forms, deciding disagreements through discussion with a third author (KB). We did not calculate formal measures of agreement to describe agreement between reviewers. The Cochrane Collaboration recommends against doing this, and instead recommends exploring reasons for any disagreement early on in the review process, which we did through discussion.

Figure 5  Random effects meta-analysis of HR for legacy (post-trial) effects of statins on cardiovascular disease mortality for four trials. Note: Within primary and secondary prevention subgroups, studies are ordered by duration of follow-up.

Figure 6  Random effects meta-analysis of HR for legacy (post-trial) effects of statins on all-cause mortality for four trials. Note: Within primary and secondary prevention subgroups, studies are ordered by duration of follow-up.
We extracted separate data on all-cause mortality and CVD-specific mortality for the within-trial and post-trial periods; the number of people at risk of each type of event at the start of the trial and at the start of the post-trial follow-up; the proportion of people taking statins within trial and post-trial; and the duration of follow-up within trial and post-trial. We attempted to extract differences in mean total cholesterol, but these were missing for at least one of the periods in the majority of studies. Further data on the original trials were obtained from Cholesterol Treatment Trialists Collaboration (CTTC).

**Figure 7** Random effects meta-analysis of HR for legacy (post-trial) effects of statins on cardiovascular disease mortality for three primary prevention trials. Note: Within primary and secondary prevention subgroups, Studies are ordered by duration of follow-up.

**Figure 8** Random effects meta-analysis of HR for legacy (post-trial) effects of statins on all-cause mortality for three primary prevention trials. Note: Within primary and secondary prevention subgroups, Studies are ordered by duration of follow-up.

**Statistical methods**

Summary statistics and plots for individual trials were generated using SAS V.9.4.

Meta-analytic models of post-trial data were built using STATA (V.14.2).

We built meta-analytic models for CVD mortality and all-cause mortality using reported number of events and number at risk for the post-trial period. Our principal summary measures for the models were relative risk and HR. We used adjusted relative risks where these were
reported, and calculated unadjusted relative risks where they were not. We built random effects models for the analysis. We assessed the heterogeneity of results using visual inspection of forest plots and I² statistics, and we conducted exploratory subgroup analysis using meta-regression to compare primary and secondary prevention trials. For the subgroup analysis, we tested for subgroup differences using a permutation test with 1000 permutations.22

We also built HR meta-analytic models for CVD mortality and all-cause mortality where these were reported in the primary studies. We undertook sensitivity analysis by restricting the model to primary prevention trials.

**Patient and public involvement**

Patients and or public were not involved in this meta-analysis of published data.

**RESULTS**

We identified 21 placebo-controlled RCTs of statins included in the CTTC for forward citation searching23–43 which retrieved 1520 abstracts (restricting search to reports which included an original trialist as an author). We identified a further 828 abstracts from Medline and Embase (searches to 16 June 2016), and after excluding duplicates, 1520 titles and abstracts were screened. We retrieved 61 papers for full-text review, 47 of which did not meet our selection criteria (see online supplementary eFigure 2). Reference searching of the remaining 14 studies identified one further study. Seven of the 15 studies used overlapping data: for each set of potentially overlapping reports, we chose the most recent report. This resulted in eight studies finally included in our review (table 1).

The original RCTs ranged in mean duration from 3.2 to 5.2 years, included trials of simvastatin, pravastatin, fluvastatin and atorvastatin, and their primary results were published between 1994 and 2003. Of the randomised participants in each trial, 0% to 52% were women, the mean age ranged from 50 to 75 years and 1% to 35% were diabetic. Between 8% and 100% had pre-existing CVD: three predominantly primary prevention/asymptomatic populations, and five predominantly secondary prevention/symptomatic populations. The difference in the proportion of people taking a statin in the randomised groups within the trial period (statin—placebo) ranged from 51% to 89%. HR (or relative risk ratio estimates when HR were unknown) for all-cause mortality and CVD-specific mortality within the trial period ranged from 0.70 to 1.02 and 0.64 to 0.96, respectively (see online supplementary eTable 1).

The post-trial follow-up ranged in mean duration from 1.6 to 15.1 years. The difference in proportion of people taking a statin in the post-trial period (for those originally randomised to statin minus those randomised to placebo) ranged from 0% to 4% (unknown for two studies). Collectively, the included studies reported on post-trial follow-up of 55732 people with 13781 deaths which occurred after the trials ended, of which 6685 were attributed to CVD. The HR (or relative risk estimates) for all-cause mortality and CVD-specific mortality ranged from 0.85 to 1.03 and 0.82 to 1.14, respectively (see online supplementary eTable 2).

**Individual trials: comparison of within-trial and post-trial effects**

The results for CVD-specific mortality for the individual trials are presented in supplementary eTable 1 and 2 and figure 1. Of the eight included trials, the six which demonstrated significant reductions in CVD mortality within the trial period (WOSCOPS, ALERT, SSSS, PROSPER, HPS and LIPID) showed less benefit in the post-trial period than in the trial period. The two trials without significant reduction in CVD mortality within the trial period (ALLHAT-LLA and ASCOT-LLA) showed a similar lack of evidence for benefit post-trial. In only one of the eight trials was there a significant reduction in CVD mortality for the post-trial period (WOSCOPS).

The results for all-cause mortality for the individual trials are presented in online supplementary eTables 1 and 2 and figure 2. Of the eight included trials, the four which demonstrated significant reduction in all-cause mortality within the trial period (WOSCOPS, SSSS, HPS and LIPID) showed less benefit in the post-trial period than in the trial period. Three trials without a significant reduction in mortality within the trial period (ALLHAT-LLA, ALERT and PROSPER) showed a similar lack of evidence for benefit post-trial. One trial (ASCOT-LLA) without a significant reduction in mortality within the trial period demonstrated more benefit in the post-trial period. In only two of the eight trials was there a significant reduction in all-cause mortality in the post-trial period (WOSCOPS and ASCOT-LLA).

**Post-trial meta-analysis**

The relative risk random effect meta-analysis using post-trial data from all eight studies is presented in figure 3 (CVD mortality) and figure 4 (all-cause mortality). Although there was no evidence overall of a post-trial (legacy) effect on CVD (p=0.15), there was some evidence of a legacy effect on all-cause mortality (p=0.01). In the exploratory subgroup analysis, there appeared to be a difference in the post-trial (legacy) effect of statins for primary prevention compared with secondary prevention studies for both CVD and for all-cause mortality. The pooled relative risk of CVD death post-trial for those originally allocated statin compared with placebo was 0.91 (0.84–0.98) for primary prevention trials, and 0.99 (0.94–1.05) for secondary prevention trials (permutation test p value for subgroup difference=0.15) (figure 3). The pooled relative risk of all-cause death post-trial for those originally allocated statin compared with placebo was 0.92 (0.88–0.96) for primary prevention trials and 0.99 (0.95–1.03) for secondary prevention trials (permutation test p value for subgroup difference=0.02) (figure 4).
The HR meta-analysis, using post-trial data from the four studies reporting HR, is presented in figure 5 (CVD mortality) and figure 6 (all-cause mortality). Similar to the meta-analysis of relative risks, there was no definite evidence of a post-trial (legacy) effect on CVD (p=0.09), but some evidence of a legacy effect on all-cause mortality (p=0.02). Pooling data from all four studies resulted in substantial heterogeneity between studies (I²=40.7% for CVD mortality and 42.3% for all-cause mortality). Restricting meta-analysis to the three primary prevention trials resulted in very low heterogeneity between studies (I²=0.0% for CVD mortality and 8.1% for all-cause mortality), and these results are presented in figure 7 (CVD mortality) and figure 8 (all-cause mortality). In the three primary prevention trials, the pooled HR for CVD death post-trial for those originally allocated statin compared with placebo was 0.87 (0.79 to 0.95, p=0.003), and for all-cause death, it was 0.90 (0.85 to 0.96, p=0.001).

DISCUSSION

We identified eight large randomised trials which had usable post-trial data to assess legacy effects on mortality outcomes. The direct effects of the statins on mortality reduction observed during the trials were much larger than potential legacy effects observed post-trial, which suggests the rhetoric on legacy effects for statins in general may not reflect the empirical evidence. WOSCOPS was the only trial to show a possible post-trial legacy effect on all-cause and CVD-specific mortality. When we pooled data from all eight studies, we found no evidence overall of legacy effects on CVD mortality, but some evidence of possible legacy effects on all-cause mortality. In the exploratory subgroup analysis, there was some evidence of a difference in results for primary prevention compared with secondary prevention. Considering these subgroups separately, we found no evidence of legacy effects following secondary prevention trials, suggesting the importance of long-term/lifelong prevention in these patients. We found evidence of possible post-trial legacy effects only when statins were started for primary prevention—these effects were observed on both CVD mortality (HR=0.87, p=0.003) and all-cause mortality (HR=0.90, p<0.001) (figure 3C,D). Participants originally randomised to placebo in two of the primary prevention trials (WOSCOPS and ASCOT-LLA) had 4% lower rates of using a statin in the first years post-trial, which will exaggerate the estimated legacy effect (bias away from the null), but this difference is unlikely to account for all the observed post-trial benefit (whether there was a difference in statin use post-trial in ALLHAT is not known). The observed post-trial reductions in CVD and all-cause mortality may potentially represent real legacy effects of statins for populations similar to those at the time of recruitment into these studies. There may be a higher likelihood of observing legacy effects for statins when this is started for primary prevention, rather than for secondary prevention.

Our sensitive search strategy means this study is likely to have included all published follow-up reports of the major placebo-controlled statin trials, including recent follow-up reports for two of the studies (WOSCAPS and LIPID). However, we did not assess for publication bias and it is possible that unpublished follow-up reports may exist that we are unaware of. We did not assess risk of bias for the included studies, but this has been assessed by others for the original trial reports, including very recently, and the included studies were generally found to be high quality. Although we believe the post-trial period is the best period to analyse for detection of legacy effects, these data are no longer a randomised comparison: some patients randomised to the statin would have been saved from dying, whereas some patients in the placebo group were not. Hence, there are additional survivors in the statin group at the beginning of post-trial follow-up who are also likely to be at higher risk of CVD than survivors in the placebo group. These differences would tend to bias our results towards the null, and mean that legacy effects may be larger than we estimated. The main limitation of our report is that because our findings are based on aggregate data, we are unable to assess the effects of whether or not an individual was treated with statins during the post-trial period, and for how long, as well as their cardiovascular risk factor levels and other potential confounders. For example, although we found evidence of possible legacy effects in primary prevention, these are largely driven by WOSCOPS which was undertaken in all male participants. If there are sex-specific effects for legacy effects, it may be the fact that all participants in WOSCOPS were male, and not that they had no history of CVD, that is the more important determinant. Similarly, participants in WOSCOPS had the lowest percentage taking statins in the post-trial period out of all the studies where this was measured (39% of active and 55% of placebo participants were taking statins at 5 years post-trial). This comparative absence of direct statin treatment effects in the post-trial period may be the more important determinant for observing a legacy effect.

We did not examine evidence of possible legacy effects on other outcomes such as non-fatal CVD, or for different post-trial follow-up times within each study, or for the same post-trial follow-up times between studies. We are aware of four other meta-analyses of data from long-term follow-up after placebo-controlled trials of lipid-lowering treatment. In three of these reports, the focus appears to have been on persistence of survival benefit, with comparison of event rates from time of randomisation, rather than post-trial legacy effect. The other meta-analysis reported separate results for the post-trial period using data from earlier follow-up reports of six of our included trials. That report found evidence of post-trial reduction in CVD mortality and all-cause mortality at 2 years, and evidence for a reduction in major coronary events at both 2 years and over the total post-trial periods. The authors did not explore possible causes of
heterogeneity for the post-trial models such as whether the primary trial was for primary or secondary prevention.

Published trial evidence supports the hypothesis that lowering cholesterol with a statin drug reduces cardiovascular events. Currently, the principle of using absolute risk to guide treatment decisions (as recommended by guidelines) is that treatment is prioritised for those at highest short-term risk, and people at low short-term risk are not treated. Data on the efficacy and safety of statins has led to treatment thresholds being lowered: in the UK, the threshold was lowered from >20% to >10% 10-year risk of CVD; in the USA, the threshold is 10% 10-year risk of CVD, with statins also recommended for selected patients with 7.5%–10% 10-year risk. However, as short-term risk is largely driven by age, younger people are unlikely to qualify for statins even with these lowered thresholds. For example, a recent report found that in the absence of smoking or raised blood pressure, a 10-year risk of CVD above 5% was infrequent in women younger than 50 and men younger than 40 years resident in the USA. Exploratory subgroup analysis in our study found evidence of possible legacy effects of statins following the primary prevention trials, which warrant further investigation. However, we note that the participants in WOSCOPS, ALLHAT and ASCOT-IIA had elevated levels of CVD risk factors (see table 1). Indeed, the majority of these people were likely to have been well above current treatment thresholds at the time of trial entry, and people with similar risk levels would now be recommended to start lifelong lipid-lowering treatment. For example, the proportion of people who had died of CVD by the end of the trial in the placebo group after 3.3 years in ASCOT, 4.8 years in ALLHAT and 4.9 years in WOSCOPS was 3%, 11% and 2%, respectively. Legacy effects in these settings serve to emphasise the benefits of starting long-term primary prevention treatment early rather than later among people at high short-term risk. It does not provide evidence to support earlier treatment for people who have lower short-term risk than current treatment thresholds.

Advocates of early intervention argue that people who are at risk of disease in the long term, but currently displaying no symptoms or signs of disease at low calculated short-term risk, should also be started on long-term treatment at an early age. But deciding when, and if, to intervene in these people is much less straightforward. The legacy effect hypothesis for statins—that the earlier you start, the lower your risk of a CVD event in the long term—has not been tested directly in an RCT comparing statins commencement at an earlier versus later age, and such a trial is unlikely to eventuate. Indirect evidence from post-trial follow-up after the large statin trials is likely the next best way to investigate this.

CONCLUSION

In this analysis of eight long-term randomised trials, we found possible post-trial legacy effects of statins on CVD mortality and all-cause mortality for primary prevention. Although the post-trial relative benefits were clearly smaller than those observed within trials, the increasing risk with age may mean that the absolute benefits are similar. Analysis of individual patient data from follow-up studies after placebo-controlled RCTs in lower-risk populations may provide more definitive evidence on whether early treatment of subclinical atherosclerosis is likely to be beneficial.


