The benefits of statins in people without established cardiovascular disease but with cardiovascular risk factors: meta-analysis of randomised controlled trials

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ABSTRACT

Objectives To investigate whether statins reduce all cause mortality and major coronary and cerebrovascular events in people without established cardiovascular disease but with cardiovascular risk factors, and whether these effects are similar in men and women, in young and older people, and in people with and without diabetes mellitus.

Design Meta-analysis of randomised trials.

Data sources Cochrane controlled trials register, Embase, and Medline.

Data abstraction Two independent investigators identified studies on the clinical effects of statins compared with a placebo or control group and with follow-up of at least one year, at least 80% or more participants with cardiovascular risk factors, and outcome data on mortality and major cardiovascular disease events. Heterogeneity was assessed using the Q and I² statistics. Publication bias was assessed by visual examination of funnel plots and the Egger regression test.

Results 10 trials enrolled a total of 70,388 people, of whom 23,681 (34%) were women and 16,078 (23%) had diabetes mellitus. Mean follow-up was 4.1 years. Treatment with statins significantly reduced the risk of all cause mortality (odds ratio 0.88, 95% confidence interval 0.81 to 0.96), major coronary events (0.70, 0.61 to 0.81), and major cerebrovascular events (0.81, 0.71 to 0.93). No evidence of an increased risk of cancer was observed. There was no significant heterogeneity of the treatment effect in clinical subgroups.

Conclusion In patients without established cardiovascular disease but with cardiovascular risk factors, statin use was associated with significantly improved survival and large reductions in the risk of major cardiovascular events.

INTRODUCTION

Cardiovascular disease is the leading cause of death and disability in the Western world and contributes substantially to healthcare budgets. Several clinical trials and meta-analyses have shown the beneficial effects of lipid lowering treatment using hydroxymethyl glutaryl coenzyme A reductase inhibitors (statins) in reducing mortality and cardiovascular morbidity in patients with established cardiovascular disease. Statins therefore have a place in the secondary prevention of cardiovascular disease.

The use of statins in patients without established cardiovascular disease (that is, primary prevention) and at relatively low risk has important public health implications. To date research has provided ambiguous answers. In addition, the reliability of treatment in older people (>65 years), women, and those with diabetes mellitus is uncertain, mainly because of a lack of data or inconsistent findings within these clinically defined groups. Most meta-analyses have been carried out on published tabular data and failed to provide consistent answers on treatment effect in these subgroups.

We carried out a meta-analysis of randomised trials that focused on primary prevention to determine whether statins reduce all cause mortality and the incidence of major coronary and cerebrovascular events in people without established cardiovascular disease but with cardiovascular risk factors. We also assessed whether these effects differed by sex, age, and the presence of diabetes.

METHODS

We followed the quality of reporting of meta-analysis guidelines. We searched the Cochrane Central Register of Controlled Trials, Medline (1990-November 2008), Embase (1980-November 2008), DARE, and the ACP Journal Club for randomised clinical trials that compared statins with a control group in people without established cardiovascular disease but with cardiovascular risk factors. We identified relevant studies using the MeSH terms “HMG-CoA reductase inhibitor”, “atorvastatin”, “simvastatin”, “pravastatin”, “fluvastatin”, “rosvastatin”, or “lovastatin”, and “cardiovascular disease”, “coronary heart disease”, “cerebrovascular disease”, or “myocardial infarction”.
and “cholesterol”, “LDL” [low density lipoprotein], “HDL” [high density lipoprotein], or “triglycerides”, and primary prevention restricted to randomised controlled trials or meta-analyses. In addition we examined the reference lists and related links of retrieved articles in PubMed to detect studies potentially eligible for inclusion.

**Study selection**

We included studies if they were randomised trials of statins compared with controls (placebo, active control, or usual care), had a mean follow-up of at least one year, reported on mortality or cardiovascular disease events as primary outcomes, and included at least 80% of people without established cardiovascular disease or reported data separately on a sole primary prevention group and provided specific numbers for patients and events in that group. Eight studies were excluded that primarily investigated statin related non-clinical and intermediate surrogate end points such as changes in the thickness of the carotid intima media and lipid levels that collectively contributed fewer than 50 clinical events.26-29 We also excluded one study in patients with renal transplants because of the specific nature of that population,24 and three studies with design problems, fewer than 20 events overall, and insufficient follow-up.25-27 Our study therefore focused on people without established cardiovascular disease but with cardiovascular risk factors.

**Validity assessment**

Our search identified 1230 studies, of which 10 fulfilled our inclusion criteria.1-10 Figure 1 summarises the results of the search. We evaluated suitable trials for concealment of treatment allocation, performance of the analysis according to the intention to treat principle, and completeness of follow-up. The Jadad scale was used to score study quality (range 0-5, higher scores indicating better quality).28 Study quality was sufficient (≥4) for all included randomised clinical trials.

**Data abstraction**

From each study two investigators separately extracted information on trial characteristics, patient data, outcome measures, and study quality using a standardised protocol and reporting document. Disagreements were resolved by consensus.

**Subgroup analysis**

We searched the papers for data on clinically defined subgroups. The Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial (ALLHAT)19 presented data on our prespecified subgroup. The other studies did not publish results stratified by age (<65 or ≥65 years), sex, or diabetes. To obtain data for these stratified groups we sent an electronic sheet with data fields to the principal investigators of these studies, requesting the number of events and number of patients in the treatment and placebo groups. We obtained data on subgroups for five trials.1-3,6,8 Subgroup analyses were therefore done in six studies.1-3,6,8 Not all end points were recorded in these studies.

**End points**

The primary end point of our meta-analysis was all cause mortality. Secondary end points were the composite of major coronary events defined as death from coronary heart disease and non-fatal myocardial infarction, and the composite of major cerebrovascular events defined as fatal and non-fatal stroke. We also assessed death from coronary heart disease, non-fatal myocardial infarction, revascularisations (percutaneous coronary intervention or coronary artery bypass graft), and cancer (fatal and non-fatal). The clinical outcomes evaluated in the subgroup analysis (data should be reported in two or more studies) were all cause mortality, major coronary events, major cerebrovascular events, and cancer.

**Quantitative data synthesis**

For each trial we calculated the summary odds ratios and 95% confidence intervals for the clinical outcomes. We pooled studies using both fixed effect and random effects models.29 A random effects model makes the assumption that individual studies are estimating different treatment effects. Our conclusions were drawn from the results of the random effects model. We were unable to exclude a small proportion of secondary prevention patients from the West of Scotland Coronary Prevention Study (1069/6395; WOSCOPS),19 ALLHAT (1470/10 355),19 and the Anglo-Scandinavian Cardiac Outcomes Trial lipid lowering arm (1906/10 305; ASCOT-LLA).10 and these therefore constitute about 6% of our study population [4445/70 388].10 In a separate analysis we verified whether our results remained consistent after exclusion of these studies. We also investigated whether our results differed when we used the original study results from ASCOT without extended follow-up.10-12

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**Fig 1 | Flow of article selection in trial**
We assessed the results for heterogeneity in the main analysis and subgroup analysis by examining the forest plots and then calculating a Q statistic, which we compared with a χ² distribution, and the I² index. The Q test indicates the statistical significance of the homogeneity hypothesis and the I² index measures the extent of the heterogeneity. We considered the results for heterogeneity to be significant at P<0.10 (two sided). Publication bias was assessed for the main end points by visually examining for funnel plot

Table 1 | Characteristics of included trials

| Characteristic | PROSPER* 2002w7 | ALLHAT-LLT 2002w7 | ASCOT-LLA 2003w10 | HPS* 2003w10 | CARDS 2004w4 | ASPEN* 2006w7 | MEGA 2006w12 | JUPITER 2008w13 |
|----------------|------------------|--------------------|--------------------|-------------|-------------|--------------|--------------|---------------|
| Target population | Elderly people with cardiovascular risk factors | People with hypertension, average or lower cholesterol levels, and at least one coronary heart disease risk factor | People with hypertension, average or lower cholesterol levels, and at least three other risk factors | People with diabetes | People with diabetes and low density lipoprotein cholesterol (no history of cardiovascular disease) | People with diabetes and low density lipoprotein cholesterol levels below guideline targets | People with hypercholesterolaemia and no history of coronary heart disease or stroke | People without vascular disease, low density lipoprotein cholesterol <130 mg/dl, and high sensitivity C reactive protein > 2.0 mg/l |
| Design | Randomised double blind placebo controlled trial | Randomised double blind placebo controlled trial | Randomised double blind placebo controlled trial | Randomised double blind placebo controlled trial | Randomised double blind placebo controlled trial | Randomised double blind placebo controlled trial | Randomised double blind placebo controlled trial | Randomised double blind placebo controlled trial |
| No of participants (statin/control) | 6595 (3302/3293) | 6605 (3304/3301) | 3239 (1585/1654) | 10355 (5170/5185) | 10305 (5168/5137) | 2912 (1455/1457) | 2838 (1428/1410) | 1905 (959/946) | 7832 (3866/3966) | 17802 (8901/8901) |
| Mean follow-up (years) | 4.9† | 5.2 | 3.2 | 4.8 | 5.5† | 4.8 | 3.9† | 4.0† | 5.3 | 1.9† |
| Drug | Pravastatin | Lovastatin | Pravastatin | Pravastatin | Atorvastatin | Simvastatin | Atorvastatin | Atorvastatin | Pravastatin | Rosuvastatin |
| Dose (mg/day) | 40 | 20-40 | 40 | 20-40 | 10 | 40 | 10 | 10 | 10 | 20 |
| Mean age (range) (years) | 55.3 (45-64) | 58 (45-73) | 75 (70-82) | 66.4 (51-81) | 63.1 (40-79) | NA (40-80) | 61.5 (40-75) | 60.5 (40-75) | 58.3 (40-70) | 661 (60-71) |
| Women (%) | 0 | 15 | 58‡ | 49 | 18.9 | NA | 32 | 38 | 68.4 | 37.9 |
| With diabetes (%) | 1 | 3.8 | 12.2‡ | 34.4 | 24.3 | 100 | 100 | 100 | 21 | 0 |
| Current smoker (%) | 44 | 13 | 33.4‡ | 23.3 | 33.2 | NA | 22 | 12 | 21 | 16 |
| Hypertension (%) | 16 | 22 | 71.6‡ | 89.9 | 80.3 | NA | 84 | 52 | 42 | 0 |
| Mean body mass index | 26 | 26.8 | 27‡ | 29.9 | 28.6 | NA | 28.7 | 28.9 | 23.8 | 28.4‡ |
| Mean systolic blood pressure (mm Hg) | 136 | 138 | 156.6‡ | 145 | 164.2 | NA | 144 | 133 | 132 | 134‡ |
| Mean diastolic blood pressure (mm Hg) | 84 | 78 | 85.2‡ | 84 | 95 | NA | 83 | 77.1 | 78.4 | 80‡ |
| Baseline lipid levels (mmol/l) (% change): | | | | | | | | | | |
| Total cholesterol | 7.0 (–20.0) | 5.7 (–19.3) | 5.7 (NA) | 5.9 (–9.6) | 5.5 (–18.2) | NA | 5.4 (–21.8) | 5.0 (–19.8) | 6.3 (–11.0) | 4.8 (NA) |
| Low density lipoprotein cholesterol | 5.0 (–26.0) | 3.9 (–26.5) | 3.8 (NA) | 3.8 (–16.7) | 3.4 (–27.6) | NA | 3.0 (–33.9) | 3.0 (–30.5) | 4.0 (–18.0) | 2.8 (NA) |
| High density lipoprotein cholesterol | 1.1 (5.0) | 1.0 (4.8) | 1.3 (NA) | 1.2 (0.9) | 1.3 (1.5) | NA | 1.4 (4.0) | 1.2 (1.9) | 1.5 (5.0) | 1.3 (NA) |
| Triglycerides | 1.8 (–12.0) | 1.7 (–12.7) | 1.5 (NA) | 1.7 (0.0) | 1.7 (–12.6) | NA | 2.0 (–15.9) | 1.6 (–4.7) | 1.4 (–7.0) | 1.3 (NA) |

NA=not available; WOSCOPS=West of Scotland Coronary Prevention Study; AFCAPS/TexCAPS=Air Force/Texas Coronary Atherosclerosis Prevention Study; PROSPER=Prospective Study of Pravastatin in the Elderly at Risk; ALLHAT-LLT=Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial; ASCOT-LLA=Anglo-Scandinavian Cardiac Outcomes Trial—Lipid Lowering Arm; HPS=Heart Protection Study (diabetic subgroup publication); CARDS=Collaborative Atorvastatin Diabetes Study; ASPEN=Atorvastatin Study for Prevention of Coronary Heart Disease Endpoints in Non-Inulin-Dependent Diabetes Mellitus; MEGA=Management of Elevated Cholesterol in the Primary Prevention Group of Adult Japanese; JUPITER=Justification for the Use of Statins in Prevention: an Intervention Trial Evaluating Rosuvastatin.

*Primary prevention subgroup data used.
†Median; in ASCOT-LLA data were from extended observations trial.‡Data from baseline characteristics publication of PROSPER.¹³
Mortality, coronary events, and cerebrovascular events

During a mean follow-up of 4.1 years, 5.7% (1925/33 793) of participants died in the control group compared with 5.1% (1725/33 683) in the statin group. Statin therapy was therefore associated with a 12% risk reduction in all cause mortality compared with the control (odds ratio 0.88, 95% confidence interval 0.81 to 0.96; fig 2 and table 2). The annual rate for all cause mortality with placebo in our study was 1.4% (fig 2).

Overall, 5.4% (1266/23 946) of participants in the control group had a major coronary event compared with 4.1% (996/23 823) in the statin group, a 30% risk reduction (odds ratio 0.70, 95% confidence interval 0.61 to 0.81). The annual rate for major coronary events with placebo in our study was 1.4% (fig 2).

Overall, 2.3% (767/33 793) of participants in the control group had a major cerebrovascular event compared with 1.9% (627/33 683) in the statin group, a 19% risk reduction.
The current meta-analysis totalled 70388 participants without established cardiovascular disease but with cardiovascular risk factors who were randomised to statin therapy or control. Statin therapy was associated with a significant risk reduction in all cause mortality of 12%, in major coronary events of 30%, and in major cerebrovascular events of 19%. Moreover, statin use was not associated with an increased risk of cancer. These results are in line with those previously published on the effects of statins in secondary prevention.5,6

Our meta-analysis differs from earlier analyses in several ways.13,14 We were able to include several recently published studies targeted at primary prevention that enrolled a large number of women and people with diabetes.6,15-21 For example, the Management of Elevated Cholesterol in the Primary Prevention Group of Adult Japanese trial (MEGA)22 comprised a large number of women (68%, 5356/7832), and we were able to obtain subgroup data. Additionally, the Atorvastatin Study for Prevention of Coronary Heart Disease Endpoints in Non-Insulin-Dependent Diabetes Mellitus study (ASPEN)23 was carried out in a large group of people with type 2 diabetes (n=1905) who did not have established cardiovascular disease. We also included data from the recently published JUPITER trial,24 totalling 17802 participants with no apparent vascular disease, low density lipoprotein cholesterol levels less than 3.4 mmol/l, and increased levels of high sensitivity C reactive protein (>2.0 mg/l).25 As our study is based on such large numbers, this meta-analysis, including the subgroups, has significant statistical power. Previously, only the JUPITER trial showed improved survival associated with statin therapy or control. Statin therapy was associated with a significant risk reduction in all cause mortality of 12%, in major coronary events of 30%, and in major cerebrovascular events of 19%. Moreover, statin use was not associated with an increased risk of cancer. These results are in line with those previously published on the effects of statins in secondary prevention.5,6

Table 2 | Treatment effects of statin therapy. Values are odds ratios (95% confidence intervals)

| Source                  | All cause mortality | Major coronary events | Major cerebrovascular events | Coronary heart disease mortality | Non-fatal myocardial infarction | Revascularisations | Fatal or non-fatal cancer |
|-------------------------|---------------------|-----------------------|-----------------------------|---------------------------------|---------------------------------|--------------------|--------------------------|
| WOSCOPS5,6              | 0.78 (0.60 to 1.01) | 0.68 (0.56 to 0.83)   | 0.90 (0.60 to 1.34)         | 0.73 (0.48 to 1.11)             | 0.69 (0.55 to 0.85)             | 0.63 (0.44 to 0.90) | 1.09 (0.84 to 1.43)     |
| AFCAPS/TexCAPS5,6       | 1.04 (0.76 to 1.43) | 0.61 (0.45 to 0.83)   | 0.82 (0.40 to 1.67)*        | 0.73 (0.34 to 1.60)             | NR                             | 0.69 (0.55 to 0.86) | 0.97 (0.81 to 1.16)     |
| PROSPER6                | 0.98 (0.79 to 1.21)*| 0.90 (0.70 to 1.15)   | 1.03 (0.72 to 1.47)         | NR                             | NR                             | 0.82 (0.54 to 1.25)* | NR                       |
| ALLHAT-LLA7             | 0.99 (0.88 to 1.11) | 0.90 (0.78 to 1.04)   | 0.90 (0.75 to 1.09)         | 0.99 (0.79 to 1.24)             | NR                             | NR                 | 1.03 (0.89 to 1.19)     |
| ASCOT-LLA8              | 0.85 (0.73 to 0.97) | 0.64 (0.52 to 0.78)   | 0.77 (0.63 to 0.95)         | NR                             | NR                             | NR                 | NR                       |
| HPS5,6                  | NR                  | 0.57 (0.41 to 0.79)*  | NR                          | NR                             | NR                             | NR                 | NR                       |
| CARDS5,6                | 0.72 (0.51 to 1.02) | 0.65 (0.44 to 0.97)   | 0.52 (0.31 to 0.90)         | 0.74 (0.40 to 1.36)             | 0.59 (0.36 to 0.98)             | 0.69 (0.41 to 1.17) | 0.65 (0.37 to 1.16)     |
| ASPEN5,6                | 1.06 (0.69 to 1.64) | NR                    | 0.92 (0.54 to 1.56)         | NR                             | NR                             | 0.92 (0.60 to 1.40) | NR                       |
| MEGA2                   | 0.71 (0.50 to 1.00) | 0.55 (0.33 to 0.91)   | 0.83 (0.57 to 1.20)         | 0.55 (0.22 to 1.38)             | 0.55 (0.30 to 1.00)             | 0.60 (0.40 to 0.90) | 0.97 (0.75 to 1.25)     |
| JUPITER8                | 0.80 (0.66 to 0.96) | NR                    | 0.51 (0.34 to 0.78)         | NR                             | 0.35 (0.22 to 0.58)             | 0.54 (0.40 to 0.72) | 0.89 (0.77 to 1.04)     |
| All trials fixed effects model | 0.90 (0.84 to 0.96) | 0.74 (0.68 to 0.81)   | 0.82 (0.74 to 0.91)         | 0.88 (0.73 to 1.05)             | 0.61 (0.52 to 0.73)             | 0.67 (0.59 to 0.76) | 0.97 (0.89 to 1.05)     |
| All trials random effects model | 0.88 (0.81 to 0.96) | 0.70 (0.61 to 0.81)   | 0.81 (0.71 to 0.93)         | 0.88 (0.73 to 1.05)             | 0.56 (0.41 to 0.76)             | 0.67 (0.59 to 0.76) | 0.97 (0.89 to 1.05)     |

Heterogeneity:

| Q statistic | 0.20† | 0.02‡ | 0.23 | 0.49 | 0.11 | 0.48 | 0.61 |
|-------------|-------|-------|------|------|------|------|------|
| I² index    | Low (27%) | Moderate (60%) | Low (24%) | Low (0%) | Moderate (50%) | Low (0%) | Low (0%) |

NR; not reported. See footnote to table 1 for full titles of studies.

*Data from Thavendiranathan et al.15 Fixed effect and random effect models in meta-analysis gave almost identical results, making important statistical heterogeneity unlikely.

†No data in primary prevention group (n=9323).16

‡Significant heterogeneity; however, a positive trend of statin therapy is observed in all trials, only of different magnitude (no neutral or negative trials).
statin use in high risk participants, but it is clear from the current analysis that a mortality benefit is a shared characteristic of long term statin use in people without previous cardiovascular disease. The currently observed benefit, a 12% risk reduction in mortality, may even be an underestimation of the true effect because subsequent death after a morbid cardiovascular event was not always considered in individual trials.

The numbers and duration of follow-up of our study allow for relatively strong inferences on risk of cancer with long term statin use. We found no evidence for an increased risk of cancer, fatal or non-fatal. One of the trials (Prospective Study of Pravastatin in the Elderly at Risk; PROSPER) did report an increased risk of cancer with use of statins among men and women older than 70. Although our results show that statins do not seem to increase the risk of cancer, longer follow-up would be helpful to determine whether new cancer events could occur with time. This is especially critical when statins are used in primary prevention. Follow-up of patients in WOSCOPS for 10 years did not show higher rates of malignancies. Concerns might remain about the higher risk of cancer in elderly patients (70-82 years) as in PROSPER, and further follow-up studies in such patients are required. Although this meta-analysis cannot fully remove that uncertainty, it confirms that the risk of cancer is not
WHAT IS ALREADY KNOWN ON THIS TOPIC
Statins are effective in patients with established cardiovascular disease (secondary prevention) but whether the benefits apply to primary prevention is unknown.
Research has provided ambiguous answers on statin use in people at relatively lower risk.
Furthermore, the efficacy of statins in subgroups of people aged more than 65, women, and those with diabetes mellitus is debated.

WHAT THIS STUDY ADDS
Statins improve survival and reduce the risk of major cardiovascular and cerebrovascular events in people without established cardiovascular disease.
No significant differences in treatment effect of statins were observed in clinically defined groups for age, sex, and diabetes status.
People at increased risk for cardiovascular disease should not be denied the relative benefits of long term statin use.

increased in middle aged patients. Tolerance to statins is also important to tackle in primary prevention. Side effects such as an increase in creatine kinase levels and myopathy have been reported relatively frequently, but rhabdomyolysis and hepatotoxicity are rare.5

Lastly, by contacting principal investigators of each trial we were able to obtain data on clinically defined subgroups. This allowed us to draw meaningful inferences on treatment effects in large numbers of women, older people, and people with diabetes. Although there is little reason to suspect different treatment effects between such groups from a pathophysiological standpoint, it is reassuring that no significant treatment heterogeneity was found between the sexes, in elderly and young people, and between people with and without diabetes.

Limitations of the study
Some limitations of our study need to be mentioned. Firstly, we included three trials in the analyses that had recruited a small proportion of patients (about 6%) with clinical cardiovascular disease.30w7 w9 Exclusion of these trials did not affect the outcome of our analyses. Secondly, the dose and type of statin differed between included trials. Depending on the statin and the dose, some treatment regimens may be more effective in lowering lipid levels. However, according to guidelines from the Adult Treatment Panel III, the statins included in our meta-analysis at their respective doses have similar clinical efficacy.9 Thirdly, the included trials represented participants with a clinically heterogeneous level of risk (although statistical heterogeneity was low). The benefit observed in the pooled estimate of treatment effect could be of different magnitude depending on the level of risk. However, exclusion of the studies with a small proportion of patients at higher risk did not influence the outcome of the analysis because our subgroup analysis indicated no heterogeneity in clinically defined groups such as elderly participants or those with diabetes mellitus who are at relatively higher risk. Such a risk dependent effect seems unlikely.

Clinical implications
Our meta-analysis shows that the relative risk reduction from long term statin use in a primary care setting is comparable to that observed in secondary prevention. Our findings confirm the results of JUPITER5 regarding the beneficial effect of statins on survival across a broader range of patients (n=70 388) at different levels of risk, and show that there is no significant difference in treatment benefit across a range of clinically defined groups (men and women, elderly people, and those with diabetes). Although our study population comprised participants without established cardiovascular disease, the pooled risk was high. The overall annual mortality was in the range of 1.4%, and fatal as well as non-fatal cardiac and cerebrovascular events occurred at an annual rate of about 1.1% and 0.6%, respectively. This is not too different from the event rates reported in trials of patients at relatively low risk in secondary prevention—for example, the European trial on reduction of cardiac events with perindopril in stable coronary artery disease and the Prevention of Events with Angiotensin-Converting Enzyme inhibition trial (PEACE).16 37 Statin based secondary prevention is considered mandatory in participants in PEACE. Still, the absolute overall treatment benefit observed in the current study population would certainly be less than 1%, and significant numbers of participants would need to be treated to prevent one event. From the currently pooled data it is not possible to exactly define one group of people who would benefit most from long term statin use. From current risk scoring systems, as well from current data, it is obvious that older men (>65 years) with risk factors, or older women with diabetes and risk factors, constitute the highest risk group. In view of the large treatment effects described here, it is likely that a considerable number of such people would benefit from long term statin use at reasonable costs. The correct identification of such people remains a challenge and, in addition to the assessment on the future cardiovascular risk based on standard cardiovascular risk factors, auxiliary diagnostic or prognostic assessments to improve risk prediction could be useful to identify these men and women more accurately. Given the favourable effects of long term statin treatment it would be wrong to deny these benefits to people at increased risk for cardiovascular disease.

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Competing interests: AMG is a consultant for Genentech, Kowa, Martek, Merck, and Merck/Schering-Plough, and serves on the board of directors for Aegeron and Arisaph. He is a member of DuPont’s health advisory board and serves on the data safety monitoring board for Novartis. JS carried out consultancy work and receives support for research from Bristol-Myers Squibb. RJW receives support for research from Bristol-Myers Squibb. HN has received travel grants and speaking honorariums from Sanofi. RHK has received research fees and speaking honorariums from Pfizer. PR has received research grant support from the National Heart and Lung Blood Institute, the National Cancer Institute, the Donald W Reynolds Foundation, the Leducq Foundation, Astra-Zeneca, Novartis, Merck, Abbott, Roche, and Sanofi-Aventis; consulting fees and lecture fees from Astra-Zeneca, Novartis, Merck-Schering Plough, Sanofi-
Aventis, ISIS, and Vascular Biogenics, and is listed as a co-inventor on patents held by the Brigham and Women's Hospital that relate to the use of inflammatory biomarkers in cardiovascular disease. These patents have been licensed to Siemens and Astra-Zeneca.

Ethical approval: Not required.

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