Beneficial effects of fluvastatin following percutaneous coronary intervention in patients with unstable and stable angina: results from the Lescol intervention prevention study (LIPS)

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Notes
Beneficial effects of fluvastatin following percutaneous coronary intervention in patients with unstable and stable angina: results from the Lescol intervention prevention study (LIPS)

C H Lee, P de Feyter, P W Serruys, F Saia, P A Lemos, D Goedhart, P R Soares, V A W M Umans, M Ciccone, M Cortellaro

Aims: To investigate the effect on risk of major adverse cardiac events (MACE) of lipid lowering treatment with fluvastatin 80 mg/day after a first percutaneous coronary intervention in patients with stable and unstable angina.

Method and results: This prespecified subgroup analysis of the LIPS (Lescol intervention prevention study) analysed 1658 patients with documented diagnosis; 824 had unstable angina (417 randomly assigned to fluvastatin, 407 to placebo) and 834 had stable angina (including silent ischaemia; fluvastatin, 418; placebo, 416). Median follow up was 3.9 years. There was no significant effect of anginal status on long term risk of MACE. Fluvastatin treatment reduced the risk of MACE by 28% compared with placebo (p = 0.03) among patients with unstable angina, with no difference between patients with unstable and patients with unstable angina (relative risk 1.07, 95% confidence interval 0.87 to 1.30, p = 0.53). Fluvastatin reduced coronary atherosclerotic events (MACE excluding restenosis) by 36% (p = 0.006) among patients with unstable angina and 31% (p = 0.02) among patients with stable angina. Fluvastatin caused similar reductions in total cholesterol and low density lipoprotein cholesterol concentrations in both patient groups.

Conclusion: Treatment with fluvastatin 80 mg/day produced significant reductions in MACE and coronary atherosclerotic events after percutaneous coronary intervention in patients with average cholesterol concentrations. The beneficial effects of fluvastatin are observed in patients with unstable or stable angina alike.

Recent evidence suggests that coronary artery disease is a systemic disorder. Although percutaneous coronary intervention (PCI) effectively improves symptoms by restoring coronary perfusion, the underlying systemic inflammatory and prothrombotic substrates are not attenuated and so life long medication is necessary after revascularisation. In particular, lipid lowering treatment with 3-hydroxy-3-methylglutaryl coenzyme A reductase inhibitors (statins) has been proved to improve survival in patients with coronary artery disease. The underlying mechanisms for the beneficial effects of statins in patients with coronary artery disease are not fully understood but are likely to involve reductions in lipid concentrations, improvement in endothelial function, anti-inflammatory action, or antithrombotic actions, or a combination of these effects.

The recent LIPS (Lescol intervention prevention study) showed that early initiation of lipid lowering treatment with fluvastatin at hospital discharge significantly reduces the incidence of major adverse cardiac events (MACE) among patients after PCI with average cholesterol concentrations. Fluvastatin treatment was well tolerated and associated with no significant side effects. In the present study, patients were classified at entry as having either unstable angina or stable angina, conditions that exhibit major differences in pathophysiology, prognosis, and response to treatment. In contrast to stable angina, which is caused by a fixed stenotic coronary lesion that impairs myocardial perfusion, unstable angina is usually caused by the rupture or erosion of an atherosclerotic plaque, leading to a reaction cascade that results in a thrombotic milieu. There is relatively little information regarding the effect of statin treatment in patients with unstable angina; hence, this study provides important insight into the potential benefits of statins in this patient group.

The objective of the present study was thus to investigate the effects of fluvastatin on the incidence of long term adverse events among patients with unstable angina or stable angina enrolled in LIPS.

METHODS

Patients

Patients were recruited into LIPS from 57 centres in 10 countries between April 1996 and October 1998. The study population was composed of 1677 patients (1406 men and 271 women, age 18–80 years). All patients had successfully undergone a first PCI (index procedure) in the native coronary arteries. Patients qualifying for enrolment in LIPS had diagnosed unstable angina (according to the Braunwald classification), stable angina (according to the Canadian Cardiovascular Society classification), or silent ischaemia.

Patients were eligible for enrolment in the study if they had a total cholesterol concentration of 3.5–7.0 mmol/l (135–270 mg/dl) with fasting triglyceride concentrations of less than 4.5 mmol/l (400 mg/dl) before the index procedure.

Abbreviations: CI, confidence interval; LIPS, Lescol intervention prevention study; MACE, major adverse cardiac events; PCI, percutaneous coronary intervention; RR, relative risk
Exclusion criteria were sustained systolic blood pressure > 180 mm Hg and diastolic blood pressure > 100 mm Hg despite drug treatment; left ventricular ejection fraction < 30%; a history of previous PCI or CABG, severe valvar disease, idiopathic cardiomyopathy, or congenital heart disease; severe renal dysfunction (defined as serum creatinine concentration > 160 μmol/l (1.8 mg/dl)); obesity (defined as a body mass index > 35 kg/m²); or the presence of malignant or other disease with a life expectancy of less than four years.

All patients provided informed written consent and the ethics committee at each participating centre approved the trial.

**Study design**
After inclusion, patients were randomly assigned to receive either fluvastatin 40 mg twice daily (Lescol, Novartis Pharma AG, Basel, Switzerland) or matching placebo for a period of at least three years and no longer than four years. The study design of LIPS has been described in more detail elsewhere.24

**End points**
The primary end point was MACE, defined as cardiac death, non-fatal myocardial infarction, or repeat intervention procedure. Prespecified secondary end points were MACE excluding repeat intervention procedures occurring in the first six months for lesions treated at the index procedure (MACE without restenosis or coronary atherosclerotic events) and cardiac death/non-fatal myocardial infarction. Secondary end points also included treatment effects on measured lipid concentrations throughout the trial, as well as the safety and tolerability of fluvastatin. Details and results of the main trial have been provided elsewhere.6

Laboratory measures, including serum total cholesterol, low density lipoprotein cholesterol, high density lipoprotein cholesterol, and fasting triglyceride concentrations, were determined from fasting blood samples and assessed at a central laboratory (Analytico Medinet, Breda, the Netherlands). Laboratory values were measured at baseline, at a clinical visit six weeks after discharge, and at six month intervals thereafter until the conclusion of the study. Study personnel and patients were not informed of the lipid results.

**Statistical analysis**
All analyses were carried out on an intention to treat basis. Continuous variables are expressed as mean (SD) and were compared by Student’s unpaired t test. Wilcoxon scores were used for categorical variables with an ordinal scale. Discrete variables are expressed as counts and percentages and were compared in terms of relative risks (RR) (for unstable angina compared with stable angina) with 95% confidence interval (CI). All statistical tests were two tailed. Event-free survival distribution was estimated according to the Kaplan-Meier method. MACE-free survival time was tested with the log rank test. Patients lost to follow up were considered at risk until the date of last contact, at which point they were censored. The Cox proportional hazards model and the Cochran-Mantel-Haenszel test were used to assess risk reduction and to compare the incidences of the primary and secondary clinical end points, respectively. The treatment by anginal status interaction was calculated in a multivariate approach with treatment, anginal stability, and an interaction factor specifying the simultaneous presence or absence of both previous factors in the model. This enabled us to calculate the fluvastatin effect in patients with stable angina separate from the same effect in patients with unstable angina, as well as to calculate the anginal status effect in the patients given placebo and those given fluvastatin.

The lipid profiles were analysed in an analysis of covariance model incorporating the baseline (at visit 1) as covariate, adding the factors treatment, visit number (visits > 1), and subgroup with all possible interaction terms.

**RESULTS**

**Patient characteristics**
Of the 1677 patients enrolled in LIPS, data regarding qualifying diagnosis were complete for 1658 (98.9%) patients. These patients were classed as having either unstable angina (total, 824; fluvastatin, 417; placebo, 407) or stable angina (total, 834; fluvastatin, 418; placebo, 416). Patients with silent ischaemia were included in the stable angina group.

Table 1 shows baseline characteristics (demographic and angiographic). There were no notable differences in the baseline and procedural characteristics between the groups. The left anterior descending artery was the affected vessel in a larger proportion of patients with unstable angina than in the stable angina group. The unstable angina group also had a higher incidence of American Heart Association classification type B2 lesions and stent implantation. In contrast, patients with diagnosed stable angina had a higher incidence of American Heart Association classification type C lesions and balloon angioplasty. In the unstable angina group, patients randomly assigned to fluvastatin had a higher incidence of diabetes mellitus and a lower incidence of previous stroke than did patients randomly assigned to placebo. Overall, patients randomly assigned to fluvastatin or placebo in the stable angina group were well matched.

The median time between index PCI and randomisation was two days in both groups (range 0–22 days in the unstable angina group and 0–21 days in the stable angina group). The median time from index PCI to initiation of study medication was also two days. The median follow up was 3.9 years in both groups.

Up to the time of the first MACE or completion of follow up among patients without MACE, 17.6% (unstable angina group) and 17.5% (stable angina group) of patients were taking the study treatment regimen less than 80% of the time and were not taking other lipid lowering drugs. It is worth noting that 16.9% of patients in the unstable angina group and 17.6% of patients in the stable angina group were taking other lipid lowering drugs (generally other statins).

The proportion of patients taking other cardiovascular drugs during the course of the study was similar between the groups. Concurrent treatments included aspirin, β blockers, calcium antagonists, nitrates, angiotensin converting enzyme inhibitors, and diuretics.

**Long term risk of unstable versus stable angina after PCI**
Among patients randomly assigned to placebo, the overall incidence of MACE was similar in patients with unstable angina (27.3%) and patients with stable angina (26.2%) (RR 0.88, 95% CI 0.66 to 1.17, p = 0.37). The overall incidence of MACE in the two angina categories was also similar in patients randomly assigned to receive fluvastatin (unstable versus stable angina 20.9% v 22.3%; RR 1.0, 95% CI 0.73 to 1.37, p = 1.0). Overall, there was no significant effect of anginal status on long term risk of MACE (RR 0.94, 95% CI 0.75 to 1.17, p = 0.56). Similarly, there was no significant difference in risk between patients with unstable angina and those with stable angina in other end points, including cardiac death/non-fatal MI or coronary atherosclerotic events (MACE without restenosis).
Effect of fluvastatin in unstable and stable angina after PCI

Table 2 shows the effect of fluvastatin on the time to first end point. In patients with unstable angina, fluvastatin treatment produced a significant risk reduction for MACE of 28% (p = 0.03). The treatment-status interaction analysis showed that the effect of fluvastatin treatment did not differ significantly between patients with stable angina and patients with unstable angina (RR 1.07, 95% CI 0.87 to 1.30, p = 0.53). The Kaplan-Meier curves for fluvastatin and placebo groups started to separate at around 1.5 years and the benefits of fluvastatin treatment appeared to increase with time (fig 1).

Table 2 shows the effects of fluvastatin on other prespecified cardiovascular outcomes and fig 1 compares these outcomes. In patients with unstable angina, fluvastatin treatment significantly lowered the risk of coronary atherosclerotic events (incidence 23.1% with placebo v 16.3%, RR = 0.64, 95% CI 0.46 to 0.88; p = 0.006) but not the risk of cardiac death/non-fatal myocardial infarction (incidence

Table 1 Patient demographics and characteristics at baseline

<table>
<thead>
<tr>
<th></th>
<th>Unstable angina</th>
<th>Stable angina</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fluvastatin</td>
<td>Placebo</td>
</tr>
<tr>
<td></td>
<td>(n = 417)</td>
<td>(n = 407)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>60.7 (10.2)</td>
<td>60.2 (9.9)</td>
</tr>
<tr>
<td>Men</td>
<td>344 (82.5%)</td>
<td>336 (82.6%)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>26.9</td>
<td>26.3</td>
</tr>
<tr>
<td>Risk factors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Smoking (previous and current)</td>
<td>303 (72.7%)</td>
<td>297 (73.0%)</td>
</tr>
<tr>
<td>Diabetes</td>
<td>65 (15.6%)</td>
<td>34 (8.4%)</td>
</tr>
<tr>
<td>Hypertension</td>
<td>157 (37.6%)</td>
<td>158 (38.8%)</td>
</tr>
<tr>
<td>Family history of IHD</td>
<td>113 (27.4%)</td>
<td>119 (29.3%)</td>
</tr>
<tr>
<td>Previous MI</td>
<td>184 (44.1%)</td>
<td>172 (42.3%)</td>
</tr>
<tr>
<td>Previous stroke</td>
<td>10 (2.4%)</td>
<td>7 (1.7%)</td>
</tr>
<tr>
<td>Multivessel disease</td>
<td>148 (35.5%)</td>
<td>138 (339%)</td>
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<tr>
<td>Braunwald classification</td>
<td></td>
<td></td>
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<tr>
<td>IB</td>
<td>53 (12.7%)</td>
<td>49 (12.0%)</td>
</tr>
<tr>
<td>IIIB</td>
<td>50 (12.0%)</td>
<td>195 (47.9%)</td>
</tr>
<tr>
<td>IC</td>
<td>82 (19.7%)</td>
<td>79 (19.4%)</td>
</tr>
<tr>
<td>IIIC</td>
<td>11 (2.6%)</td>
<td>10 (2.5%)</td>
</tr>
<tr>
<td>IIIC</td>
<td>49 (11.8%)</td>
<td>34 (8.4%)</td>
</tr>
<tr>
<td>IIIC</td>
<td>36 (8.6%)</td>
<td>40 (9.8%)</td>
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<tr>
<td>CC classification (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>II</td>
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<td>NA</td>
</tr>
<tr>
<td>III</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>IV</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Silent ischaemia</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Baseline lipid concentration (mmol/l)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total cholesterol</td>
<td>5.17 (0.81)</td>
<td>5.16 (0.84)</td>
</tr>
<tr>
<td>LDL cholesterol</td>
<td>3.43 (0.75)</td>
<td>3.42 (0.79)</td>
</tr>
<tr>
<td>HDL cholesterol</td>
<td>0.95 (0.28)</td>
<td>0.95 (0.26)</td>
</tr>
<tr>
<td>Triglycerides</td>
<td>1.75 (0.66)</td>
<td>1.75 (0.71)</td>
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<tr>
<td>Systolic BP (mm Hg)</td>
<td>128</td>
<td>127</td>
</tr>
<tr>
<td>Diastolic BP (mm Hg)</td>
<td>74</td>
<td>75</td>
</tr>
<tr>
<td>Ejection fraction (%)</td>
<td>62.0</td>
<td>62.9</td>
</tr>
<tr>
<td>Lesion type (AHA class)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>20.3</td>
<td>18.6</td>
</tr>
<tr>
<td>B1</td>
<td>34.1</td>
<td>35.6</td>
</tr>
<tr>
<td>B2</td>
<td>36.5</td>
<td>36.2</td>
</tr>
<tr>
<td>C</td>
<td>9.1</td>
<td>9.6</td>
</tr>
<tr>
<td>Type of PCI*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stent</td>
<td>61.7</td>
<td>58.8</td>
</tr>
<tr>
<td>Others</td>
<td>3.4</td>
<td>4.5</td>
</tr>
</tbody>
</table>

*Percutaneous coronary intervention (PCI) categories are not mutually exclusive; hence, percentage values may not total 100%.
AHA, American Heart Association; BMI, body mass index; BP, blood pressure; CCS, Canadian Cardiovascular Society; HDL, high density lipoprotein; IHD, ischaemic heart disease; LDL, low density lipoprotein; MI, myocardial infarction; NA, not applicable.

Table 2 Effects of fluvastatin on prespecified cardiovascular end points

<table>
<thead>
<tr>
<th>End point</th>
<th>Unstable angina</th>
<th>Stable angina</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fluvastatin</td>
<td>Placebo</td>
</tr>
<tr>
<td></td>
<td>(n = 417)</td>
<td>(n = 407)</td>
</tr>
<tr>
<td>MACE without restenosis</td>
<td>68 (16.3%)</td>
<td>94 (23.1%)</td>
</tr>
<tr>
<td>Cardiac death/non-fatal MI</td>
<td>22 (5.3%)</td>
<td>35 (8.6%)</td>
</tr>
<tr>
<td>Time to first MACE</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

NA, not applicable.
8.6% with placebo vs 5.3%, RR = 0.60, 95% CI 0.34 to 1.05; p = 0.06). Fluvastatin also significantly decreased the risk of coronary atherosclerotic events in patients with stable angina (incidence 22.1% with placebo vs 15.8%, RR = 0.69, 95% CI 0.50 to 0.95; p = 0.02), although the reduction in cardiac death/non-fatal myocardial infarction was not significant (incidence 5.8% with placebo vs 4.8%, RR = 0.85, 95% CI 0.46 to 1.55; p = 0.59).

The proportion of patients with unstable angina with no MACE did not vary significantly according to Braunwald classification in either the fluvastatin (class I, 75%; class II, 80%; class III, 80%) or the placebo group (class I, 81%; class II, 73%; class III, 68%).

**Changes in lipid concentrations**

Baseline lipid concentrations were similar in the unstable angina and stable angina groups, as well as in the fluvastatin and placebo arms within each group (table 1). Figure 2 and table 3 show the changes in lipid concentrations with fluvastatin analysed on an intention to treat basis. Fluvastatin produced a significant reduction in total and low density lipoprotein cholesterol concentrations in both the unstable and stable angina groups, with comparable efficacy in the two groups. Reductions in lipid concentrations were first observed at visit 2 (six weeks after discharge) and were maintained throughout the study period. Although high density lipoprotein cholesterol concentrations increased above baseline concentrations in both the unstable and stable angina groups, similar increases were observed in patients randomly assigned to placebo; hence, there was no net benefit in favour of fluvastatin. Similarly, no significant changes in triglyceride concentrations attributable to fluvastatin were observed in either group.

**Safety and tolerability**

Full safety data have been reported previously.16 No significant adverse events were observed with fluvastatin.

**DISCUSSION**

LIPS was the first prospective study of patients with average lipid concentrations after successful PCI, with a significant reduction of 22% in fatal or non-fatal MACE as a result of early statin treatment with fluvastatin 80 mg/day. These results support the use of early lipid lowering treatment with fluvastatin for patients after PCI regardless of baseline lipid concentration.

Most previous studies investigating the long term effects of statin have focused on patients after an infarction or with stable angina.14 Information regarding the effects of statin in patients with unstable angina, particularly after PCI, is scarce. LIPS therefore offered the opportunity to investigate the effect of fluvastatin in patients with unstable angina who had successfully undergone PCI, a group that made up approximately half of the LIPS population.

The results of the present study show that patients classed as having both unstable and stable angina in LIPS had an

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Figure 1  Kaplan-Meier curves showing survival time free of adverse coronary atherosclerotic events (cardiac death, non-fatal myocardial infarction, and all reinterventions not caused by coronary restenosis) in patients with stable or unstable angina randomly assigned to receive either fluvastatin 80 mg/day or placebo.

Figure 2  Mean change in lipid concentrations throughout follow up among patients with stable or unstable angina randomly assigned to receive either fluvastatin 80 mg/day or placebo. (A) Total cholesterol; (B) low density lipoprotein (LDL) cholesterol; (C) high density lipoprotein (HDL) cholesterol; (D) triglyceride concentrations.
impaired long term prognosis, with up to one quarter to one third of patients experiencing one or more MACE within four years of PCI. These findings emphasise the importance of additional treatment after successful revascularisation. Importantly, fluvastatin treatment was of clear benefit both for patients with stable angina and for patients with unstable angina and was associated with clinically relevant reductions in the incidence of coronary atherosclerotic events in both patient groups. The benefits of fluvastatin treatment in reducing the incidence of MACE and prolonging the time to first MACE were significant in patients with unstable angina. A treatment-status interaction analysis showed no significant difference in the effects of fluvastatin between patients with unstable angina and patients with stable angina.

There was also no significant difference in outcome between patients with stable angina and those with unstable angina in the placebo group. This is somewhat surprising but may be a result of the successful PCI that took place before randomisation, which not only treats the obstructive lesion but also stabilises any initially unstable culprit plaque.

Previous studies have shown that, after successful coronary angioplasty, patients with unstable angina have a higher event rate than do patients with stable angina. This has generally been attributed to a higher repeat revascularisation rate. In contrast, LIPS showed that unstable and stable angina had comparable long term outcomes after successful revascularisation. There are several possible reasons for the difference. Firstly, this difference may reflect the more widespread use of coronary stent, a technique that has been shown to reduce restenosis in LIPS. Secondly, the inherited bias of recruiting solely patients with unstable angina who have undergone successful angioplasty may have effaced the risk imposed by the disease entity. Thirdly, the administration of new adjunctive medications such as glycoprotein IIb/IIIa inhibitors for unstable angina may also have contributed to lowering the risk of this patient group.

For patients with unstable coronary syndrome, early initiation of statin before hospital discharge has been independently associated with a reduction in MACE in the short term, as well as over long term follow up. However, successful revascularisation was not an inclusion criterion in these studies. It has also been reported that early initiation of statin for patients with unstable angina undergoing stent implantation is associated with an improvement in six month outcomes. Long term event rates, however, were not reported. Giving the positive effects of fluvastatin on long term outcomes in both patients with unstable angina and those with stable angina in our analysis, extrapolation of previous findings to all patients with coronary artery disease who have undergone a first successful PCI seems reasonable.

The beneficial effects of statin among patients with unstable angina may be attributed to its effect on vulnerable plaques, as vulnerable plaque rupture is the major triggering event for unstable coronary disease. Experimental data suggest that statin, in addition to its lipid lowering effects, may stabilise vulnerable plaques by converting lipid rich plaques (which are at high risk of rupture) into more stable fibrotic plaques.

### Study limitations

Because the number of patients with silent ischaemia was small, this group was included in the stable angina group for analysis. Patients with previous PCI or coronary bypass surgery were excluded; hence, the conclusions of the present study cannot be extrapolated to these patients. As this study is an analysis of a subgroup, albeit prespecified, it suffers from inherent limitations, such as the lack of sufficient power, because of the limited number of patients in each subgroup.

### Conclusions

The results of the present study provide evidence for the long term benefits of fluvastatin treatment initiated at hospital discharge for all patients with angina after successful PCI. It is worth noting that evidence from previous studies has shown that early initiation of lipid lowering treatment before hospital discharge also improves compliance. On the basis of these findings, we suggest that early lipid lowering treatment should be initiated for all patients after PCI, regardless of baseline lipid concentrations and clinical presentation.

### ACKNOWLEDGEMENTS

This study was supported by a grant from Novartis Pharma AG. Novartis provided the fluvastatin and matched placebo used in the present study.

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M Cortellaro, Istituto Policlinico S Donato, San Donato Milanese, Italy

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FROM BMJ JOURNALS

Systematic review of interventions to reduce delay in patients with suspected heart attack

Objectives: To evaluate the effectiveness of interventions aiming to reduce time from onset of signs and symptoms of an acute myocardial infarction (AMI) to seeking medical help/arrival at hospital.

Methods: A systematic review was conducted. Fifteen electronic databases, the internet, and bibliographies of included studies were searched, and experts in the field of cardiac care were contacted. Randomised controlled trials (RCTs), controlled trials, and before and after studies conducted in any setting that assessed an intervention aimed at reducing time from onset of signs and symptoms of an AMI to seeking medical help and/or arrival in hospital were eligible for inclusion.

Results: Eleven media/public education intervention studies met the inclusion criteria. Five (one controlled and four before and after studies) reported the intervention to have a statistically positive effect on delay time and six (two RCTs and four before and after studies) reported no statistically significant effect. Three (one RCT and two before and after studies) of five studies evaluating the effect of the intervention on emergency department visits reported an increase in this outcome as a result of the intervention, and both studies (one RCT and one before and after study) examining calls made to emergency switchboards reported an increase in this outcome after the intervention.

Conclusions: There was little evidence that media/public education interventions reduced delay. There is some evidence that they may result in an increase in emergency switchboard calls and emergency department visits. Despite substantial expenditure of time and effort, methodological deficiencies of the studies mean that it is not possible to make definitive recommendations. ▲ Emergency Medicine Journal 2004;21:506–508.