Lack of clopidogrel–CYP3A4 statin interaction in patients with acute coronary syndrome

D Mukherjee, E Kline-Rogers, J Fang, K Munir, K A Eagle

Objective: To assess a clinically significant interaction between cytochrome P450 3A4 (CYP3A4) metabolised statin and clopidogrel.

Design: Prospective single centre cohort study.

Setting: Academic teaching hospital in the USA.

Patients: 1651 patients presenting with acute coronary syndromes between January 1999 and February 2003 were studied. Data on baseline demographics, co-morbidities, and in-hospital management were collected.

Main outcome measure: Association of CYP3A4 metabolised statin and clopidogrel use with in-hospital and six month mortality. The impact of the combined use of a CYP3A4 statin and clopidogrel on six month mortality and major adverse cardiac events was analysed by a risk adjusted logistic regression model.

Results: The odds ratios for six month mortality were: for CYP3A4 statin, 0.43 (95% confidence interval [CI] 0.27 to 0.71, \( p = 0.0009 \)); for CYP3A4 statin plus clopidogrel, 0.36 (95% CI 0.23 to 0.60, \( p < 0.001 \)); for non-CYP3A4 statin, 0.22 (95% CI 0.08 to 0.59, \( p = 0.002 \)); and for non-CYP3A4 statin plus clopidogrel, 0.22 (95% CI 0.06 to 0.73, \( p = 0.016 \)).

Conclusions: Use of a combination of a CYP3A4 statin plus clopidogrel was associated with lower six month mortality and morbidity in patients with acute coronary syndromes. There was no significant difference in clinical benefit between a CYP3A4 statin and a non-CYP3A4 statin when used in conjunction with clopidogrel. This suggests that the proposed interaction is probably an ex vivo phenomenon and may not be clinically relevant.

Methods

Patients

From 1 January 1999 to 28 February 2003, 1651 patients were admitted to or discharged from inpatient services at our institution with a diagnosis of unstable angina or acute myocardial infarction (MI). All patients were identified by admission or discharge diagnoses and then the charts were reviewed to screen for entry criteria. Inclusion in the study required symptoms consistent with ACS and ECG changes suggestive of ischaemia (ST segment elevation or depression of \( \geq 1 \) mm, T wave inversion, or increased cardiac biomarkers). A final diagnosis of MI required increased creatine kinase MB (CK-MB) fraction or troponin as described in the American College of Cardiology guidelines. Reinfarction was defined as recurrent chest pain with new ECG changes (ST elevation of at least 1 mm in two contiguous leads, new Q waves) or new enzyme increases—that is, a re-increase of CK-MB after reaching a plateau or trough or a \( > 20\% \) increase of the previous value of already increased CK-MB. Stroke was identified clinically by a neurological deficit persisting \( > 24 \) hours with or without confirmation by computed tomography or magnetic resonance imaging. The study protocol was approved by the institutional review board at the University of Michigan and informed consent was obtained from all patients.

Data collected

Clinical, demographic, treatment, and outcome data were abstracted from medical charts by trained abstractors (cardiology fellows and cardiology research nurses). Definitions were based on those recommended by the American College of Cardiology data standards committee. ECG changes and initial laboratory data were recorded. Data describing patient management covered use of \( \beta \) blockers, aspirin, clopidogrel, angiotensin converting enzyme inhibitors, angiotensin receptor blockers, CYP3A4 and non-CYP3A4 statins, and percutaneous coronary interventions or coronary artery bypass grafting. Data on particular statin use was not collected prospectively but rather retrospectively from medical chart records. All other data were collected prospectively for our institutional ACS database. Six month mortality, MI, and stroke data were obtained on 100% of the patients from a health system record review or telephone call interview.

Statistical analysis

Baseline characteristics were summarised by the use of frequencies and percentages for categorical factors and mean (SD) for continuous factors. Multivariable logistic regression analysis was performed for six month death and major...
adverse cardiac events (MACE; composite of death, MI, and stroke) in patients with ACS treated with the statin plus clopidogrel adjusted for age, sex, positive biomarker, new ST elevation, history of diabetes, renal failure, heart failure, and revascularisation. Both a c-index (a measure of model discrimination) and Hosmer-Lemeshow test (a measure of model calibration) were used to determine the performance of the multivariate models. All data were analysed with SAS.

RESULTS

Table 1 shows the baseline characteristics of the study cohort. Of 1651 patients, 331 patients received neither clopidogrel nor statin, 180 patients received clopidogrel only, 387 used CYP3A4 statin only, 561 received CYP3A4 statin plus clopidogrel, 101 received non-CYP3A4 statin only, and 91 received non-CYP3A4 statin plus clopidogrel. Patients with ACS receiving a CYP3A4 metabolised statin were slightly older, less likely to have a prior history of angina, and more likely to have hypertension and hyperlipidaemia. There were no significant differences in history of prior MI, diabetes, smoking, increased biomarkers, renal function, or clinical presentation between patients stratified by type of statin use.

Table 2 shows cumulative six month clinical event rates for patients treated with a combination of CYP3A4 and non-CYP3A4 statin and clopidogrel. There were no significant differences in clinical outcomes stratified by use of CYP3A4 statin. For example, with atorvastatin as the CYP3A4 statin and pravastatin as the non-CYP3A4 statin, there were no differences in clinical outcome between these agents when used in combination with clopidogrel (fig 1).

The multivariate risk adjusted odds ratios for six month death were: for CYP3A4 statin alone, 0.43 (95% confidence interval (CI) 0.27 to 0.71, p = 0.0009); for clopidogrel alone, 0.48 (95% CI 0.26 to 0.89, p = 0.01); for CYP3A4 statin plus clopidogrel, 0.36 (95% CI 0.23 to 0.60, p < 0.001); for non-CYP3A4 statin, 0.22 (95% CI 0.08 to 0.59, p = 0.002); and for non-CYP3A4 statin plus clopidogrel, 0.22 (95% CI 0.06 to 0.75, p = 0.016) (table 3). The c-index for this model was 0.80, suggesting excellent model discrimination. The Hosmer-Lemeshow test statistic was 0.71, suggesting good model calibration and goodness of fit. Table 3 shows the risk of death, MI, stroke, and MACE in six months stratified by use of each type of statin.
adjusted odds ratios for MACE and shows a significant reduction in MACE with the combination of clopidogrel and a CYP3A4 statin (odds ratio 0.53, 95% CI 0.36 to 0.79, p = 0.001).

In a separate model, we compared the association between non-CYP3A4 statin plus clopidogrel and mortality with CYP3A4 statin plus clopidogrel as the reference group. The odds ratio was 0.56 (95% CI 0.15 to 2.1, p = 0.38) suggesting no significant differences in mortality with type of statin used in conjunction with clopidogrel.

**DISCUSSION**

Several recent ex vivo studies have suggested that statins metabolised by CYP3A4 such as atorvastatin may significantly attenuate platelet aggregation inhibition by clopidogrel. Such potential drug interactions with clopidogrel may be particularly important to recognise in patients with ACS who may be prescribed clopidogrel for 9–12 months. We systematically assessed the association of CYP3A4 metabolised statin and clopidogrel use with in-hospital and six month clinical event rates to ascertain whether a clinically significant interaction exists. There was no significant difference in clinical benefit between a CYP3A4 statin and a non-CYP3A4 statin when used in conjunction with clopidogrel. The results are in agreement with the reports by Saw and colleagues, who found no clinically relevant interaction between CYP3A4 metabolised statin and clopidogrel in a post hoc analysis of a placebo controlled study, and by Wienbergen and colleagues, who also found no significant difference between atorvastatin and other statins in the clinical outcomes of patients with ACS receiving clopidogrel. The results are also consistent with the report by Muller and colleagues, who used optical aggregometry for an ex vivo analysis and showed that concomitant use of CYP3A4 statins with clopidogrel did not significantly inhibit antiplatelet activity when clopidogrel was administered at a higher loading dose of 600 mg.

In our analysis of patients who received CYP3A4 metabolised statin, clopidogrel was associated with decreased six month mortality and a reduction in MACE. The discordance between the ex vivo data and clinical results may have several potential explanations. One concern is that the sample size in the initial ex vivo studies was very small, increasing the likelihood of detecting a spurious association based on biological variability. Moreover, platelet function in ex vivo tests may not adequately reflect the in vivo milieu when agents are administered together. In the only ex vivo analysis that used one of the ideal platelet function measurements, optical aggregometry, there was no significant interaction. Another explanation is that the non-platelet aggregation inhibition benefit of clopidogrel in reducing inflammation may result in clinical benefit in these patients.

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Significant multivariable risk adjusted predictors of six month mortality and MACE after acute coronary syndrome</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><strong>Adjusted OR for death (95% CI)</strong></td>
</tr>
<tr>
<td>Age</td>
<td>1.04 (1.02 to 1.06)</td>
</tr>
<tr>
<td>Positive biomarker</td>
<td>1.75 (1.11 to 2.75)</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>1.42 (0.95 to 2.09)</td>
</tr>
<tr>
<td>Renal insufficiency</td>
<td>2.21 (1.42 to 3.44)</td>
</tr>
<tr>
<td>Heart failure at presentation</td>
<td>3.88 (2.61 to 5.76)</td>
</tr>
<tr>
<td>Clopidogrel</td>
<td>0.48 (0.26 to 0.89)</td>
</tr>
<tr>
<td>CYP3A4 statin</td>
<td>0.43 (0.27 to 0.71)</td>
</tr>
<tr>
<td>CYP3A4 statin + clopidogrel</td>
<td>0.36 (0.23 to 0.60)</td>
</tr>
<tr>
<td>Non-CYP3A4 statin</td>
<td>0.22 (0.08 to 0.59)</td>
</tr>
<tr>
<td>Non-CYP3A4 statin + clopidogrel</td>
<td>0.22 (0.06 to 0.76)</td>
</tr>
</tbody>
</table>

C-index for the mortality model was 0.80; c-index for the MACE model was 0.72. CI, confidence interval; OR, odds ratio.
A limitation of our study is its non-randomised nature. We also assumed that patients who were prescribed statins and clopidogrel actually took their medications and we did not confirm compliance. However, we used a multivariate logistic regression analysis to adjust for differences in baseline demographics, co-morbidities, and revascularisation, and the model was robust in its overall ability to predict risk of both death and MACE.

We showed that a combination of a CYP3A4 statin plus clopidogrel was associated with lower six month mortality and MACE in patients with ACS. Although there are some differences in the odds ratios and risk reduction there was no significant difference in clinical benefit between a CYP3A4 statin and a non-CYP3A4 statin when used in conjunction with clopidogrel. The results from this large prospective registry of patients suggest the absence of any significant adverse clinical interaction of these two potent and effective agents in the management of patients with ACS.

Authors’ affiliations
D Mukherjee, E Kline-Rogers, J Fang, K Munir, K A Eagle, University of Michigan, Ann Arbor, Michigan, USA.

REFERENCES

Images in Cardiology

Wire artefact—in reverse!

A 75 year old woman with symptomatic hypertrophic cardiomyopathy and a left ventricular outflow tract gradient under stress exceeding 100 mm Hg was admitted for alcohol septal ablation. Panel A shows an apparent stenosis in the proximal right coronary artery identified at coronary angiography (left anterior oblique [LAO] 20°, cranial 20°) immediately pre-procedure. After the ablation procedure, it was elected to perform percutaneous coronary intervention on this lesion. On passage of a Balloon middleweight wire to the distal vessel, however, the “stenosis” resolved (panel B). The lesion reappeared when the wire was withdrawn and resolved again when the wire was passed back down the vessel. It was concluded that this apparent lesion represented a kink in the vessel and no intervention was performed.

It is well recognised that conformational change of a vessel wall by an angioplasty wire can cause appearances suggestive of coronary stenosis or dissection. Such a “wire artefact” occurs most commonly when the wire results in straightening of a tortuous vessel causing rucking of the vessel wall. These images demonstrate quite the reverse—an apparent stenosis that is abolished, rather than caused, by passage of the wire.

R A Archbold
C J Knight
andrew.archbold@bartsandthe london.nhs.uk