Influence of previous aspirin treatment and smoking on the electrocardiographic manifestations of injury in acute myocardial infarction

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Abstract

Objective—To examine demographic and clinical characteristics of patients with acute myocardial infarction in order to identify factors affecting the electrocardiographic evolution of injury.

Methods—Prospective cohort study of 1399 consecutive patients with a first myocardial infarction. Baseline clinical data associated with ST elevation and Q wave development were identified and 12 month survival was estimated.

Results—Smoking had complex effects on the evolution of injury, increasing the odds of ST elevation (odds ratio (OR) 1.61; 95% confidence interval (CI) 1.08 to 2.36), but reducing the odds of Q wave development (OR 0.69, 95% CI 0.49 to 0.96). The effects of previous aspirin treatment were more consistent with reductions in the odds of ST elevation (OR 0.57, 95% CI 0.35 to 0.94) and Q wave development (OR 0.53, 95% CI 0.34 to 0.84). ST elevation and Q wave development were both associated with an adverse prognosis, with estimated 12 month survival rates of 80.6% (95% CI 78.2% to 83.1%) and 80.0% (95% CI 77.5% to 82.5%), respectively, compared with 86.5% (95% CI 81.2% to 91.9%) and 89.9% (95% CI 86.2% to 93.7%) for patients without these ECG changes.

Conclusions—The thrombogenicity of the blood may be a major determinant of infarct severity. Smoking increases thrombogenicity and the likelihood of ST elevation, but because coronary occlusion is relatively more thrombotic in smokers, responses to both endogenous and exogenous thrombolysis are better, reducing the risk of Q wave development. Previous aspirin treatment reduces thrombogenicity, protecting against ST elevation and Q wave development.

Keywords: myocardial infarction; smoking; aspirin
infarction was diagnosed if any two of the following criteria were fulfilled:
- cardiac chest pain lasting at least 30 minutes;
- ST elevation of $\geq 0.1$ mV in at least one standard lead or $\geq 0.2$ mV in two or more contiguous chest leads;
- creatine kinase $\geq 400$ IU/l (upper limit of reference range 200 IU/l).

Thrombolysis was recommended for any patient with regional ST elevation who presented within 12 hours of the onset of continuous chest pain.

**DATA COLLECTION**

Baseline clinical characteristics, including demographic, clinical, and biochemical data, were collected prospectively by one of three study investigators and entered into a purpose built electronic database. The hospital diagnostic of a qualifying event (ST elevation or new Q wave development) was determined by one of two attending cardiologists (KR and ADT).

Thrombolysis was recommended for any patient with regional ST elevation who presented within 12 hours of the onset of continuous chest pain.

**FOLLOW UP**

The purpose of the follow up was to define all cause mortality and recurrent ischaemic events (readmission with unstable angina or acute myocardial infarction) from the admission date until censoring in July 1998. The patients were flagged by the Office of National Statistics to obtain follow up mortality data. In patients known to be alive at the censoring date and in whom information about non-fatal ischaemic events was not already available from outpatient follow up or readmission to the coronary care unit, postal questionnaires were used to obtain this information, backed up by telephone inquiry for non-responders. In this way, follow up data beyond discharge were obtained in 98% of the study group.

**STATISTICAL ANALYSIS**

Survival was calculated by the Kaplan–Meier method, and survival probabilities were expressed as percentages with 95% confidence intervals (CI). Comparison of discrete variables was by $\chi^2$ analysis and continuous variables by the Mann–Whitney U test. In order to evaluate their independent influence,
variables showing significant differences or pronounced trends in univariate analysis or believed to be of clinical or biological relevance were entered into a logistic regression analysis. Improvements in model fit were based on comparison of likelihood ratios. Odds ratios (OR) are quoted together with 95% CI.

Results

BASELINE CHARACTERISTICS: ST ELEVATION
These data are given in table 1. Comparison of baseline characteristics in patients with and without ST elevation showed no significant differences with regard to age, sex distribution, or race. At the time of admission, significantly more patients with ST elevation were current smokers but fewer were taking aspirin. Patients with ST elevation presented earlier after the onset of chest pain than those without, and tended to have higher glucose but lower potassium concentrations in their admission blood samples. Significantly more patients with ST elevation received thrombolytic treatment in accordance with treatment guidelines.

BASELINE CHARACTERISTICS: Q WAVES
Comparison of baseline characteristics in patients with ST elevation who did and did not develop pathological Q waves showed no significant differences as regards age, sex distribution, and racial group (table 2). Significantly fewer patients who developed Q waves were current smokers and fewer were taking aspirin at the time of admission. Proportions who received emergency treatment with aspirin and thrombolysis were similar in both groups. Comparison of smokers who did (n = 516) and did not (n = 134) receive thrombolytic treatment showed similar rates of Q wave development (76.9% vs 78.4%).

PREDICTORS OF ST ELEVATION AND Q WAVE DEVELOPMENT
These data are given in tables 3 and 4. Smoking increased the odds of ST elevation by 60% but reduced the odds of Q wave development by 31%. Being on aspirin at the time of admission reduced the odds of ST elevation and Q wave development by 43% and 47%, respectively. Diabetes also reduced the odds of Q wave development. Admission within two hours of the onset of chest pain more than doubled the odds of ST elevation but did not influence Q wave development.

RELATIONS BETWEEN PREVIOUS ASPIRIN TREATMENT, SMOKING, ST ELEVATION, AND Q WAVE DEVELOPMENT
ST elevation occurred most commonly in smokers and least commonly in patients on previous aspirin treatment (table 5). Q wave development occurred most commonly in patients who neither smoked nor took aspirin, and least commonly in those who both smoked and took aspirin. For other groups, rates of ST elevation and Q wave development were intermediate.

SURVIVAL ANALYSIS
ST elevation and Q wave development were both associated with an increase in all cause
mortality as well as recurrent ischaemic events in the form of non-fatal myocardial infarction and readmission with unstable angina (fig 1). The estimated 12 month survival rates were 80.6% (95% CI 78.2% to 83.1%) for patients presenting with ST elevation and 80.0% (95% CI 77.5% to 82.5%) for patients who developed Q waves, compared with 86.5% (95% CI 81.2% to 91.9%) and 89.9% (95% CI 86.2% to 93.7%), respectively, for patients without these ECG changes. Similarly, estimated 12 month event free survival rates were 71.0% (95% CI 67.4% to 74.6%) and 69.6% (95% CI 66.7% to 72.5%) compared with 76.0% (95% CI 69.3% to 82.8%) and 74.7% (95% CI 69.1% to 80.3%).

Discussion
This prospective cohort study of patients with acute myocardial infarction has shown important independent effects of smoking and previous aspirin treatment on ST segment change and Q wave development that have not been reported previously. The effects of smoking were complex, associations with ST elevation being consistent with its prothrombotic activity but apparently inconsistent with the non-development of Q waves. The effects of previous aspirin treatment were generally favourable and likely to reflect its antiplatelet activity.

ST ELEVATION
ST elevation in people with chest pain is a reliable marker of coronary occlusion and usually predicts Q wave development.12 In this study, previous aspirin treatment reduced the incidence of ST elevation, whereas smoking was predictive of ST elevation. This presumably reflects first, the antiplatelet effects of aspirin, which shift the endogenous balance of thrombotic and thrombolytic activity in favour of early coronary patency, and second, the prothrombotic effects of smoking,13–15 which exaggerate the thrombotic response to plaque rupture and increase the risk of total rather than partial coronary occlusion. This is consistent with earlier studies of acute coronary syndromes in which smokers were more likely to present with myocardial infarction than unstable angina, while previous aspirin treatment favoured presentation with unstable angina.9–11

Q WAVE DEVELOPMENT
The reduced incidence of Q wave development in patients on previous aspirin treatment is consistent with the antiplatelet effect of aspirin. In contrast, the association between smoking and non-Q wave infarction appears paradoxical. However, it is well established that high grade reflow following thrombolysis is more common in smokers than in non-smokers; this is thought to reflect a thrombotic rather than an atherosclerotic coronary occlusion, which is more susceptible to thrombolysis.16–18 It is plausible, therefore, that the prothrombotic effects of smoking account not only for its association with ST elevation but also for its association with non-Q wave infarction, by providing a substrate for successful thrombolysis. It may also explain why smokers with acute myocardial infarction often have a relatively good short term prognosis, as absence of Q waves usually denotes more limited myocardial injury.16–21 Thrombolytic treatment clearly has beneficial effects on prognosis, but a lower mortality in smokers was documented in studies in the prethrombolytic era22 and, in common with other investigators,23 we found that neither thrombolytic treatment nor aspirin given after hospital arrival influenced Q wave development, even though the median arrival time for this inner city population was only just over two hours after the onset of chest pain. However, if smokers are also more susceptible than non-smokers to endogenous lytic mechanisms shortly after the plaque event, recanalisation would occur sufficiently early to protect against Q wave development. Thus previous studies have shown that in acute myocardial infarction coronary patency at the time of presentation commonly predicts non-Q wave infarction,24 even in patients with ST elevation on the presenting ECG.5 7 This emphasises how the haemostatic environment at the time of plaque rupture, which is modified both by previous aspirin treatment and smoking, can have more profound effects on its clinical consequences than antithrombotic treatment given after hospital arrival.

PAIN DURATION AND ADMISSION BIOCHEMISTRY
The duration of symptoms was a major predictor of ST change. Thus ST elevation was more common in patients presenting within two hours after the onset of chest pain than in those presenting later. This is consistent with the work of DeWood and colleagues, who showed that as time elapses after the onset of myocardial infarction the inci- dence of complete coronary occlusion declines, presumably reflecting endogenous thrombolytic activity.12 Increased pain severity may also have contributed to the earlier presentation of patients with ST elevation and this would predict a heightened sympathoadrenal response, which no doubt explains the tendency for blood glucose to be higher and potassium lower in patients with ST elevation.22–23 Not surprisingly, however, admission biochemistry was unrelated to the later development of Q waves, although patients who were diabetic were more likely to have non-Q-wave infarction. This has been reported previously24 and may reflect the diffuse and chronic nature of diabetic coronary disease which encourages collateralisation of the myocardium and protects against ischaemic injury in the event of thrombotic coronary occlusion.25

PROGNOSIS
The effects of smoking and previous aspirin treatment on the evolution of ECG changes in acute myocardial infarction are likely to have important clinical consequences, particularly regarding longer term prognosis. We and others26 27 have previously reported that patients with ST elevation have a worse prognosis than those without, an observation supported
by the data in the present study. The high proportion of patients presenting with ST elevation in this study may reflect not only the inner city location of our hospital, which ensures that patients with myocardial infarction present early (almost half presented within two hours after the onset of pain), but also the protective effect of myocardial collateralisation. 29–30 The extent of myocardial collateralisation is dependent upon the severity and duration of ischaemic heart disease, and other studies with a lower proportion of patients presenting with ST elevation 31–32 have not excluded those with previous myocardial infarction. There are conflicting data on the relevance of Q wave development for prognosis. Our data, however, are in agreement with those of Barbagalata and colleagues, who have also reported a lower one year mortality for non-Q wave infarction in a large cohort of patients presenting with their first myocardial infarction. 30 Of course, any prognostic benefits of smoking that are related to the non-development of Q waves must be minimal when set against the overall increase in coronary risk. However, the effects of previous aspirin treatment in protecting against ST elevation and Q wave development in acute myocardial infarction may contribute to its benefit in coronary artery disease.

CONCLUSIONS

The thrombogenicity of the blood at the time of a plaque event may be a major determinant of infarct severity, as reflected by ST segment change and Q wave development. Our study suggests that smoking, which increases thrombogenicity, has complex effects, increasing the likelihood of ST elevation while at the same time providing a substrate for successful thrombolysis, which reduces the risk of Q wave development. The net effect on prognosis may be beneficial. On the other hand, previous aspirin treatment may, by reducing thrombogenicity, reduce infarct severity, protecting against ST elevation and Q wave development more effectively than aspirin given after hospital arrival.