Glycoprotein Ia C807T gene polymorphism and increased risk of recurrent acute coronary syndromes: a five year follow up

A M Leone, V De Stefano, F Buzzotta, P Chiusolo, I Casorelli, K Paciaroni, E Rossi, A Sciahbasi, L Testa, G Leone, F Crea, F Andreotti

Despite aggressive control of modifiable risk factors, recurrent acute coronary syndromes are still common, and the underlying mechanisms largely elusive. Glycoprotein (Gp) Ia/IIa is one of several collagen receptors involved in platelet adhesion. A C807T gene variant of Gpla is related to Gp Ia/IIa receptor density on the platelet surface, with the density proportionate to the number of T alleles. Previous case-control studies in white subjects have associated this variant with acute ischaemic heart disease. As it is known whether the 807TT allele may also predict recurrent acute coronary syndromes, this was investigated in 117 survivors of a first acute coronary event; patients were followed for up to five years.

METHODS

Between 1996 and 1998 we enrolled 195 patients below the age of 65 years; these patients were discharged from our coronary care unit with a diagnosis of acute myocardial infarction (MI), according to World Health Organization criteria, (MI, n = 141) or severe Braunwald class IIIB unstable angina (UA, n = 54) at first presentation of disease. After informed consent for genetic analyses, a final number of 117 patients (79 MI, 38 UA) accepted follow up assessment which took place at six months, two years, and five years. Clinical evaluation was blinded to the results of genotyping. Major cardiovascular risk factors, angiographic extent of disease, and left ventricular ejection fraction (LVEF) were assessed at enrolment. Antithrombotic drugs during follow up consisted of aspirin or ticlopidine if aspirin was contra-indicated. Following percutaneous coronary interventions, aspirin and ticlopidine were both prescribed for four weeks. Gp IIIb/IIa inhibitors were not administered.

The primary end point was the composite of fatal and non-fatal MI and UA, but additional separate analyses were carried out for incident MI or UA alone. During follow up, fatal MI was defined as death preceded by symptoms, or signs, indicative of prolonged myocardial ischaemia, while non-fatal MI and UA were defined as the index event by WHO criteria and as Braunwald class IIIB, respectively. Genotyping was completed in December 2001 by operators blinded to the clinical data, as previously described. The Kaplan-Meier survival curves free of MI and UA started to differ at two months (p = 0.034) and remained significantly different at six months, two years, and five years (p = 0.0009, p = 0.014 and p = 0.036, respectively) (fig 1A). The difference in event rates was still significant at six months, two years, and five years for incident MI (p = 0.005, p = 0.075 and p = 0.048) (fig 1B), but not for UA alone (p = 0.07, p = 0.10 and p = 0.37, power 73%) (fig 1C). Comparing the three genotypes (CC v CT v TT), there was a trend towards a worse outcome with increasing number of T alleles (p = 0.054, log rank test). These results were confirmed in the subgroup of 79 patients who had an initial diagnosis of MI, in whom T allele carriers showed a worse outcome at five years for the composite end point (p = 0.04) and for MI (p = 0.04), but not for UA (p = 0.48, power 57%).

RESULTS

Genotyping revealed 807CC in 49 patients (41.9%), 807CT in 51 patients (43.6%), and 807TT in 17 patients (14.5%). This distribution did not differ significantly from that found in the original population of 195 patients (CC 37.4%, CT 47.7%, TT 14.9%). T allele carriers did not differ significantly from CC homozygotes for the major cardiovascular risk factors, extent of coronary disease, percentage undergoing surgical or percutaneous coronary revascularisation, antithrombotic drugs, and LVEF at baseline (table 1). There was also no significant difference in the use of β blockers and angiotensin converting enzyme inhibitors between T allele carriers (42% and 51%, respectively) and CC homozygotes (45% and 48%, respectively). Mean follow up was 46 months with a maximum of five years. All 117 patients were assessed at 30 days; at this time three patients (all carriers of the T allele) had a recurrence; 115 patients (98.3%) participated in follow up at six months, 86 patients (73.5%) at two years, and 81 patients (69.2%) at five years.

T allele carriers suffered significantly more events during follow up compared to patients homozygous for the C allele. The Kaplan-Meier survival curves free of MI and UA started to differ at two months (p = 0.034) and remained significantly different at six months, two years, and five years (p = 0.0009, p = 0.014 and p = 0.036, respectively) (fig 1A). The difference in event rates was still significant at six months, two years, and five years for incident MI (p = 0.005, p = 0.075 and p = 0.048) (fig 1B), but not for UA alone (p = 0.07, p = 0.10 and p = 0.37, power 73%) (fig 1C). Comparing the three genotypes (CC v CT v TT), there was a trend towards a worse outcome with increasing number of T alleles (p = 0.054, log rank test). These results were confirmed in the subgroup of 79 patients who had an initial diagnosis of MI, in whom T allele carriers showed a worse outcome at five years for the composite end point (p = 0.04) and for MI (p = 0.04), but not for UA (p = 0.48, power 57%).

Abbreviations: Gp, glycoprotein; LVEF, left ventricular ejection fraction; MI, myocardial infarction; UA, unstable angina
Table 1. Clinical characteristics of the study population

<table>
<thead>
<tr>
<th></th>
<th>All patients</th>
<th>T allele carriers</th>
<th>CC genotype</th>
<th>p Value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>117 (100%)</td>
<td>68 (58.1%)</td>
<td>49 (41.9%)</td>
<td></td>
</tr>
<tr>
<td>Mean age (years)</td>
<td>54.3 (8.2)</td>
<td>55.1 (7.5)</td>
<td>52.9 (1.3)</td>
<td>0.27</td>
</tr>
<tr>
<td>Females</td>
<td>16 (13.7%)</td>
<td>10 (14.7%)</td>
<td>6 (12.2%)</td>
<td>0.79</td>
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<tr>
<td>Smokers</td>
<td>71 (60.7%)</td>
<td>36 (52.9%)</td>
<td>35 (71.4%)</td>
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<tr>
<td>Systemic hypertension</td>
<td>64 (54.7%)</td>
<td>37 (54.4%)</td>
<td>27 (55.1%)</td>
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<tr>
<td>Hypercholesterolemia</td>
<td>52 (44.4%)</td>
<td>31 (45.6%)</td>
<td>21 (42.9%)</td>
<td>0.85</td>
</tr>
<tr>
<td>Family history of ischaemic heart disease</td>
<td>44 (37.6%)</td>
<td>23 (33.8%)</td>
<td>21 (42.9%)</td>
<td>0.34</td>
</tr>
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<td>Diabetes mellitus</td>
<td>19 (16.2%)</td>
<td>13 (19.1%)</td>
<td>6 (12.2%)</td>
<td>0.32</td>
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<tr>
<td>Mean left ventricular ejection fraction</td>
<td>50.1% (10.2)</td>
<td>49.4% (10.4)</td>
<td>51.1% (9.9)</td>
<td>0.42</td>
</tr>
<tr>
<td>Percutaneous coronary interventions</td>
<td>30 (25.6%)</td>
<td>17 (25.0%)</td>
<td>13 (26.5%)</td>
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<td>Coronary artery bypass grafting</td>
<td>12 (10.3%)</td>
<td>8 (11.8%)</td>
<td>4 (8.2%)</td>
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<td>Coronary angiography</td>
<td>90 (100%)</td>
<td>54 (79.4%)</td>
<td>36 (73.5%)</td>
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<tr>
<td>Single vessel disease</td>
<td>34 (37.8%)</td>
<td>23 (42.6%)</td>
<td>11 (30.6%)</td>
<td>0.28</td>
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<td>Multi vessel disease</td>
<td>51 (56.7%)</td>
<td>29 (53.7%)</td>
<td>22 (61.1%)</td>
<td>0.82</td>
</tr>
</tbody>
</table>

*p Values compare T allele carriers to CC genotype.

Hypercholesterolemia was defined as: total cholesterol >240 mg/dl; diabetes as fasting glycaemia >126 mg/dl on at least two occasions; hypertension as blood pressure >140/90 mm Hg; smoking as a smoker of > 1 cigarette per day at the time of admission; family history as documented acute coronary syndrome before 60 years of age in at least one first degree relative.

Figure 1. Kaplan-Meier event-free survival curves (A to C) in T allele carriers (continuous curves) and in CC homozygotes (broken curves). Compared to CC homozygotes, significantly fewer T allele carriers were free of any event during the five year follow up and free of MI at both six months and five years. Survival free of UA did not differ significantly between groups. Recurrences (D) occurred sooner in T allele carriers compared to CC homozygotes (horizontal bars indicating median time to recurrence).
In the entire group, after adjusting for age, sex, diabetes mellitus, LVEF < 40%, type of index event, and use of revascularisation procedures, the relative risk for the primary composite end point among T allele carriers compared to non-carriers was 7.6 (95% confidence interval (CI) 1.68 to 33.8, \( p = 0.007 \)) at six months, 2.4 (95% CI 1.19 to 4.83, \( p = 0.014 \)) at two years, and 1.9 (95% CI 1.05 to 3.43, \( p = 0.035 \)) at five years. Median event-free survival was shorter in T allele carriers compared to CC homozygotes (27 v 57 months) and, among patients who experienced adverse events, these occurred sooner in the presence of the T allele (median 6 v 18 months, \( p = 0.015 \)) (fig 1D).

**DISCUSSION**

Our findings suggest that carriers of the 807T allele of the GpIa gene among patients suffering a first acute coronary syndrome before the age of 65 years may significantly increase the risk and anticipate the timing of a recurrent event over the next five years, even after correction for established prognostic factors. At six months, having the T allele increased the adjusted risk of recurrence by approximately sevenfold compared to homozygosity for the C allele. The less favourable outcome in T allele carriers appears to be attributable to a higher incidence of fatal and non-fatal MI rather than UA (although the power to detect differences in the rates of UA was below 80%). The 807T allele may thus confer susceptibility to more severe recurrences.

The 33.3% recurrence rate of MI at five years seen in our cohort is comparable to that previously reported in survivors of acute coronary syndromes.1 Moshfeg and colleagues5 first described a threefold risk of MI among carriers of the 807TT genotype compared to C allele carriers. Several, though not all, subsequent cross sectional studies have confirmed the association between the C807T polymorphism and acute ischaemic heart disease.1 To our knowledge this is the first longitudinal study to investigate and identify a relation between a GpIa gene polymorphism and the risk of recurrences in patients presenting with acute coronary syndrome. Further studies are needed to confirm these preliminary findings and to assess whether the prognostic role of this polymorphism may be incremental over that of markers of myonecrosis and inflammation.

**Authors’ affiliations**

A M Leone, V De Stefano, F Burzotta, P Chiusolo*, I Casorelli*, K Paciaronti*, E Rossi*, A Sciabbbasi, L Testa, G Leone*, F Crea, F Andreotti, Institute of Cardiology and Institute of Haematology, Catholic University, Rome, Italy

Correspondence to: Dr Antonio Maria Leone, Istituto di Cardiologia, Universita` Cattolica del Sacro Cuore, Largo A. Gemelli 8, 00168 Rome, Italy; antonionomarialeone@libero.it

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