Lack of association between haemostatic variables and the presence or the extent of coronary atherosclerosis

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SUMMARY Blood samples were taken for haemostatic analysis from 225 patients with angina pectoris who were admitted to hospital for coronary angiography. β thromboglobulin, platelet factor 3, platelet factor 4, factor VII:C, factor VIII:C, von Willebrand factor antigen, activated partial thromboplastin time, fibrinogen, antithrombin III, protein C:Ag, plasminogen, and antiplasmin were measured before angiography. Patients who had had a myocardial infarction in the two months before the investigation were excluded from the study. Multiple linear regression analysis showed that none of the haemostatic variables contributed independently to the prediction of an angiographic score that indicated the extent of coronary atherosclerosis. History of myocardial infarction, male sex, worsening of angina pectoris, serum triglycerides, and ejection fraction were independently associated with the angiographic score. There were some significant correlations between haemostatic variables and conventional risk factors for coronary heart disease.

Thus data obtained from haemostatic analyses of peripheral venous blood do not permit the presence or the extent of atherosclerosis in coronary arteries to be predicted.

There is increasing evidence that coronary heart disease is associated with an activation of the clotting system. Coronary thrombosis occurs in most patients with acute myocardial infarction, and there is also epidemiological evidence indicating the pathogenetic importance of haemostatic function in coronary heart disease. The precise role of haemostatic factors in the presence of coronary atherosclerosis, however, is still unknown.

Although recent clinical data support the proposition that activation of the coagulation and the platelet system is closely associated with myocardial ischaemia, there is little information on the relation between the development of coronary atherosclerosis and the haemostatic system. Nichols et al did not detect increased concentrations of platelet factor 4, β thromboglobulin, and fibrinopeptide A in a group of patients with abnormal coronary angiograms without previous myocardial infarction. We have measured several haemostatic variables in blood samples from 225 patients with angina pectoris. All of them also had coronary angiography within the next two days. We graded the extent of coronary atherosclerosis according to an angiographic score and studied the associations of this score with different clinical variables as well as the haemostatic tests. Thus the aim of this study was to find out, in a cross sectional survey, whether there was a relation between stenotic disease and clotting activity.

Patients and methods

Patients
We studied men and women of all ages with angina...
who were admitted to hospital for coronary angiography. We excluded patients with non-cardiac diseases likely to cause death within one year, right heart failure with peripheral oedema, valve defects, or acute myocardial infarction within the preceding two months. Angina was described as “worsening” if the patient said that the attacks of chest pain had increased in frequency or in severity within recent weeks. Blood was drawn in the morning from patients at rest who had fasted for at least eight hours. A record was kept of any drugs, particularly anticoagulants and platelet inhibitors, taken within 10 days before blood sampling.

CORONARY ANGIOGRAPHY
Left ventricular catheterisation and coronary angiography were performed percutaneously from the femoral artery according to the Judkins’ technique.7 Coronary atherosclerosis on the arteriograms was scored from 1 to 5 as (1) no changes, (2) less than 50% stenosis (diameter reduction) in one or more vessels, (3) at least 50% stenosis in one vessel, (4) at least 50% stenosis in two vessels, and (5) at least 50% stenosis in three vessels. The ejection fraction was estimated according to Dodge et al10 as modified by Kennedy et al.9 Akinesis and dyskinesia were defined according to Herman et al.10

BLOOD SAMPLES
Samples were drawn by venupuncture with a 19 gauge butterfly system by specially trained staff. The first 5 ml was not used for haemostatic analyses. Blood for analysis was collected into precooled tubes containing Thrombotect. (Abbott, North Chicago, USA) reagent (9 + 1 v/v). These blood samples were used for the assays of platelet factor 4 and β thromboglobulin. Blood samples were centrifuged at 0°C for 60 min at 1900 g. Plasma was quick frozen and kept at −70°C until assay. Blood for coagulation assays was mixed (9 + 1 v/v) with tri-sodium citrate solution (0-130 mol/l, pH 7-5). Blood was centrifuged for 30 min at 2500 g (20°C) within 60 minutes of venupuncture. Plasma was snap frozen and kept at −70°C before analysis. Samples that were used for assays of activated partial thromboplastin time and platelet factor 3 were not frozen and were used immediately.

HAEMOSTATIC ASSAYS
Platelet factor 4 in plasma was measured by the radioimmunoassay kit produced by Abbott and β thromboglobulin by a radioimmunoassay kit supplied by Amersham (Buchler, Braunschweig, West Germany).

Factor VII:C and factor VIII:C were measured by a one-step assays with reagents and factor VII-deficient plasma from Behringwerke AG (Marburg, West Germany). Standard human plasma (Behringwerke) and a plasma pool prepared from 25 male donors (aged 20–40 years) were used as a reference plasma for the standard curves for factor VII:C and factor VIII:C respectively. Results were expressed as percentages of the standards. Fibrinogen was measured according to the method of Claus,11 with instructions and reagents from Behringwerke. Platelet factor 3 was measured in freshly prepared platelet rich plasma by a microcoagulation assay described in detail elsewhere.12

Activated partial thromboplastin time was measured semiautomatically with a coagulometer (Amelung, Lemgo, West Germany) and a cephalin reagent provided by the National (UK) Reference Laboratory for Anticoagulant Reagents and Control, Manchester. The biological activities of the protease inhibitors antithrombin III and antiplasmin were measured with the synthetic substrates S2238 and S2251,14 respectively, supplied by KabiVitrum (Munich, West Germany). Fibrinogen was measured by means of the synthetic substrate S2251 supplied by Kabi. von Willebrand factor antigen was measured by the electroimmunoassay described by Laurell.16 Heterologous antiserum against human von Willebrand factor:Ag (Behringwerke) was used. The results were expressed as a percentage of the standard prepared from normal human blood donors as described above. Protein C antigen was assayed by an enzyme linked immunosorbent assay kit17 (Boehringer Mannheim, Mannheim, West Germany).

STATISTICAL ANALYSIS
Multiple linear regression with forward stepwise selection of variables with a critical p value for entry of 0.05 was applied to test for possible relations between the angiographic score and the other variables. Logarithmic transformation of some variables was used to achieve an approximately normal distribution. p values, given to the nearest significant number if <0.0001, were derived from the multiple regressions. Independent associations with each of the individual haemostatic variables were also investigated by multiple regression analyses. In analyses of activated partial thromboplastin time, factor VII:C, or protein C:Ag we excluded the 40 patients on anticoagulant treatment. Because the number of comparisons was large results of only borderline statistical significance may have arisen by chance.

Results
CHARACTERISTICS OF THE PATIENTS ADMITTED TO THE STUDY
Two hundred and sixty three patients were admitted
to the study. Thirty eight were excluded because angiograms or other relevant documents were lost (n = 12), blood sampling was inadequate (n = 14), or incompatibilities with the entry criteria were detected after entry to the trial (n = 12).

Table 1 summarises the main clinical data of the 225 patients. Ninety nine patients of this group described themselves as ex-smokers and 42 were active smokers. The men were 52-4 (7·1) years old, and the women 52·9 (6·6) (mean (SD)).

**INDEPENDENT ASSOCIATIONS WITH THE ANGIOGRAPHIC SCORE**

Regression analysis showed that the following were independently associated with increases of the angiographic score: history of myocardial infarction (p < 0·0001, table 1), male sex (p < 0·0001, table 1), worsening of angina pectoris (p < 0·0001, table 1), increased serum triglyceride concentration (p = 0·003, table 2), and lower ejection fraction (p = 0·02, table 2). Although white blood cell count (table 2), current or past smoking (table 1), and the presence of akinesia or dyskinesia (table 1) were positively related to the angiographic score, these associations did not remain statistically significant when the above variables had been taken into account. The same applies to serum cholesterol concentrations (table 2), which were positively correlated with serum triglycerides (r = 0·36). Among those with a past history of myocardial infarction, there was no significant association between the interval since this event and the angiographic score. None of the haematological or haemostatic variables (table 2) contributed independently to the prediction of this score. In addition, no significant differences were seen when the haemostatic variables measured in the patients without angiographic changes were compared with the corresponding values in the patients with coronary atherosclerosis.

**INDEPENDENT ASSOCIATIONS WITH HAEMOSTATIC VARIABLES**

There were several significant independent associations between haemostatic variables and relevant clinical or laboratory variables. Patients with a history of myocardial infarction had a higher average of von Willebrand factor:Ag than those without (123% vs 102%, table 2, p = 0·003). The ejection fraction was associated with concentrations of antiplasmin (r = 0·18, p = 0·009) and inversely correlated with concentrations of β thromboglobulin (r = −0·15, p = 0·03) and fibrogen (r = −0·18, p = 0·006). Triglyceride concentrations correlated with plasminogen (r = 0·21, p = 0·003) and factor VII:C (r = 0·40, p = < 0·0001). Increases in cholesterol concentration were associated with higher plasminogen (r = 0·17, p = 0·02), antiplasmin (r = 0·22, p = 0·0009), and protein C:Ag (r = 0·53, p < 0·0001).

**Discussion**

In the present study we found no associations between the extent of coronary atherosclerosis and an activation of the haemostatic system, either for activation of the platelet and coagulation system or for depression of the fibrinolytic system. These results may seem to be in conflict with recent experimental, angiographic, and histological studies of coronary thrombosis in unstable angina pectoris, acute myocardial infarction, and sudden cardiac death. But none of our patients had an acute myocardial infarction and only a few of them had unstable angina. It is conceivable that myocardial ischaemia is a more important determinant of the activation of the haemostatic system than the extent of anatomical obstruction. Earlier studies did not demonstrate a relation between the extent of coronary atherosclerosis and concentrations of platelet
factor 46 19 20 or fibrinopeptide A.5 21 The last study found that the extent of coronary artery disease and β thromboglobulin concentrations were related in a group of 40 patients.21 Nichols et al., who studied 82 patients with ischaemic heart disease, only saw increased concentrations of platelet factor 4 and β thromboglobulin in those patients in whom myocardial infarction had occurred (more than six months previously, other cases of myocardial infarction being excluded).6 The results of Nichols et al.'s study were interpreted as indicating that release of platelet factors in patients with ischaemic heart disease results from the reaction of platelets with the previously infarcted ventricular wall rather than with the arteriosclerotic coronary arteries.

In our study the inverse correlation between lower ejection fractions (which were generally associated with akinesia or dyskinesia of the left ventricle) and β thromboglobulin concentrations, supports this interpretation. The strong positive correlation of antiplasmin and the ejection fraction and the negative correlation of fibrinogen with the ejection fraction that we found have yet to be explained.

We confirmed some other independent associations of haemostatic variables with established risk factors for coronary heart disease. Factor VII:C was strongly related to triglyceride concentration,3 22 and plasminogen correlated directly with serum concentrations of triglycerides and cholesterol.23 Protein C:Ag was not raised in our patients (unlike other reports24), but it was independently associated with cholesterol. The close relations between factor VII and some components of the fibrinolytic system25 on the one hand and different indices of hyperlipoproteinemia on the other may well reflect an important pathogenetic factor in coronary heart disease. In contrast, an association between cholesterol or triglyceride concentrations and platelet aggregability could not be confirmed.26

The results obtained in the present study do not support the hypothesis that an activation of the haemostatic system associated with atherosclerotic coronary arteries is detectable in venous blood. Rather, they show that correlates of fibrin formation or platelet release are detectable when processes that are secondary to coronary atherosclerosis occur and lead to vessel obstruction and ischaemic injury.

The cross sectional design of the present study means that the question of whether occlusive coronary events are preceded by a long term activation of the haemostatic system remains unresolved. The results of our study do not conflict with previous
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epidemiological work that suggests that individual clotting factors influence the development of coronary heart disease.

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