Antithrombin III in patients on long-term oral anticoagulants

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SUMMARY The antithrombin III (AT III) concentration in plasma was measured in 63 patients on oral anticoagulant treatment (mean age 57±7 years), 26 healthy laboratory controls (mean age 28 years), and 21 patients attending the hypertensive clinic who had never been on oral anticoagulants (mean age 50 years). Three methods were used to measure AT III: a coagulation assay, a chromogenic substrate assay, and an immunological assay. In patients on oral anticoagulants, the mean values for AT III in the three assays were: 124%, 107%, and 96% respectively. The mean AT III concentration in laboratory staff was 103±4%, 94%, and 104±1% for the three assays; patients attending the hypertensive clinic had AT III concentrations indistinguishable from those in patients on oral anticoagulants: 117±9%, 110±5%, and 93±9%. The difference between both patient groups and laboratory staff was statistically highly significant, but no difference was demonstrated between patients on anticoagulant treatment and those not receiving it. Our results show that the increase in the functional AT III concentration (measured by coagulation and chromogenic assays) observed in patients on oral anticoagulants is probably due to the effects of age and underlying disease rather than to the anticoagulant treatment itself.

Decreased plasma levels of AT III are associated with an increased incidence of thromboembolic disease, as seen in congenital deficiency, in women taking oral contraceptives with high oestrogen content, and during the postoperative period (for recent review see Barrowcliffe et al.1 and Davies and McNicol2).

Marciniak et al.3 reported an increase in plasma AT III concentration in congenitally deficient individuals when they were given small doses of oral anticoagulants. O'Brien and Etherington4 found that patients on long-term oral anticoagulant therapy for thromboembolic conditions had higher AT III plasma concentrations than healthy controls.

In contrast, Wessler et al.5 reported that plasma AT III was similar in patients on oral anticoagulants and controls, but that the Xa inhibitory activity was significantly higher in those on warfarin. To establish whether or not the AT III concentration rises in patients on oral anticoagulants we have investigated 63 patients on long-term oral anticoagulants and compared the values obtained with those of two control groups: healthy laboratory staff and hospital patients not on oral anticoagulants. If an increase in plasma AT III concentration occurs regularly with the administration of oral anticoagulants, an entirely different dose schedule and pattern of administration may be effective and justifiable in the prophylaxis of deep venous thrombosis.

Methods and subjects

Controls

Two control groups were studied. The first group consisted of 13 men and 13 women. They were members of the laboratory staff and their mean age was 28 years (Table 1). Five women were taking low oestrogen contraceptive pills; their AT III values did not differ from those in the women not taking the pill and they were included in the control group. The second control group comprised 21 patients attending the hypertensive clinic. There were 12 men and nine women and their mean age was 50 years. These patients received a variety of anti-hypertensive drugs, including β blockers, hydralazine, methyl dopa, and diuretics, but they had never taken oral anticoagulants.

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Table 1  
Details of subjects studied

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<th>Group</th>
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<td></td>
<td>No. Age, mean</td>
<td>No. Age, mean</td>
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<tr>
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<td>13 28</td>
<td>13 29</td>
<td>26 28</td>
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<tr>
<td>Patients not on oral anticoagulants</td>
<td>12 53</td>
<td>9 46</td>
<td>21 50</td>
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<td>Patients on oral anticoagulants:</td>
<td></td>
<td></td>
<td></td>
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<tr>
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<td>15 58</td>
<td>30 56</td>
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<td>Group 2</td>
<td>11 62</td>
<td>15 58</td>
<td>26 59</td>
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<tr>
<td>Group 3</td>
<td>7 60</td>
<td>—</td>
<td>7 60</td>
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Patients
A total of 63 patients stabilised on oral anticoagulants were studied. The patients had been on warfarin for at least three months since the last thromboembolic episode. They comprised three groups of individuals: group 1—30 patients who had had deep venous thrombosis and/or pulmonary embolism; group 2—26 patients with mitral valve disease; and group 3—seven men who had been anticoagulated after myocardial infarction. The details of age and sex are shown in Table 1.

Methods
AT III concentration was measured using three techniques:
1 Progressive antithrombin activity (PAT) by the coagulation assay of Abildgaard et al. Bovine fibrinogen (Diagnostic Reagents Ltd, Thame, Oxon) was used as substrate and Parke Davis bovine thrombin as enzyme.
2 Amidolytic assay of AT III (Chrom AT III) with the chromogenic substrate S 2238 (a generous gift of Kabi-Vitrum Ltd) using Fibrindex, human thrombin (Ortho Diagnostics Ltd, High Wycombe, Bucks), according to the method of Odegaard et al.
3 AT III antigen (AT III Ag) was determined by rocket immunoelectrophoresis using Behringwerke antiserum (Hoechst UK Ltd, Hounslow).

Blood samples were collected into 3.13% citrate, and the plasma were separated and stored for up to two weeks at —20°C before being assayed in batches. Pooled fresh normal plasma (10 donors) was assigned 100% potency and was used as a standard in PAT and AT III Ag assays. The standard for the Chrom AT III was the freeze-dried normal plasma supplied by Kabi Vitrum Ltd.

Thrombotest was performed on citrated venous blood according to the method recommended by Nyegaard and Co. A/S Oslo. The results were analysed using Student's t test and the linear coefficient of correlation.

Results
The results obtained with the PAT assay are shown in Figure 1. The mean values were 103.4% for laboratory controls, 117.9% for patients not on oral anticoagulants, 130.2% for group 1 anticoagulated patients, 117.7% for group 2, and 117.1% for group 3. The PAT values were significantly higher in hypertensive patients and in groups 1 and 2 than in the laboratory controls. Group 3 was too small for analysis.

AT III levels measured by the chromogenic assay are shown in Figure 2. Again laboratory controls showed lower values than the patients, and there was no difference between different patient groups. Mean Chrom AT III values were 94.2% for laboratory controls, 110.5% for hypertensive patients, 106.5% for group 1 patients on anticoagulants, 105.6% for group 2, and 111.5% for group 3.

Fig. 1  AT III levels measured using PAT assay in laboratory controls, hypertensive patients, and patients on oral anticoagulants. Each dot represents one measurement in one patient; the dotted line denotes the mean for each group.
The results of the immunological assay are shown in Figure 3. Laboratory controls had higher mean AT III antigen (104.1%) than hypertensive patients (93.9%), anticoagulated patients group 1 (98.6%), group 2 (96.2%), and group 3 (90.2%). These differences were significant (Table 2). There was no correlation between Thrombotest values and any of the AT III modalities measured.

Discussion

In comparison with healthy young controls, our patients on oral anticoagulants had higher mean PAT and Chrom AT III levels and lower AT III Ag concentrations. Surprisingly, similar high PAT and Chrom AT III values and low AT III Ag were obtained in a group of hypertensive patients not on oral anticoagulants. This suggests that the changes observed are due to the effects of age and/or disease rather than to warfarin therapy.
Meade and North\(^9\) reported that AT III Ag decreases with age, but no changes in AT III functional activity were observed in the industrial population studied. Increased levels of functional AT III have been described in ischaemic heart disease,\(^10\) after myocardial infarction,\(^11\) and as part of acute phase reaction.\(^1\) Our results emphasise the need for careful selection of controls for AT III assays.

Direct comparison of AT III values was possible only for PAT and AT III Ag assay where the same standard was used throughout. The two values were in good agreement in laboratory controls but showed a marked discrepancy in all patient groups where high PAT values were often associated with low AT III Ag values.

Poor correlation between functional and immunological AT III is well recognised\(^12\) and thought to be an index of AT III consumption. The discrepancy may also arise as a result of endothelial damage and disease, as Chan and Chan\(^13\) have recently demonstrated that AT III is present in the endothelial cells.

A fall in plasma AT III levels during heparin therapy is well documented, as is the subsequent rise while patients are given oral anticoagulants alone.\(^9\)\(^14\) Our observations on three patients confirm these findings and may explain the rise in AT III level seen during the early stages of anticoagulant therapy; this may well be the return to high pretreatment values after the heparin-induced fall. A larger series of patients tested regularly before, during, and after heparin and warfarin treatment is required to clarify this problem.

References

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