Mesenteric venous thrombosis and antithrombin III deficiency

C WILSON,† I D WALKER,* J F DAVIDSON,* C W IMRIE†

From the †Division of Surgery and *Department of Haematology, Royal Infirmary, Glasgow

SUMMARY Of the 123 patients with acute mesenteric infarction treated over the past 12 years, 16 (13%) had mesenteric venous thrombosis. Eight of the patients with mesenteric venous thrombosis survived the initial episode; two have since died. The remaining six patients were studied for evidence of haemostatic deficiencies or abnormalities. Antithrombin III deficiency, which is known to be associated with recurrent venous thrombosis, was found in three patients. It is recommended that all patients with mesenteric venous thrombosis should be screened for antithrombin III deficiency as treatment with coumarin anticoagulants may be indicated, providing effective prophylaxis against further thrombotic episodes.

Mesenteric venous thrombosis accounts for over 7% of patients with mesenteric infarction.1 Conditions predisposing to mesenteric venous thrombosis include portal hypertension, intra-abdominal sepsis, haematological disorders (such as polycythaemia rubra vera), neoplasms, surgery (particularly splenectomy) and use of oral contraceptives. Most patients have primary mesenteric venous thrombosis, where no underlying cause is found.

Abnormalities of the haemostatic mechanism are increasingly recognised: antithrombin III (ATIII), protein C, and protein S deficiencies and plasminogen abnormalities have been described in recurrent venous thrombosis.2–5 Mesenteric venous thrombosis was first reported in association with familial ATIII deficiency in 1975.6

Patients and methods

We reviewed the records of all 123 patients with acute mesenteric infarction presenting between January 1973 and August 1984, of whom 16 (13%) had mesenteric venous thrombosis. Thirteen of these patients were diagnosed at laparotomy and three at necropsy. Mean age was 55 years (range 32–74); eight patients (50%) were male. Mesenteric venous thrombosis was considered to be secondary to other underlying disease in five patients (31%), four of whom died. Three of these patients had underlying gastrointestinal carcinoma, one patient was receiving stilboestrol for prostatic carcinoma, and the fifth patient developed superior and inferior mesenteric vein and splenic vein thrombosis secondary to acute necrotising pancreatitis. In the remaining patients no recognised underlying cause of thrombosis could be implicated although one developed mesenteric venous thrombosis three weeks after giving birth and another had evidence of secondary polycythaemia on admission; both factors were likely to have been of importance. Thrombosis was not associated with use of oral contraceptives. Three patients (19%) had a history of thrombotic episodes, one had had mesenteric venous thrombosis nine years before and the other two each had had an episode of deep vein thrombosis.

Eight patients (50%) survived the initial episode, and two of them have since died. The remaining six patients were studied for evidence of haemostatic deficiencies or abnormalities.

ATIII and plasminogen antigen were measured by radial immunodiffusion (NOR and M Partigen plates, Behring Diagnostics). ATIII (functional) was measured by amidolytic assay7 and protein C by a modified enzyme linked immunosorbent assay (ELISA) method (Boehringer Mannheim). The functional assay of plasminogen was done according to the amidolytic method of Friberger.8 The normal ranges for all tests were established in over 40 healthy volunteers.

Case reports

CASE 1
A 32 year old man who presented with abdominal pain in 1984 was found at laparotomy to have venous congestion of the entire small bowel with multiple infarcted areas. The spleen appeared enlarged but the
Mesenteric venous thrombosis and antithrombin III deficiency

Table Results of haemostatic assessment

<table>
<thead>
<tr>
<th>Case No</th>
<th>Antithrombin III</th>
<th>Protein C</th>
<th>Plasminogen</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Immunological (29–42 mg/dl)</td>
<td>Amidolytic (85–129%)</td>
<td>Immunological (60–133%)</td>
</tr>
<tr>
<td>1</td>
<td>23</td>
<td>69</td>
<td>50*</td>
</tr>
<tr>
<td>2</td>
<td>20</td>
<td>61</td>
<td>59</td>
</tr>
<tr>
<td>3</td>
<td>25</td>
<td>72</td>
<td>113</td>
</tr>
<tr>
<td>4</td>
<td>36</td>
<td>107</td>
<td>100</td>
</tr>
<tr>
<td>5</td>
<td>29</td>
<td>89</td>
<td>87</td>
</tr>
<tr>
<td>6</td>
<td>34</td>
<td>116</td>
<td></td>
</tr>
</tbody>
</table>

*Patient receiving warfarin.

Case 3
A 76 year old man admitted in 1974 was found at laparotomy to have venous congestion of the entire small bowel, which was maximal in the mid jejunum. The bowel was viable and therefore not resected. He was not investigated postoperatively but treated empirically with heparin, making an uneventful recovery. He has since had one episode of deep vein thrombosis. As part of this survey he was studied for haemostatic abnormality, which showed mild ATIII deficiency (ATIII 72% of normal). Estimation of protein C, plasminogen, and coagulation screen and liver function tests all yielded normal results.

Three other survivors were reviewed and screened for ATIII, protein C, and plasminogen abnormalities two, three, and 11 years, respectively, following their mesenteric venous thrombosis. None has had further thrombotic episodes and all had normal haemostatic values at the time of writing.

Discussion

Antithrombin III is the most important naturally occurring antithrombin. It inhibits thrombin and also factor Xa, and thus regulates thrombin formation. ATIII is an α-2-globulin with a molecular weight of around 65,000. It is synthesised in the liver and has a narrow physiological range in plasma, values below this range being associated with an increased tendency to venous thrombosis. Deficiency of ATIII may result from impaired or faulty synthesis in congenital or acquired disorders, increased consumption or catabolism, loss from the intravascular compartment, or from unknown causes.9

Hereditary ATIII deficiency is usually transmitted as an autosomal dominant and such families have a high incidence of venous thrombosis. Over 85% of those aged 50 or older have had at least one thrombotic episode, often precipitated by pregnancy, parturition, surgery, trauma, or use of oral contraceptives.9 The mesenteric veins are a rare but characteristic site of thrombosis in familial ATIII
deficiency, affecting 8% of reported cases.10

Biochemical and haematological evidence suggested liver dysfunction as a possible cause for ATIII deficiency in case 2. ATIII concentrations may be reduced in chronic liver disease, but because of a simultaneous decrease in the procoagulants, there may be no increased risk of thrombosis.9 In case 2, despite the reduced procoagulants, the ATIII deficiency and thrombocythaemia presumably favoured thrombosis.

The aetiology of ATIII deficiency in the two other patients remains unclear, although in case 1 there may have been a liver abnormality. We were unable to show the deficiency in the other family members who were available for testing. Patients did not have abnormal plasminogen values and none had protein C deficiency, although this is thought to be commoner than ATIII deficiency.11

Moderate ATIII deficiency seems to have been a factor in three of the six survivors from mesenteric venous thrombosis tested, and it may explain the mechanism of thrombosis in some patients who would previously have been classified as having primary mesenteric venous thrombosis. Familial ATIII deficiency is a recognised but rare cause of mesenteric venous thrombosis. Our study suggests that even acquired ATIII deficiency may be associated with this condition, and we would therefore recommend investigating all patients with mesenteric venous thrombosis for ATIII deficiency as soon as the condition is suspected. In the presence of clinically important thrombosis ATIII activity may be decreased due to ATIII consumption and therefore results in the acute phase of the illness should be interpreted with caution. ATIII activity may fall further in the postoperative period if the patient remains unwell due to infection, sepsicaemia, or disseminated intravascular coagulation.9 Careful follow up with repeated ATIII estimation is therefore essential to avoid mislabelling patients whose ATIII activity subsequently returns to normal. Families of those discovered to have ATIII deficiency should also be studied to exclude familial deficiency because of their very high risk of thrombotic disease.

Treatment with coumarin anticoagulants provides effective prophylaxis further thrombosis in ATIII deficiency12 and is indicated in familial cases, in those with a low ATIII activity associated with previous thrombotic episodes, and particularly in situations likely to predispose to thrombosis such as surgery. In pregnancy, due to the teratogenic risks associated with coumarin anticoagulants, prophylactic anticoagulation with heparin is preferable at least during the first trimester.10 Immediately after surgery anticoagulation with heparin alone may be ineffective in preventing further thrombosis in patients with ATIII deficiency. The concurrent administration of ATIII concentrate and heparin permits immediate, full anticoagulation, although the doses of both require careful laboratory monitoring,12 which for ATIII, should be by a functional amidolytic assay.13

We thank Dr J Conkie and the staff of the Thrombosis Research Laboratory for performing the assays.

References

Mesenteric venous thrombosis and antithrombin III deficiency.

C Wilson, I D Walker, J F Davidson and C W Imrie

doi: 10.1136/jcp.40.8.906

Updated information and services can be found at:
[http://jcp.bmj.com/content/40/8/906](http://jcp.bmj.com/content/40/8/906)

These include:

**Email alerting service**

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Notes**

To request permissions go to:
[http://group.bmj.com/group/rights-licensing/permissions](http://group.bmj.com/group/rights-licensing/permissions)

To order reprints go to:
[http://journals.bmj.com/cgi/reprintform](http://journals.bmj.com/cgi/reprintform)

To subscribe to BMJ go to:
[http://group.bmj.com/subscribe/](http://group.bmj.com/subscribe/)