Mesenteric arteritis

J G Mosley, A Desai, I Gupta

Abstract

Four patients are presented with small bowel infarction secondary to a vascular arterial. In three patients there was a history of rheumatoid arthritis. In each patient infarcted bowel was resected and a primary anastomosis performed. In one patient the anastomosis broke down and she subsequently died. One patient died from a disseminated rectal tumour three years later. The remaining patients remain well. If operated on early, intestinal infarction due to arteritis has a good prognosis.

Acute intestinal infarction is a surgical emergency. Except in patients with mechanical bowel obstruction, it is caused by arterial or venous occlusion, or a vasculitis, or may arise in a low flow state when mesenteric perfusion is inadequate.

The precise cause is difficult to define, even at laparotomy, which accounts for the variation in incidence reported. Nevertheless, the consensus indicates that superior mesenteric arterial occlusion is responsible for most cases of intestinal ischaemia, while mesenteric venous thrombosis and non-occlusive ischaemia together represent 30% of cases. The rarer cause of intestinal ischaemia is due to mesenteric vasculitis. Early recognition and treatment is important as intestinal infarction may progress unless the involved bowel is resected. We report the clinical features and outcome for four adults with small bowel infarction secondary to vasculitis.

Patients

A short history of non-specific abdominal pain and lack of clear physical signs was characteristic of each patient (Table). Three of the patients had a history of severe rheumatoid arthritis and were taking 5–10 mg prednisolone/day on presenta-

<table>
<thead>
<tr>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
<th>Case 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>65</td>
<td>67</td>
<td>74</td>
</tr>
<tr>
<td>Sex</td>
<td>Female</td>
<td>Male</td>
<td>Male</td>
</tr>
<tr>
<td>Relevant medical</td>
<td>Rheumatoid arthritis for 20 years. Taking prednisolone and gold</td>
<td>Rheumatoid arthritis for five years. Taking prednisolone and gold</td>
<td>Rheumatoid arthritis for 10 years. Taking prednisolone. Antkylosing spondylitis</td>
</tr>
<tr>
<td>Symptoms</td>
<td>One week severe diffuse abdominal pain</td>
<td>One week abdominal pain and vomiting</td>
<td>Two days’ abdominal pain. Necrotic skin lesions. Livedo reticularis. Tender in RIF</td>
</tr>
<tr>
<td>Signs</td>
<td>Generalised tenderness</td>
<td>Generalised tenderness Rheumatoid factor</td>
<td>Rheumatoid factor</td>
</tr>
<tr>
<td>Operation</td>
<td>Multiple segments of gangrenous small bowel 400 cm of jejunum and ileum resected. Primary anastomosis</td>
<td>Infarcted 90 cm ileum resected. Primary anastomosis</td>
<td>Infarcted 150 cm small bowel with 4 cm perforation. Resection and primary anastomosis</td>
</tr>
<tr>
<td>Outcome</td>
<td>Anastomotic dehiscence. Started on parenteral nutrition but subsequently abandoned. Died eight weeks</td>
<td>Well at two years</td>
<td>Well for three years then presented with disseminated rectal tumour</td>
</tr>
</tbody>
</table>

Discussion

Mesenteric vasculitis is a rare cause of intestinal infarction representing about 2% of cases. Indeed, this presentation is the largest reported series. Intestinal vasculitis represents a group of
Mesenteric arteritis

Histological appearance of mesenteric vasculitis, showing loss of columnar epithelial lining of the bowel, thrombosis of vessels, and inflammation around involved vessels.

disorders sharing the histological features of inflammation and necrosis of blood vessels. Classification is based on a combination of clinical, radiological, and pathological features. The intestinal involvement may occur in isolation (as in the patient in case 4) or multiple systems may be involved.2

A variety of different vasculitic complications have been implicated as the cause of intestinal ischaemia. The reported diagnostic categories have been rheumatoid arthritis.3 This is generally regarded as a rare cause of intestinal infarction, though it was probably responsible for three cases in this series. Scleroderma4 has been described and the patient in case 4 had certain symptoms that may indicate this as an underlying abnormality. Polyarteritis nodosa may rarely cause intestinal ischaemia and infarction,5 and the patient in case 3 had histological evidence of fibrinoid necrosis of the vessel wall which would be consistent with polyarteritis nodosa. In addition, he had necrotic areas of skin.

In those cases with an isolated mesenteric vasculitis it is often impossible to identify a specific clinical syndrome and such patients have been described as having an atypical collagen vasculitis.6 The patient in case 4 would probably be regarded in this category. Indeed, difficulties arise in the classification of arteritis. There is often considerable overlap between the different syndromes7 which will not be resolved until the pathogenesis of vasculitic disease is better understood.

Should a patient present with vasculitis it may be possible to prevent the development of mesenteric vasculitis by aggressive treatment with cyclophosphamide.

Preoperative diagnosis of intestinal ischaemia is notoriously difficult. It has been found that a high white cell count is frequently found in this condition.8 Interestingly, the expected leucocytosis was present in these patients, three of whom were taking prednisolone.

In all cases the infarcted bowel was resected and a primary anastomosis performed. In the patient in case 1, however, this subsequently broke down and ultimately was responsible for her death. It may be important that in the three patients who did well there was an isolated ischaemic segment. The patient in case 1 had many separated areas of necrosis and about three quarters of her small bowel had to be resected which had both necrotic and viable segments. In the patient in case 1 the vasculitis may have involved the suture line, and in retrospect we would not recommend primary anastomosis for anyone with diffuse small bowel involvement. Rather, we favour resection of the ischaemic bowel and an end jejunostomy and ileostomy, which would be closed after two weeks if the mucosal surface was healthy. In addition, this patient also showed us the difficulties patients with severe arthritis experience when they have to manage an ostomy bag.

Mesenteric vasculitis causing bowel ischaemia is a rare condition; nevertheless, if operated on promptly it has a prognosis considerably better than that in patients with infarction due to superior mesenteric artery occlusion.9 Indeed, three of the patients made a good postoperative recovery. We therefore urge those responsible for the management of patients with abdominal symptoms and a connective tissue disorder to have a high index of suspicion as prompt operative intervention can be life saving.

We thank Dr D R Swinson for his advice.