Ulceraive colitis complicated by disseminated intravascular coagulation

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Summary: We report a case of chronic ulcerative colitis complicated by clinical evidence of disseminated intravascular coagulation and pathological evidence of intestinal ischaemia secondary to venular and capillary fibrin thrombi. This may well represent an example of univisceral Shwartzman reaction occurring in a sensitized target organ.

Introduction

Disseminated intravascular coagulation (DIC) occurs in association with a variety of conditions. It is characterized by the formation of micro-thrombi in small vessels producing ischaemic damage to many organs including lung, heart, kidney, liver and, less commonly, to the gastro-intestinal tract.¹ Vascular damage to the intestine has been described in a number of circumstances² and may be part of a generalized coagulopathy.³ However, the relationship of ulcerative colitis to DIC is not well documented and we describe a case which terminally resulted in fulminant ischaemic colitis with perforation.

Case report

A 59 year old man presented with a 4-month history of bloody diarrhoea and weight loss. One month before admission, rectal biopsy showed an active colitis with gland distortion, goblet cell depletion, focal ulceration and crypt abscesses (Figure 1). A barium enema performed 4 days before admission showed proctosigmoid colitis.

On examination he was apyrexial, dehydrated and clinically anaemic. Initial investigation revealed a haemoglobin of 6.6 g/dl, white cell count 13.2 x 10⁹/l, platelets 150 x 10⁹/l, sodium 129 mmol/l, potassium 2.7 mmol/l, urea 5.5 mmol/l, creatinine 114 μmol/l, albumin 21 g/l. A plain abdominal film showed mucosal oedema of the left colon and caecal dilatation. He was re-hydrated, transfused and given gentamycin and metronidazole. Blood and stool cultures were negative. Over a period of 48 hours his rectal bleeding increased and a clotting screen showed a prolonged prothrombin time of 18 seconds (control 13) and partial thromboplastin time (PTT) 49 seconds (control 37); the platelet count fell to 58 x 10⁹/l. Fibrin degradation products (FDPs) were greater than 20 but less than 40 μmol/l and a clinical diagnosis of DIC was made. He was treated with fresh frozen plasma, platelet transfusion and parenteral hydrocortisone. Radiologically colonic appearances improved. However, his girth increased and rectal bleeding persisted in spite of correction of the thrombocytopaenia. Two days later he developed signs of peritonitis. At laparotomy there was a perforation in the transverse colon with evidence of sub-total colitis.

Figure 1 Rectal biopsy showing active chronic colitis with goblet cell depletion, crypt abscess formation, a heavy inflammatory infiltrate and occasional bifid glands (H & E × 64)

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Accepted: 3 February 1987
Panproctocolectomy was performed and an ileostomy was fashioned. The colon showed large areas of ulceration and necrosis predominantly along the antimesenteric border, together with a 1.5 cm perforation in the transverse colon. The terminal ileum was normal.

The most striking histological feature was the large numbers of recent fibrin thrombi in many capillaries and veins within all layers of the colonic wall, mesentery, and sinusoids of mesenteric lymph nodes. The large mesenteric arteries and terminal ileal vessels were strikingly normal. All sections were stained to identify and confirm fibrin thrombi. The ulcers extended into the submucosa, with adjacent less severe inflammatory changes superimposed on chronic architectural and glandular damage (Figure 2). In the transverse colon there was transmural ischaemic necrosis with perforation. There was no fibrosis or any other feature to suggest chronic ischaemic damage.

Post-operatively his clotting remained deranged and he developed Gram-negative septicema, hypotension and renal failure. He died 18 days post-operatively.

At autopsy there was evidence of left ventricular failure and ischaemic heart disease with an old healed anterior myocardial infarct. Large mesenteric vessels were patent but there was recent portal vein thrombosis and a small liver infarct.

Histology confirmed multi-organ failure with ‘shock lung’, hepatic centriflobular necrosis, renal tubular necrosis and cerebral micro-haemorrhage. The small bowel showed mucosal autolysis, but no fibrin thrombi were to be seen here or in any other organ.

Discussion

This case illustrates several interesting features with respect to the role of ischaemia in patients presenting with fulminant colitis, and its relationship to ulcerative colitis.

Submucosal fibrin thrombi are rarely found in inflammatory bowel disease and do not appear to be related to the severity of the colitis. Vascular thrombosis and necrosis are not features of fulminant ulcerative colitis, but may be seen in the small or large bowel following periods of hypotension. The term ‘ischaemic enterocolitis’ has been used to describe this phenomenon. Previous studies have shown that it is usually of less than 20 days duration. In our case the insidious onset and relatively long history (3 months) together with the rectal biopsy appearances of chronic colitis and the sharp cut-off point at the ileo-caecal valve favour an underlying inflammatory bowel disease rather than an ischaemic enterocolitis.

Macroscopically, ischaemic enterocolitis may resemble fulminant ulcerative colitis although, in the former, the lesions are predominantly left-sided and rectal involvement is uncommon. The histological features are generally different, although on recovery from ischaemic colitis the changes may resemble those of a quiescent inflammatory bowel disease. In this case, there was extensive venous thrombosis with subsequent ischaemic necrosis and perforation superimposed on an established ulcerative colitis, but no features of chronic ischaemic damage such as muscle atrophy and fibrosis. Furthermore, the recent nature of the fibrin thrombi make low-grade chronic DIC unlikely.

The association of ulcerative colitis and DIC has only been reported infrequently. Lo describes a young female with a 3-day history of diarrhoea and Gram-negative septicemia complicated by DIC. On review, the history and histology are more suggestive of an acute infective colitis and the DIC could well have been secondary to Gram-negative sepsis. The two Japanese cases have no haematological evidence of DIC, and no histological evidence of fibrin thrombi involving the colon.

The haematological evidence of DIC in our case is presumptive with a normal platelet count on admission falling rapidly to 58 × 10⁹/l. This was associated with prolongation of prothrombin time and P I I , and a slight elevation of FDPs. Although in active DIC FDPs are usually greater than 40 μmol/l, levels can fluctuate widely. Severe acute DIC rarely causes significant organ damage unless pre-existing local disease is also present. In this case, the large intestine was already the site of a chronic inflammatory condition and once DIC supervened, ischaemia compounded damage to the colon.

DIC is known to produce ischaemic intestinal necrosis in dogs and in man. The latter may be one of

Figure 2 Colon showing distorted and branching glands, residual mucosal chronic inflammation and multiple submucosal fibrin thrombi (Picro-Mallory × 40)
the multi-system manifestations of DIC. However, fibrin thrombi have been producing enterocolitis in animals following endotoxaemia. As well as the generalized form, a univisceral or single organ Shwartzman reaction has been described which may be implicated in the pathogenesis of ischaemic enterocolitis. One mechanism postulated for the univisceral Shwartzman reaction is that of a 'generalized provocation' acting on an already damaged target organ. In our case, the initial thrombotic stimulus provided by DIC acts focally on the inflamed colon producing the univisceral damage seen in the colectomy specimen.

Persistence of fibrin thrombi may result from the secondary antithrombin III deficiency which has been demonstrated in patients with ulcerative colitis. The mechanisms producing reduced antithrombin III levels include increased catabolism and increased loss in a protein-losing enteropathy. The combined effects of these two mechanisms may result in significantly reduced antithrombin III activity and subsequent failure to neutralize thrombin formation.

Acknowledgements
The authors thank Mr R.T.J. Holl-Allen for allowing us to report this case, Dr A.B. Price for helpful comments on the histology, and Mr M.J. Chard, Mr C. Davies and Miss E. White for technical and secretarial assistance.

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doi: 10.1136/pgmj.63.742.689

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