Non-steroidal anti-inflammatory drug-induced jejunal and colonic diaphragm disease: a report of two cases

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Abstract

Two patients with intestinal submucosal diaphragm disease associated with non-steroidal anti-inflammatory drugs are presented. The jejunum was affected in one, and in the other the ascending colon. Most previously reported cases have shown ileal disease and no case to date has shown involvement of the colon. The reasons for the localisation of these diaphragms remain unclear.

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It is clear that non-steroidal anti-inflammatory drugs (NSAIDs) can cause a range of abnormalities in the small intestine including, rarely, stenoses caused by submucosal fibrosis. However, all the seven patients reported by Lang et al showed stenoses of the ileum. In the present study, two patients are presented in whom NSAID resulted in stenoses of the jejunum and ascending colon, respectively.

Case reports

CASE I

A 69 year old woman presented with a five week history of anorexia, weight loss of 6 kg, nausea, vomiting, and abdominal bloating. Seven years previously, she had been diagnosed as having seropositive rheumatoid arthritis and Sjogren’s syndrome. She began treatment with ibuprofen and later diclofenac, and within one year developed an iron deficiency anaemia of 8.3 g/dl. This was treated with iron tablets and later intramuscular injections of iron but she remained moderately anaemic. Gastroscopy showed a benign prepyloric ulcer but the patient declined a barium enema and further investigation was not performed. Over the succeeding seven years, she received fenbufen, ibuprofen, tiaprofenic acid, and flurbiprofen sequentially, using the latter as the sole NSAID for five years. There was no overt gastrointestinal bleeding. Physical examination showed a thin woman of 40 kg with obvious rheumatoid arthritis. Abdominal examination, rectal examination, and sigmoidoscopy to 20 cm were normal. A barium enema was normal. Gastroscopy showed a small inflammatory ulcer at the junction of the first and second parts of the duodenum, confirmed by biopsy.

The patient’s haemoglobin concentration was 10.3 g/dl, mean cell volume 61.5 (normal 80–100), white cell count 6.6×10⁹/l, platelets 616×10⁹/l, and ferritin 19 μg/l (normal 23–300). Serum vitamin B12 and red cell folate concentrations were normal. Serum glucose, electrolyte, urea, and creatinine concentrations were normal as were biochemical liver function tests.

Fine bore nasogastric tube feeding was begun but within one week, she developed vomiting, abdominal distension with visible peristalsis and constipation. Plain abdominal radiology showed dilated small intestinal loops. A provisional diagnosis of small bowel obstruction due to NSAID induced stenosis was made. After a period of no oral intake, intravenous feeding and parenteral iron therapy, during which the signs of intestinal obstruction resolved, a laparotomy was performed. This showed four strictures of the jejunum (Fig 1), which were excised with

Figure 1: Segment of resected jejunum showing four diaphragm strictures.

Figure 2: Section through the tip of one of the jejunal diaphragms. (Haematoxylin and eosin, original magnification ×20.)
primary anastomosis of the bowel. The patient made an uneventful recovery. NSAIDs were stopped and four months later she was well with a haemoglobin concentration of 12.5 g/dl.

Macroscopical examination of the strictures showed the typical appearances of diaphragm disease. The mucosal folds of the diaphragm were supported by proliferation of longitudinal bands of thickened and splayed muscularis mucosae extending into the submucosa. Bands of smooth muscle also extended into the lamina propria at the tip of the stricture (Fig 2). In places the overlying glands showed pyloric gland metaplasia caused by previous ulceration. Broader based mucosal folds were also seen. There was no evidence of coeliac disease.

CASE 2
A 77 year old woman complained of loss of appetite and weight loss of 12 kg in 6 months. She had not suffered abdominal pain but had been constipated, opening her bowels every 3–7 days for about two years. Rarely, she noted a little red blood on the paper after defecation.

She had suffered from polyarthritis for about 20 years and had taken slow release diclofenac 100 mg once daily for five years. Physical examination showed koilonychia and deformity of the hands consistent with rheumatoid arthritis. Abdominal examination showed a surgical scar of vagotomy and pyloroplasty for duodenal ulceration 16 years previously. The descending colon was palpable because of faeces. Rectal examination and rigid sigmoidoscopy were normal.

She had developed a hypochromic microcytic anaemia of 8.8 g/dl, which responded to iron injections given by her general practitioner. Serum urea, electrolytes, creatinine, and liver function tests were normal. Barium enema showed a few sigmoid diverticula and a short stricture of the ascending colon (Fig 3). Laparotomy confirmed a short stricture of the ascending colon and a right hemicolecotomy was performed. The patient made an uneventful recovery.

The right hemicolecotomy specimen showed several strictures at the origin of the ascending colon characterised by irregular, partly annular, mucosal folds. Microscopical examination showed that these broad based mucosal folds were almost identical to the jejunal lesions in the previous case with hyperplastic splayed muscularis mucosae extending into the lamina propria and submucosa (Fig 4). These features have not previously been described in the colon.

Discussion
The two patients presented had suffered iron deficiency anaemia and had taken NSAIDs for protracted periods. The proliferation of muscularis mucosae into the lamina propria of the mucosal folds and extension into the submucosa in these two patients was similar to that described as caused by NSAIDs but the location, in the jejunum and ascending colon, respectively contrasts with the ileal involvement in the patients of Lang et al.

Similar strictures, mainly affecting the jejunum, were reported in a patient with coeliac disease, although it is not known whether the patient had taken NSAIDs. Mucosal diaphragms in the duodenum and jejunum of patients with coeliac disease have also been described. However, there was no evidence of coeliac disease in patient 1 in the present study. Jejunal stricture was reported in one patient taking oral indomethacin for five years and distal jejunal strictures were reported in NSAID users.

Large intestinal effects of NSAIDs include inflammation, bleeding, ulceration, and perforation. Although the latter authors cite colonic strictures as a side effect of NSAIDs, they provide no specific references for this observation. The effects of NSAIDs on the large intestine are largely based on anecdotal cases. De Vos et al. found mild acute colitis in colonoscopic biopsy specimens of only 1 of 27 patients with rheumatoid arthritis submitted to ileocolonos-
copy. They stated most of these patients were taking NSAIDs but did not provide any other details of drug consumption and excluded any effect of NSAIDs in colonic inflammation. Recently, Jenkins et al. showed that NSAIDs increased colonic permeability to chromium labelled ethylenediamine tetra-acetate. The mechanisms underlying the induction of intestinal damage by NSAIDs remain unclear, although the amelioration of the NSAID-induced increase in small intestinal permeability by misoprostol implicates effects on prostaglandin synthesis. Other factors such as increased lysosomal fragility, local irritation, and impaired oxidative phosphorylation, may also be involved. Similarly, the factors determining the location of NSAID induced submucosal diaphragms are unclear. The patients presented in this paper indicate that the disease is not confined to the ileum but can affect the jejunum and colon.