Rectal lymphoma after colectomy for ulcerative colitis

J P Teare, S M Greenfield, S Slater

Abstract
Primary colonic lymphoma is an increasingly recognised complication of ulcerative colitis. We report the first known case of rectal lymphoma occurring after colectomy and ileorectal anastomosis in ulcerative colitis.

Although colonic carcinoma is the major long-term complication of ulcerative colitis, primary colonic lymphoma has also been shown to be associated with ulcerative colitis. Patients present with features similar to carcinoma, with an alteration in bowel habit, passage of blood rectally, or an abdominal/rectal mass. Biopsy of the tumour found at sigmoidoscopy or colonoscopy usually confirms the diagnosis. Surgical excision of lymphoma remains the mainstay of treatment and although chemotherapy or radiotherapy is often given as adjuvant treatment, the value of these treatments cannot be assessed from published reports. We report a patient who presented with a rectal lymphoma many years after ileorectal anastomosis.

Case report
A 74 year old woman presented in 1956 with severe bloody diarrhoea. At that time sigmoidoscopy showed an acutely inflamed, granular mucosa and a rectal biopsy showed features of an acute proctitis. Repeat stool cultures at presentation were negative. She failed to respond to medical treatment and underwent colectomy (an ileorectal anastomosis was fashioned three months later), having developed toxic megacolon secondary to acute ulcerative colitis. The slides from the original colectomy specimen have been reviewed by Dr Slater and an independent histopathologist, Dr S Rasbridge. These show features typical of ulcerative colitis with pseudopolyp formation, distortion of crypts, crypt abscesses, and a severely inflamed and congested mucosa. In addition, no giant cells, granulomas, or fissuring ulceration were seen and there were no features of ischaemic or pseudomembranous colitis.

During the intervening period she remained well and underwent regular sigmoidoscopy and rectal biopsy. These biopsy specimens merely showed features of chronic and occasionally acute proctitis, but no dysplasia was ever seen (further confirmed by Drs Slater and Rasbridge). This patient was lost to follow up in 1988 but she presented again in August 1990 with a change in bowel habit, passing mucus and blood rectally.

Examination showed a 5 kg weight loss, no lymphadenopathy, abdominal masses, or organomegaly. At sigmoidoscopy a thickened irregular rectal wall with contact bleeding was found. Biopsy of this showed infiltration and complete replacement of the rectal mucosa by mitotically active pleomorphic cells; these were leucocyte common antigen positive and carcino-embryonic antigen and epithelial membrane antigen negative consistent with a high grade non-Hodgkin's lymphoma (Figs 1 and 2). Further immunocytochemistry confirmed this to be a B cell lymphoma. A small bowel meal showed a narrowed rectum distal to the ileorectal anastomosis, while a chest x ray film, total and differential white cell count, and bone marrow examination were normal. In addition, computed tomography of the thorax failed to show mediastinal lymph nodes.

Since the tumour was found to be fixed in the pelvis at surgery, it was left in situ with the formation of an ileostomy and closure of the rectum; the liver and spleen were normal.

The tumour therefore fulfilled Dawson et al and Richards's criteria for primary colonic lymphoma. Postoperatively the patient made an uneventful recovery and began combination chemotherapy as an outpatient.

Figure 1: Many mitotically active pleomorphic cells infiltrating the rectum.

Figure 2: Cell surface membranes of the infiltrating cells stained darkly with leucocyte common antigen.
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Discussion

Colonic adenocarcinoma is a recognised complication of ulcerative colitis with a 9% risk at 25 years in an extensive colitis. Less is known about the association between this disease and primary colonic lymphoma, a rare condition constituting only 0-2% of all colonic malignancies. There are now approximately 30 reported cases of colonic lymphoma in association with ulcerative colitis. Eight of these reports described lymphoma involving the rectum. Our patient is unique in that she developed primary lymphoma many years after colectomy and to our knowledge is the first reported case of rectal lymphoma after an ileorectal anastomosis. There has been one report of lymphoma in an ileostomy stoma after colectomy for ulcerative colitis.

The patient we described showed similar features to those patients in one of the largest series of colonic lymphoma in inflammatory bowel disease. These patients tended to have longstanding colitis, with left sided and often multiple tumours which were B cell, high grade, and at a late stage at diagnosis, being modified Dukes B and C. An explanation for this delayed presentation is that symptoms may be confused both clinically and radiologically with a disease flare up. Thus patients with longstanding colitis who notice a change in their symptoms should be thoroughly investigated.

The small numbers of reported cases makes it difficult to prove a definite association between ulcerative colitis and colonic lymphoma, although various mechanisms of pathogenesis have been postulated. These include repeated episodes of lymphoid hyperplasia and abnormalities of the reticuloendothelial system. The incidence of carcinoma of the rectum increases with time after an ileorectal anastomosis in ulcerative colitis and tends to occur in those patients with total colonic involvement before surgery. We believe our report further emphasises the importance of continued surveillance to detect rectal lymphoma as well as carcinoma.

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