Collagenous colitis: Histologic progression, extraintestinal features and lack of response to 5-ASA — A case report

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ABSTRACT: Collagenous colitis is a clinicopathologic syndrome which presents as chronic, watery, nonbloody diarrhea. Colonic biopsies characteristically show a thick subepithelial collagen band. Barium enema and colonoscopy generally fail to establish the diagnosis. This case report provides evidence for histologic progression over time, pancolonic involvement and the presence of extraintestinal features of collagenous colitis. The reported patient showed no response to 5-aminosalicylic acid. Colonoscopic biopsies in cases of chronic diarrhea can save the patient repeated, uncomfortable and costly investigations. Can J Gastroenterol 1990;4(1):19-22

Key Words: 5-aminosalicylic acid, Collagenous colitis, Chronic diarrhea, Extraintestinal features

La colite du collagène: Évolution histologique progressive, manifestations extraintestinales et manque de réponse à 5-ASA

RESUME: La colite du collagène est un syndrome anatomoclinique caractérisé par une diarrhée chronique aqueuse non sanglante. De manière caractéristique, les biopsies coliques montrent une épaisse bride de collagène sous-épithéliale. Le lavement baryté et la colonoscopie ne parviennent généralement pas à établir le diagnostic. Le cas présenté montre une évolution histologique progressive, une atteinte pancolique et des manifestations extraintestinales propres à la colite du collagène. Le patient concerné n'a pas répondu à l'acide 5-aminosalicylique. Dans les cas de diarrhée chronique, le recours aux biopsies colonscopiques peut épargner au patient des investigations répétées, désagréables et coûteuses.

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THE CLINICOPATHOLOGIC ENTITY of collagenous colitis has become increasingly recognized since its first description 13 years ago (1), with at least 50 cases reported in the English literature.

A case of collagenous colitis is reported which not only demonstrates the difficulty in diagnosis but also adds information with respect to mucosal progression, possible extraintestinal manifestations and treatment of this condition.

CASE PRESENTATION

A 71-year-old female presented with a 16 year history of intermittent diarrhea. The diarrhea was characterized by up to 15 loose to watery bowel movements per day, which often awakened her at night. It was preceded by bilateral lower abdominal cramps and associated with tenesmus. She denied the passage of blood but did notice mucus. These episodes occurred several times per year lasting up to three to four weeks and were followed by more regular soft to loose bowel movements num-
hering two to three per day. Over the past year prior to admission she began having more frequent bouts of diarrhea often leading to fecal incontinence. In addition, she complained of mild anorexia, nausea and severe heartburn that was relieved by H2 blocker therapy.

At age 56 she was found to have a slightly raised erythrocyte sedimentation rate, but stool cultures, sigmoidoscopy and barium x-rays of her upper and lower gastrointestinal tract were normal. An exploratory laparotomy revealed cholelithiasis and her gallbladder was removed. A jejunal biopsy was reported as normal.

At age 64 she was readmitted because of worsening diarrhea. She was found to have seronegative deforming arthritis involving the small joints of her hands and knees. Stool cultures, sigmoidoscopy, barium enema and liver-spleen scan were negative and she was treated with a fibre-enriched diet and psyllium hydrophilic mucilloid.

One year later she required readmission to hospital because of persistence of her symptoms despite the administration of codeine and loperamide. Repeat barium studies, a D-xylene excretion test, Schilling test, serum gastrin levels and gastric analysis were all normal as was the serum cortisol and a 24 h urine sample for 5-hydroxyindoleacetic acid (5-HIAA). A 72 h stool collection yielded an average of 165 g of feces with 1.8 g of fat per 24 h. A colonoscopy and repeat small bowel biopsy were within normal limits.

Further investigations during the following year again revealed negative stool cultures as well as a negative test for Clostridium difficile cytoxin. A small bowel aspirate and string test failed to recover Giardia lamblia cysts or trophozoites. A sigmoidoscopy was normal but a random rectal biopsy demonstrated 'nonspecific colitis' (Figure 1). She was treated with total parenteral nutrition for 11 days which led to improvement of her diarrhea and weight gain. She was discharged on codeine but experienced intermittent relapses of cramps and diarrhea over the next five years.

She presented again recently and was found to have a slightly elevated blood glucose level which was treated by diet. She complained of periodic depression particularly since her husband's death two years earlier. She denied the use of laxatives. Other past medical history included recurrent pneumonias over the past 15 years, a hysterectomy, bilateral ligation of varicose veins and a hemorrhoidectomy.

There was no family history of inflammatory bowel disease or other intestinal disorders.

On examination the patient appeared depressed. Her head and neck were normal and there was no thyromegaly. The chest and cardiovascular system examination were unremarkable apart from a grade II/VI systolic ejection murmur radiating to the base. There were two old scars on her abdomen, which was slightly distended and tympanitic. There was mild diffuse tenderness throughout her abdomen but no evidence of organomegaly or mass. Rectal examination was normal. She had joint deformities affecting her proximal and distal interphalangeal joints, wrists and knees.
but no actively inflamed joints were found. She had several patches of vitiligo over her forearms.

Further investigations in hospital demonstrated a normal complete blood count and erythrocyte sedimentation rate. Serum electrolytes, calcium and proteins were normal but the fasting blood glucose and glycosylated hemoglobin were mildly elevated. A 72 h stool collection resulted in a stool weight of 225 g per 24 h and 14 g of fat per day while on a 100 g fat containing diet. Stool electrolytes were potassium 99 mmol/L, sodium 61 mmol/L, chloride 68 mmol/L, with a stool osmolarity of 468 mmol/L. The pH was 8 and stool was negative for occult blood and reducing substances.

Upper endoscopy revealed grade II ulcerative esophagitis and esophageal biopsy was consistent with this. Small bowel biopsy was normal. A colonicoscopic examination was unremarkable but multiple biopsies obtained at 10 cm intervals from cecum to rectum showed the presence of collagenous colitis in all specimens (Figure 2). The patient was started on 5-aminosalicylic acid (5-ASA) (600 mg tid), a high fibre diet and psyllium mucilloid. Although she improved initially her diarrhea recurred after three weeks while on the medication and repeat biopsies showed no histologic improvement.

Subsequent therapy with sulfasalazine and oral prednisone was poorly tolerated and did not lead to symptomatic improvement. Currently, she is being treated with steroid enemas and she passes three to five semiformal bowel movements per day.

DISCUSSION
Collagenous colitis was first described by Lindström in 1976 (1) and has been the subject of recent reviews (2,3). This clinicopathologic syndrome characteristically presents with a chronic watery diarrhea and colonic mucosal biopsies reveal the presence of a linear subepithelial fibrous thickening and a chronic inflammatory infiltrate in the lamina propria. As in the patient described, radiologic and even colonoscopic findings may be entirely normal or nonspecifically abnormal and the diagnosis is eventually made only by mucosal biopsy (2,4).

The disease appears to be most common in middle-aged to elderly women. The diarrhea is often intermittent, nonbloody, and associated with abdominal cramps. It is frequently misdiagnosed as a manifestation of irritable bowel syndrome (5). However, the presence of nocturnal diarrhea, fecal incontinence and weight loss, should suggest an organic cause for diarrhea.

The patient in this case report suffered from a seronegative peripheral polyarthritis; its association with collagenous colitis has repeatedly been reported (6-8). In addition, the occurrence of glucose intolerance, thyroid disease and the finding of antinuclear antibodies in some patients with collagenous colitis has been cited as evidence for an autoimmune pathogenetic mechanism (9). The presence of vitiligo has not been described previously.

The reported patient initially had a rectal biopsy five years prior to the finding of a thickened subepithelial collagen table in the same anatomic location. This initial biopsy revealed an active colitis consistent with the recently recognized entity of microscopic colitis (10). Hence, this case adds further support to the hypothesis that collagenous colitis and microscopic colitis are but different aspects of the same condition (9,11,12). The pathogenesis of the formation of the collagen band is not completely understood, but likely requires the presence of chronic subepithelial inflammation (13). In addition, the existence of a local abnormality of collagen synthesis has been postulated (2).

This case demonstrates the pancolonic distribution of collagenous colitis (3,14). Characteristically, jejunal biopsies are normal except in rare cases in which celiac disease was found to be associated with collagenous or microscopic colitis (12,14-17). However, there was no evidence of small bowel villous atrophy in this patient and her mild steatorrhea remains unexplained.

The treatment response to various drugs is unpredictable and often disappointing (14). Both sulfasalazine and corticosteroids have induced remission in several cases (2,9) but have been ineffective in others (14). Anecdotal therapeutic success has been achieved with a number of other compounds including mepacrine (13), metronidazole (18), and bismuth subsalicylate (19). Sulfasalazine has been shown in several cases to lead not only to clinical but also histologic improvement seen in subsequent biopsies (9,12). Whether this effect was brought about by the antimicrobial (sulfonamide) or anti-inflammatory (5-ASA) moiety, however, is not clear. This patient was treated with 5-ASA and despite a good therapeutic dose and duration of therapy there was no clinical or histologic response. Furthermore, sulfasalazine did not lead to improvement of diarrhea. Rams et al (2) proposed that collagenous colitis is an inflammatory disorder, possibly of infectious origin, which is initially characterized by an acute inflammatory process which progresses over time and results in a gradual increase of collagen which may act as a diffusion barrier, further contributing to diarrhea. Conceivably, neither anti-inflammatory nor antimicrobial agents are effective at this later stage of the disease.

Finally, this case report underlines the importance of colonic biopsies even in the absence of colonic findings in patients with unexplained chronic diarrhea. If positive, the diagnosis is established, and repeated, uncomfortable and costly investigations can be avoided.

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REFERENCES


