An asymptomatic male with coexisting celiac disease and primary biliary cirrhosis

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Both primary biliary cirrhosis and celiac disease may be asymptomatic in many patients for extended periods of time. Rarely the disorders have been reported to occur together, in which case symptomatic malabsorption and diarrhea are virtually always present. An unusual case is presented of an asymptomatic male patient who had both primary biliary cirrhosis and celiac disease. The literature describing this coexistence, and theories regarding a common etiological link between the two disorders, are reviewed.

Key Words: Celiac disease, Malabsorption, Primary biliary cirrhosis

Patient asymptomatique porteur d'une maladie coeliaque et d'une cirrhose biliaire primaire

RESUME: La cirrhose biliaire primaire et la maladie coeliaque peuvent étre toutes deux asymptomatiques chez de nombreux patients et sur de longues périodes. Mais, d'après les rapports, ces deux affections se manifestent rarement ensemble – auquel cas une malabsorption et une diarrhée symptomatiques sont virtuellement toujours présentes. On rapporte le cas inhabituel d'un patient asymptomatique, porteurs des deux maladies; on examine la littérature faisant état de cette coexistence et des théories relatives au lien étiologique qui existerait entre les deux affections.

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Received for publication July 12, 1991. Accepted September 13, 1991

A number of autoimmune diseases have been reported in association with primary biliary cirrhosis, and in the presence of diarrhea and malabsorption, concomitant celiac disease should be considered in the differential diagnosis (1).

An unusual case is described of a 51-year-old male who was found to have both primary biliary cirrhosis and celiac disease, in the absence of symptoms.

CASE PRESENTATION

A 51-year-old male was referred in January 1989 because of an elevated serum alkaline phosphatase. He had been originally investigated in December 1987 for transient chest pain after lifting heavy objects. Cardiac evaluation was negative but the serum alkaline phosphatase was 1039 U/L (normal 41 to 133). Bone scan was normal. Repeat serum alkaline phosphatase six months later was 1080 U/L, with the hepatic isoenzyme fraction elevated.
The patient had been previously well except for a tonsillectomy and rheumatic fever at the age of nine years. He gave no history of alcohol or drug use. He never travelled outside Canada and denied weight loss, abdominal pain, diarrhea, gastrointestinal bleeding or a prior history of jaundice or anemia. There was no history of pruritus and the patient had no specific complaint. He denied any family history of gastrointestinal or hepatobiliary diseases.

Physical examination was completely normal and the patient lacked stigmata of chronic liver disease. Hemoglobin was 104 g/L (normal 117 to 166) with a hypochromic microcytic blood film showing numerous target cells. The white blood cell and platelet counts were normal. Serum iron studies and ferritin level confirmed the presence of iron deficiency. Serum alkaline phosphatase was 1195 U/L, gamma-glutamyltransferase 733 U/L (normal 9 to 76) and aspartate aminotransferase 112 U/L (normal 8 to 40).

Ultrasound of the abdomen and a liver-spleen scan were normal. An air contrast barium enema suggested a cecal filling defect, but colonoscopy ruled out any colonic lesion. Endoscopic retrograde cholangiopancreatography was completely normal and, in particular, the intrahepatic biliary tree was well visualized and free from any abnormality (Figure 1). However, the duodenal mucosa looked atrophic with loss of Kerckring’s folds. Two small bowel biopsies taken from the third part of the duodenum confirmed the presence of total villous atrophy (Figure 2). Further malabsorption workup revealed an abnormal D-xylose test and reduced serum and red cell folate levels. Serum B12, calcium, albumin, partial thromboplastin and prothrombin time were normal.

An upper gastrointestinal series and small bowel follow-through showed prolonged transit and mild flocculation of barium; there was also mild dilation of small bowel loops. Hepatitis B serology, antismooth muscle and antinuclear antibody titres were negative. The antimitochondrial antibody was present at a titre of greater than 1:640. A percutaneous liver biopsy showed expansion of portal tracts with replacement of the bile ductules by a
lymphocytic aggregate (Figure 3) and focal piecemeal necrosis in some portal triads. The patient was placed on a strict gluten-free diet and was given a three month course of oral iron and folic acid therapy.

Within a few weeks the patient reported significant improvement of energy, general well-being and weight gain, despite not having had any earlier complaints.

Re-evaluation six months later included a normal upper endoscopy. Small bowel biopsies taken from the distal third part of the duodenum showed restoration of the villous architecture with lack of inflammation and crypt hyperplasia (Figure 4). Hemoglobin was 159 g/L and folic acid and iron studies were normal. Liver enzyme profile showed elevated alkaline phosphatase (805 U/L) and aspartate aminotransferase (134 U/L), but normal bilirubin levels. A repeat antimitochondrial antibody titre was again greater than 1:640. His human lymphocyte antigen (HLA) haplotype was determined to be HLA-A1, B8, DR3.

A repeat liver enzyme profile in October 1990 showed a persistently elevated alkaline phosphatase of 762 U/L, an aspartate aminotransferase of 155 U/L and normal serum bilirubin levels.

At present, the patient continues to feel well on a gluten-free diet.

**DISCUSSION**

Celiac sprue is characterized by a structurally abnormal small intestinal mucosa leading to malabsorption which improves while the patient is on a gluten-free diet. Re-introduction of dietary gluten usually leads to clinical and histological relapse. The reported patient had evidence of malabsorption caused by inflamed atrophic small bowel mucosa. The clinical improvement and weight gain, the correction of iron and folate deficiencies, and the marked improvement in small intestinal biopsy after the institution of a gluten-free diet, all support the diagnosis of celiac disease in this patient.

Various liver diseases have been described in conjunction with celiac disease (2): the spectrum ranges from hepatic steatosis (3) and 'nonspecific' hepatitis to chronic active hepatitis and cirrhosis (4). In a retrospective analysis of 74 celiac patients, Hagender et al (2) documented histological evidence of liver injury in 16% and abnormal liver function tests in 39%, indicating that hepatic injury is not rare in adult celiac disease. More recently there has been a report of three celiac patients affected by primary sclerosing cholangitis (5).

Primary biliary cirrhosis, another autoimmune disorder characterized by a distinct histological picture – biochemical evidence of cholestasis in the presence of a normal biliary tree and the finding of antimitochondrial antibodies – has previously been described in patients with celiac disease (6). Up to the present time fewer than 20 fully documented cases of coexisting primary biliary cirrhosis and celiac disease have...
appeared in the literature (7-18). Although the majority of these patients were female, the present patient is the sixth reported male subject affected by both disorders. This might suggest that the large female preponderance of 10:1 seen in primary biliary cirrhosis (1) may be offset in the presence of gluten-sensitive enteropathy.

It is well recognized that both primary biliary cirrhosis and celiac disease may be relatively asymptomatic for extended periods of time. Indeed, a recent Scandinavian study suggests that about 50% of patients with adult celiac disease have no gastrointestinal symptoms (19). In primary biliary cirrhosis the prevalence of asymptomatic patients ranges from 13 to 48% (20). However, most of the reported cases in which both diseases coexisted were symptomatic, with weight loss, diarrhea and dermatitis herpetiformis most frequently reported (7-18).

The present patient was somewhat unusual in that he was virtually asymptomatic. Subclinical steatorrhea might have been present, as quantitative stool fat determinations were not performed. Although the patient had no specific complaints prior to treatment, he noticed an improved sense of well-being after starting a gluten-free diet, a phenomenon frequently seen in treated celiacs (21).

The present patient’s serum alkaline phosphatase decreased from 1195 U/L prior to treatment to 762, after he was on a gluten-free diet for slightly more than two years, while his aspartate aminotransferase level did not improve significantly. It is difficult to know whether this reflects an improvement in his liver disease, and a repeat liver biopsy would not necessarily be able to settle this issue because of the possibility of sampling error. Alternatively, improvement of an associated subclinical metabolic bone disease could have been reflected in a drop in serum alkaline phosphatase.

In none of the previously reported cases was there significant improvement of primary biliary cirrhosis after treatment with a gluten-free diet. Similarly, primary sclerosing cholangitis does not appear to be affected by dietary
treatment in the three recently reported patients with celiac disease (5). However, other types of liver injury in celiac patients may be reversible, and progressive liver disease may be preventable by a gluten-free diet in such patients (2).

There appears to be no obvious reason for the association between primary biliary cirrhosis and celiac disease. Jilfe and Owen (9) estimated the likelihood of both diseases occurring in the same patient to be on the order of one in 90 million; a chance association thus seems improbable.

Several theories have been put forward to explain a common etiology between the two diseases (14). A genetic predisposition appears unlikely, as the close association between celiac disease and the HLA loci B8 and DR3, seen in the present patient, is not shared with primary biliary cirrhosis (20). Circulating immune complexes derived from the normal gut mucosa could conceivably be deposited in portal tracts in the liver and thus play a direct role in tissue injury (7). This is also a proposed mechanism for other autoimmune disorders associated with celiac disease (22). Circulating immune complexes are found in some patients with primary biliary cirrhosis, but their role in the pathogenesis of the liver disease remains unclear, and current data suggest that it is unlikely that immune complexes play a significant role in primary biliary cirrhosis (20).

A common defect in the cellular immune response may be a possible etiological link between celiac disease and primary biliary cirrhosis. Hence, in celiac disease abnormal suppressor T cell function may permit cytotoxic reactions directed against enterocytes to which gluten is bound (23). Diminished T suppressor cell function has also been shown to occur in primary biliary cirrhosis, and this may play a part in the autoimmune process, allowing, for example, cytotoxic T lymphocyte responses against native cholangiocyte antigens or neoantigens (23).

Irrespective of the underlying pathogenetic mechanism, it is important for the clinician to be aware of the possible coexistence of both disorders, which may even be asymptomatic in the early stages, as was the case in the patient presently described. Biochemical hepatic evaluation should be included in the work-up of patients suffering from celiac disease.

REFERENCES
