Occult celiac disease associated with lymphocytic sclerosing cholangitis

HUGH J FREEMAN MD, WC PETER KWAN MD

Previous reports have described hepatobiliary tract abnormalities in celiac disease, including chronic cholestatic syndromes (1-3). Indeed, the coexistence of celiac disease and primary biliary cirrhosis is already well described in over 20 patients, including previous reports from two different Canadian teaching hospitals (4,5). Perhaps less well appreciated are the very rare reports of primary sclerosing cholangitis associated with celiac disease (3,6) or lymphoma (7), a condition also known to occur with increased frequency in patients with celiac disease (8,9). Weight loss, malabsorption, bone disease, steatorrhea and elevated alkaline phosphatase activities may be seen as common features in both celiac disease and either of these hepatobiliary tract disorders so that at an early stage of their coexistence, diagnosis of one or the other condition may be missed.

The pathological features in the small intestine of celiac disease are well known. These include blunting of the small intestinal mucosal villi, expansion of the lamina propria by lymphocytes and plasma cells, crypt hyperplasia and marked infiltration of the villous epithelium by lymphocytes with accompanying epithelial cell vacuolation and flattening (10). These in-
In June 1992 for further evaluation of abnormal liver chemistry tests and diarrhea for at least one year. In 1970, dermatitis herpetiformis was diagnosed; he was treated with intermittent dapsone. In 1976, an enlarged right inguinal lymph node was removed; a localized nodal lymphoma was detected and he was treated with chemotherapy and radiotherapy. No recurrent lymphoma was seen during follow-up at the British Columbia Cancer Agency in Vancouver. In May 1991, however, abnormal serum chemistry tests were first seen: alkaline phosphatase, 313 IU/L (normal 29 to 133); alanine aminotransferase, 61 IU/L (normal 7 to 35); and aspartate aminotransferase, 67 IU/L (normal 5 to 52). Because of diarrhea (one to four semiformal or loose stools per day), barium radiographs of the upper gastrointestinal tract were done in August 1991. These were normal. An abdominal ultrasound was also normal. Serum chemistry tests were repeated in March 1992 (alkaline phosphatase, 417 IU/L; aspartate aminotransferase, 85 IU/L) and May 1992 (alkaline phosphatase 555 IU/L; aspartate aminotransferase, 124 IU/L).

At the patient’s initial review at University Hospital in Vancouver, British Columbia, in June 1992, physical examination was normal. There was no lymphadenopathy and his liver and spleen were not enlarged. There were no peripheral stigmas of chronic liver disease. Laboratory investigations revealed: hemoglobin, 118 (normal 140 to 180 g/L); white blood cell count 4.4x10^3 (normal 4.0 to 11.0x10^3); and normal serum iron, iron binding capacity, ferritin, carotene, folic acid and vitamin B12. A bone marrow aspirate showed iron but no lymphoma. Upper gastrointestinal endoscopic evaluation in June 1992 was normal. There were no varices and the stomach and small intestine were macroscopically normal. Small bowel biopsies, however, showed a severe ‘flat’ lesion with crypt hyperplastic villous atrophy characteristic of celiac disease. Gastric biopsies were normal with no lymphocytic gastritis (16). Flexible sigmoidoscopy was normal but a colonic biopsy showed features of epithelial lymphocytosis (15). Serum chemistry tests were more abnormal (alkaline phosphatase, 630 IU/L; aspartate aminotransferase, 173 IU/L). Antinuclear and antimicrosomal antibodies were negative. Immunoglobulin G, A and M quantitation was normal. Fecal cultures and parasite examinations were negative. Abdominal ultrasound, abdominal and pelvic computed tomography scans were normal; lymphadenopathy was not detected. Percutaneous needle liver biopsy (Figures 1,2) revealed normal liver lobules with bile-duct centred inflammatory change, especially with lymphocytic infiltrates. The limiting plates were intact and there was no lobular inflammation. Focal damage to reactive bile duct epithelium with bile ductular proliferation was present. Larger portal tracts showed inflammation with some fibrosis present in a concentric lamellar arrangement; severe reactive atypia of bile duct epithelial cells was also present. Lymphoma was not present. Following liver biopsy, a gluten-free diet was initiated. By July 1992, his bowel habit was normal with one formed stool daily. Serum chemistry tests had improved: alkaline phosphatase, 465 IU/L; and aspartate aminotransferase, 168 IU/L. Endoscopic retrograde cholangiography (Figure 3), however, showed multiple strictures within the intrahepatic biliary tree associated with areas of proximal dilatation. A long tubular fixed narrowing with irregular margins extended throughout the length of the common bile duct but contrast was seen to flow into the duodenum. The appearances suggested sclerosing cholangitis involving both the intrahepatic and extrahepatic biliary tree. A duodenal biopsy done at the time of the cholangiogram showed histological improvement with reappearance of villi. After introduction of the gluten-free diet, serum chemistry tests in November 1992 (alkaline phosphatase, 287 IU/L; aspartate aminotransferase, 117 IU/L; alanine aminotransferase, 93 IU/L) and March 1993 (alkaline phosphatase, 224 IU/L; aspartate aminotransferase, 51 IU/L; alanine aminotransferase, 53 IU/L) were improved but not normal; an abdominal ultrasound repeated in January 1993 was normal.
DISCUSSION

The presented patient had histologically defined occult celiac disease that was preceded by a long-standing history of dermatitis herpetiformis and a successfully treated lymphoma, a relationship previously described elsewhere (17). Studies in this patient, however, also revealed the presence of lymphocytic colitis (15) and an antimitochondrial antibody negative form of chronic cholestatic liver disease. Further investigations showed a bile duct centred inflammatory process characterized by fibrosis, a predominance of lymphocytes in duct epithelium and cholangiographic features typical of primary sclerosing cholangitis involving the intrahepatic and extrahepatic bile ducts. A number of hepatobiliary tract disorders have been recorded in celiac disease including cholestatic liver diseases such as primary biliary cirrhosis (4,5). Although a common immunological basis may be present, extensive studies to date have failed to identify a common genetic predisposition or common immunological alteration in celiac disease and primary biliary cirrhosis (5). Changes of primary sclerosing cholangitis have been only very rarely recorded in patients with celiac disease (3,6), and the precise relationship between celiac disease and primary sclerosing cholangitis has never been systematically explored (19). Either small bowel biopsy studies of patients with primary sclerosing cholangitis or cholangiographic studies of patients with celiac disease are needed to define this potentially intriguing relationship more precisely. As in celiac disease, many patients with primary sclerosing cholangitis are asymptomatic. Indeed, as is well known in celiac disease, our patient had only limited symptoms despite histological detection of both small and large intestinal disease as well as eventual cholangiographic documentation of rather extensive intrahepatic and extrahepatic radiological changes. These findings further emphasize that biliary tract epithelial changes may be more frequent in celiac disease than is appreciated.

Additional studies are needed to elucidate the natural history of the biliary tract abnormalities in celiac disease and possible biliary tract responses in these patients to a gluten-free diet. Although at least two previous studies indicated that abnormal liver chemistry tests (ie, aspartate aminotransferase, alanine aminotransferase, alkaline phosphatase activities) in celiac disease patients with various forms of hepatobiliary disease improve with gluten restriction (2,3) – as was apparent in this present patient with sclerosing cholangitis – it is premature to suggest that these improved blood tests were indicative of a convincing biliary tract re-
address conclusively this issue. Previous pathological studies have emphasized the predominately lymphocytic nature of the portal inflammatory process in celiac associated liver disease (1). Detailed quantitative studies of intraepithelial lymphocyte numbers in treated celiac disease have shown a persistent elevation in the intestine despite a gluten-free diet (20,21). Thus, the biliary tract response to a gluten-free diet might be difficult to document if the defining pathological end-point consists of lymphocyte counts in ductal epithelium. Increased clinical awareness of this 'more than chance' (19) association of sclerosing cholangitis with celiac disease, however, may eventually permit a more detailed exploration of a possible biliary response to gluten restriction.

Previous studies have documented a wide array of immunological abnormalities in celiac disease as well as sclerosing cholangitis (22-24), and various theories have been noted to explain the concomitant presence of celiac disease and hepatobiliary tract disease. Possibly, immune complexes are formed with a common antigenic basis resulting in tissue damage; no specific antigen, however, has been detected. Alternatively, diminished suppressor T cell function might permit effector cytotoxic lymphocytes to alter a modifying antigen, like gluten. These effector cells might then recognize and attack a patient's own histocompatibility antigens, such as human leukocyte antigen B-8 and deoxyribose-3 phosphates recognized in high incidence in both diseases (19), and possibly present in high concentrations in biliary and intestinal epithelial cells. More immunological studies are needed to elucidate further this intriguing relationship between celiac disease and lymphocytic sclerosing cholangitis.

REFERENCES
