Celiac disease associated with primary biliary cirrhosis in a Coast Salish native

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Celiac disease associated with primary biliary cirrhosis in a Coast Salish native. Can J Gastroenterol 1994;8(2):105-107. A 41-year-old Coast Salish female was initially diagnosed with typical features of classical adult celiac disease. Clinical and pathological features of primary biliary cirrhosis were also present, along with a familial history of insulin-dependent diabetes. Later, childhood celiac disease was detected in a male first-degree relative with diabetes. These patients are the first reported natives in Canada with celiac disease, a disorder believed to be genetically based but dependent on environmental factors for its clinical expression. The recognition of a 'new' disease in the setting of an aboriginal population may reflect geographical and climatic factors that permitted subsistence of this culturally complex food-gathering society up until most recent historical times, followed by adaptation of this society to European-based agricultural methods, particularly wheat cultivation.

Key Words: Antimitochondrial antibodies, Celiac disease epidemiology, Liver disease, Malabsorption, Native celiac disease, Primary biliary cirrhosis

Maladie cœliaque associée à une cirrhose primaire chez une autochtone Salish de la côte ouest

RÉSUMÉ : Une patiente Salish, de la côte ouest, a d’abord reçu un diagnostic de maladie cœliaque de l’adulte. Des caractéristiques cliniques et pathologiques de cirrhose biliaire primaire étaient également présentes, de même que des antécédents familiaux de diabète insulinodépendant. Par la suite, une maladie cœliaque infantile a été décelée chez un parent du premier degré atteint de diabète. Ces deux sujets sont les deux premiers autochtones du Canada chez qui le diagnostic de maladie cœliaque est déclaré; cette affection avait longtemps été considérée avant tout génétique, quoique dépendante de facteurs environnementaux pour son expression clinique. L’observation d’une « nouvelle » maladie chez une population aborigène pourrait signaler le rôle de facteurs géographiques et climatiques dans la survie de cette complexe société encore non agraire, jusqu’à très récemment, et qui a dû s’adapter aux méthodes agricoles d’origine européenne, particulièrement à la culture du blé.

Epidemiological studies on the geographical distribution of celiac disease suggest that it occurs almost exclusively in Europeans and their descendants, particularly those who have emigrated to North America and Australia (1). In Ireland, for example, the prevalence of clinically apparent celiac disease is estimated to be as high as one in 300 (2). Interestingly, there is a very high frequency of human leukocyte B8 antigen in western Ireland, a genetic marker known to be linked to a number of disorders, including celiac disease. In addition, there is a very long cultural history of wheat cultivation in Europe dating from at least 1000 BC (3). Such genetic and environmental factors appear to have played a critically important role in the clinical expression of celiac disease in different populations (4).

While recent epidemiological studies of celiac disease in Europe and other Mediterranean countries have been reported (5), this disorder has also been widely recognized in inhabitants from a number of other non-European countries (1) and has occasionally been detected in other non-Caucasian races, including possibly Blacks (6,7), but not in Asians (8). Although both genetic and environmental factors are important (4), the very low case fatality rate of celiac disease, its relative rarity in most populations and wide variations in case ascertainment methods for dif-
fferent countries have made epidemiological studies of celiac disease more difficult (9). It has been estimated, for example, that up to one-half of all adult celiac patients identified in some populations have few or no symptoms (9); obviously, then, case ascertainment would substantially influence differences in prevalence rates for celiac disease that might be reported from different centres. Additional studies are required, therefore, to map more precisely the epidemiological topography of celiac disease on a global scale.

This report documents an adult Coast Salish female with celiac disease, and explores possible environmental factors that may be significant historically in the appearance of a 'new disease' in the native peoples of British Columbia.

CASE PRESENTATION

A 41-year-old Coast Salish female was referred because of weight loss of 18 kg. She also had pruritus, and her family physician found altered liver chemistry results including: alkaline phosphatase, 906 IU/L (normal 23 to 103); total bilirubin, 21 µmol/L (normal 2 to 17); and aspartate aminotransferase, 186 IU/L (normal 5 to 52). Other laboratory studies showed a mild hypochromic, microcytic anemia with: hemoglobin, 113 g/L (normal 118 to 154); erythrocyte sedimentation rate, 43 mm/h (normal 0 to 20); ferritin, 9 nmol/L (normal 25 to 250); INR, 0.8 (normal 0.9 to 1.2); cholesterol, 7.74 mmol/L (normal 3.36 to 5.19); total protein, 82 g/L (normal 61 to 78); albumin, 38 g/L (normal 35 to 55). Serological studies for hepatitis A, B and C viruses were negative and an abdominal ultrasound was normal.

Further studies following referral revealed positive antimitochondrial and smooth muscle antibodies (both in titres of 1:640); an elevated serum immunoglobulin M level of 4.24 g/L (with normal levels of other immunoglobulins); normal red cell folate, 896 nmol/L (normal 400 to 1600); and normal vitamin B12, 586 pmol/L (normal 150 to 600). Although serum calcium levels were normal, 25-hydroxy-vitamin D was 35 nmol/L (normal 40 to 185) and 1,25-dihydroxy-vitamin D was 95 pmol/L (normal 36 to 119). Bone densitometry studies revealed changes consistent with early osteoporosis. A liver biopsy revealed features consistent with primary biliary cirrhosis, a condition previously associated with adult celiac disease and reported by other investigators elsewhere (10-22).

Because of diarrhea, endoscopic studies of the upper and lower gastrointestinal tracts were done and were normal. Small intestinal biopsies from the proximal small bowel, however, showed typical changes consistent with celiac disease. Gastric and colonic mucosal biopsies failed to show changes of lymphocytic gastritis (23), or collagenous or lymphocytic colitis (24,25). A gluten-free diet resulted in resolution of the patient's diarrhea, weight gain of 6 kg and normalization of her small intestinal biopsies, resulting in the typical histological features of treated celiac disease.

Additional Coast Salish familial historical data revealed a 10-year-old insulin-dependent diabetic daughter; in addition, the patient's sister and two-year-old nephew were diagnosed as diabetic. The nephew, a first-degree relative, also developed diarrhea, and small intestinal biopsies done at another teaching hospital showed features typical of childhood celiac disease.

DISCUSSION

The native patient described in the present report had both celiac disease and primary biliary cirrhosis, two conditions that have previously been observed to coexist in over 20 patients from other populations (10-22). In addition, both conditions are often associated with a range of other disorders having an immunological basis (21,26), including diabetes and thyroiditis, although evidence for a common genetic predisposition or a common immunological alteration has not yet been identified. Weight loss, malabsorption, bone disease, steatorrhea and elevated alkaline phosphatase activities are common features of both diseases so that at an early stage of their coexistence, celiac disease or primary biliary cirrhosis may not readily be recognized. In all patients reported with both conditions, regardless of geographical origin or race, restriction of dietary gluten caused cessation of diarrhea, as in the patient reported here, but abnormal liver chemistry tests were not improved despite a gluten-free diet.

A number of theories have been considered to explain the concomitant presence of celiac disease and primary biliary cirrhosis. One suggests that immune complexes are formed with a common antigenic basis, and this mechanism mediates tissue damage; no specific antigen, however, has been identified. Alternatively, diminished function of suppressor T cells in patients with both diseases might allow effector cytotoxic lymphocytes to attack a modifying antigen, such as gluten. These effector cells might then recognize and attack a patient's histocompatibility antigens, present in high concentrations in biliary as well as intestinal epithelial cells (23). Further studies are needed, however, to elucidate the very intriguing relationship between celiac disease and primary biliary cirrhosis.

The recognition of celiac disease in this native patient has particular interest as a 'new' disorder appearing for the first time in this population. The Coast Salish seemed to have survived and attained one of the highest known levels of cultural complexity, subsisting largely on fishing as well as gathering of roots and berries, rather than horticulture or herds (27). As a result of their rich, maritime temperate zone habitat, these peoples developed societies with permanent houses in villages of more than 1000 residents, social stratification with slaves and ranked nobility, elaborate ceremonies, multiple linguistic dialects and one of the world's great art styles (27). Early descriptions are reported to describe the Coast Salish tribes as a hunting and fishing peoples without knowledge of soil cultivation methods (28). Although possible sources of potato and wheat cultivation may have been Russian settlements in Alaska or some Spanish settlements at Nootka Sound in British Columbia, most historical evidence suggests that
the Coast Salish tribes acquired some cultivation techniques only after the establishment of Fort Langley in 1827 by the Hudson’s Bay Company. This was followed by very rapid diffusion of the methods elsewhere in British Columbia (29). Previously, access to Europeans was very limited. This recent and very significant change from a food-gathering society to an agricultural food-producing society apparently occurred rapidly in these aboriginal peoples. Although heritable factors for celiac disease are clearly important in the clinical expression of this disorder, recent environmental changes during the past century, including the introduction of wheat cultivation methods, may be relevant in the pathogenesis of celiac disease in this particular population.

REFERENCES
