Intestinal intussusception as an atypical presentation of celiac disease

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CASE REPORT

A 31-year-old woman was studied because of chronic ferropenic anemia and colicky abdominal pain for many years. An upper gastrointestinal endoscopy was performed that revealed no mucosal changes, with a normal villous pattern and normal duodenal biopsy. Colonoscopy yielded no findings. A CT scan revealed a thickened jejunal wall with mesenteric adenopathies and no evidence of intestinal ischemia or obstruction. Capsule endoscopy (CE) was indicated, which revealed from the proximal jejunum the characteristic, diffusely cracked mucosa of celiac disease (CD), with atrophy and a mosaic pattern (Fig. 1A). A jeuno-jejunal invagination was identified in the distal jejunum (Fig. 1B) with a transition of normal mucosa over the involved mucosa that might correspond to the image shown by CT. CE showed a preserved villous pattern in the distal small bowel. A subsequent double-balloon enteroscopy with histopathologic study confirmed the presence of patchy CD in the proximal jejunum.

DISCUSSION

CE is indicated for suspected atypical or patchy CD, where duodenal biopsy may obtain a false negative result (1,2). Furthermore, in established CD, CE may be useful to monitor mucosal healing, or in refractory cases to screen potential accompanying complications (ulcerative jejunitis, lymphoma) (3).

The cause of recurrent colicky pain in CD remains unclear. However, the invagination of the atrophic proximal mucosa into the normal distal (thicker) mucosa has been suggested to account for this manifestation.

Transient intestinal intussusception is a rare complication of CD more commonly seen in children (4). In the adult, organic disease (neoplasms) must be ruled out, but in a proper clinical setting intussusception may be highly suggestive of CD. In our patient, CE allowed a diagnosis of suspected CD and ruled out other, potentially more severe causes of invagination.

REFERENCES