“Leopard skin sign”: the use of narrow-band imaging with magnification endoscopy in celiac disease

Signo de la “piel de leopardo”: el uso de endoscopía con magnificación y narrow banding en enfermedad celiaca

Asadur J. Tchekmedyian 1, Emmanuel Coronel 2, Frank Czul 3

1 Universidad de la República, Departamento de Gastroenterología Prof. Henry Cohen, Hospital de Clínicas. Montevideo, Uruguay
2 Division of Gastroenterology, Hepatology and Nutrition, Department of Medicine, University of Chicago. Chicago, EE UU.
3 Miller School of Medicine, Division of Gastroenterology, Department of Medicine, University of Miami. Miami, EE UU.

Recibido: 05-08-2014; Aprobado: 06-10-2014

ABSTRACT
Celiac disease (CD) is an immune reaction to gluten containing foods such as rye, wheat and barley. This condition affects individuals with a genetic predisposition; it targets the small bowel and may cause symptoms including diarrhea, malabsorption, weight loss, abdominal pain and bloating. The diagnosis is made by serologic testing of celiac-specific antibodies and confirmed by histology. Certain endoscopic characteristics, such as scarring, reduction in the number of folds, mosaic-pattern mucosa or nodular mucosa, are suggestive of CD and can be visualized under white light endoscopy. Due to its low sensitivity, endoscopy alone is not recommended to diagnose CD; however, enhanced visual identification of suspected mucosal abnormalities through the use of new technologies, such as narrow band imaging with magnification (NBI-ME), could assist in targeting biopsies and thereby increasing the sensitivity of endoscopy. This is a case series of seven patients with serologic and histologic diagnoses of CD who underwent upper endoscopies with NBI-ME imaging technology as part of their CD evaluation. By employing this imaging technology, we could identify patchy atrophy sites in a mosaic pattern, with flattened villi and alteration of the central capillaries of the duodenal mucosa. We refer to this epithelial pattern as “Leopard Skin Sign”. Since epithelial lesions are easily seen using NBI-ME, we found it beneficial for identifying and targeting biopsy sites. Larger prospective studies are warranted to confirm our findings.

Key words: Celiac disease; Endoscopy; Duodenum; Narrow band imaging; Diagnosis (source: MeSH NLM).

ABSTRACT
La enfermedad celiaca (EC) es una reacción inmune a los alimentos que contienen gluten como el centeno, el trigo y la cebada. Esta condición afecta a las personas con predisposición genética, comprometiendo al intestino delgado causando síntomas como diarrea, mala absorción, pérdida de peso, dolor abdominal y meteorismo. El diagnóstico se hace con estudios serológicos de anticuerpos celiacos y es confirmado por histología. Algunas características endoscópicas tales como “scalloping”, reducción en el número de pliegues, patrón mucoso tipo mosaico o mucosa nodular, son sugestivos de EC y se pueden observar con endoscopia de luz blanca. Debido a su baja sensibilidad la endoscopia por sí sola no se recomienda para diagnosticar EC, sin embargo, una visualización cuidadosa de las anormalidades mucosas sospechosas a través de nuevas tecnologías como “Narrow Band Imaging” con magnificación (NBI-ME) puede ayudar a dirigir las biopsias y así incrementar la sensibilidad de la endoscopia. Esta es una serie de siete casos con diagnóstico serológico e histológico de EC a quienes se les realizó una endoscopia digestiva alta con NBI-ME. En ellos se pudo identificar sitios de atrofia parcelar en un patrón de mosaico, con vellosidades aplanadas y alteración de los capilares de la mucosa duodenal. Nos referimos a esta alteración como el “Signo de la Piel de Leopardo”. Como las lesiones epiteliales se ven fácilmente usando NBI-ME, lo encontramos beneficioso para identificar y dirigir los sitios donde tomar las biopsias. Estudios prospectivos más grandes deben realizarse para confirmar nuestros hallazgos.

Palabras clave: Enfermedad celiaca; Endoscopia; Duodeno; Imagen de banda estrecha; Diagnóstico (fuente: DeCS BIREME).

INTRODUCTION
Celiac disease (CD) is defined as an immune reaction to gluten containing foods such as rye, wheat and barley. This condition affects individuals with a genetic predisposition; it targets the small bowel and may cause symptoms including diarrhea, malabsorption, weight loss, abdominal pain and bloating. A CD diagnosis is made by serologic testing of celiac-specific antibodies in individuals following a gluten containing diet and is confirmed by endoscopy with multiple duodenal biopsies for histology. The mainstay of therapy is a gluten-free diet (GFD) which requires significant education and commitment (1,2).

The role of endoscopy in CD has been evaluated in multiple studies, mainly to describe the pattern of mucosal injury and test the accuracy of visualization of villous atrophy. In CD, the intestinal damage can have a patchy distribution, and the macroscopic features can be related to the degree of severity of the histological lesions. Several endoscopic features reflect the presence...
of villous atrophy; however, their sensitivity varies from 50% to 94%. Consequently, endoscopy alone is not recommended to establish a CD diagnosis. Endoscopy plays an important role, though, in the diagnosis and management of this disease. Enhanced visual identification of suspected mucosal atrophy could assist in targeting biopsies and increase the sensitivity of endoscopy, reducing the need of random biopsies.

Narrow band imaging (NBI) is a technology that uses two different light filters reducing the light wavelength to 415 and 540 nanometers; the penetration of the shorter wavelength blue light is limited to the mucosa. This corresponds to the main and secondary hemoglobin absorption improving the contrast between the sub mucosal vasculature and mucosa, enhancing the mucosal structure.

This is a case series of seven patients with a serologic and histologic diagnosis of CD who underwent evaluation with narrow-band imaging with magnification endoscopy (NBI-ME) prior to duodenal biopsy taking.

**CASE SERIES**

This case series was obtained from the Department of Gastroenterology of the “Hospital de Clinicas” in Montevideo, Uruguay. All patients had serology results consistent with CD. Two patients had established CD diagnoses, positive serologies and biopsies, and were evaluated due to refractory symptoms. The five remaining patients were initially diagnosed with CD by serology and confirmed with endoscopy and histology. Patients underwent upper endoscopies with NBI prior to duodenal biopsies as part of their evaluation. Table 1 represents patient characteristics and histologic findings using the Modified Marsh Classification.

We used a magnification gastroscope (GIF Q160Z, Olympus, Tokyo, Japan) with optical magnification of 115x and NBI technology. At the time of endoscopic evaluation with white light, it was difficult to find all the characteristic endoscopic changes suggestive of CD such as: scalloping and reduction in the number of folds, mosaic-pattern mucosa, and nodular mucosa. We used NBI-ME to target the biopsy sites and obtained multiple biopsies (at least 4). All seven patients had severe intestinal atrophy by histology, and through NBI-ME, all displayed patchy atrophy sites in a mosaic pattern, with flattened villi and alteration of central capillaries of the duodenal mucosa. We called this epithelial pattern the “Leopard Skin Sign” (Figures 1 and 2). We found NBI useful for identifying and targeting biopsy sites since the pattern of epithelial lesions is easily observed using this technology.

Table 1. Baseline characteristics of patients according to clinical presentation, serology consistent with CD, microscopic characteristics of duodenal biopsies using the Modified Marsh Classification. (GFD) Gluten free diet, (TTG) tissue-transglutaminase, (+) positive, (-) negative.

<table>
<thead>
<tr>
<th>Case</th>
<th>Patient characteristics</th>
<th>Serology</th>
<th>Modified Marsh Classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>41 y/o woman evaluated for alternating diarrhea, bloating and abdominal pain</td>
<td>Anti TTG (IgA) + Antigliadin - Antiendomisial +</td>
<td>3c</td>
</tr>
<tr>
<td>2</td>
<td>40 y/o woman with chronic diarrhea and bloating</td>
<td>Anti TTG (IgA) + Antigliadin + Antiendomisial +</td>
<td>3c</td>
</tr>
<tr>
<td>3</td>
<td>29 y/o woman with diarrhea and bloating</td>
<td>Anti TTG (IgA) + Antigliadin + Antiendomisial +</td>
<td>3b-3c</td>
</tr>
<tr>
<td>4</td>
<td>24 y/o woman with diarrhea, hair loss, malaise and skin lesions consistent with dermatitis herpetiformis</td>
<td>Anti TTG (IgA) + Antigliadin + Antiendomisial +</td>
<td>3a-3b</td>
</tr>
<tr>
<td>5</td>
<td>29 y/o woman with confirmed celiac disease, evaluated because of refractory abdominal symptoms due to non-compliance with GFD</td>
<td>Anti TTG (IgA) + Antigliadin + Antiendomisial +</td>
<td>3c</td>
</tr>
<tr>
<td>6</td>
<td>39 y/o woman with confirmed celiac disease, evaluated because of refractory abdominal symptoms due to non-compliance with GFD</td>
<td>Anti TTG (IgA) + Antigliadin + Antiendomisial +</td>
<td>3c</td>
</tr>
<tr>
<td>7</td>
<td>34 y/o woman evaluated for chronic watery diarrhea</td>
<td>Anti TTG (IgA) + Antigliadin + Antiendomisial +</td>
<td>3b</td>
</tr>
</tbody>
</table>
Figure 1. NBI-ME of the duodenum in seven patients with CD. Panels (a-g) show a patchy appearance/mosaic pattern of flattened villi and alteration of central capillaries of the duodenal mucosa. We called this epithelial pattern the “leopard skin sign”. Panel (h) leopard skin picture to compare with NBI-ME images. Narrow–band imaging (NBI), magnification endoscopy (ME), celiac disease (CD).

Figure 2. White-light and NBI-ME images of two patients with confirmed CD. Panels (a,b) are the NBI-ME pictures showing the same mucosal pattern described in figure 1. Panels (c,d) are the white-light pictures to compare. Narrow–band imaging (NBI), magnification endoscopy (ME), celiac disease (CD).
DISCUSSION

It is controversial whether CD can be diagnosed without a duodenal biopsy. The European Society of Pediatric Gastroenterology, Hepatology, and Nutrition (ESPGHAN) proposed that it may be possible to avoid any intestinal biopsy in children who meet the following criteria: characteristic symptoms of CD, Tissue Transglutaminase (TTG) IgA levels > 10 × upper limit of normal (confirmed with a positive anti-endomysial antibody in a different blood sample), and positive HLA-DQ2 (triple testing). There is no sufficient data to recommend triple testing as a diagnostic strategy since there is no standardization of TTG assays and there is not enough data in unselected populations such as family members of individuals with confirmed CD.

Endoscopic evaluation with histopathological sampling is an important step for patients undergoing evaluation for CD or as part of their management. Routinely, white light endoscopy is used to study patients with CD, but it has been found to have a low sensitivity in detecting typical mucosal changes. New imaging technologies have been used to facilitate the diagnosis of different digestive conditions such as early dysplasia of the intestinal tract. There is not sufficient data, though, to recommend its systematic use for diagnosis/confirmation of CD. Similar to a prior study that found magnification endoscopy improved the sensitivity and specificity of endoscopy in the diagnosis of CD, we found that combining NBI with this technology useful for identifying and targeting biopsy sites as well as enhancing areas of mucosal atrophy for better visualization.

Better assessment of mucosal healing in patients with CD is warranted. Recent findings described by Lebwohl et al., describe an increased risk of lymphoproliferative malignancy in patients with CD and persistent villous atrophy, hence the importance of using enhanced imaging technologies to assess mucosal architecture.

The small number of patients is a limitation of our case series; larger prospective studies are necessary to confirm our present findings.

Acknowledgements:

To: Prof. Dr. Henry Cohen, Dr. Julio Chilewski, Dr. Juan Pablo Gutierrez, Dr. Andres Taullard and Dr. Virginia Lopez (Hospital de Clinicas, Montevideo, Uruguay) who have helped by obtaining data for the preparation of this manuscript.

Conflict of interest:

None of the authors have any conflicts of interest to disclose.

REFERENCES


Correspondence:

Dr. Asadur Jorge Tchekmedyian Rambala República del Perú 1413 – Ap. 201. Montevideo 11300. Uruguay
E-mail: asadur@adinet.com.uy