A 62-year-old man had been passing small stool for 2 months. He had been diagnosed as having mantle cell lymphoma of the spleen 8 years ago, and complete remission was achieved after chemotherapy. A computed tomography (CT) scan of the abdomen, taken to evaluate the intrabdominal lymph nodes, revealed a rectal mass with perilesional lymphadenopathy. A primary rectal cancer was suspected (Fig. 1, 2). Colonoscopy revealed an ulcerative rectal mass with loss of rectal glandular structure confirmed with narrow-band imaging (Fig. 3, 4). Histological examination of the biopsy specimens showed numerous lymphocytic infiltrations (Fig. 5, 6), which were positive for cyclin D1 for B cells. Recurrent mantle cell lymphoma was diagnosed and the patient was treated with rituximab-based chemotherapy.

A large solitary ulcerative rectal mass is a typical presentation of primary rectal cancer. Intestinal involvement of mantle cell lymphoma, in contrast, typically presents with multiple lymphomatous polyposis [1, 2]. The stomach is the favored location [2]. Rates of involvement as documented in previous endoscopy reports are: esophagus 6%, stomach 74%, duodenum 34%, ileum 48%, cecum 14%, colon 57%, and rectum 48% [4]. Intestinal lesions of mantle cell lymphoma presented as multiple lesions in nearly 80%, whereas a protruding mass was found in 18% [2]. Narrow-band imaging of mantle cell lymphoma of the stomach has revealed loss of normal glandular structure and tree-like appearance of abnormal blood vessels [3]. The present rectal mantle lymphoma showed hypervascularity of the mucosa with loss of standard rectal glandular structure. Immunohistochemical staining provides a definite diagnosis, and infiltrates of small, atypical lymphocyte-like cells, which stain positive with pan B-cell marker, T-cell CD5, and cyclin D1, are

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Fig. 1  Computed tomography (CT) scan (coronal section) showing a 4-cm intraluminal polypoid mass (arrow) at the left rectal wall, near the anorectal junction.

Fig. 2  Computed tomography (CT) scan (axial section) showing several enlarged lymph nodes (arrows) along the presacral space.

Fig. 3  A central ulcerative mass, 4 cm in diameter, at the posterior wall of rectum, at 3 cm from the anal verge, demonstrating contact bleeding.

Fig. 4  Narrow-band images demonstrating loss of rectal glandular structure with increased numbers of abnormal blood vessels.

Fig. 5  Abnormal dense lymphoid infiltration in the lamina propria. There was no proliferation in the center or lymphoepithelial lesion (hematoxylin and eosin stain).
characteristic of the disease. Typically the disease is aggressive and the median survival is 3–5 years despite aggressive chemotherapy [4].

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Fig. 6 Small to medium sized lymphoid cells with hyperchromatic irregular nuclei and some scattered plasma cells. Note that the normal glandular structures have been destroyed.