Ileocecal invagination due to an inflammatory fibroid polyp. A case report

The invagination develops as a result of the dislocation of proximal bowel segment within the following distal portion. It is rarely seen in adult population. It generally develops as a result of intestinal wall neoplasia in the adult patients. Only 3-6% of gastrointestinal system tumors are seen to be localized to small bowels. Inflammatory fibroid polyp is a rarely seen benign tumor in gastrointestinal system. We present the case of 47 year-old female with ileo-cecal invagination caused by Inflammatory fibroid polyp in ileum.

**KEY WORDS:** Invagination, Intussusception, Inflammatory Fibroid Polyp

**Case Report**

47 year-old female patient having no comorbidity disease or previous abdomen surgery history applied to our emergency service with the complaints of abdominal pain for 1 week and vomiting and nausea for 2 days. There was no fecal discharge for 2 days. In abdominal examination, the distention, decrease in intestinal voice, and sensitivity throughout the abdomen were detected. In rectal examination, the rectum was empty and there is no indicator of intestinal hemorrhage. Hematocrit was 28.1, hemoglobin was 9.1, and white blood cell count was 14.8. The serum electrolyte level was normal. In direct abdominal radiograph taken in standing position, a liquid-air leveling localized to right lower quadrant was seen (Fig. 1A). In contrasted abdominal tomography (CT), ileocecal invagination was observed in approximately 8cm segment of right lower quadrant, and the jejunal anuses were dilated in proximal (Fig. 1B). Taken into surgery with these findings, the patient was found to have approx. 10 cm ileum segment (starting from the ileocecal valve) invaginated into the cecum. There was no circulatory impairment. When the bowel was re-invaginated, a 3cm x 5cm intralumen mass was observed at approx. 10cm proximal to the ileocecal valve.
The patient was undergone right hemicolectomy and extreme collateral ileotransversostomy with terminal ileum resection. The patient was discharged on 4th postoperative day, and kept in follow-up period for 3 months. No complication was observed since then. In histopathological analysis, the mixed-type inflammatory cell infiltration rich in eosinophil leucocytes and the veins were observed in tissues of small bowels. In immune staining, Vimentin and CD34 were found to be positive, whereas CD117 (C-KT) was found to be negative. The IFP diagnosis was made based on these findings (Fig. 3).

**Discussion**

Even though the invagination was observed rarely among the adults, 90% of the cases originate from the mass developing within the lumen. The development of small bowel obstruction might occur in acute, chronic, and intermittent manner. The abdominal pain, nausea, vomiting, and (especially in malign lesions) melena might be seen. In the present study, there were abdominal pain for 1 week and nausea and vomiting for 2 days but no melena. Among the adults, 65-75% of the invagination orig-
nates from the benign or malign lesions, whereas 15-25% arises from the non-neoplastic factors. Only 3-6% of GIS tumors are seen to be localized to small bowels. The small bowel tumors are very hard-to-diagnose unless they lead to hemorrhage or obstruction. They are generally diagnosed lately. In the present case, the diagnosis was made upon observing the invagination in CT taken after the obstruction and detecting the mass during the surgical procedure.

IFP is a rare lesion of GIS identified by Vanek. It is generally asymptomatic. The symptoms are seen depending on the localization. It causes invagination in 5-16% of the cases. From macroscopic aspect, it is in sessile or pedicled polypoid form. It contains inflammatory cells such as eosinophil, lymphocyte, and plasm cells. It should be microscopically distinguished from gastrointestinal stromal tumor (GIST), leiomyoma, schwannoma, and metastatic malign melanoma. IFP is the lesions having morphology similar to inflammatory myofibroblastic tumor (IMT) and GIST, and the immunohistochemical examination is required for the diagnosis. IFP is positive for CD34 and vimentin, whereas IMT is negative for CD34 and GIST is positive for CD34 and CD117 (C-KIT). In the present case, the vimentin and CD34 were found to be positive and CD117 (C-KT) to be negative. This immune profile was interpreted in favor of IFP.

In adult population, invagination generally requires the surgical treatment. Resection and primary anastomosis are recommended. In invaginations developing based on the mass, there is no accepted approach to the surgical limits in the mass-caused invaginations. In the present case, the patient was undergone right hemicolectomy due to the suspicion of malignity in the mass. In conclusion, the invagination is a cause of ileus that is rarely seen among adults and difficult-to-diagnose. The suitable treatment method is the surgical resection depending on the mass.

**Riassunto**

L’invaginazione è costituita da una dislocazione di un segmento prossimale dell’intestino nella porzione distale contigua. Si tratta di una evenienza rara nella popolazione adulta, e nel caso si verifica per la presenza di una neoplasia della parete intestinale del tratto invaginato. Soltanto il 3-6% dei tumori gastro-intestinali sono localizzati nell’intestino tenue, ed anche il polipo inflammatore fibroso benigno è di rara incidenza.

Viene presentato il caso di una donna di 47 anni affetta da invaginazione ileo-ciecale determinata appunto dalla presenza di un polipo fibroso inflammatario dell’ileo.

**References**