Small bowel diverticulum complicated by enterocutaneous fistula and abdominal wall abscess

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Abstract

Introduction: Small bowel diverticular disease is uncommon, representing 1-2% of general population. The ileum diverticulum is very rare, especially if it is complicated by enterocutaneous fistula with abdominal wall abscess.

Presentation of the case: We report a case of small bowel diverticulum, ileum diverticulum with enterocutaneous fistula in a 65-year man presented with lower abdominal pain and an irreducible swelling for 5 days.

Discussion: Diverticulosis has no exact cause, but theoretically the diverticula result from peristalsis abnormalities, intestinal dyskinesia, or high segmental intraluminal pressures. Complications of small intestinal diverticulum include bleeding, diverticulitis and its complication (e.g. fistula formation, and intra-abdominal abscess).

Conclusion: This is considered rare and serious complication of small bowel diverticulum could occur if diagnosis is delayed.

Introduction

Diverticula are sac-like protrusions of the bowel wall and occur throughout the small and large bowel, usually from mesenteric border. A small bowel diverticular disease is uncommon; the incidence is 1-2% of general population [1]. The duodenal diverticula are more common than jejunoileal diverticula [2]. Jejunal diverticula are more common and are larger than those in the ileum, most of them are acquired diverticula occurring in patients at 6th and 7th decade of life. Majority of them are asymptomatic, may be diagnosed during diagnostic imaging for unrelated symptoms or intraoperative [3], however they may present with their signs and symptoms of complications. In our case, we reported one of very rare complication of small bowel diverticulum was ileal diverticulum with enterocutaneous fistula and abdominal wall abscess, associated with intestinal wall serosal lipoma and enterolith. We reported the case to discuss the etiopathogenesis and proper management of such cases.

Case Report

A 65-year-old Russian male, not known to have chronic medical illnesses, came to the ED complaining of painful swelling in the lower abdomen which had been going on for five days. Abdominal pain was severe colicky in nature with no relieving factors, associated with nausea and vomiting multiple times. There had been no change in bowel habits, fever or change in appetite. The patient had a history of lower abdominal surgery at the age of two, but he had no medical report.

On physical examination the patient was conscious and had a normal body built. His blood pressure was 126/92, pulse was 88 and temperature was 36.2°C is symmetrically distended with a swelling in the lower abdomen 12 × 15 cm in size with negative cough impulse, erythema and tenderness on the overlying skin. The rest of the abdomen was soft on palpation with positive bowel sounds. Investigation of his hemoglobin gave 10.8 wbc’s with 11.5 sodium 139 potassium 3.2 creatinine 0.7.

The patient was admitted as a case of abdominal pain for investigation. The CT of abdomen and pelvic with IV and oral contrast was done showing thickened terminal ileum with marked luminal narrowing which appeared adherent to the urinary bladder wall with no line of cleavage. Two fistula tracts were seen superior and inferior; the superior one lead to a pocket of collection filled by contrast 36 × 20mm in size. The inferior tract was connected to an anterior abdominal wall collection measuring about 18.7 × 14.4mm with marginal enhancement denoting an abscess. There was diffuse anterior abdominal wall fat stranded with subcutaneous pockets of air denoting infection. Subcentemetric mesenteric lymphadenopathy was observed (Figures 1 and 2).

Patient was taken to the OR for exploratory laparotomy and drainage of the abscess. Upon entrance to the abdomen a large pocket of pus in subcutaneous layer was opened and evacuated and a swab was sent for culture and sensitivity. A firm mass inclosing the pelvic was dissected and found to be a large diverticulum 10 cm from the iliocecal junction. The mass was attaching to the urinary bladder and was fistulating to the subcutaneous pus collection. Urologist was called in at this point and the urinary bladder was checked by injecting methylene blue dye; there was no leak. Limited right hemicolectomy was performed with a primary ileocolic anastomosis (Figures 3 and 4).

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Figure 1. A and B, Abdomen CT with IV and oral contrast showed thickening of terminal ileum.

Figure 2. Abdominal CT with IV and oral contrast showed air pocket in bladder and ileo-cutaneous fistula.

Figure 3. Resected bowel of 62-year-old male patient diagnosed with ileum diverticulum associated with abdominal wall abscess.
Histopathology was consistent with diverticulum of the small bowel and serosa lipoma with a pocket containing multiple staghorn-type black stones (Figure 5). Patient wound culture from OR showed E. coli which was sensitive to Tigacyclin. Treatment was started with this antibiotic and patient’s condition improved. Postoperative course was uneventful except for a small dehiscence at the lower part of the abdominal wound, which was treated conservatively with VAC dressing. Patient was discharged to travel to his country, and the wound was left for secondary closing.

Discussion

Small intestinal diverticular disease is defined as outpouching of intestinal wall [4], usually from the mesenteric border [2]. It is uncommon; the incidence is 1-2% of general population [1]. Jejunoileal diverticula are less common than duodenal diverticula [2], occurring in 0.3-1.3% of all cases of small bowel diverticular disease [1].

Diverticulosis has no exact cause, but theoretically the diverticula result from peristalsis abnormalities (e.g., intestinal spasms), intestinal dyskinesia, or high segmental intraluminal pressures. There are two types: A true type that is formed of all intestinal layer, and a false type, formed of just mucosa and submucosa [4]. Majority of them are asymptomatic, may be diagnosed during diagnostic imaging for unrelated symptoms or intraoperative [3] procedures; however, they may present signs and symptoms of complications. Complications of small intestinal diverticulum include bleeding, diverticulitis and its complication (e.g., obstruction, perforation, peritonitis, fistula formation, and intra-abdominal abscess), or segmental diverticulitis (e.g., inflammation in segments of the mucosal segments of colon in between diverticula) [4]. In the case reported here, the diverticulum was on the anti-mesenteric border instead of mesenteric border. In addition, there was the possibility of diverticula adhering to the abdominal wall due to the previous abdominal surgery, which can explain enterocutaneous fistula formation leading to an abdominal wall abscess rather than an intra-abdominal formed abscess.

In the literature, abscess was drained percutaneous in Alvarez’s patient; however, resection and anastomosis of affected segment was the treatment for two Japanese patients (Eriguchi’s and Fujisawa’s) [5]. This was the treatment in present case and case of Meckel’s diverticulum reported by Oguzhan [3] which was also complicated by abdominal abscess. All of them had uneventful postoperative course.

An enterolith formation is not usually formed in a normal anatomy of bowel; however, it can develop in the setting of intestinal stasis in the presence of intestinal diverticula, surgical enteroanastomoses, blind pouches, afferent loops, incarcerated hernias, small intestinal tumours, intestinal kinking from intra-abdominal adhesions, and stenosis or stricture of Crohn’s disease and intestinal tuberculosis [6]. In our case there were two predisposing factors: history of enterostomy and present ileal diverticulum. Therefore, it is suspected to be a primary result of intestinal stasis. Its prevalence ranges from 0.3-10% in selected populations [6]. Perforation is one of the rare complications of the enteralith. In our case we suppose the enterolith made compressions on the ileum diverticulum wall which was adherent to abdominal wall as a sequel of previous surgery or diverticulitis.

Conclusion

We should consider the bowel communication in abdominal wall abscess cases and use imaging modalities, like computed tomography, to optimize the pathway of the management and decrease morbidity rate.

Consent

Written informed consent was obtained from the patient for publication of the features and accompanying images.

Conflict of interest and funding

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References

1. Koh FJ, Khor JL, Chew MH (2017) An Unusual Case of Giant Ileal diverticulum–A Case Report. Int J Surg Case Rep 41: 277–279. [Crossref]
2. Hubbard TJ, Balasubramanian R, Smith JJ (2015) Jejunal Diverticulum Enterolith Causing Perforation and Upper Abdominal Peritonitis. BMJ Case Rep 2015: bcr2015210095. [Crossref]
3. Karatepe O, Adas G, Altiok M, Ozcan D, Kamali S, et al. (2009) Meckel’s Diverticulum Manifested by a Subcutaneous Abscess. World J Gastroenterol 15: 6123–6125. [Crossref]

4. Foris LA, Bhimji SS (2018) Diverticulosis. StatPearls. U.S. National Library of Medicine. [Crossref]

5. Eriguchi N, Aoyagi S, Nakayama T, Emi Y, Saka M, et al. (1988) Ileo-Abdominal wall fistula caused by diverticulum of the ileum. J Gastroenterol 33: 272-275. [Crossref]

6. Gervits GE, Lan G (2014) Enterolithiasis. World J Gastroenterol 20: 17819–17829. [Crossref]