Enterovesical Fistula From Meckel Diverticulum

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Case Report

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INTRODUCTION

An enterovesical fistula (EVF) is an abnormal communication between the intestine and urinary bladder. An EVF can be caused by an inflammatory process, such as Crohn disease or diverticulitis; a malignancy; or surgical trauma. An EVF is a rare complication of Meckel diverticulum, with several reports in the literature [1-7], but only a few cases in Asia [2, 4]. We report this very rare complication of Meckel diverticulum in Korea.

CASE REPORT

A 51-year-old male was referred to our hospital with a 10-year history of pyuria and increased urinary frequency. Although he had been treated with antibiotics for a chronic urinary tract infection, pyuria persisted without fever or abdominal pain. The patient had hypertension but no significant past surgical history or family history of inflammatory disease.

The prostate was soft and smooth without tenderness. Initial routine blood tests showed no leukocytosis and a normal C-reactive protein level. The urine sediment revealed large numbers of white blood cells and many bacteria. Urine nitrite test was positive, and culture showed growth of Escherichia coli sensitive to a third-generation cephalosporin. Simple abdominal X-ray showed no significant abnormality in the bowel gas pattern or urinary tract. Computed tomography (CT) showed mild, diffuse wall thickening in the urinary bladder with intravesical air density (Fig. 1A). Although adjacent small bowel was observed on the dome of the urinary bladder on coronal CT (Fig. 1B), an EVF could not be identified. Cystoscopy demonstrated a bullous, mass-like lesion at the dome of the urinary bladder, with a hole suggesting a fistula (Fig. 2). Transurethral biopsy was consistent with a vesicointestinal fistula (Fig. 3).

Laparoscopic exploration was performed, and intraoperative findings demonstrated a Meckel diverticulum (8.5 × 2.4 cm) attached to the dome of the urinary bladder and omental adhesions. Segmental resection of the small bowel including the diverticulum and primary repair of the urinary bladder along with partial cystectomy were performed. The postoperative clinical course was uneventful. An enterovesical fistula is a very rare complication resulting from chronic inflammation of a Meckel diverticulum.

Keywords: Enterovesical fistula; Meckel diverticulum; Complication

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Meckel diverticulum was performed with partial cystectomy. Macroscopic findings also indicated that the tip of a Meckel diverticulum was connected to the urinary bladder (Fig. 4). The final histologic findings demonstrated a gastric-type Meckel diverticulum with a vesicointestinal fistula.

The postoperative course was uneventful, and the patient was discharged on postoperative day nine without any significant complications. The Institutional Review Board of St. Vincent Hospital approved this case report and waived the informed consent requirement (No. VC18ZOSI-0252).

DISCUSSION

A Meckel diverticulum, a remnant of the omphalomesenteric duct that connects the midgut of the fetus to the yolk sac, is one of the most common congenital malformations of the gastrointestinal tract. It is reportedly present in 2%–4% of the population [8], with symptoms appearing in only 4%–6% of cases [9]. Gastrointestinal bleeding is the most common presentation of Meckel diverticulum and usually occurs in children, especially those younger than 2 years [8]. In adults, intestinal obstruction due to intussusception or an adhesive band is the most frequent complication of Meckel diverticulum, accounting for 14%–53% of complicated cases, followed by diverticulitis, enterolith formation, and perforation [8, 10]. A neoplasm, such as a carcinoid, sarcoma, or adenocarcinoma, accounts for up to 3% of complicated cases [11].

There are few reports of an EVF associated with Meckel diverticulum. A PubMed and Embase search for articles published between 1990 and 2018 using terms of “enterovesical fistula,” “vesicoenteric fistula,” and “Meckel’s diverticulum” identified seven cases of fistula in the English literature [1-6, 12]. The causes were no coexisting bowel or bladder disease (3 cases), adenocarcinoma in ectopic pancreatic tissue (1 case), Crohn disease (1 case), enterolith (1 case), and ingested foreign body in Meckel diverticulum.
lum (1 case) (Table 1).

Preoperative diagnosis of Meckel diverticulum is difficult. One study reported that, of 776 patients diagnosed with Meckel diverticulum, only 11% had symptoms other than bleeding [13]. A preoperative diagnosis of EVF is also challenging. Magnetic resonance imaging and CT have contributed to improved diagnosis of EVF. These modalities are highly effective in determining bladder wall thickness, presence of gas in the bladder, and continuity of a bowel segment adjacent to the bladder. Cystoscopy may also help diagnose EVF through direct visualization of a fistula in 6.7%–67% of cases [7]. This wide variation in detection rate with cystoscopy may be associated with the presence of mucosal edema and hyperplasia, which can cause difficulty in identifying a small fistula orifice [1]. Before visiting our hospital, this patient had undergone cystoscopy at two other hospitals, but no EVF was detected.

Herein, we report the first case of EVF involving a gastric-type Meckel diverticulum in the Asia-Pacific region. Although such a case is very rare and is difficult to diagnose preoperatively, surgeons should be aware of this possibility to avoid a missed or delayed diagnosis.

**CONFLICT OF INTEREST**

No potential conflict of interest related to this article was reported.

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**Table 1. Published case reports of enterovesical fistula associated with Meckel diverticulum**

| Study            | Publication year (country) | Sex | Age (yr) | Chief complaint               | Type            | Preoperative confirmation of diagnosis |
|------------------|---------------------------|-----|----------|-------------------------------|----------------|----------------------------------------|
| Petros et al. [12] | 1990 (United States)      | Male | 22       | Fever, dysuria                | Gastric         | Confirmed with cystography              |
| Hudson et al. [5]  | 1992 (United States)      | Male | 23       | Abdominal pain, diarrhea      | No ectopic mucosa | Suspected on CT                         |
| Grazioti et al. [3] | 2002 (Italy)             | Male | 40       | Recurrent dysuria, fever      | -               | Confirmed with cystoscopic biopsy       |
| Boussida et al. [1] | 2013 (Tunisia)         | Female | 66      | Fever, dysuria                | No ectopic mucosa | Suspected on cystoscopy                  |
| Bourguiba et al. [6] | 2017 (Tunisia)         | Male | 35       | Fever, abdominal pain with distension | -               | No laparotomy                           |
| Fujita et al. [2]  | 2018 (Japan)             | Male | 58       | Dysuria, frequency            | Pancreatic      | Confirmed with cystoscopic biopsy       |
| Hakoda et al. [4]  | 2018 (Japan)             | Male | 46       | Hematuria                      | No ectopic mucosa | Suspected on cystoscopy, CT, MRI         |

CT, computed tomography; MRI, magnetic resonance imaging.