Perforated Meckel’s Diverticulum in a 3-day-old Neonate; A Case Report

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INTRODUCTION

Meckel’s diverticulum (MD) is the most common congenital anomaly of the gastrointestinal tract.1 Symptomatic MD in the neonatal period is quite rare. Complication rate is about 4%.2 Intestinal perforation is a less common complication of MD in children that occurs in 10% of patients.3 It rarely occurs in neonatal period and only few cases have been reported until now in the literature.4

CASE REPORT

A 3-day-old male neonate was referred to our hospital because of repeated vomiting after breastfeeding. Vomiting was bilious and was associated with abdominal distention.

The neonate’s birth weight was 3200 gr and he was born from a 28-year-old mother gravid 1 with uncomplicated pregnancy course and normal vaginal delivery with APGAR score 9 at min 1, and 10 at min 5.

On admission, careful physical exam was done, which revealed tachycardia, hypotension, and severe abdominal distention, while the neonate was lethargic. Laboratory assessment was done in the first day of admission. White blood cell count was 8600. Hemoglobin was 17.1 g/dL. Platelet count was 237000 (table 1). Blood urea nitrogen was 21 mg/dL. Due to bilious vomiting, thoraco-abdominal radiography was done (figure 1).

After physical examination and radiological evaluation the patient underwent exploratory laparotomy with the impression of bowel perforation. Free air was observed in abdomen. Bowel content was discovered
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in abdomen too and perforated MD was diagnosed. Resection and irrigation with 1 liter normal saline was done.

Pathological report of tissue specimen was inflamed MD with heterotopic gastric mucosa. Five days after operation, nasogastric tube was discontinued and oral feeding with formula was started without any complication. Two days later (7 days of hospital admission) the neonate left the hospital with good condition.

DISCUSSION:

MD is a 3-6 cm outpouching from the antimesenteric border of the ileum at 50-75 cm from the ileocecal valve. Failure of involution of the omphalomesenteric duct during the 5th and 7th week of gestation, results in MD. MD contains four layers of intestine and may have different ectopic tissues such as gastric, pancreatic, colonic, duodenal, or endometrial in about 30% to 50% of patients. The omphalomesenteric artery that arises from an ileal branch of the superior mesenteric artery provides the blood supply of diverticulum.

Although MD is the most common congenital anomaly of the gastrointestinal tract, its symptomatic manifestation in the neonatal period is rare. Common presentations of neonatal Meckel’s diverticulum that have been reported in the literature include perforation, intussusception, segmental ileal dilatation, and ileal volvulus. Perforation was one of our findings in the current report.

Most of the symptomatic MD occurs by the age of 3 years. It is more common in boys (male to female ratio 2:1). In our report, a male neonate was affected.

Heterotopic gastric mucosa was reported in our patient. In other reports, heterotopic tissue was not found in some cases.
Pneumoperitoneum was reported in the literature\(^{10,11}\). Pneumoperitoneum was seen in our case.

Bowel obstruction is the most common presenting feature in neonates\(^ {12}\). In our report, bowel obstruction with bilious vomiting was the first presentation. In the study by Bertozzi and colleagues\(^ {13}\), bowel obstruction (58.3%) and pneumoperitoneum (33.3%) were the most common clinical manifestations of symptomatic MD among neonates.

Mild hypochloremic alkalosis was reported by Bertozzi and co-workers\(^ {13}\). In our case, mild alkalosis was found in the blood gas analysis. Incidence of perforated MD in neonates is very rare and this case is the first report from our country.

It should be considered that other causes of intestinal perforation in neonatal period include necrotizing enterocolitis (NEC), Hirschsprung’s disease, meconium ileus in neonates with cystic fibrosis, intestinal atresia, and intestinal volvulus\(^ {3}\).

MD perforation can be occurred due to either diverticulitis or heterotopic mucosa within the diverticulum. This has been proven in the last 25-year overview of MD perforation, which shows that about 75% of cases have acute inflammation and/or heterotopic mucosa in their specimens. Other than the present patient only in three other patients, pancreatic heterotopic mucosa was identified, but it may occur spontaneously. In our report, inflamed MD with heterotopic gastric mucosa was found.

Other predisposing factors for MD perforation are: corticosteroid therapy in antenatal and postnatal period, maternal use of corticosteroids or cocaine, perinatal asphyxia or hypoxia, exchange transfusion, trauma due to feeding by nasogastric tube, congenital absence of muscle in gastrointestinal wall, and decreased intrauterine blood flow\(^ {14}\). None of these factors were detected in our patient.

There are only two mortality reports from perforated MD. The first one happened in 1927 when a 4-day-old neonate with abdominal distention and shock died and during autopsy evaluation perforated MD was diagnosed\(^ {10}\). The second case was a neonate with multiple congenital anomalies (esophageal atresia, tracheoesophageal fistula, and imperforate anus who died due to cardiorespiratory failure)\(^ {11}\).

Perforated MD in neonates can mimic several diseases such as NEC, perforated appendicitis, and solitary ileal perforation. Treatment is different, but confirming a preoperative diagnosis of MD in cases with signs of perforation is not necessary because prompt surgical intervention is mandatory if an intra-abdominal pathology is suspected. If diagnosis and management is done at proper time, good prognosis with about 100% survival will be expected for neonates with isolated perforated MD.

**CONFLICT OF INTEREST**

The authors declare no conflict of interest related to this work.
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