Perforation of Meckel’s diverticulum in a very low birth weight neonate with severe pneumoperitoneum and review of literature

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Perforation of Meckel’s diverticulum (MD), but it could be severe, even life-threatening for pediatric patients. There is a paucity of data on etiology of perforation, as well as clinical manifestations, management and prognosis in very low birth weight (VLBW) neonates with perforated MD. We report a rare case of spontaneously perforated MD in a VLBW neonate presenting with severe pneumoperitoneum. To our knowledge, this is one of the earliest reported VLBW cases with this rare complication. Furthermore, we review relevant publications and summarize major characteristics of all VLBW cases previously reported in order to provide some practical experience and interesting issues for pediatricians. Perforated MD should be kept in mind when VLBW neonates present with pneumoperitoneum.

Key words: Meckel’s diverticulum; intestinal perforation; very low birth weight; neonates; pneumoperitoneum.
improved after continuous positive airway pressure (CPAP) and surfactant treatment.

On the 24th hour of life, the bedside X-ray, which was initially aimed to confirm the position of venous catheters, revealed signs of free air in the peritoneum (Fig.1). Although distension of his abdomen was not remarkable, intestinal perforation was suspected. However, his parents refused surgical intervention. One hour later, his abdominal distension progressed, and plain radiograph of his abdomen demonstrated significant pneumoperitoneum with bowels gathering together (Fig. 2). On the 26th hour, the baby was taken to the operating room for emergent laparotomy immediately after his parents gave their written consent. When general anesthesia was utilized, he already had severe abdominal distension.

Upon entering the peritoneal cavity, a lot of gas came out. There were no signs of peritonitis, and we didn’t observe intraperitoneal haemorrhage or exudate accumulation. After careful examination of the entire bowel, a perforated MD was discovered, measuring up to 1.5 cm within about 20 cm of the ileocecal valve (Fig. 3). The diverticulum and

Fig. 1. Plain radiograph showing free gas on the 24th hour after birth.

Fig. 2. Severe pneumoperitoneum accompanied with bowels gathering together was revealed by plain radiograph.

Fig. 3. Meckel’s diverticulum with perforation at its tip.
its surrounding bowel were macroscopically normal without involvement of inflammation. We didn’t find any other suspected perforation area or other macroscopic pathology such as necrotising enterocolitis (NEC). Wedge bowel resection with transverse suturing was performed. Pathological examination confirmed the specimen was an MD with perforation, slightly focal infiltration of reactive inflammatory cells and no evidence of ectopic mucosa.

The patient recovered well from the surgery during the first 3 weeks. Recurrent abdominal distension and vomiting occurred approximately 2 weeks after enteral feeding was introduced. X-ray showed intestinal dilatation without signs of pneumoperitoneum. His symptoms improved quickly and significantly when enteral feeding was discontinued. The baby was discharged in stable condition well tolerating oral diet at the age of 57 days. Further follow-up is proceeding.

Discussion

Meckel’s diverticulum could be found incidentally or when causing various complications. According to Cullen et al, the lifetime risk of complications was estimated to be approximately 6.4%. Although patients born with MD could develop complications at any age, the dominate types differ between children and adults. Pediatric patients tend to present with painless intestinal hemorrhage, whereas diverticulitis and intestinal obstruction are more common in adult patients. Perforation is a rare complication of MD, evident in approximately 3-10% of patients who have symptoms. To our knowledge, only four isolated cases of spontaneous MD perforation in VLBW neonates have been described in the literature, information is summarized in Table I and II.

The mechanisms underlying the development of perforation of MD in VLBW neonates have not been fully elucidated. In adults and older children, perforation of MD is often secondary to mucosal ulceration caused by acid-producing ectopic tissue within the wall of MD. As observed in acute appendicitis, long, narrow-based diverticulum with poor self-emptying are more prone to luminal obstruction leading to subsequently inflammation, necrosis, and even perforation. According to previous studies, bowel obstruction was the most common presentation in term infants with perforated MD, which suggested that rupture of the thin diverticulum due to high pressure within lumen may play a vital role in the development of perforated MD in neonatal period. Of note, ectopic tissue appeared not to be the major cause of perforation of MD in VLBW neonates as displayed in Table I. Hypoxia, inadequate blood supply, permeability of the intestine, and early use of postnatal steroid and indomethacin are thought to increase the risk of spontaneous intestinal perforation (SIP), but the potential roles have not been

<table>
<thead>
<tr>
<th>Reference</th>
<th>Sex</th>
<th>Gestation</th>
<th>Birth weight</th>
<th>Presentation</th>
<th>Age</th>
<th>Presence of ectopic tissue</th>
<th>Pathology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aguayo et al⁹</td>
<td>Male</td>
<td>28 weeks</td>
<td>798g</td>
<td>abdominal distension, 6 days</td>
<td>6 days</td>
<td>No</td>
<td>perforation</td>
</tr>
<tr>
<td>Smolkin et al¹¹</td>
<td>Male</td>
<td>28 weeks</td>
<td>1200g</td>
<td>abdominal distension, 5 days</td>
<td>5 days</td>
<td>No</td>
<td>perforation, submucosal hemorrhages</td>
</tr>
<tr>
<td>Khan et al¹⁰</td>
<td>Male</td>
<td>29 weeks</td>
<td>650 g</td>
<td>abdominal distension, 6 days</td>
<td>6 days</td>
<td>NA</td>
<td>perforation, inflammation</td>
</tr>
<tr>
<td>Borgi et al⁷</td>
<td>Male</td>
<td>29 weeks</td>
<td>1400g</td>
<td>abdominal distension, 3 hours</td>
<td>3 hours</td>
<td>No</td>
<td>perforation, discrete inflammation</td>
</tr>
</tbody>
</table>

NA Not available
determined. It is yet to be investigated whether these factors predisposed MD to perforation in premature neonates. MD and the surrounding intestinal wall were involved with extensive inflammation in one of the four previous cases. It is unclear why the baby presented here developed a perforated MD. Both ectopic mucosa and gangrenous inflammation were absent pathologically. We infer that the perforation of MD may be mainly attributed to intestinal immaturity in this case.

Symptomatic MD has various clinical manifestations and is easily misdiagnosed preoperatively. Perforated MD often mimic acute complicated appendicitis with a sudden onset of peri-umbilical or right lower quadrant pain. Conventional radiographs have limited value in detecting perforated MD, and clinicians often don’t have sufficient time to approach various diagnostic measures under the condition of acute abdomen. Although pneumoperitoneum is a vital radiological sign of the presence of intestinal perforation, it is often unremarkable when perforated MD occurs in adults and children. Conversely, pneumoperitoneum appears to be the most common symptom in VLBW neonates with perforated MD. We speculate that it may be associated with the use of noninvasive mechanical ventilation and fast conditions of these babies.

In neonates with intestinal perforation and abdominal distension, NEC is an important disease for differential diagnosis due to its similar clinical manifestations with SIP and other gastrointestinal disorders. NEC rarely occurs during the first week after birth. In comparison, SIP mainly affects infants of lower birth weight, more immaturity and at an earlier postnatal age. The median age of all the five VLBW cases with perforated MD at symptoms onset were 5 days (range: 3 hours to 6 days). Of note, intraoperative findings of the case we reported suggested that his severe abdominal distension appeared to be associated with the accumulation of free gas. Limited content of his peritoneal cavity could be a possible explanation, but mechanical ventilation may also partly contribute to the rapid progression of pneumoperitoneum. If the hypothesis did exist, it needs to be emphasized for clinicians to prohibit general condition of these patients form worsening and strive time for possibly delayed surgical intervention.

It is generally recommended that symptomatic MD should be removed and in some cases surgical intervention is urgent. Histological state of MD and the surrounding bowel and morphological characteristics are proposed to be taken into consideration when surgeons chose a surgical method. Complete resection of ectopic tissue is a vital step of the surgical procedure to minimize the risk of recurrent symptoms. Previous studies have provided evidence that the external appearance could indicate the distribution of ectopic mucosa within the wall of MD. Ectopic tissue were found to have a wide distribution

<table>
<thead>
<tr>
<th>Reference</th>
<th>Surgical procedure</th>
<th>Clinical Outcome</th>
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<tbody>
<tr>
<td>Aguayo et al</td>
<td>bowel segment resection and ileostomy; reanastomosis was performed two months later</td>
<td>had gastrointestinal dysfunction related to prematurity and discharged in stable condition</td>
</tr>
<tr>
<td>Smolkin et al</td>
<td>bowel segment resection with end-to-end anastomosis</td>
<td>NA</td>
</tr>
<tr>
<td>Khan et al</td>
<td>bowel segment resection with end-to-end anastomosis was performed when abdominal distention occurred again after pulling out of draining tube</td>
<td>recovered well from surgery, but demised following a suspected aspiration episode</td>
</tr>
<tr>
<td>Borgi et al</td>
<td>bowel segment resection with end-to-end anastomosis</td>
<td>recovered well and discharged on day 16 of life</td>
</tr>
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NA Not available
from tip to base in short MD (HDR less than 1.6 or 2.0), and wedge resection of MD or segmental resection of the ileum containing the diverticulum was safer to avoid ectopic tissue being left. Transverse suturing is recommended to decrease the risk of narrowing the ileal lumen after wedge bowel resection. We performed wedge resection with transverse suturing for the baby presented here, considering the normal appearance of the surrounding bowel without signs of inflammatory involvement and shortening operative time.

The overall prognosis of SIP is better than perforation secondary to NEC, but few studies focus on further intestinal problems after primary SIP. Drewett et al. found that neonates with SIP are at significant risk of further intestinal complications such as new spontaneous perforation and NEC. Indeed, these issues could occur as long as 3 months after the initial perforation and may require surgery. One death after suspected aspiration was reported in the four previous VLBW cases. The patient we reported was discharged in stable condition well tolerating oral diet. It is unclear whether his recurrent abdominal distension during hospitalization was attributed to his prematurity or surgery. Regular follow-up is necessary.

Perforated MD in VLBW neonates is very rare, but it should be kept in mind when these patients present with pneumoperitoneum in the neonatal period. The underlying mechanisms have not been fully elucidated and ectopic mucosa appeared to have little association with the development of perforation in VLBW neonates. Intestinal immaturity may be predominate in the case we reported, and the role of mechanical ventilation in the progress of abdominal distension needs to be further investigated. After prompt surgery, these patients could have a good clinical outcome, but long-period follow-up is needed with awareness of possible intestinal issues in their later life.

REFERENCES


