CASE REPORTS

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Chylolymphatic cyst as a rare cause of acute intestinal obstruction in infants



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Abstract

Background Chylolymphatic cysts are benign pathology in lymphatic vessels due to obstruction in the lymphatic system. Their incidence is still unknown. However, they account for approximately 3 to 9% of all pediatric lymphangiomas. Most of these lesions are asymptomatic and are discovered accidentally. They can be presented with nonspecific abdominal pain, vomiting, and changes in bowel habits. Rarely, chylolymphatic cysts can as intestinal obstruction, volvulus or even torsion.

Case presentation A 3-month-old male presented to our unit with manifestations of acute intestinal obstruction. Abdominal examination revealed signs of generalized peritonitis and intestinal obstruction. Abdominal ultrasound revealed minimal turbid intraperitoneal collections and no motility of bowel loops. Plain erect abdominal radio-graph revealed multiple small bowel air fluid levels. Abdominal exploration revealed a large chylolymphatic cyst causing acute intestinal obstruction. Excision of the cysts was done with preservation of adjacent bowel loop. The patient was discharged in good health on the 6th day postoperative. A chylolymphatic cyst with a diameter of 5 cm was identified by histological investigation.

Conclusions Most chylolymphatic cysts in infants are asymptomatic, but they can cause fatal complications such as intestinal obstruction, volvulus, or even torsion.

Keywords Chylolymphatic cysts, Mesenteric cysts, Intestinal obstruction, Infants

Background

Chylolymphatic cysts are benign pathology in lymphatic vessels due to obstruction in the lymphatic system. Their incidence is still unknown [1]. However, they account for approximately 3 to 9% of all pediatric lymphangiomas [2]. Most of these lesions are asymptomatic and are discovered accidentally. They can be presented with nonspecific abdominal pain, vomiting, and changes in bowel habits. Rarely, chylolymphatic cysts can as intestinal obstruction, volvulus, or even torsion [3].

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Case description

A 3-month-old male presented to our unit with his mother complaining of repeated bilious vomiting, abdominal distension, and absolute constipation for 2 days. General examination revealed toxic look of the infant and moderate dehydration. He had a temperature of 38.5 °C, respiratory rate of 34 breaths per minute, and pulse of 160 bpm, and his blood pressure was 70/40 mmHg. Abdominal examination revealed stepladder appearance of bowel loops, diffuse guarding, and empty rectum. The rest of the examination was unremarkable. After resuscitation with intravenous fluids and empirical antibiotics, a 10-French nasogastric tube was inserted. A white blood cell count showed a leukocytosis of 18,000 cells/mm³ with a neutrophilic predominance of 15,500 cells/mm³. We requested urgent abdominal ultrasound, which revealed minimal turbid intraperitoneal collections and no motility of bowel loops. Plain erect



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abdominal radiograph revealed multiple small bowel air fluid levels with no free air under diaphragm (Fig. 1). We prepared the infant for urgent abdominal exploration. Via right transverse supra umbilical incision, we performed formal abdominal exploration that revealed highly turbid fluid collection and a large multilocular chylolymphatic cyst adjacent to a jejunal loop 30 cm far from duodenojejunal junction causing intestinal obstruction (Fig. 2). Excision of the cysts was done with preservation of adjacent bowel loop (Fig. 3). Repair of a small mesenteric defect at the bed of the cyst was done with serosal repair of mesenteric border of adjacent jejunal loop (Fig. 4a, b). Peritoneal lavage was done, and a pelvic drain was inserted. The patient was discharged in good health on the 6th day postoperative. A chylolymphatic cyst with a diameter of 5 cm has been identified by histological investigation.

Discussion

Mesenteric cyst is a rare intra-abdominal tumor with prevalence of about 1:20,000 in pediatric age group. Chylolymphatic cysts are rare variants of mesenteric cysts; they account for 3 to 9.2% of all pediatric lymphangiomas [2].

They are benign cystic pathology lined with a thin endothelium or mesothelium that occurred due to gross proliferation of isolated or ectopic lymphatics in mesentery that lack communication with the rest of the lymphatic system. The accumulation in these channels results from an imbalance between the inflow and outflow of fluid [1]. They can occur in any part of the mesentery of the gastrointestinal tract but most commonly



Fig. 2 Large multilocular chylolymphatic cyst (arrow) adjacent to a jejunal loop 30 cm far from duodenojejunal junction

are localized in the mesentery of the small intestine, large intestine, and retroperitoneum. They can be single or multiple and unilocular or multilocular and can have serous, chylous, hemorrhagic, or mixed fluid [4]. Our case had a single multilocular chylolymphatic cyst in the mesentery of proximal jejunum.

Based on their etiology, Beahrs et al. [2] classified mesenteric cysts into embryonic or developmental, traumatic, or acquired, neoplastic, or non-neoplastic and infective or degenerative. Based on their contents,



Fig. 1 Plain erect radiograph on abdomen showing multiple small intestinal air fluid levels



Fig. 3 Large multilocular chylolymphatic cyst after excision



a: Mesenteric defect (Arrow) after excision of the mesenteric cyst.

b: Repair of small mesenteric defect at the bed of the cyst was done with serosal repair of mesenteric border of adjacent jejunal loop (Arrow).

Fig. 4 a Mesenteric defect (arrow) after excision of the mesenteric cyst. b Repair of small mesenteric defect at the bed of the cyst was done with serosal repair of mesenteric border of adjacent jejunal loop (arrow)

mesenteric cysts can be divided into serous, chylous, hemorrhagic, and chylolymphatic cyst. Regarding pathology, mesenteric cysts are classified into four types including type 1 (pedicled) and type 2 (sessile), which are limited to the mesentery, hence can be excised completely with or without resection of the involved gut. Type 3 and type 4 are multicentric, so they require complex surgery and often sclerotherapy because of their extension into retroperitoneum [3]. This case is type 2 (sessile) chylolymphatic cyst.

Most of mesenteric cysts may present as asymptomatic abdominal mass or incidental finding on imaging or laparotomy for other abdominal conditions. In rare occasions, mesenteric cysts may cause acute abdomen from cyst rupture, infection, hemorrhage, intestinal obstruction, and volvulus [5]. In our case, a large chylolymphatic cyst presented as acute intestinal obstruction.

Preoperative diagnosis of mesenteric cyst in most cases is difficult as they have no definite characteristic features. Abdominal ultrasound which is the imaging modality of choice can reveal mesenteric cysts as cystic tumors [6]. Plain X-ray abdomen may show gasless, homogenous mass displacing bowel loops. Multiple air fluid levels may be seen in erect film [5]. Computed tomography can be also used to confirm the diagnosis [3]. In our case, preoperative plain radiograph revealed multiple air fluid levels due to compression and inflammatory adhesion caused by a large multilocular chylolymphatic cyst. Postoperative diagnosis is confirmed with histopathology, which can differentiate the chylolymphatic cyst from other mesenteric cysts [3]. Surgical removal of the cyst is the standard treatment for chylolymphatic cysts. The cyst is removed along with the adjacent bowel loop. Enucleation and cyst drainage were two more techniques mentioned [7]. We just remove the cyst in our case.

Conclusions

Most chylolymphatic cysts in children are asymptomatic, but they can cause fatal complications as intestinal obstruction, volvulus, or even torsion.

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Authors' contributions

AKA contributed to management and wrote the manuscript. TAS contributed to acquisition of data and revision of manuscript. AKA and TAS were the surgeon who operated this case. AHM was the anesthesiologist. AKA and IMA, critically revised the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials

Data sharing is not applicable to this article.

Declarations

Ethics approval and consent to participate

Ethical approval no. E9854 from Assiut University Ethics Committee was taken. Written informed consent was obtained from the patient for publication of this case report and accompanying images. A copy of the written consent is available for review by the editor in chief of this journal.

Competing interests

The authors declare that they have no competing interests.

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