Epiploic Cystic Lymphangioma Rupture as a Result of Acute Abdomen in a 4-year-old Patient. Case Report

Department General Surgery, Autonomous University of Nuevo León, “Dr. José E. González” University Hospital, Nuevo León, Mexico

Abstract

Introduction: Cystic lymphangiomas are rare, benign, and congenital lesions which result from malformations of the lymphatic system. These malformations occur more frequently in pediatric patients, with diverse clinical symptoms. Imaging studies such as ultrasounds, computed tomography scans, and magnetic resonance imaging (MRI) have proven to be of great use in their diagnosis, and with the surgical removal of the lesion being the treatment of choice. Clinical case: A 4-year-old male with a history of hiporexia and early satiety since he was 2 years old. Furthermore, 72 h before consultation had suffered blunt abdominal trauma, presented tegument paleness and abdominal pain as a result of this trauma. The pain decreased after analgesics. Afterward, there was abdominal distention. We were able to observe a cystic lesion with data gained through an abdominal MRI along with free fluid and active bleeding. During the surgical procedure, a cystic lesion-dependent of the greater omentum is found and verified with a histopathological report. The post-operative evolution of the patient was positive. Conclusion: Abdominal cystic lymphangiomas are rare; the definitive treatment is the removal of the lesion, either with minimally invasive surgery, by laparoscopy or open technique. We ought to have clinical suspicion to offer timely treatment, with the support of imaging studies since clinical presentation varies within the pathology.

Key words: Cystic lymphangioma. Greater omentum. Rupture.

Introduction

Abdominal cystic lymphangiomas are rare, benign, and congenital tumors which result from malformations of the lymphatic system usually located in the mesentery, followed by the omentum, mesocolon, and retroperitoneum. An epiploic cystic lymphangioma can occur in the greater or lesser omentum and occurs more frequently in pediatric patients (under 10 years of age). Reports show that 60% of these cases occur in patients under 5 years of age.

Its clinical presentation can be very diverse, from a palpable symptomatic/asymptomatic mass to an acute abdominal mass, which can be life-threatening. Imaging studies are fundamental for its diagnosis since its clinical picture is unspecified. Among the radiological techniques are ultrasounds, CT scans, and/or magnetic resonance imaging (MRIs). The treatment of choice is complete surgical removal.

Case report

The patient, a 4-year-old male without a significant background (with a history of hiporexia and early satiety since he was 2 years old) began his condition 3 days before his admittance, having suffered blunt abdominal

Correspondence:
Roberto Pineda-Quinonez
E-mail: droberto@live.com
Available online: 01-10-2018
Medicina Universitaria, 2018;20(2):88-90
www.medicinauniversitaria.org
1665-5796© 2018 Universidad Autónoma de Nuevo León. Published by Permanyer México SA de CV. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).
trauma (a kick). After the first 24 h, he presents as a result of this trauma, tegument paleness, and 48 h later, moderate abdominal pain with a predominance in the right lower quadrant, which improved with analgesics. On the 3rd day after the trauma, the patient presented abdominal distention with increasing intensity of pain, for this reason, they go to the emergency room at our hospital, with generalized abdominal pain and tachycardia. A thoracoabdominal X-ray is taken showing displacement of bowel loops as well as radiopacity in the anterior compartment. In his pre-operative studies, the patient presents hemoglobin at 7.29 g/dL and hematocrit at 21.5%. An abdominal ultrasound was conducted, where we were able to see free fluid in the splenorenal fossa, the right parietocolic gutter, and both iliac fossae and the pelvic cavity (Fig. 1). Further studies were conducted with an abdominal MRI, showing a septated multicystic image and hypointense material in the T2 sequence, suggesting a hemorrhage (Figs. 2 and 3). Reanimation of the patient is performed using intravenous fluids. During the surgical approach, 400 cc of hemoperitoneum are found, in addition to an epiploic cyst of approximately 20 cm × 15 cm × 8 cm, ruptured with hematic content inside (500 cc), dependent of the greater omentum (Fig. 4). Cyst removal and omentectomy are performed. An erythrocyte concentrate is transfused in the transoperative. The patient presents a satisfactory evolution and is discharged on day 3 of his post-operative stay. In his histopathological report, we observed cystic lymphangioma, edema, and a recent hemorrhage (Fig. 5).

Discussion

Cystic lymphangioma is a rare benign tumoral pathology. It may be single, multiple, unilocular, or multilocular, from a few centimeters to 40 cm in diameter. In general, located in the subcutaneous tissue in the head, neck, or armpits and can, rarely, be intrabdominal (2-5%). It is more common in male pediatric patients, with a reported incidence of approximately 1 in 140,000 patients. Its pathophysiology originates from a malformation in lymph vessels, characterized by dilated cysts of lymphatic content, pus or blood as a result of bleeding. They generally occur in a multiloculated and multiseptated form. Its clinical presentation is very variable, from asymptomatic lesions all the way up to intestinal occlusions, which may occur suddenly.
Despite the advances in diagnostic techniques, pre-operative diagnosis is a challenge due to its presentation and form, and can sometimes be incidentally discovered when performing a laparotomy procedure or during an autopsy. Ultrasound, tomography, and MRIs are currently valuable tools in the diagnosis of this pathology. There is rarely a recurrence of the tumor following complete surgical removal. A follow-up of the lesion is recommended in the post-operative with an imaging study, an ultrasound being the golden choice. Within its differential diagnosis of an abdominal cystic mass, there are - cystic teratoma, mucinous cystadenoma, ovarian cysts, bile duct cysts, pancreatic pseudocysts, complicated ascites, renal cysts, and splenic cysts, among others.

Definitive treatment of abdominal cystic lymphangioma involves complete resection. Surgical removal can be conducted using laparoscopic techniques if possible, and lower recurrence rates have been reported in complete removal versus partial removal, aspiration, or sclerosing therapy.

The case we are presenting is interesting since the patient arrived at the emergency room for an assessment with acute abdominal pain (with a background of early satiety and hiporexia. This may very well have been caused by the growth of the lesion) accompanied by signs and symptoms of hypovolemia after the trauma, which lead us to conduct imaging studies to rule out differential diagnoses, and thus obtaining a rare diagnosis.

Conclusion

Abdominal cystic lymphangioma is rare, definitive treatment is lesion removal, with a rare rate of recurrence. The treatment, while depending on the state of the patient, ought to be in an early manner. The complicated aspect of these cases is the timely diagnosis due to the varied symptomatology.

Ethical disclosures

Protection of human and animal subjects. The authors declare that no experiments were performed on humans or animals for this study.

Confidentiality of data. The authors declare that they have followed the protocols of their work center on the publication of patient data.

Right to privacy and informed consent. The authors declare that no patient data appear in this article.

References


Figure 4. Lymphangioma adhered to omentum before resection

Figure 5. (A-D) Large lymphatic channels containing eosinophilic proteinaceous material are shown in a loose stroma of fibroconnective tissue. Disorganization of the muscular wall can be seen in some foci