Huge Mesenteric Lymphangioma – A Rare Cause of Acute Abdomen

Abstract

Lymphangiomas are benign congenital masses which occur most commonly in head and neck of children and incidence of mesenteric lymphangiomas is very rare. We report such a case of huge mesenteric lymphangioma in a 20 year old male who presented to us with acute abdomen. Pre-operative diagnosis is difficult due to silent clinical course and lack of awareness of the clinical and morphological features of this disease.

Keywords: Benign tumours; Congenital masses; Lymphangioma.

Introduction

Lymphangiomas are benign lesions of vascular origin that show lymphatic differentiation. They occur in many anatomic locations and may have a pediatric or adult clinical presentation. Most (95%) occur in the head and neck and axillary regions. Traditionally, lymphangiomas are classified as simple, cavernous, or cystic. The simple type is usually situated superficially in the skin and composed of small thin-walled lymphatic vessels. Cavernous type is composed of dilated lymphatic vessels and lymphoid stroma, and has a connection with various normal adjacent lymphatics while the cystic type consists of lymphatic spaces of various sizes that contain fascicles of smooth muscle and collagen bundles, but has no connection with adjacent normal lymphatics [1].

Although benign in nature, mesenteric lymphangiomas may cause significant morbidity or mortality due to their large size and critical location, when they compress the adjacent structures. The most common mode of acute presentation in children is a small bowel obstruction, sometimes associated with volvulus and intestinal infarction [2]. Intestinal obstruction related to the mesenteric cysts can be induced by compression or traction by the mass [3].

Abdominal radiograph shows a gasless, homogeneous, water-dense mass that displaces bowel loops around it [4]. Ultrasonography often
reveals a sharply-defined cystic or multicystic mass with internal septations [5]. The typical CT scan feature of mesenteric lymphangioma is a multiloculated fluid-filled mass. Characteristic thin walls and septa also can be seen on a CT scan. MDCT with thin slice multiplanar reformation is helpful to not only diagnosis the lesion, but also to assess the complications such as volvulus. CT may add important preoperative information regarding size, anatomical location, adjacent organ involvement, and complications. One report suggested that fine needle aspiration of a milky fluid containing lymphoid cells could confirm a preoperative diagnosis of lymphangioma [6].

The differential diagnosis of a cystic lesion in the pelvic cavity includes reactive ascites, duplication cysts arising from the bowel or bladder, and cystic lesions of the seminal vesicle.

The optimal treatment is radical excision, even when asymptomatic [7]. It may cause infiltration of the intestine, or involvement of the main branch of mesenteric arteries or adjacent organs that necessitate segmental resection of the intestine as was done in our case [8]. Sometimes, radical resection might be technically impossible. The prognosis is excellent if resection is complete. USG is the modality of choice for follow-up, especially in children.

Case Report

A 20 year male presented with central abdominal dull aching pain for the last 15 days, abdominal distension and constipation for 5 days and two episodes of bilious vomiting and low grade fever one day prior to visit to the hospital and was admitted into the ward.

General physical examination and vital signs were normal. On examination, abdomen was distended predominantly in epigastric and left hypochondrium and no lump was palpable. There was a dull note on percussion all over the abdomen except in left hypochondrium and epigastrium and bowel sounds were also appreciable only in above described region. Per rectal and routine blood investigations were normal. X-ray Flat plate abdomen revealed multiple air fluid levels suggestive of small intestinal obstruction. Ultrasonography was suggestive of septic ascites. CT scan (Figure 1) revealed a large lobulated mass in right subhepatic space, involving mesentery, extending up to pelvic cavity and right paracolic gutter and adjacent to ascending colon and caecum, encasing superior mesenteric vessels. Provisional diagnosis of lymphangioma was made and exploratory laprotomy was planned.

On opening abdomen a huge grey-brown multicystic mass in mesentery was discovered (Figure 2). This was surrounding terminal ileum, ascending and proximal transverse colon and encasing superior mesenteric vessels.

Figure 1: CT scan showing huge mass displacing small bowel.

Figure 2: Per operative picture showing mesenteric lymphangioma
Right hemicolecotomy with excision of lymphangioma was done and the part adherent to superior mesenteric vessels was left unresected. Histopathology confirmed the mass as a lymphangioma measuring 20x8x7 cm. The mass was multiloculated and the cut surface was spongy (Figure 3).

Figure 3: Cut surface of the excised mass

Discussion

This report is a case of a huge mesenteric lymphangioma which is a very rare entity. Lymphangiomas are benign lesions of vascular origin that show lymphatic differentiation. About 95% of cases occur in the head, neck and axillary regions; the remaining 5% are located in the mesentery, retroperitoneum, abdominal viscera, lung, and mediastinum [9]. The “classical presentation” is that of a low grade, partial-intestinal obstruction combined with a palpable, freely-mobile abdominal mass [10]. A definite mass may be difficult to palpate because of its large size, fluid consistency, and great mobility. This case presented to us with acute intestinal obstruction without palpable lump.

The gold standard for the diagnosis of the abdominal lymphangioma in an adult is CT scan which was performed in our case. Otitmal treatment is radical excision. We performed right hemicolecotomy with excision of lymphangioma and a part adherent to superior mesenteric vessels was left unresected. The tumor was infiltrating into main trunk of superior mesenteric artery which is a rare entity. We could find very little literature about ways to deal with such a situation which motivated us to report this case so as to throw more light to such a scenario. Post-operative course in our case was un-eventful. In our procedure, Ryle’s tube was removed on day 4 and patient was allowed orally. The patient recovery was remarkable and was discharged on 9th post operative day. A follow up CT scan after 1 year was normal without any recurrence.

Conclusion

A rare case of huge mesenteric lymphangioma has been reported in a 20 year old male. This case was successfully managed surgically with the patient making remarkable recovery.

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Conflict of Interest

No conflict of interest associated with this work.

Contribution of Authors

We declare that this work was done by the authors named in this article and all liabilities pertaining to claims relating to the content of this article will be borne by the authors.

References


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