Lymphangioma of the Spleen—A Rare Tumor Rarely seen in an Adult: A Case Report and a Comprehensive Literature Review

ABSTRACT

Lymphangioma of the spleen, a cystic, benign and slow growing tumor, is rarely seen in children and in adults and usually found only incidentally. We would like to report a case of a splenic lymphangioma in a 33-year-old female that presented to our hospital. To our knowledge this is the first case of its kind being reported from the Indian subcontinent in over 35 years. The previous case was reported in 1974 by Devi et al. In addition to this, our case is also unique because of the age of the patient and the size of the lymphangioma being one of the largest reported so far. This case report will be followed with a detailed literature review of splenic lymphangioma and its management.

INTRODUCTION

Lymphangioma of the spleen is cystic, benign and slow growing tumor that is rarely seen in children and even rarer in adults and is usually found only incidentally. To our knowledge this is the first case of its kind being reported from the Indian subcontinent in over 35 years. The previous case was reported in 1974 by Devi et al. Lymphangiomas were first described by Rodenber in 1828 and the first case of lymphangioma of spleen was reported by Frink in 1885. Primary benign tumors of the spleen account for only 0.007% of all operations and autopsies. Since, 1939 to 1990 around 180 cases of splenic lymphangioma have been reported. The majority of those cases was cystic lymphangioma and included cases which developed as a part of systemic lymphangiomatosis. Another feature of the same disease is isolated splenic lymphangiomatosis and is a much rarer form; from 1990 to 2010, only nine cases have been described in the literature. Spleenic lymphangioma with a solid gross appearance is rare, and only a few cases have been reported so far.

CASE REPORT

A 33 years old, unmarried female patient reported to the surgical department with a history of pain in abdomen and early satiety since 1 month. The pain was more in the epigastric region with occasional radiation to the left upper quadrant. She described the pain as a burning type of pain which aggravated with intake of food. She also complained of fullness of the abdomen and early satiety on taking food. She had no other complaints. She reported having similar complaints intermittently in the past which used to resolve with over the counter antacids. She had no history of weight loss. She had no previous history of trauma to the abdomen or any intra-abdominal infection.

Physical examination revealed no acute distress with stable vital signs. General examination revealed no pallor, no icterus and no lymphadenopathy. On abdominal examination, spleen was palpable till the umbilicus. It was firm and nontender. Rest of the systemic examination was normal.

Her blood investigations were reported as follows; hemoglobulin: 11 gm/dl, white blood cell count: 6,100 cells/mm³, differential count: neutrophils 60%, lymphocytes 36%, eosinophils 4%, erythrocyte sedimentation rate (ESR): 16 mm/hr, platelet count: 2.57 lakhs/dl, bleeding time: 2 minutes, Clotting time: 8 minutes, prothrombin time: 16 seconds (control: 14 seconds), international normalized ratio (INR): 1.11, activated partial thromboplastin time (APTT): 30 seconds (control: 28 seconds), random blood sugar: 91 mg/dl, blood urea: 19 mg/dl, serum creatinine: 0.6 mg/dl, sodium: 133 mEq/l, potassium: 4.7 mEq/l, chloride: 95 mEq/l. Chest X-ray was normal. Urine pregnancy test was negative.

Patient then underwent an abdominopelvic ultrasound to further evaluate the mass. It was reported as a focal, fairly well defined multiloculated cystic mass (12-14 cm) with...
multiple thin internal septations seen at left hypochondriac region causing posteromedial displacement of spleen abutting upper pole of left kidney. Upper gastrointestinal endoscopy was done which revealed antral gastritis and extrinsic compression of stomach along the greater curvature of stomach. Patient was treated with oral pantoprazole 40 mg once a day for the gastritis and it was decided to further evaluate the patient. To further delineate the lesion an abdominal computed tomographic (CT) scan with oral and intravenous contrast was done which showed an enlarged spleen with a 18 × 13 cm fluid density lesion with multiple septa (Figs 1 and 2). The lesion showed an imperceptible wall and was seen impressing the upper pole of left kidney.

A diagnosis of lymphangioma of the spleen was made with a differential of pseudo cyst of the pancreas and it was decided to perform splenectomy as the size of the mass was large and to rule out any malignant changes.

Patient was given pneumococcal vaccine 14 days before the planned surgery. Nasogastric tube was inserted prior to the surgery for gastric aspiration and to prevent post-splenectomy acute dilatation of stomach. Patient was given injection ceftriaxone 1 gm chemoprophylaxis prior to the induction of surgery.

Patient underwent midline laparotomy and splenectomy was done in conventional method and the spleen measuring 19 × 13 × 8 cm weighing approximately 1,740 gm with large multiple cysts and was sent for a histopathological evaluation (Fig. 3). Few accessory splenic masses (4 splenunculi) were also identified and removed during the surgery.

Histopathological examination on cut section revealed multiple cystic spaces filled with gelatinous material with thin rim of splenic parenchyma on the periphery. Microscopy (Fig. 4) showed multiple dilated lymphatic channels lined by endothelial cells with walls showing disorganized bundle cells and lymphocytes with the remaining parenchyma showing congested sinuses with features consistent with lymphangioma of spleen. Since, our hospital is situated in rural part of Kerala, India we were not able to do immunohistochemical study of the specimen. Patient was kept nil per oral for 2 days, and later started on oral diet. Post splenectomy platelet counts were done on day and day 14 were normal. Patient had an uneventful hospital stay and was discharged 4 days postsurgery. Skin sutures were removed on 7th day postsurgery. On subsequent reviews the patient had no further complaints.

**DISCUSSION**

Cystic lesions of the spleen have numbers of different classifications. Currently, the most accepted classification is whether cystic lesions can be classified as primary (true) or secondary (false), based on the presence of a cellular or brous lining. Primary cysts can be further divided into nonparasitic or parasitic (i.e. echinococcal). True nonparasitic cysts include congenital (i.e. epithelial) and neoplastic cysts (lymphangioma, metastases, hemangioma). False cysts may develop secondary to trauma, hemorrhage, infarction-degeneration and inflammation. Histologically, it is classified into three subtypes: Simple (capillary), cavernous and cystic. Cystic type is the most common.

Lymphangiomas are benign malformations composed of endothelial-lined cysts containing lymph. The most commonly involved sites are the neck (75%) and axilla (20%). Lymphangioma is less commonly encountered in the mediastinum, adrenal gland, kidney, bone, omentum, gastrointestinal track, retroperitoneum, spleen, liver and pancreas.
Cystic, benign, slow-growing tumors are usually seen in children, where it is discovered incidentally. These tumors occur more frequently in females, and 80 to 90% are detected before the end of the second year of life and rarely manifests itself after the age of 20 years. Lymphangioma of the spleen can involve the spleen alone either in the form of a single cyst or multiple cysts, or it can be a part of multivisceral involvement also known as systemic cystic angiomatosis. They are generally considered to be a developmental malformation in which obstruction or agenesis of lymphatic tissue results in lymphangiectasia, which is caused by a lack of normal communication of the lymphatic system. It could be due to bleeding or inflammation of the lymphatic system which causes an obstruction leading to additional lymphangiomas. Sexual hormones can influence the growth of lymphangioma. This may explain why it is more commonly seen in females and the sudden increase in the size of lymphangioma during pregnancy.

Isolated splenic lymphangioma can present with different manifestations. It is asymptomatic in the majority of cases. Large cystic lesions may attain sufficient size to cause significant splenomegaly and left-upper quadrant symptoms. Symptoms are usually related to the splenic size. The clinical manifestations of lymphangioma are left upper quadrant pain, abdominal distension, loss of appetite, nausea, vomiting and a palpable mass. These are usually nonspecific and are mostly due to compression of adjacent organs, such as stomach, diaphragm or kidney. The complications associated with more extensive or larger lymphangiomas of the spleen include bleeding, consumptive coagulopathy, hypersplenism and portal hypertension. The effect of abdominal mass produced by the lymphangioma when it exceeds 3,000 or 4,000 gm can occasionally lead to diaphragmatic immobility and consequent atelectasis or pneumonia. Rarely, reversible hypertension due to renal artery compression may be seen. Acute abdominal pain, or a rapid increase in size of the cyst, may occur because of infection or rupture of a cyst.

The radiologic differential diagnosis of primary vascular tumors of the spleen is often difficult to achieve, because of the frequent overlap of their features. Because the lesions are cyst-like, their appearance on ultrasound, CT and magnetic resonance imaging (MRI) is similar to and indistinguishable from one another. Radiographic findings of the cystic form have been well described, but a radiological diagnosis of the capillary and cavernous forms is more difficult. Solitary splenic lymphangiomas are traditionally described as subcapsular, multicystic proliferations that often have identical findings. USG commonly shows hypoechoic spaces which may contain internal echoes. Although CT is nonspecific for distinguishing between many types of retroperitoneal cysts, clinical history and certain details seen at CT can assist in making the correct diagnosis. CT scans usually demonstrate low density, multiple thin-walled, sharply marginated subcapsular cysts, which may contain mural calcifications, thus suggesting a diagnosis of cystic lymphangioma.

On MRI, the mass is shown as multiloculated hyperintensity areas on the T2-weighted images, thus corresponding to the dilated lymphatic spaces whereas T1 imaging is only slightly increased. This is usually due to proteinaceous or hemorrhagic content. The septa are demonstrated as hypointensity bands, corresponding to an abundant amount of fibrous connective tissue. The MR findings typically correlate well with the histologic findings. The angiographic findings include well-defined avascular lesions of varying size scattered throughout the spleen, stretching of the intraparenchymal arterial branches, and an absence of neovascularity, arterial shunting, or
venous pooling. A characteristic ‘Swiss cheese’ appearance of the spleen has been considered pathognomonic. Due to financial constraints of the patient we were unable to do a MRI of her abdomen which would have helped us further delineate the lesion.

Fine needle aspiration biopsy in splenic lymphangioma is contraindicated because of the bleeding risk and limited amount of tissue for accurate diagnosis.

The correct diagnosis depends on histopathologic examination after removal of the spleen. At gross examination, these cysts have a thick fibros wall with an internal morphology that is characterized by fibros trabeculae. Since, lymphatics are found only in the subcapsular region or in large trabeculae, these are the areas of the spleen where lymphangiomas are usually found. Cut section of the lymphangioma reveals a honeycombing of large and small thin-walled cysts usually containing a clear fluid composed of protein. As seen at histologic analysis, capillary, cavernous, and cystic lymphangioma each consists of a single layer of flattened endothelium-lined spaces between fat, fibrotic and lymphatic structures, which are filled with eosinophilic proteinaceous material. Hyalinization and calcification of the fibrous connective tissue may be present.

Histochemical staining of the endothelium demonstrates reactivity with CD31, CD34, factor VIII-related antigen and keratin to varying degrees. Since, our hospital is situated in a rural set up in Kerala, India, we do not have the required facilities required for the immunohistochemical studies. But the histopathological studies done on the specimen were typical of a splenic lymphangioma with all of the characteristic features.

Management of a splenic lymphangioma usually depends on the size. Incidental findings or small lesions do not warrant a splenectomy. Larger lesions that are symptomatic have generally been treated by a splenectomy, but more recently, with the advent of partial resection techniques, a partial splenectomy has also been employed.

For solitary single cysts, this is the preferred treatment. Leaving a splenic remnant in the diffusely involved spleen involves the risk of further growth and enlargement of the spleen, necessitating a possible second operation. Aspiration has proven of little value.

Laparoscopic splenectomy (LS) is emerging as the procedure of choice for splenic removal in patients with a normal to moderately enlarged spleen, benign tumors and hypersplenism. Successful laparoscopic complete excision of a splenic lymphangioma was first reported by Kwan et al in 2001.

LS recommended when a splenic tumor is suspected to be either benign or borderline. However, massive splenomegaly has been considered a contraindication for LS. During surgery, both open and laparoscopic; the search for accessory spleens is an important step. These must be removed even if they appear macroscopically normal, because they could be involved in the pathological process.

The postoperative recurrence rate and the rate of transformation into malignancy is low and the prognosis is good. There have been very few reports of a splenic lymphangioma developing into malignant lymphangiosarcoma. Complications following surgery may include peritonitis, bleeding, abscess and torsion.

Differential diagnosis that always must be borne in mind include but are not limited to hydatid cysts, pancreatic pseudocysts, splenic cysts, cystic hamartomas and hemangiomas, coelomic cysts and cystic intestinal remnants.

In a young patient with splenic lymphangioma, the diagnostic evaluation should be extended to include extrasplicic organs as it has been found that the younger the patient the more the likelihood of multiorgan involvement.

REFERENCES

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