CASE REPORT

Mesenteric calcified cystic lymphangioma in an adult patient

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Abdominal cystic lymphangiomas are rare congenital benign malformations of the lymphatic system. To the best of our knowledge, only 6 mesenteric calcified cystic lymphangiomas have ever been reported. We herein describe a woman who presented to our hospital with stomachache that had been continuous for approximately 8 months. An abdominal computed tomography showed a cystic lesion. In the exploration, the cyst was totally excised. Based on the histomorphological data, a case of ‘calcified cystic lymphangioma’ was diagnosed. Although mesenteric lymphangiomas are rare, especially in adults, they should be considered as a possible cause of abdominal pain. Treatment is surgery with resection of the mass, sometimes including resection of adjacent bowel.

Key words: Calcified cystic lymphangioma, CD31, mesentery

INTRODUCTION

Mesenteric cystic lymphangiomas (MCLs) are rare congenital benign malformations of the lymphatic system that are commonly located on the small bowel mesentery and less typically on the omentum, mesocolon and retroperitoneum (1). Most lymphangiomas are found in the head and neck; intra-abdominal locations are very unusual. Abdominal cystic lymphangiomas are more frequent in boys (5:2), with a mean age at presentation of 2 years (2). The common presentations of intra-abdominal cystic lymphangiomas are abdominal mass and distension, and most have cystic and septal components (3). To the best of our knowledge, only six mesenteric calcified cystic lymphangiomas in adults have ever been reported (4). We describe an extraordinary case of mesenteric calcified cystic lymphangioma in a 32-year-old woman.

CASE REPORT

A 32-year-old female patient presented to our clinic with stomachache that had been continuous for approximately eight months. Her arterial blood pressure was 120/80 mmHg, pulse rate 85 beats/minute and respiration 19/minute, and her surface temperature was normal. An approximately 10 cm in diameter mass was palpable in her...
abdomen. The rest of the abdomen and a digital rectal examination were unremarkable. Laboratory results were within normal limits. The chest X-ray was normal. An abdominal computed tomography showed an 8x7x7 cm cystic lesion with sharp boundaries and lobular contours in the middle abdominal region between the jejunal loop, starting at the L2 vertebra level and ending at the L4 vertebra level. This was thought to be a mesenteric hydatid or diverticular cyst, when evaluated together with the abdominal ultrasonography taken in correlation (Figure 1).

Surgery was planned for the patient. In the exploration, a white cystic lesion of approximately 8 cm in diameter, with increased vascularization and lobular contours, was observed in the intestinal mesentery, 200 cm proximal to the ileocecal valve. A decision was made to excise the lesion (Figure 2). The cyst ruptured during excision and a milky white liquid drained out. It was totally excised.

The pathological study of slices taken from the macroscopically 7x5x3 cm, grayish yellow, irregular surfaced tissue sample of the excised cyst showed a cystic structure with 1-2 mm thick walls filled with a milky liquid. The histopathological study was carried out on hematoxylin and eosin (H&E)-dyed slices prepared from tissue samples taken from the cyst wall after the follow-up. Large numbers of lymphatic veins covered with flattened endothelial cells, generally having smooth muscle bundles in their walls and showing cystic expansion in places, were observed. In some areas of the vein walls, lymphoid aggregates and dystrophic calcification were noted. The immunohistochemical CD31 test showed immune reactivity in the endothelial cells. Based on the histomorphological and immunohistochemical data, a case of “calcified cystic lymphangioma” was diagnosed (Figure 3).

**DISCUSSION**

Mesenteric cystic lymphangiomas (MCLs) occur at all ages, although most (65%) are present at birth. In general, 90% become symptomatic before the second year of life, and nearly 60% are diagnosed before the fifth year of life (5). MCLs are encountered only rarely in surgical practice; the frequency is less than 1 per 100,000 hospital admissions (6).

Losanoff et al. (3) defined four different types of MCLs: pedicled, sessile, retroperitoneal extended,
and multicentric. The etiology of lymphangiomas is still a matter of discussion. A well-established theory suggests that lymphangiomas arise from sequestrations of lymphatic tissue during embryonic development \((1,7,8)\). On the other hand, Godar postulated that premature lymphatics appear as mesenchymal slits, which coalesce and normally communicate with the venous system. Failure to establish this communication may lead to a lymphangioma \((8)\). Fifty percent of cases involve the head and neck, with only 10% occurring in internal organs. Intra-abdominal cystic lymphangiomas may arise from the retroperitoneum, the mesentery and the visceral organs \((2)\). There are only a few case reports of mesenteric lymphangioma in adolescence \((3,7)\).

The common presentations of intra-abdominal cystic lymphangiomas are abdominal mass and distension, and most have cystic and septal components \((2)\). Most MCL patients are initially asymptomatic, with vague and obscure abdominal symptoms emerging early or late, depending on the size and location of the cyst. The clinical symptoms are protean and include pain, nausea, vomiting, or alterations in bowel habits \((9)\). Our patient had chronic stomach pains that had been ongoing for eight months. Known complications that may associate with mesenteric cysts include torsion, rupture, hemorrhage, intestinal obstruction due to volvulus, extraneous pressure on nearby organs, and very rarely, malignant transformation – usually sarcoma, although adenocarcinoma has also been reported \((10,11)\). To our knowledge, only six mesenteric calcified cystic lymphangiomas have been reported previously \((4)\). Ultrasonography and computed tomography are considered the most appropriate radiodiagnostic modalities to evaluate cysts of the mesentery, although usually ultrasonography alone will suffice \((2,12)\).

The differential diagnosis of abdominal cystic lymphangiomas must include other fluid-filled lesions such as pseudocysts, dermoid cysts, enteric duplications, lymphoceles, or neoplasms like mesotheliomas, pancreatic tumors, lipomas, teratomas, leiomyosarcomas, neurofibromas, or liposarcomas \((13)\). The treatment of choice is radical excision, since incomplete resection may lead to recurrence \((14)\). We completely excised the cystic lesion present in the intestinal mesentery of our patient. Although mesenteric lymphangiomas are rare, especially in adults, they should be considered as a possible cause of abdominal pain. Treatment is surgical with resection of the mass, sometimes including resection of the adjacent bowel. After surgery, recurrence rates are low and are usually the result of incomplete excision.

**REFERENCES**