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**Experience in the diagnosis and treatment of mesenteric lymphangioma in adults: A case report and review of literature**

Chen J *et al*. Mesenteric lymphangioma in adults

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**Abstract**

***BACKGROUND***

Mesenteric lymphangioma (ML) in adults is a very rare disease. We reported six hospitalized adult patients with ML in our hospital between January 2013 and July 2018 to investigate the characteristics and prognosis of ML in adults.

***CASE SUMMARY***

The male-to-female ratio was 3:3 and the median age at diagnosis was 55.2 years. Clinical manifestations varied; however, most were acute cases (5/6). No history of trauma was reported．None (0/6) of the patients were accurately diagnosed with ML in the emergency and outpatient departments. Mesenteric cyst was identified in four patients (66.7%) by abdominal ultrasound and in 5 patients (83.3%) by computed tomography．ML was confirmed by pathology postoperatively．Most MLs (4/6) were associated with infection of other systems. ML was located in the mesentery of small intestinal (*n* = 4), ileum (*n* = 1) and rectum (*n* = 1). Cyst fluid was clear (*n* = 4), chylous (*n* = 1) and bloody (*n* = 1). Surgical procedures included complete tumor removal and partial intestinal excision (*n* = 6). Recurrence and adhesive intestinal obstruction were not observed during the 3-12 mo follow-up period.

***CONCLUSION***

ML in adults is a rare benign acquired disease which can be cured by surgical treatment. Infection may be a cause of ML.

**Key words:** Mesenteric lymphangioma; Mesenteric cyst; Adults; Acute abdominal pain; Case report

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**Core tip:** Mesenteric lymphangioma (ML) is a rare congenital lymphangioma occurs predominantly in children. We reported six cases of adult patients with ML and reviewed the literature. The report is helpful in comprehensively understanding the characteristics and prognosis of ML in adults and arousing the clinician's attention to this disease.

Chen J, Du L, Wang DR. Experience in the diagnosis and treatment of mesenteric lymphangioma in adults: A case report and review of literature. *World J Gastrointest Oncol* 2018; In press

**INTRODUCTION**

Mesenteric lymphangioma (ML) is a rare congenital lymphangioma of uncertain etiology, and occurs predominantly in children. ML lacks specific clinical signs and symptoms, and patients often attend hospital due to complications, such as abdominal pain, abdominal distension, intestinal obstruction and other acute abdominal manifestations. In adults, it is often found by accident during auxiliary examinations or even [exploratory](javascript:;) [laparotomy](javascript:;) of an acute abdomen, and leads to passive surgery or surgical preparation, and even missing the best surgical opportunity due to delayed diagnosis. Cases and misdiagnosed cases have been reported in previous studies[1-3]. However, these studies mainly focused on imaging. Here, we retrospectively analyzed six adults with ML confirmed by pathological examination, admitted between January 2013 and July 2018. Our findings may improve our understanding of this disease and provide more clinical references for early and correct treatment.

**CASE PRESENTATION**

***Chief complaints***

**Case 1:** A 45-year-old man with abdominal pain for 16 h.

**Case 2:** A 59-year-old man with severe abdominal pain, no defecation and exhaust for 3 d.

**Case 3:** A 62-year-old man with abdominal pain, nausea and fever for 3 h.

**Case 4:** A 71-year-old woman with abdominal pain, fever, diarrhea, and gastrointestinal bleeding for 1 wk.

**Case 5:** A 42-year-old woman with abdominal distension for 2 years ruccurent 1 d.

**Case 6:** A 52-year-old woman with mild abdominal pain and distension, nausea and fever for more than one day.

***History of present illness***

None of the six patients had a significant history of trauma. Most MLs (4/6) were associated with infection of other systems. Case 1: cholecystitis; Case 2: no other systemic co-infection was found; Case 3: cholangitis; Case 4: urinary tract infection; Case 5: no other systemic co-infection was found; Case 6: Gastrointestinal tract infection, acute gastritis and colitis.

***History of past illness***

The history of symptoms ranged from 3 h to 2 years. Case 3 had a medical history of cholangiolithiasis and Endoscopic retrograde cholangiopancreatography, Case 5 had an abdominal cyst 2 years previously, which was untreated and diabetes for more than 5 years, and was admitted to our hospital due to acute abdominal distension. All other cases were admitted to the emergency department for acute abdominal pain without history of other chronic diseases.

***Physical examination***

Case 2: middle and upper abdominal tenderness and rebound tenderness, no muscle guarding; bowl sound was high pitched tinkling. Case 4: periumbilical tenderness, no rebound tenderness and muscle guarding, a palpable mass of about 10 cm in diameter can be detected when abdominal pain occurs, and disappeared when abdominal pain is relieved. All other cases showed tenderness in different parts of the abdomen, no rebound tenderness and muscle guarding, bowel sound were normal.

***Laboratory testing***

Tumor markers were normal in all cases, and laboratory tests indicated increased white blood cell to different degrees, while Case 2 and 3 showed slight increase in alanine aminotransferase and γ-glutamyl transpeptidase. Case 4 and Case 6 indicated fecal occult blood (+), and fecal bacteria culture were negative.

***Imaging examination***

All patients underwent abdominal ultrasound and abdominal computed tomography/magnetic resonance imaging (CT/MRI) examination after admission (Figures 1 and 2), and some patients had abdominal X-ray examination. In one patient with acute intussusception and one patient with diarrhea, abdominal X-ray showed fluid and an incomplete intestinal obstruction. In other patients with acute abdominal pain, abdominal X-ray showed no obvious abnormalities. Five patients were found to have an intraperitoneal cystic or solid cystic mass by CT/MRI, Abdominal ultrasonography failed to detect an abdominal cyst in Case 3 and Case 4. Preoperative ultrasound and CT failed to detect an abdominal cyst in Case 3.

**FINAL DIAGNOSIS**

All cysts were examined by pathology after operation, and all were ML (Figures 3).

**TREATMENT**

Case 6 of a suspected rectal tumor and Case 5 of an intra-abdominal benign cyst chose elective surgery, and the remaining four patients underwent emergency surgery within 48 h of hospital admission. Intraoperative ML was identified in the jejunum in four cases, the ileum in one case, and the rectum in one case. Four cases had clear cystic fluid, one case had chylous fluid and one case had bloody fluid. Tumor invasion was noted in the bile duct in one case, the duodenum in one case, the transverse colon in one case, and the rectum in one case. All six cases underwent complete removal of tumor and partial intestine excision, ranging from 6 to 50 cm.

**OUTCOME AND FOLLOW-UP**

Six patients were included in this study. Patient information and clinical manifestations are shown in Table 1. Of the six patients, three were male and three were female, aged 42-71 years, with an average age of 55.2 years. The clinical manifestations of ML included acute abdominal pain, acute intestinal obstruction (vomiting, abdominal distension, no defecation and exhaust), fever, diarrhea, and gastrointestinal bleeding. The initial diagnoses included: one case of acute intussusception, three cases of abdominal tumors (mesenteric lipoma, duodenal and rectum tumor), one case of acute cholangitis, and one case of acute hemorrhagic enteritis. All six patients were diagnosed with ML by pathology following surgery. The accuracy of initial diagnosis was zero (0/6). The diagnostic accuracy of ultrasound for mesenteric cyst was 66.7% (4/6). The diagnostic accuracy of CT/MRI for mesenteric cyst was 83.3% (5/6).

All the patients had good postoperative recovery, the abdominal mass disappeared, appetite, defecation and urine output became normal. All patients were followed up for 3 mo by abdominal CT, and no recurrence or adhesive intestinal obstruction occurred. Five cases (83.3%) were followed for up to one year and no recurrences were observed.

DISCUSSION

Lymphangiomas are uncommon congenital malformations of the lymphatic system. They can occur at any site in the body, but are most commonly found in the neck and head area as well as the abdominal wall, but rarely in the mesentery[4]. It is reported that the incidence of ML is approximately 5%, and the male-female ratio is about 1.5-3:1[5,6]. In addition, ML has been described in less than 1% of all lymphangiomas[1,7,8].

The exact etiology of ML is unknown. It is likely to be a developmental anomaly of the lymphatic system, as 65% of MLs are present at birth and 90% of all patients are diagnosed before the age of 2 years[7]. However, they are also formed due to an inflammatory process, lymphatic obstruction, surgery, radiation and abdominal trauma[9,10]. ML is rarely seen in adults. So it's not clear what the incidence of ML is in adults. Only a few case reports of ML in adults are available in the published literature[11]. In adults, lymphangiomas mostly occur on the body surface or in the abdominal cavity, and the incidence of ML is 1/100000[2], mostly in the small intestine, followed by the omentummajus, mesentery and retroperitoneum. ML is a benign lesion, with a relatively asymptomatic onset, slow growth and a long disease course. Most MLs are initially asymptomatic, and so are usually discovered incidentally. However, when the tumor is large, it can compress the surrounding viscera or block the intestine producing corresponding symptoms. The clinical symptoms of ML vary depending on location. ML can manifest as abdominal pain, abdominal distension, diarrhea, hematochezia, constipation, hypoproteinemia, intussusception, and decreased physical quality[12]**.** Due to the lack of specific clinical signs and symptoms, ML is easily missed and misdiagnosed. Abdominal ultrasonography or CT can be used to identify an early abdominal cystic mass, especially in patients with recurrent abdominal pain, abdominal distension, and stubborn constipation. Emergency doctors should be vigilant, and early abdominal ultrasound or CT examination should be carried out to detect this disease as soon as possible. Ultrasonography is of high diagnostic value in detecting the location, size, division of the cyst, cyst fluid, cyst wall and its relation to surrounding tissues[13]. In small cysts, abdominal CT is more sensitive and helps to differentiate from other related abdominal pelvic cysts, such as greater omentum cysts, intestinal repetitive malformations, ovarian cysts, common bile duct cysts, and kidney cysts. CT is useful for further understanding the relationship between the cyst and the surrounding tissues and organs, especially large blood vessels and the bowel, which ultimately aids treatment decisions and surgical approaches in these patients[14]. MRI is more sensitive in patients with intracavitary hemorrhage. In the case of intrathecal hemorrhage, the imagingmay show solid cystic signs[15,16]. Pathological examination is the gold standard for the diagnosis of ML, and provides strong evidence for postoperative identification of other types of cysts. Microscopy shows dilated lymphatics, and the thin wall lining epithelial cells in the lymphatic cavity gap, and a small amount of smooth muscle tissue can be seen. During infection, the infiltration of lymphocytes, plasma cells, eosinophils and other inflammatory cells are also visible. MLs are classified as simple, cavernous and cystic. The simple type of ML is mainly situated superficially in the skin and is composed of small thin-walled lymphatic vessels. The cavernous type has dilated lymphatic vessels and has connections with normal adjacent lymphatics. Cystic lymphangioma is composed of large macroscopic lymphatic spaces surrounded by collagen and smooth muscle, and does not have connections with adjacent normal lymphatics (CD34 and CD31 positive)[17]. ML needs to be differentiated from peritoneal abscess, hematoma, malignant tumor center necrosis liquefaction, malignant tumor cystic adenoma and some solid masses such as sarcomas derived from mesenchymal tissue. Simple abdominal ultrasonography may be difficult to distinguish, and abdominal CT/MRI can provide more information. However, the diagnosis of ML mainly depends on pathological diagnosis. When necessary, ultrasound-guided diagnostic puncture can further differentiate the characteristics of cyst, abscess and hematoma. Ultrasound-guided needle aspiration cytology can identify the benign and malignant, then providing a basis for differential diagnosis. To treat ML, most doctors recommend radical surgical excision, as ML can grow very large and invade adjacent structures, develop complications and the risk of sarcoma transformation on irradiation[9]. After excise a mass that involves the whole mesentery, internal herniation is likely to occur due to the presence of skeletonized vessels. A biological collagen implant can be used to repair the mesenteric defect after excision of a large ML and monitoring for recurrence during follow-up is necessary[18]. But there is also a different opinion on treatment. A published literature[19] shows that, for asymptomatic or mild lymphangioma patients, conservative treatment and close follow-up are recommended. In a case report, colorectal lymphangioma can spontaneously disappear. In my opinion, if ML is relatively large, it is possible to infiltrate the surrounding organs and cause tissue ischemic necrosis, even develop life-threatening complications, such as traumatic rupture, anaemia secondary to intraabdominal or intra-cavitary bleeding, intestinal gangrene secondary to volvulus and intermittent intestinal obstruction[2,16]. It is better to have it removed as soon as possible.

In the present study, the ratio of men to women was 3:3, with no significant gender difference. The rate of accurate initial diagnosis was zero. This indicated that clinicians lack of awareness of ML in adults, and the average age at diagnosis in these patients was 55.2 years. All of these patients had received routine physical examination in the past few years, but no abdominal cysts were found. Five of these six cases had no previous history of abdominal cysts, but all patients had acute onset, mainly presenting with acute abdominal pain, incomplete intestinal obstruction, and mucinous bloody diarrhea. The remaining patient was found to have abdominal cysts 2 years previously, which were untreated. The patient’s acute abdominal distension was not treated with anti-inflammatory therapy, and was diagnosed and cured after surgery. Following symptom onset, the abdominal cysts were not identified by abdominal color Doppler ultrasound and CT. However, the diagnosis of abdominal cysts was confirmed during laparotomy. It is speculated that adult ML may be an acquired disease, and infection may be a risk factor. Of these six cases, two had biliary tract infection, one had urinary tract infection, one had digestive tract infection, and two patients had no obvious infection. The pathogenesis of ML may include the formation of secondary cysts caused by lymphatic obstruction due to infection. Moreover, we found that the detection rate of abdominal cysts by ultrasound was lower than that of abdominal CT (66.7% *vs* 88.3%). This may have been due to the following possibilities: (1) due to the rarity of ML in adults, ultrasound clinicians lack awareness of ML, resulting in missed diagnosis; (2) misdiagnosed cystic lesions such as ascites or dilated bowel may be caused by intestinal obstruction; and (3) ML may be an acquired disease. As clinicians are accustomed to prescribing abdominal ultrasound first and then CT, an abdominal cyst may not yet have formed when ultrasonography is performed. Pathological examination is the gold standard in the diagnosis of ML. An accurate and thorough surgical technique is an effective method of treating ML. No recurrence was found during the follow-up period of 3-12 mo in the six patients included in this study. However, timely surgery is essential. In a patient with mucinous diarrhea, infiltration of the transverse colon was found during surgery, leading to transverse colonic ischemic necrosis, and partial transverse colectomy was performed.

In summary, ML in adults is an extremely rare benign disease, which may be an acquired disease, different from that in children which is due to congenital lymphatic dysplasia. Infection may be involved in the pathogenesis of ML in adults. However, the exact etiology should be confirmed in a large sample study. In some cases, ML can be fatal, as it may cause tissue ischemic necrosis due to infiltration of surrounding viscera, compression of the intestine or peritoneal vessels, resulting in serious complications. Timely and effective radical surgery is necessary, which requires increased awareness of the disease in clinicians to avoid misdiagnosis and missed diagnosis.

**EXPERIENCES AND LESSONS**

Although ML occurs more frequently in children, it also occurs in adults, the exact etiology in adults needs some further studies .Although ML is benign, it can also lead to serious and fatal consequences. Timely and effective radical surgery can cure this disease. Clinicians should raise awareness of ML in adults.

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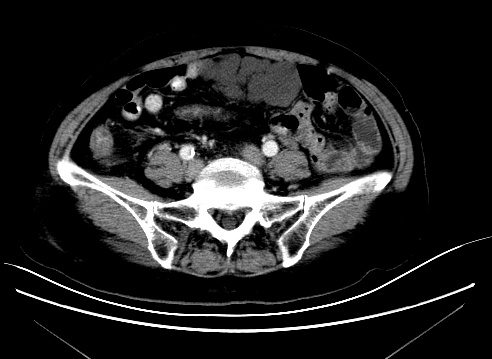
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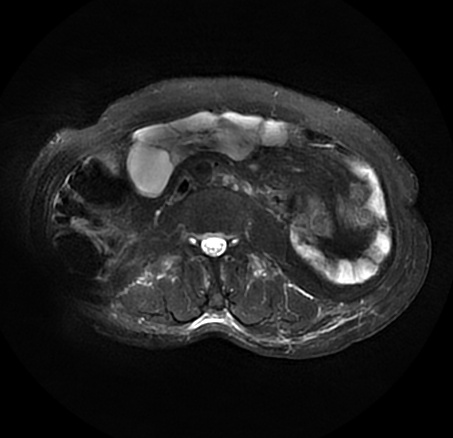
**Table 1 Information and clinical manifestations in the six patients**

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| Cases | 1 | 2 | 3 | 4 | 5 | 6 |
| Gender | M | M | M | F | F | F |
| Age (yr) | 45 | 59 | 62 | 71 | 42 | 52 |
| Abdominal pain | + | + | + | + | - | + |
| Nausea | - | - | + | - | - | + |
| Vomiting | - | + | - | - | - | - |
| Diarrhea | - | - | - | + | - | - |
| GI bleeding | - | - | - | + | - | - |
| Fever | - | - | + | + | - | + |
| Concurrent infection | Cholecystitis | UK | Cholangitis | Urinary tract | UK | GI tract |
| Abdominal distension | - | - | + | + | + | + |
| Intestinal obstruction | - | + | - | - | - | - |
| Medical history | - | - | Cholangiolithiasis,  ERCP | - | Diabetes | - |
| Trauma | - | - | - | - | - | - |

UK: Unknown; ERCP: Endoscopic retrograde cholangiopancreatography; GI: Gastrointestinal tract.



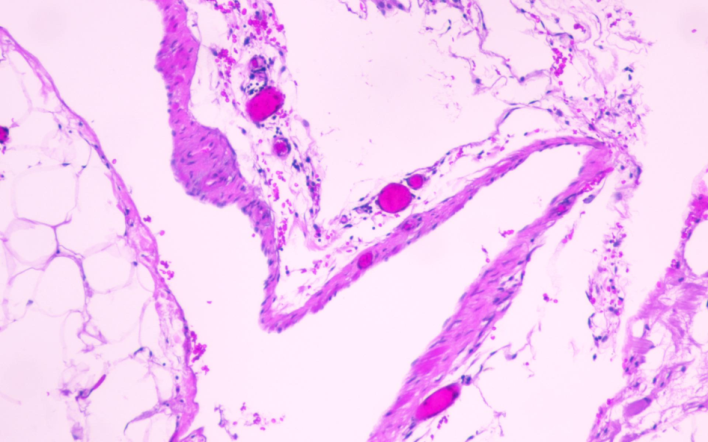
**Figure 1 Abdominal computed tomography scan reveals lumpy low-density shadows around the upper middle intestine, 86 mm × 42 mm in size, and no enhancement was observed (Case 4).**



B

A

**Figure 2 Abdominal magnetic resonance imaging shows cystic long T1 and long T2 signal masses in the anterior wall of the middle abdomen, 110 mm× 40 mm in size.** No abnormal enhancement was observed (Case 4). A: Coronal position; B: Axial position.



**Figure 3 Histopathology shows that the cyst wall was composed of fibrous adipose tissue, and a few lymphatic endothelial proliferations.** Case 4, hematoxylin and eosin, 100 ×.