Case report

Lymphangiomatosis of the ileum with perforation: A case report and review of the literature

Antonio Giuliani\textsuperscript{a}, Lucia Romano\textsuperscript{a,}\textsuperscript{*}, Gino Coletti\textsuperscript{b}, Mohammad Walid A Fatayer\textsuperscript{b}, Giuseppe Calvisi\textsuperscript{b}, Francesco Maffione\textsuperscript{a}, Chiara Muolo\textsuperscript{a}, Vincenzo Vicentini\textsuperscript{a}, Mario Schietroma\textsuperscript{a}, Francesco Carlei\textsuperscript{a}

\textsuperscript{a}Department of Surgery, University of L’Aquila, Italy
\textsuperscript{b}UOC Anatomia Patologica, ASL1 Abruzzo, Ospedale San Salvatore, L’aquila, Italy

\textbf{ARTICLE INFO}

\textbf{Keywords:}
Lymphangiomatosis
Bowel perforation
Small intestine
Bowel resection
Acute abdomen

\textbf{ABSTRACT}

Lymphangiomatosis is a benign proliferation of lymph vessels. Lymphatic diseases can vary from small lymphangioma to generalized lymphangiomatosis, which is a rare condition and can have several clinical manifestations. The gastrointestinal tract may be affected, but the incidence in the intestinal wall is very low. We propose in our study a case of ileal lymphangiomatosis presenting with perforation, in which the diagnosis was made after the pathological analysis of the resected intestinal tract. Although rare and not described in the literature, intestinal lymphangiomatosis could manifest itself with acute abdomen and could be a surgical urgency. This disease should be considered when intestinal perforation is observed.

1. Introduction

Lymphangiomatosis is a benign proliferation of lymph vessels. Lymphatic diseases can vary from small lymphangioma to generalized lymphangiomatosis, which is a rare condition and can have several clinical manifestations \cite{1-6}. Although the underlying pathogenetic mechanism is unknown, it is generally considered as congenital malformation of the lymphatic system associated with alterations in the circulatory dynamics of the lymph \cite{7}. It can occur anywhere in the body and also abdominal lymphangiomatosis is reported, but in many cases it involves the mesentery, omentum, mesocolon and retroperitoneum. The gastrointestinal (GI) tract may be affected, but the incidence in the intestinal wall is very low \cite{8,9}. In the few cases described in the literature, the symptomatology was characterized mainly by abdominal pain and bleeding. Aggressive surgery should be avoided in symptomless cases, because it is now known that these lesions are benign \cite{7}.

We propose in our study a case of ileal lymphangiomatosis presenting with perforation, in which the diagnosis was made after the pathological analysis of the resected intestinal tract. In addition, the relevant medical literature on intestinal lymphangiomatosis was reviewed. This work has been reported in line with the SCARE criteria \cite{8}.

2. Presentation of case

A 41-year-old male presented to the Emergency Department with significant diffuse abdominal pain, nausea, vomiting and inability to pass gas or stool (constipation); the patient reported that these symptoms were present for 10 hours. He denied any significant family history of disease and he did not refer to major diseases or prior surgical interventions in his own medical history. During the physical examination, the subject was in discrete general conditions, collaborative, the sensory was intact, the decubitus indifferent, the breath eupneic and the pulse rhythmic. The vital parameters were preserved, and temperature was within normal limits. The abdominal examination revealed a flat abdomen, tender to palpation, painful to deep palpation on all quadrants, liver size appears within normal limits, Murphy’s sign was negative, Blumberg’s sign was positive, bowel sounds were absent. Rectal exploration indicated nothing significant. Laboratory values upon admission reported hemoglobin value of 15.5 g/dL, White Blood Cells 12.000/mmc, PCR 0.04 mg/dl. Upright abdominal films, then confirmed with computed tomography (CT) enhancement scan, revealed bowel distension and the presence of multiple gas-fluid levels. This framework suggested the presence of small bowel obstruction. As the etiology of the obstruction remained unidentified, the decision was made to perform a diagnostic laparoscopy. On entering the peritoneal cavity, the small bowel was examined: the intestinal loops were vital.

\textsuperscript{*}Corresponding author. University of L’Aquila, Dipartimento di Scienze Chirurgiche, Ospedale San Salvatore, 67100, Coppito, AQ, Italy.
E-mail address: lucia.romano1989@libero.it (L. Romano).

https://doi.org/10.1016/j.amsu.2019.03.010
Received 14 February 2019; Accepted 24 March 2019
2049-0801/© 2019 The Author(s). Published by Elsevier Ltd on behalf of IJS Publishing Group Ltd. This is an open access article under the CC BY license (http://creativecommons.org/licenses/BY/4.0/).
and vascularized. At approximately 80 cm from the ileocaecal valve, a volvulus was identified, and in close proximity there was a long tubular structure, which proved to be a Meckel’s diverticulum, with adhesions to parietal peritoneum. A lysis of adhesions was performed, the structure, which proved to be a Meckel’s diverticulum, with adhesions volvulus was identified, and vascularized. At approximately 80 cm from the ileocaecal valve, a microperforation of the bowel. The perforation site seemed to be distant above the previous diverticulum. About 20 cm of ileal resection was performed and side-to-side mechanical anastomosis was made. Two surgical drains were also placed. The postoperative course was regular, and the patient was discharged one the 10th post-operative day.

Surgical drain took out enteric material and blood. The body temperature was 38 °C. Laboratory values reported hemoglobin value of 12.7 g/dL, White Blood Cells 7.000/mmc, PCR 10.22 mg/dl. The patient again underwent urgent surgery. The intervention was an exploratory laparoscopy, then converted to open ileal resection for a perforation is reported.

Histological examination showed stratification of fibrin and granulocytes on the serosa and presence of diffuse lymphangiomatosis (positive immunohistochemistry of D2-40 marker) involving the submucosa and, in some parts, the full-thickness muscular wall (Figs. 1 and 2).

3. Discussion

Lymphangiomatosis is a multisystemic disorder, characterized by congenital malformation of the lymphatic system with channels and cystic spaces of varying size. It may affect all the areas of the body; therefore, symptoms and complications are related to localization. Usually in the course of the disease patients are asymptomatic at first, but then the abnormally proliferating lymphatic channels are capable of massive expansion and infiltration into surrounding tissues. Abdominal lymphangiomatosis is quite often reported, but in many cases it arises in the mesentery, omentum, mesocolon and retroperitoneum. The GI tract may be affected, but the incidence of lymphangioma of the intestinal wall is very low, and it is even rarer in the small bowel (less than 1%) [9,10]. To the best of our knowledge, all cases reported in the literature with GI tract involvement are summarized in Table 1 [1,2,7,10–18].

There is only one case report of lymphangiomatosis involving the oesophagus [18] and one case report involving fundus of the stomach [2]. Lymphangiomatosis of the colon is described in four papers [7,13–15]. In these patients, the main symptoms were abdominal discomfort, bleeding and anaemia.

Lymphangiomatosis of small bowel is described in 6 works [1,10–12,15,16]. Patients had abdominal pain, bleeding, melena, anaemia, pedal edema, weakness and weight loss, but no case of perforation is reported.

In the majority of patients described in the literature, diagnosis was suggested by colonoscopy and biopsy, which mainly showed protruding submucosal lesions. The diagnostic workup was sometimes integrated by contrast-enhanced CT, showing marked thickening of the walls of the bowel loops.

In five papers [10–12,15,16], patients were treated with surgical resection of the affected part of intestine, whereas one patient improved on conservative management and was put on low-fat and high-protein diet [1]. In all reported cases, follow-up was of short duration. In two cases patients continued to have symptoms after surgery [2] or after endoscopic mucosectomy [13].

The histological features of the lymphangiomatosis are non-specific, and the definitive diagnosis requires the demonstration of an hyperproliferation of normal lymphatic vessels with normal endothelium, predominantly in the context of submucosa, with disruption of the muscular layer and sometimes of the serosa. This condition creates a locus minoris resistentiae, and this may explain the pathogenesis of the

Fig. 1. Microscopic findings. (A) Hematoxylin and eosin staining of numerous dilated lymphatic vessels (4× magnification). (B) Immunohistochemical D2-40 expression (brown color) in dilated lymphatic vessels of the submucosa (10× magnification). (C) Subserous dilated lymphatic vessels with discontinuity of the muscular layer and serositis (H&E, 10× magnification). (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

Fig. 2. (A) D2-40 immunostaining shows positive reactivity for lining endothelial cells of lymphatic spaces in the muscular layer (10× magnification). (B) Numerous submucosal dilated lymphatic vessels (H&E, 10× magnification). (C) Lymphatic vessels that interrupt muscular layer (D2-40 stain, 10× magnification).

A. Giuliani, et al. Annals of Medicine and Surgery 41 (2019) 6–10
### Table 1

Papers about lymphangiomatosis-related disease.

| Reference        | Age and gender | GI tract organs involved | Type of lesions | Histopathologic findings                                                                 | Clinical features                                                                 | Diagnostic workup                              | Treatment                                                                 | Follow-up and outcome                                                                 |
|------------------|----------------|--------------------------|-----------------|------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------|--------------------------------------------------------------------------------|---------------------------------------------------------------------------------|----------------------------------------------------------------------------------|
| Valakada J, Madhusudhan KS et al. [1] | 59-years-old woman | duodenum, jejunum, mesentery and retroperitoneum | marked thickening of the small bowel loops in the duodenum and jejunum and multiple tubular channels in the mesentery and retroperitoneum hyperintense on T2-weighted images. | lymphangectasia | recurrent abdominal pain, multiple episodes of melena, pedal edema, pallor and mild hepatosplenomegaly | abdominal magnetic resonance imaging (MRI), abdominal ultrasonography, double-balloon enteroscopy and biopsy | conservative management: low-fat and high-protein diet |                                                                                     |
| Lin RJ, Zou H et al. [2] | 38-year-old female | fundus of the stomach, peripancreatic area, mesenteric area, retroperitoneal space of the spleen, right upper quadrant of the greater omentum | multiple small cystic lesions without enhancement (TC), multiple cystic dark areas (abdominal ultrasonography) | submucosal microcystic cysts of lymphatic channels with walls composed of thin fibrous tissue | melena for 3 months, weakness for 10 days, hemoptysis for 4 months | computed tomography, abdominal ultrasonography, biopsy | distal gastric resection and Billroth II-type anastomosis | she continued to present melena, iron deficiency anemia and hypoproteinemia after the surgery |
| Jung SW, Cha JM et al. [7] | 31-years-old woman | ascending colon, from the cecum to the hepatic flexure | multiple thumbprint-like lesions on the air contrast barium enema; clusters of round submucosal tumors with smooth surface, without ulcerations or erosions and positive to the cushion sign on the colonoscopy; the EUS showed echo free cysts with a clear border and septal walls in the submucosal layer | submucosal cysts lined by endothelial cells, serous liquid resembling lymphatic fluid, with occasional multinucleated cells and without fat or blood cell components | air contrast barium enema, colonoscopy, EUS and endoscopic biopsy | the patient was not treated with invasive treatment because she was asymptomatic |                                                                                     |
| Rai P, Rao RN et al. [10] | 31-years-old man | small bowel and small bowel mesentery starting from mid-jejunum to ileocecal junction | protruding submucosal lesions on the colonoscopy, cystic lesions on the CT | multiple irregular dilated space lined by endothelial cells with lymphoid aggregates, filled with acellular proteinaceous material and no evidence of malignant cells | recurrent melena for the last 8 years and iron deficiency | colonoscopy, capsule endoscopy, contrast-enhanced CT, laparotomy with intraoperative endoscopy and endoscopic biopsy | limited ileocolic resection, end ileostomy and distal mucus fistula. After few days continuity was restored. Complete surgical resection of the segment involving the lesions | no gastrointestinal bleed, haemoglobin and albumin were normalised |
| Hwang SS, Choi HJ et al. [11] | 71-year-old man | jejunal and adjacent mesentery | multiple nodular mesenteric masses infiltrating into the jejunum and adjacent mesentery; multiloculated cystic lesion from the mucosa to the subserosa diffuse wall thickness (CT) | numerous multiloculated, cystically dilated spaces lined by attenuated endothelial cells that appeared to dissect through the muscularis propria of the small intestine with inside fluid containing lymphocytes | weakness, swelling in leg, weight loss, preterminal edema and recurrent upper respiratory infections | computed tomography, 18FDG PET/CT, biopsy | no gastrointestinal bleed, haemoglobin and albumin were normalised |                                                                                     |
| Ilhan M, Oner G et al. [12] | 43-years-old woman | ileum and jejunum | | | | | |                                                                                     |
| Chung WC, Kim HK et al. [13] | 48-years-old man | proximal transverse colon | several protruding mucosal lesions covered | | abdominal discomfort and anemia | colonoscopy, abdomen CT, biopsy | after 1 month surgery preterminal edema was not seen, protein and albumin increased | the patient had abdominal pain and (continued on next page) |
| Reference | Age and gender | GI tract organs involved | Type of lesions | Histopathologic findings | Clinical features | Diagnostic workup | Treatment | Follow-up and outcome |
|-----------|----------------|--------------------------|----------------|--------------------------|------------------|------------------|-----------|----------------------|
| Lee JS, Kim GW et al. [14] | 38 year-old man for a general check-up | mid-portion of the ascending colon up to the proximal portion of the transverse colon | variably sized cystic mass lesions | with normal mucosa on the colonoscopy | recurrent melena for more than 2 months | chest and abdominal radiography, esophagogastroduodenoscopy, colonoscopy, abdominal ultrasonography, CT and biopsy | several incisions and excisional biopsies | anemia when he was followed up 3 month after musectomy |
| Fang JF, Qiu LF et al. [15] | 57-years-old woman | small intestine, 30 cm distal to the flexor tendon | mass with ulcers and erosion approximately of 5 cm × 4 cm | intrinsic layer of dilated lymphatic vessels and a small amount of interstitial neutrophil, eosinophil, plasma cell infiltration | gastroscopy, enteroscopy, and biopsy | partial resection of the small intestine | during the follow-up no recurrence was observed |
| Dong A, Zhang L et al. [16] | 22-years-old female | mesentery and ileum | mass involving mesentery and ileum with nodules in the mass | proliferation and dilation of the mucosal lymphatic, containing a large amount of red blood cells. The cells were positive for CD31, CD34 and D2-40. Ki-67 was about 1%. | 9-month history of intermittent melena, weakness and palpitation | abdominal MR, abdominal CT, PET-CT and biopsy | resection of the abdominal mass and a segment of 60 cm of the ileum invaded by the abdominal mass | after surgery symptoms improved and follow-up laboratory tests showed normal red blood cell count and hemoglobin level |
| Lu G, Li H et al. [17] | 79-year-old man | sigmoid colon | multiple cystic masses (colonoscopy), with septal walls in the submucosal layer | cysts located in the submucosal layer surrounded by flat endothelial cells that were positive for D2-40 at the immunohistochemistry multiple dilated lymphatic vessels of different sizes filled with pink beneath squamous epithelium | intermittent attacks of bowel bleeding and abdominal discomfort for 3 months | colonoscopy, endoscopic ultrasound and biopsy | laparoscopy-assisted partial sigmoid colon resection | in the 2-year follow-up after the operation, no bleeding or other complications were noticed |
| Xue L, Guo WG et al. [18] | 58-year-old man | lower esophagus | longitudinally protruding mass covered with normal esophageal mucosa and a lesion outside but adjacent to the wall of the esophagus | dysphagia of 7 months | esophagogastroscopy, esophageal ultrasonography, chest CT and biopsy | a right lateral thoracotomy was performed for the resection of the cysts, first the lesion outside and than that protruding in to the esophageal lumen | the postoperative course was uneventful and at the patient was discharged on th 10th postoperative day |
perforation, reported in our case. The involvement can be continuous or, more frequently, segmental. The impairment of the muscular layer and of the submucosal nerve plexus could also have contributed to the development of the intestinal obstruction, which was the symptom because our patient came to our attention. This suggests that also lymphangiomatosis should be taken into account among other rare causes of intestinal obstruction [19,20].

4. Conclusion

In conclusion, lymphangiomatosis of the small bowel is a rare disease that has no specific clinical features, so it is easy for a clinician to make a misdiagnosis or to miss diagnosis. In some cases, surgical resection may be required to provide definitive histological diagnosis, as occurred in our cases. We want to share our experience about this, because, although rare and not described in the literature, intestinal lymphangiomatosis could manifest itself with an acute abdomen and surgical urgency. This disease should be considered when intestinal perforation is observed. In particular, the pathologist should keep it in mind in the differential diagnosis, when he analyses a case of perforation whose cause is not very clear and specified.

Ethical approval

Ethical approval was not required.

Sources of funding

No funding

Author contribution

Antonio Giuliani, Lucia Romano: Writing.
Gino Coletti, Mohammad walid A Fatayer, Giuseppe Calvisi: Images and contribution to the text.
Francesco Maffione, Chiara Muolo, Vincenzo Vicentini: Data collection.
Mario Schietroma, Francesco Carlei: Study design and review.

Conflicts of interest

No conflict of interest.

Research registration number

None.

Guarantor

Prof. Francesco Carlei.

Consent

Written informed consent was obtained from the patient for publication of this case report and accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal on request.

Provenance and peer review

Not commissioned, externally peer reviewed.

Acknowledgments

All authors disclose the absence of any financial and personal relationships with other people or organizations that could inappropriately influence (bias) their work.

References

[1] J. Valakada, K.S. Madhusudhan, g Ranjan, et al., Abdominal lymphangiomatosis with intestinal lymphangiectasia diagnosed by magnetic resonance lymphangiography: a case report. Curr. Probl. Diagn. Radiol. 47 (3) (2018 May - Jun) 200–202.
[2] R.Y. Lin, H. Zou, T.-Z. Chen, et al., Abdominal lymphangiomatosis in a 38-year-old female: case report and literature review, World J. Gastroenterol. 20 (25) (2014) 8320–8324.
[3] K.C. Woon, B.J. Jang, T.N. Kim, et al., A case of primary intestinal lymphangiectasia, Korean J. Intern. Med. 8 (1) (1993) 51–56.
[4] D.H. Yang, H.W. Goo, Generalized lymphangiomatosis: radiologic findings in three pediatric patients, Korean J. Radiol. 7 (4) (2006) 287.
[5] A.S. Puri, R. Aggarwal, R.K. Gupta, et al., Intestinal lymphangiectasia: evaluation by CT and scintigraphy, Gastrointest. Radiol. 17 (2) (1992) 119–121.
[6] F. Blei, Lymphangiomatosis: clinical overview, Lymphatic Res. Biol. 9 (2011) 185–190, https://doi.org/10.1089/lrb.2011.0020 PMID: 22196283.
[7] S.W. Jung, J.M. Cha, J.I. Lee, et al., A case report with lymphangiomatosis of the colon, J. Korean Med. Sci. 25 (1) (2010 Jan) 155–158.
[8] Riaz A. Agha, Mimi R. Borrelli, Reem Farwana, Kiron Koshy, Alexander J. Fowler, Dennis P. O’Regan, For the SCARE Group, The SCARE 2018 statement: updating consensus Surgical Case Report (SCARE) guidelines, Int. J. Surg. 60 (2018) 132–136.
[9] T. Matsuda, T. Matsutani, Y. Tsuchiya, et al., A clinical evaluation of lymphangioma of the large intestine: a case presentation of lymphangiomatosis of the descending colon and a review of 279 Japanese cases, J. Nippon Med. Sch. 68 (2001) 262–265.
[10] P. Rai, R.N. Rao, S.B. Chakraborthy, Caecal lymphangiomatosis: a rare cause of gastrointestinal blood loss, BMJ Case Rep. (2013 Apr 19) 2013.
[11] S.S. Hwang, H.J. Choi, S.Y. Park, Cavernous mesenteric lymphangiomatosis mimicking metastasis in a patient with rectal cancer: a case report, World J. Gastroenterol. 15 (31) (2009 Aug 21) 3947–3949, PMID: 19607144.
[12] M. Ilhan, G. Onur, A. Alibeyoglu, et al., Primary intestinal lymphangiomatosis of the ileum in an adult: the role of surgical approach, J. Surg. Case Rep. (8) (2016 Aug 17) 2016 pii: rjw133.
[13] W.C. Chung, H.K. Kim, J.Y. Yoo, et al., Colonic lymphangiomatosis associated with anemia, World J. Gastroenterol. 14 (37) (2008 Oct 7) 5760–5762.
[14] Y.S. Lee, G.W. Kim, H.J. Cho, et al., Colonic lymphangiomatosis resolved after excisional biopsy, Clin. Endosc. 48 (1) (2015 Jan) 81–84.
[15] Y.F. Fang, L.F. Qiu, Y. Du, et al., Small intestinal hemolympangiomia with bleeding: a case report, World J. Gastroenterol. 18 (17) (2012 May 7) 2145–2146.
[16] A. Dong, L. Zhang, Y. Wang, et al., Abdominal kaposiform hemangioendothelioma associated with lymphangiomatosis involving mesentery and ileum: a case report, J. Korean Med. Sci. 73 (3) (2016 Apr) 8320–8324.
[17] G. Lu, H. Li, Y. Li, Lymphangiomatosis of the sigmoid colon - a rare cause of lower gastrointestinal bleeding: a case report and review of the literature, Oncol. Lett. 13 (1) (2017 Jan) 339–341.
[18] L. Xue, W.G. Guo, J. Hou, et al., Huge lymphangiomatosis of the esophagus, Ann. Thorac. Surg. 93 (6) (2012 Jun) 2048–2051.
[19] G. De Santis, F. Sista, A. Giuliani, et al., Idiopathic intramura hematoma of sigmoid colon. A case report, Ann. Ital. Chir. 82 (5) (2011 Sep-Oct) 395–397.
[20] J.J. de Vries, J.M. Vogten, P.C. de Bruin, et al., Mesenterical lymphangiomatosis causing volvulus and intestinal obstruction, Lymphatic Res. Biol. 5 (4) (2007) 269–273.