Massive Chylous Ascites and Chylothorax Secondary to Chronic Pancreatitis: A Novel Surgical Option

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Chylous ascites is a debilitating condition characterized by milky, triglyceride-rich fluid accumulating in the peritoneum due to disruption of the intraabdominal lymphatic system. Medical management includes low-fat diets, somatostatin analogues, and therapeutic paracentesis, but is unsuccessful in one-third of patients.

Case Report

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The patient had severe gallstone pancreatitis in September 2014, which required an intensive care unit stay at an outside hospital, during which time he underwent a laparoscopic
cholecystectomy. He recovered well, but in April 2015, he developed dyspnea and was found to have a large right-sided pleural effusion. He underwent thoracentesis, with removal of 3 L of chyle. He had no abdominal pain; however, axial imaging demonstrated extensive pancreatic necrosis consistent with his prior pancreatitis; portal vein, superior mesenteric vein, and splenic vein thrombosis; extensive ascites; a possible defect in the right hemidiaphragm; and a large right pleural effusion with associated collapse of the right lower lobe (Fig. 1). There was no pancreatic ductal dilatation to suggest malignancy or obstruction.

Attempts were first made to drain the chyle that had already collected in his peritoneum and pleural cavity, and slow the production of chyle. He was admitted to the MGH in September 2015, where he underwent percutaneous right-sided chest tube placement and was started on a nonfat clear liquid diet, fat-free total parenteral nutrition (TPN), and octreotide. Triglyceride levels in the pleural fluid decreased from 244 to 17 mg/dL, and cytology demonstrated no malignant cells. There was almost complete resolution of his ascites. His chest tube was removed on the day of discharge. However, though he continued a low-sodium, nonfat diet supplemented with medium-chain triglycerides after discharge, the chylothorax and chylous ascites recurred.

In October 2015, he underwent a lymphangiogram, which opacified the cisterna chyli and demonstrated no evidence of a leak. Several transabdominal attempts were made to access the cisterna chyli, in addition to attempts to cannulate the thoracic duct, without success. He was admitted in December 2015 and again underwent chest tube placement, and was started on a nonfat diet, TPN, and octreotide. Again, there was temporary improvement, but the ascites and chylothorax recurred soon after discharge.

The hepatology service suggested that his ascites and effusion may be due in part to cirrhosis and portal hypertension. Transjugular liver biopsy demonstrated no cirrhosis. In March 2016, he underwent portal vein recanalization and stent placement. The accumulation of ascites slowed but continued, and he required paracenteses and thoracenteses with increasing frequency. Repeat axial imaging in August 2016 demonstrated partial in-stent thrombosis. During this time, he developed increasingly large and uncomfortable bilateral inguinal hernias and an umbilical hernia, as well as worsening dyspnea due to repeated reaccumulation of the chylothorax. These symptoms were particularly debilitating given that the patient was recently widowed and the sole provider for two young twin boys. His nutritional and functional status began deteriorating, and his overall health was reaching a point of chronic critical illness, with his serum albumin declining to 2.5 g/dL.

At this point, the decision was made to attempt repair of the inguinal and umbilical hernias along with a novel lymphatic cable flap to prevent reaccumulation of chylous ascites. This flap, proposed by Chen et al, is designed to divert lymph away from the leak by transferring the superior epigastric vessels with surrounding lymphatic fatty tissue into the abdominal cavity and anastomosing them to mesenteric vessels, thereby permitting chyle to drain superiorly through the lymph node chain of the flap and into the thoracic duct (Fig. 2). The patient was admitted in January 2017, 1 week prior to scheduled surgery. He underwent paracentesis with removal of 8.4 L of chylous ascites, along with right chest tube placement. He was started on a nonfat clear liquid diet, octreotide, and TPN to improve his nutritional status prior to surgery. He was taken to the operating room and undergone exploratory laparotomy via a small midline incision, with removal of 6 L of ascites. No chyle leak was identified, even with a heavy cream challenge the night before and via orogastric tube in the operating room. Further exploration of the retroperitoneum was deferred due to the significant inflammatory mass surrounding the pancreas and extensive venous collaterals from portal venous thrombosis. There was no evidence of a diaphragmatic defect.

The plastic surgery team performed the lymphatic cable flap. They dissected the right deep inferior epigastric artery and vein free from the posterior rectus muscle after lateral exposure of the muscle. Next, they raised the vascularized lymph node flap, which consisted of the right deep inferior epigastric artery and vein pedicle and its surrounding fat and lymphatic tissue. The deep inferior epigastric vessels were ligated distally just before they enter the iliac vessels to obtain length for mobility of the flap. The flap, which was technically now based on the superior epigastric vessels as its pedicle (which technically also included the choke vessels connecting the superior epigastric vessels and deep inferior epigastric vessels), was then swung medially and cephalad and placed into the abdomen via an opening in the posterior rectus sheath. The recipient vessels in the mesentry of the flap were ligated and divided to obtain length. Several attempts were made to cannulate the thoracic duct via the cisterna chyli, in addition to attempts to access the cisterna chyli and inferior vena cava with a Rosen catheter and thoracic duct catheter. Transjugular liver biopsy demonstrated no cirrhosis. In March 2016, he underwent portal vein recanalization and stent placement. The accumulation of ascites slowed but continued, and he required paracenteses and thoracenteses with increasing frequency. Repeat axial imaging in August 2016 demonstrated partial in-stent thrombosis.

Fig. 1 Computed tomography images of the (A) abdomen and (B) chest obtained prior to surgical intervention, demonstrating severe ascites, right chylothorax, and right lung collapse.
ileum were prepared for microsurgical anastomosis approximately 10 cm proximal to the ileocolic junction. They then performed microsurgical anastomoses of the artery and vein in standard fashion, and confirmed patency with Doppler signals. At this point, the right inguinal hernia was repaired with a mesh plug through the right groin incision made for the lymphatic flap, and the left inguinal hernia was repaired with a mesh plug through a new left groin incision. Three peritoneal drains were placed, and the umbilical hernia was repaired primarily with closure of the midline incision.

The patient tolerated the procedure well, though required aggressive fluid resuscitation. He recovered in the intensive care unit and quickly weaned off pressor support. He was advanced to a low-fat diet, and received octreotide and TPN.

Fig. 2  Diagram depicting the lymphatic cable flap, which diverts lymph away from the leak by transferring the superior epigastric vessels (including the choke vessels connecting the superior epigastric vessels and deep inferior epigastric vessels) with surrounding fat and lymphatic tissue into the abdominal cavity and anastomosing them to mesenteric vessels, thereby permitting chyle to drain superiorly through the lymph node chain of the flap and into the thoracic duct.
Chylous ascites is a debilitating condition associated with mortality rates as high as 43 to 71%, though rates have been reported to be lower in cases of postoperative chyle leaks. The leakage of chyle prevents the return of protein to the systemic circulation, leading to severe protein-losing enteropathy and malnutrition. Patients also often become immunocompromised due to loss of lymphocyte-rich fluid, in addition to developing hypogammaglobulinemia from protein losses, thus increasing their susceptibility to infections. Respiratory embarrassment may occur from diaphragmatic dysfunction as ascites builds and also from chyle accumulation in the pleural spaces, presumably from natural diaphragmatic channels that may exist.

The management of chylous ascites begins with accurate diagnosis, treatment of the underlying cause, and preservation of the patient’s nutritional status. Diagnosis of chylous ascites is variably reported as an ascitic triglyceride concentration of >110 or >200 mg/dL. Medical management of chylous ascites initially comprises dietary adjustments to reduce the production of chyle. Patients are placed on a low-fat, high-protein diet with supplementation of medium-chain triglycerides (which are absorbed directly into portal venous circulation, bypassing lymphatics).

In addition to dietary modifications, many patients are started on octreotide or other somatostatin analogs, which are believed to decrease portal pressure and reduce intestinal fat absorption. Other therapies have been suggested, such as glue embolization, surgical ligation, and embolization of disrupted lymphatic channels; however, data are limited and results are mixed. Surgical identification and ligation of the source of chyle leak is often unsuccessful due to extensive inflammatory or fibrotic changes and because patients often have multiple chyloperitoneal fistulas. Peritoneovenous shunting has been used in the past but is no longer suggested due to high rates of complications including sepsis, disseminated intravascular coagulation, and early occlusion due to the high viscosity of chyle. Unfortunately, one-third of patients with chylous ascites do not respond to medical therapies. Palliation is achieved by intermittent thoracentesis and paracentesis. Surgical or chemical pleurodesis may be attempted but may be ineffective or result in loculated effusions in the setting of massive chyle production, as in our patient.

In this setting, Chen et al harnessed their experience with using vascularized lymph node flaps for extremity
lymphedema, and created a vascularized lymphatic cable flap based on the superior epigastric vessels and their surrounding fatty, lymphatic-rich tissue. This flap addresses the underlying lymphatic obstruction by creating an extraperitoneal bypass of the damaged lymphatics, enabling lymph and chylous ascites to drain via the superior epigastric vein and surrounding lymphatics in the thoracic wall into the thoracic duct and subclavian vein. Their operative technique consists of identifying the deep inferior epigastric vessels on the posterior aspect of the rectus abdominis muscle, and dissecting the vascular bundle with its surrounding fat and lymphatic tissue down to the vessels’ origin at the external iliac. They next enter the peritoneum, isolate the fourth jejunal artery and vein, and perform end-to-end anastomoses between the deep inferior epigastric vessels and the fourth jejunal vessels. They performed this procedure in three patients with chylous ascites, with resolution of ascites, thoracic reexpansion, normalized serum albumin, and no perioperative complications.

Our patient is an additional compelling example of the success of a lymphatic cable flap for cases of intractable chylous ascites. This technique proposed by Chen et al may be a promising surgical option to offer to patients suffering from this debilitating condition.

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Conflict of Interest
None declared.

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