Percutaneous Transhepatic and Translumbar Sclerotherapy of a Thoracic Duct Cyst: A Case Report

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A 58-year-old male presented with chronic abdominal pain lasting for 5 years and a 15 × 6-cm multicystic thoracic duct cyst with three compartments, located in the right retrocrural and retrocardiac regions from T5/T6 to T12/L1. A transhepatic route was selected to sclerose the middle and lower compartments. Subsequent contrast injection showed minimal contrast passage into the upper compartment through a narrow neck, but the guidewire could not pass into this compartment. The contrast-filled upper compartment was punctured with a 22-g Chiba needle using a translumbar approach under cone-beam computed tomography (CT) guidance and ethanol sclerotherapy was performed. Six-month follow-up CT revealed decreased thoracic duct cyst size (5×3 cm) and no pain. This case illustrates successful percutaneous transhepatic and translumbar sclerotherapy for retrocardiac and retrocrural thoracic duct cysts, which are very difficult to remove surgically.

Abstract

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Keywords

► sclerotherapy
► thoracic duct cyst
► ethanol
► percutaneous

Introduction

Although uncommon, thoracic duct cysts are lymph-filled collections that may arise from any portion of the thoracic duct.1,2 Such cysts may be accompanied with intermittent abdominal pain.1

To accurately diagnose and prevent potential complications such as spontaneous or traumatic cyst rupture, surgery and subsequent histological examination are required.2-4 Here, we describe percutaneous transhepatic and translumbar sclerotherapy of a thoracic duct cyst located in the right retrocrural and retrocardiac regions, which are difficult to access using conventional surgery. To the best of our knowledge, percutaneous sclerotherapy of a thoracic duct cyst in this challenging location has not been previously described.

Case Presentation

A 58-year-old male presented with chronic vague abdominal pain lasting for 5 years. The pain was aggravated by respiration and movement. There was no fever, weight loss, or other gastrointestinal symptoms. He had no history of trauma, previous surgery, or congenital abnormalities. Magnetic resonance imaging (►Fig. 1A) and enhanced computed tomography (CT) (►Fig. 1B) revealed a 15×6 cm, nonenhancing, and multicystic lesion in the right retrocrural and retrocardiac regions, representing a thoracic duct cyst with three compartments and ranging from T5/T6 to T12/L1. There was no ascites, pleural effusion, or lymphedema. After a multidisciplinary discussion with vascular and cardiothoracic teams, it was decided to perform percutaneous sclerotherapy.
The intranodal lymphangiogram was not performed because it was thought that there was no communication with the normal cystic duct due to calcification at the distal part of the cyst. There appeared to be no safe route other than the transhepatic route; therefore, this route was selected to puncture the lower compartment under ultrasound guidance, using a 22-g Chiba needle under local anesthesia. Approximately 50 mL of yellowish chylous fluid was aspirated (Fig. 2). Contrast injection in the lower compartment showed a very narrow passage into the middle compartment, and a stiff 0.035” guidewire barely passed through the connecting region. An 8.5-Fr pigtail catheter was inserted into the middle compartment. Subsequent contrast injection showed minimal contrast passage into the upper compartment through a very narrow neck, but guidewires of various diameters failed to reach into the upper compartment. After excluding any connections with surrounding structures, sclerotherapy was first performed in the middle compartment where the pigtail catheter was located; this was followed by lower compartment sclerotherapy after pulling the catheter so that it was located in the lower compartment. Each compartment was washed three times with 3 mL of 99% ethanol to minimize ethanol dilution, followed by indwelling of 11 mL of 99% ethanol for 7 minutes, while the patient was positioned from the right to the left lateral decubitus to allow the ethanol to contact the entire cyst wall. Next, the ethanol was aspirated and the catheter removed. The upper compartment had a retrocardiac location and was not accessible from the middle compartment. On prone position, the contrast-filled upper compartment was punctured with a 22-g Chiba needle, using a translumbar approach under cone-beam CT guidance. The needle path passed between the aorta and the spine, followed by the placement of a pigtail catheter over a guidewire, and a similar fluid was aspirated and the contrast injection showed no communication with the surrounding structures. Then the 99% alcohol sclerotherapy technique was applied similarly as before for the lower and middle compartments.

The patient’s pain began to subside over time. A 6-month follow-up CT scan revealed a significant decrease in thoracic duct cyst size (5 × 3 cm) (Fig. 3), and the patient’s pain had disappeared, with no complications related to the procedure during the follow-up period.
Discussion

The cyst was considered as a thoracic duct cyst because it was difficult to regard as macrocystic lymphatic malformation because the path of the cyst was very consistent with the thoracic duct. Thoracic duct cysts can be observed anywhere along the course of the thoracic duct, from its origin in the cisterna chyli in the abdomen all the way up to the left part of the neck where the duct is inserted to the junction of the left subclavian vein and internal jugular vein. The pathogenesis of thoracic duct cysts is not clearly known; however, it is assumed to be linked to the congenital or degenerative weakness of the thoracic duct wall, lymphatic flow obstruction, or iatrogenic injuries, or blunt traumas. Cysts cause symptoms by applying pressure to adjacent structures. Symptoms such as dysphagia, dyspnea, abdominal pain, spontaneous chylothorax, and supraclavicular or neck masses may develop.

These cysts are usually recommended to be removed surgically. The surgical procedure should be undertaken to validate the diagnosis by histological examination and to avoid possible complications such as spontaneous or traumatic cyst rupture. The most common complication following surgery is chylothorax that needs reoperation or embolization of the thoracic duct.

Transcatheter sclerotherapy of cystic lesions has become a safe, effective, minimally invasive treatment option, especially in cases where surgery is difficult and/or results in an incomplete resection, such as this case, where the retrocardiac and retrocrural location of the thoracic duct cyst made it unsuitable for surgical removal. Among various sclerosing agents, ethanol remains the most cost-effective agent. Ethanol induces denaturation of proteins, cell death, and inflammatory fibrosis after contact with the cyst wall; however, repeated sessions may be necessary to achieve sufficient inflammatory changes and cyst wall scarring. Therefore, contact with the whole cyst wall is essential during treatment. In adults, the maximum ethanol amount should be limited to 100 mL to avoid alcohol toxicity, and patient monitoring is important for signs of alcohol toxicity during sclerotherapy. Potential complications of alcohol ablation include pain, as well as bleeding, infection, and injury to adjacent organs.

Translumbar access can be utilized when other endovascular or endoluminal approaches appear infeasible or risky. The translumbar approach is performed under both fluoroscopic and cone-beam CT scanning, and the needle path to the target and possible complications like bleeding or erroneous punctures can be monitored.

Conclusion

In the retrocardiac and retrocrural areas, this case illustrates successful percutaneous catheter-directed sclerotherapy for a thoracic duct cyst, which would be difficult to remove surgically.

Conflict of Interest

None.

References