Isolated Mediastinal Lymphangioma: Prenatal Diagnosis and Thoracoscopic Treatment

Varlet F.1, Guye E.1, Varlet M. N.2, Tronchet M.2, Mariat G.3, Chene G.2
1Departments of Pediatric Surgery, Centre Hospitalier Universitaire, Saint-Etienne, France
2Obstetrics and Gynecology, Centre Hospitalier Universitaire, Saint-Etienne, France
3Anesthesiology Centre Hospitalier Universitaire, Saint-Etienne, France
E-mail: chenegautier@yahoo.fr
Received June 12, 2010; accepted July 13, 2010

Abstract

Isolated mediastinal lymphangiomas are uncommon. We report a case of a 14 × 8 mm right paracardiac cyst diagnosed at 20 weeks’ gestation. The prenatal evolution was uneventful and a magnetic resonance imaging at 31 weeks showed the limited extension of the cyst into the anterior mediastinum. At birth, the baby was asymptomatic, but the size of the lesion increased steadily (48 × 29 mm). At 7 months of life, he underwent a thoracoscopic resection of the cyst without intra or postoperative complications. Histological examination showed a lymphangioma. This case is remarkable for its prenatal diagnosis, the thoracoscopic treatment and the 8 years of follow-up without recurrence.

Keywords: Mediastinal Tumor; Mediastinal Lymphangioma; Thoracoscopic Treatment; Prenatal Diagnosis

1. Introduction

Isolated anterior mediastinal lymphangiomas (ML) are uncommon, with an occurrence less than 1% of all the lymphangiomas [1], and most of them are asymptomatic during childhood. They can lead to compression of vital structures, even life-threatening airway compromise. A prenatal diagnosis is now possible, but several pathologies can be evoked when a paracardial cystic lesion is discovered. Once diagnosed, they should be resected, typically by thoracotomy or median sternotomy. We report a case of ML with prenatal diagnosis and thoracoscopic treatment.

2. Case Report

An 8 months old boy presented a 14 × 8 mm an echo-genic right-sided anterior mediastinal cyst, which had been diagnosed at 20 week’s gestation by ultrasonographic examination (Figure 1) and confirmed by magnetic resonance imaging (MRI) at 31 weeks’ gestation (Figure 2). At 34 weeks, the cyst was heterogeneous and measured 27 × 23 mm (Figure 3), but no complications were observed during the pregnancy and the baby was delivered at 37 weeks, weighing 2870 g, without respiratory distress. In the first week of life, sonography and MRI showed a 33 × 26 mm cyst and it was decided to delay resection for a few months. By 7 months, the cyst had enlarged to 48 × 29 mm, without respiratory complications, and the baby was operated on account of this evolution. In the operating room, the patient was placed in left lateral decubitus position and 4 ports were necessary. The cyst was to the right of the thymus, close to the phrenic nerve. The posterior parietal pleura was opened over the cyst from its lower part and easily dissected off the thymus; the dissection was performed cephalad along the right phrenic nerve and the pedicle was ligated close to the superior vena cava. Pathological examination showed a typical lymphangioma. The postoperative course was uneventful and no phrenic palsy or pleural effusion was noted. The patient remains asymptomatic 8 years after surgical excision, without recurrence of the lymphangioma.

3. Discussion

Lymphangiomas are benign hamartomatous tumors of the lymphatic system and less than 1% of all cystic lymphangiomas are purely mediastinal in origin [1]. They constitute about 3% of all mediastinal masses in children [2]. A prenatal diagnosis has already been reported in 8 cases for a single isolated ML [3-10]. It may be suspected when the sonographic examination shows a single or multilocated paracardiac anterior mediastinal cystic
mass. Sometimes, the lymphangioma was described in the posterior mediastinum [4,10,11]. In our case, a fetal MRI was also performed and showed the exact location of the lesion and its extension. As the intracystic septations are not always visible on fetal ultrasound, other diagnoses have to be proposed: pericardial cyst, bronchogenic cyst, thymic cyst, teratoma, esophageal duplication and neurenteric cyst [8,12,13]. A poor outcome is possible with fetal hydrops and hypoplastic lungs, and prenatal thoracocentese may be discussed [3].

For the 8 cases with prenatal diagnosis, the evolution during the pregnancy was variable with 1 spontaneous disappearance, 3 stable lesions and 4 increases of the ML. Among the 4 last cases, 3 fetal hydrops occurred with 1 neonatal death [3], 1 prematurity at 35 weeks’ gestation [7] and the third underwent drainage of the cyst at 24 weeks with success [5]. The ML may be associated with a cervical cyst [9,14] and sometimes with an abdominal extension [15]. A termination of the pregnancy was performed for one fetus pre sented a cervico-mediastino-retroperitoneal lymphangioma [11].

After birth, most ML are not diagnosed because they are asymptomatic; among the patients presenting symptoms, the most common are respiratory, cough or stridor by extrinsic compression of the airway as a result of hemorrhage or inflammation, sometimes with acute respiratory distress [15,16]. Less common symptoms are dysphagia, superior vena cava syndrome, dysrythmia, Horner’s syndrome or phrenic nerve paresis [17]; a fatal outcome in a 12 year-old boy has been described [18]. Chest radiograph may show an anterior mediastinal mass and sonography may establish its cystic aspect with septations; however, computerized tomography and especially MRI are useful for the diagnosis and the extension of the lesion [14-16]. Calcifications have been described in ML, although this is more characteristic of teratomas [19].

Mediastinal lymphangiomas, as other mediastinal masses, must be removed to avoid complications. Among the
Figure 3. Sonography at 34 weeks' gestation with a 27 × 23 mm heterogeneous right anterior mediastinal cyst.

4. Conclusions

Isolated mediastinal lymphangiomas must be suspected when a cystic mass is noted on prenatal sonography in the anterior mediastinum, differential diagnosis including especially pericardial cyst or thymic cyst. The evolution is variable from spontaneous disappearance to fetal hydrops or life-threatening complications. A thoracoscopic approach is now possible, even in infant.

Author Disclosure Statement: there is no conflict of interest for each author.

5. References


