Recurrent thoracic duplication cyst with associated mediastinal gas

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Mediastinal cysts are not uncommon in the pediatric age group. Presentation varies from an abnormality found on routine chest radiograph to severe respiratory distress and even respiratory failure. Presentation depends on the age of the patient, the location of the lesion, the extent and the size of the mass, and what structures are involved. The case of a six-year-old boy who presented with recurrence of a mediastinal mass associated with gas two years after surgical removal of an infected esophageal duplication cyst is described. No connection between the cyst and the esophagus to explain the presence of gas was documented. This appears to be the first reported case of esophageal duplication cyst associated with mediastinal gas.

Key Words: Children, Mediastinal cysts

CASE PRESENTATION

A four-year-old boy developed cough and fever followed by respiratory distress, which progressed over few days to severe respiratory failure requiring admission to the pediatric intensive care unit of the Winnipeg Children’s Hospital in June 1994. He was found to have a mediastinal mass on chest radiograph (Figure 1) that was further delineated by an esophagogram and chest computed tomography (CT) scan (Figure 2). The mass was compressing the trachea and accounted for the severe respiratory distress and carbon dioxide retention. He was operated on within three days of admission and found to have an infected duplication cyst that was drained and excised. Microscopic examination of the cyst showed...
stratified squamous epithelium, and the lumen contained a large number of polymorphonuclear cells. Culture from the purulent cyst fluid grew *Bacteroides fragilis*. Postoperatively he was treated with cefoxitin and gentamycin. He recovered uneventfully from the surgery and was discharged home after five days.

Before this presentation there was a history of noisy breathing with colds. He also had a mild right hemiparesis that was noted at about age one year when he started walking, but which had remained stable since that time with no progression. The etiology of the hemiparesis remained obscure, and no further investigations, such as imaging of the brain or spinal cord, were completed. Intellectual function was normal. There was no history of foreign body aspiration or ingestion, and no history of trauma to the chest. He was born at full term with normal spontaneous vaginal delivery, and Apgar score was three and nine at 1 and 5 mins, respectively. He had jitteriness noted after birth, which was attributed to withdrawal symptoms secondary to the anticonvulsive medication that the mother was taking during pregnancy; blood glucose concentration was normal.

Two years later (June 1996) at age six years he again presented at the Winnipeg Children’s Hospital with a six-month history of shortness of breath with exercise, resolving without treatment after 30 mins of rest. There was no history of wheeze or nocturnal cough. Physical examination was normal except for mild right-sided weakness.

Pulmonary function tests were completed and were normal, with no evidence of airway obstruction: forced vital capacity (FVC) 1.56 L (89% predicted), forced expiratory volume in 1 s (FEV₁) 1.41 L (89% predicted), FEV₁/FVC 90%, forced expiratory flow 25% to 75% 1.73 L/s (87% predicted). Because of the history of the excised mediastinal duplication cyst two years previously, a chest radiograph was completed that showed a superior mediastinal mass with lateral deviation and compression of the trachea to the right (Figure 3). A chest CT scan showed a soft tissue density situated between the esophagus and trachea at the thoracic inlet, similar to that noted two years earlier. In addition, a small amount of gas was seen in the mediastinum that had not been present on the earlier CT scans (Figure 4). A gastrografin swallow followed by a barium swallow in the supine position did not demonstrate any leak from the esophagus into the cyst.

A 6 min run at 6 km/h was completed at a grade of 12°, and no fall in FEV₁ dyspnea, wheeze or stridor was detected. Because he was now asymptomatic, no further therapy was undertaken. At follow-up five months later (October 1996), he denied any further symptoms, and chest radiograph revealed slight deviation of the trachea to the right and minimal soft tissue density in the superior mediastinum but no gas evident. The patient has remained asymptomatic, and no further clinical or radiological studies have been completed.

**DISCUSSION**

The nomenclature of duplication cysts of the gastrointestinal tract remain confusing with various terms used, such as enteric, enterogenous, and gastroenteric cyst or enterocystoma (2). In studying mediastinal cysts Pokorny and Sherman (6) and Snyder et al (7) reported that bronchogenic cysts were more common than esophageal duplication cysts. In contrast, Haller et al (8) found that esophageal duplication cysts were more common than bronchogenic cysts. In any case, the appropriate management of these benign cysts is surgical excision. Symptomatology depends on the degree of compression of neighbouring structures by the mass and can
range from no symptoms to severe respiratory distress, stridor, wheezing, dysphagia, vomiting and regurgitation. Chest pain may be reported by older children. Age at the time of presentation varies. Symptoms may start as early as the newborn period (3,4,7) or may be delayed until adulthood (5,9), but most esophageal duplication cysts are discovered in childhood. Vertebral anomalies may be associated with duplication cysts, especially those in the posterior mediastinum (1,2,10). Mediastinal masses may also present as diagnostic dilemmas, and operation may be delayed (7) unless there is high index of suspicion.

In our case a four-year-old child presented with severe respiratory distress secondary to an esophageal duplication cyst. What is puzzling and novel is the recurrence of the mass and the presence of mediastinal gas two years after surgical excision. Radiological studies did not demonstrate an esophageal leak that would explain the presence of gas. Stringel et al (11) reported esophageal duplication cyst containing a foreign body, demonstrated by CT scan of the chest, and found the foreign body to be a bingo chip at time of operation. They could not demonstrate a communication between the cyst and the esophagus by barium esophagography or at time of surgery. Obviously, a communication between the esophagus and the cyst must have existed at some stage and presumably closed spontaneously. In our case, although there was no foreign body, the situation may be similar in that gas entered the cyst. Another possible explanation is that the gas was introduced at the time of the initial surgery.

Although thought to have been completely excised during the original operation, some remnant of the cyst may have remained, and this could explain the recurrence of the mediastinal mass. To our knowledge, this is the first reported case of recurrent esophageal duplication cyst in association with mediastinal gas.

REFERENCES

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