Benign teratoma in the left pleural space with amylase in the pleural fluid

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CASE REPORT


The case of a 28-year-old man who experienced repeated episodes of pleuritic chest pain is reported. Eventual discovery of amylase in the pleural fluid and a supradiaphragmatic mass on computed tomography led to exploratory thoracotomy. A benign cystic teratoma was discovered in the pleural space overlying the left hemidiaphragm and connected by a peduncle to the thymus.

Key Words: Benign teratoma, Pleural amylase Subpulmonary tumour

Tératome bénin dans l’espace pleural gauche et présence d’amylase dans le liquide pleural

RÉSUMÉ : Le cas d’un homme âgé de 28 ans qui souffrait d’épisodes répétés de douleur thoracique d’origine pleurétique est rapporté. La découverte d’amylase dans le liquide pleural qui s’ensuivit ainsi qu’une masse supradiaphragmatique révélée par tomodensitométrie ont nécessité une thoracotomie exploratoire. On a découvert un tératome kystique bénin dans l’espace pleural recouvrant l’hémidiaphragme gauche et relié au thymus par un pédoncule.

A 28-year-old Caucasian man from western Canada presented to a tertiary care hospital with sudden onset of left-sided chest pain concentrated in the lower anterior region. His pain had a classical pleuritic quality. There were no other respiratory or systemic symptoms. He was an infrequent binge drinker and smoked a half pack of cigarettes daily. He had been drinking for several days before this episode of chest pain.

Physical examination revealed an afebrile, fit male in significant distress. He had shallow breathing and guarded movement as he ‘splinted’ his left side. The left chest wall was tender in the lower anterolateral region with dullness to percussion at the left base. The remainder of the physical examination was unremarkable.

Laboratory investigations included a peripheral leukocyte count of $13.5 \times 10^9/L$ with a marked granulocytosis and no bands. Arterial blood gases were within normal limits. Chest radiograph demonstrated a density in the left lower lung field (Figure 1), interpreted as being either within or below the left lower lobe. A pleural effusion of moderate size was con-
firmed on a lateral decubitus film. Thoracentesis revealed straw-coloured, turbid fluid with pH of 7.42, leukocyte count 44x10^9/L (differential 79% polymorphs, 21% monocytes), protein 41 g/L (serum protein 63 g/L, normal range 60 to 84 g/L), glucose 4 mmol/L (serum glucose 4.9 mmol/L, normal range 3.6 to 6.1 mmol/L), lactate dehydrogenase (LDH) 465 U/L (serum LDH 110 U/L, normal range 71 to 183 U/L) and amylase 927 U/L (serum amylase 74 U/L, normal range 25 to 115 U/L). Amylase isoenzyme fractionation was not available.

In the preceding year this unfortunate man had experienced four similar episodes, each involving admission to a rural hospital. The clinical and radiological picture was similar on each occasion. Thoracentesis was not carried out on any of these admissions. Each time the patient was started on intravenous antibiotics with rapid resolution of symptoms in the first 48 h and early discharge without follow-up. There was never a clear diagnosis of an infectious process. Previous radioisotope lung scan for pulmonary embolism had been indeterminate.

Further investigations were carried out on admission to hospital. A computed tomographic (CT) scan of the chest demonstrated a small left-sided pleural effusion and a subpulmonic density interpreted as representing either loculated fluid or tumour (Figure 2). Flexible bronchoscopy failed to demonstrate an endobronchial lesion. Gastroesophagogram and gastroscopy were normal. Cultures of blood, pleural fluid and bronchial washings failed to grow pathogenic bacteria or mycobacteria.

The patient was started on clindamycin empirically on admission. Symptoms resolved completely within 48 h and antibiotics were discontinued. A chest radiograph three days after admission demonstrated reduction in the size of the density and effusion (Figure 3).

Since the diagnosis remained unclear, exploratory thoracotomy was performed. At surgery, a tumour was found overlying the left hemidiaphragm within the pleural space. A long pedunculated portion of the mass was dissected to the region of the left pulmonary artery and thymus. Following excision of the tumour, the patient remained asymptomatic during follow-up over nine months.
Histopathology of the tumour revealed this to be a benign cystic teratoma measuring 15 cm in greatest diameter. There were multiple cysts which were lined with squamous and bronchial tissue. Well differentiated bowel and pancreatic tissue were identified, including exocrine and endocrine cells (Figure 4). There was a suggestion of residual thymic tissue and no evidence of malignancy.

**DISCUSSION**

Germ cell neoplasms are the third most common mediastinal tumour, following thymic lesions and lymphoma (1). They account for approximately 8% of all tumours of the mediastinum (2). Approximately 80% of germ cell tumours are benign teratomas (3).

The literature describes a varied clinical presentation for benign teratomas. In one series, the diagnosis was made at a mean age of 28 years, with a range of seven months to 65 years (4). In another study, up to 62% of patients were asymptomatic, with the tumour being discovered on routine chest radiograph (5). The most common symptoms were pain in the chest, back and shoulder, followed less commonly by dyspnea and cough (4).

This case demonstrated several unique characteristics. First, the subpulmonic location of this tumour has never been reported in the literature, to the best of our knowledge. The majority of mediastinal teratomas are located in the anterior compartment, with 3% to 8% arising from the posterior compartment (2,6). One series described extension into the right or left hemithorax in 48% of cases (4). Our case demonstrated an extreme migration to the subpulmonic region of the left pleural space.

Second, our patient had a unique clinical presentation of recurrent chest pain with transient pleural effusion. Diagnosis was delayed due to rapid resolution of symptoms and a near-normal chest radiograph between episodes (Figure 3).

Third, our case demonstrated a high amylase level in pleural fluid. Rare reports of amylase in the pleural fluid associated with benign teratomas may be found in the literature (7-9). Histopathology in those cases demonstrated either pancreatic (7,8) or salivary (9) glandular tissue. We suspect that the amylase discovered in our patient in the pleural fluid originated in pancreatic exocrine tissue proven to be contained within the tumour (Figure 4). Amylase levels from cystic fluid and isoenzyme fractionation would have been helpful, but were unavailable.

It is interesting to speculate what pathophysiological process may have contributed to our patient’s clinical course. Sommerlad et al (7) reported a cystic teratoma with amylase activity and surrounding inflammation. In that report, it was suggested that leakage of amylase and other digestive en-
zymes from a cyst might contribute to inflammation. In our patient, intermittent leakage from a cyst to the pleural space seems to be the most plausible explanation. We speculate that the patient’s ‘binge drinking’ may have precipitated an equivalent of pancreatitis within the teratoma, leading to the clinical events. The episodes of chest pain appear to have been preceded by excessive alcohol intake, and this may have caused inflammation, pain, pleural effusion and amylase excretion. Another plausible explanation could be intermittent torsion and ischemia, although there was no histopathological evidence for this.

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REFERENCES
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