Hemoptysis caused by an endobronchial lipoma

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Benign tumors of the lung and endobronchial tree are uncommon and need to be differentiated from malignant lesions. Among benign lesions, endobronchial lipoma (EL) is extremely uncommon, accounting for only 0.1% to 0.5% of all bronchial tumors. We present a case in which an EL caused hemoptysis, bronchial obstruction, and distal pulmonary fibrosis. Surgical resection with a right lower lobectomy was accomplished.

Clinical Summary
A 62-year-old man who smoked heavily had hemoptysis sporadically for 3 years. Six months before admission he had cough and shortness of breath, which was treated as asthma. Finally, he was admitted to our department because of increasing hemoptysis with fever and purulent secretions.

A chest radiograph demonstrated a right lower pulmonary infiltrate. Thoracic computed tomographic scan showed damage to the right lower lobe and a fat density mass in the right lower bronchus (Figure 1). A bronchoscopic study confirmed a smooth round polypoid endobronchial lesion in the right lower bronchus (Figure 2); histologic and cytologic studies (biopsy and brushing) found no neoplastic cells. Bronchoscopic resection was not considered feasible because it was impossible to clearly identify the tumor’s endobronchial origin. A right thoracotomy was thus performed.

At bronchotomy, a yellow smooth fatty lesion occluding the lower bronchus was found and a frozen section confirmed no neoplastic tissue. Pus was found in the distal pulmonary parenchyma and lobectomy was necessary. A less extensive resection was at risk for postoperative severe infective complications.

Pathologic examination showed mature adipose tissue growing in the submucosal layer, completely covered with ciliate respiratory epithelium. A diagnosis of endobronchial lipoma was made. The resected lung tissue showed important inflammatory infiltrates, pus, and fibrotic areas.

The postoperative course was characterized by fever and purulent bronchial secretions for 6 days, controlled with broad-spectrum antibiotics.

Discussion
We present an uncommon case of EL in which surgical resection of the tumor was accomplished with a right lower lobectomy because of the pulmonary damage following long-term obstructive pneumonia.

Figure 1. Thoracic computed tomographic scan (particular) with a right lower endobronchial lesion.

Figure 2. Bronchoscopic view of the lesion.
ELs are rare benign tumors of the bronchial tree; only 80 cases have been previously reported in the English literature. Most ELs arise in the submucosal layer of the bronchus; fatty tissue is normally found in both the mucosa and the tissue external to the cartilage. Almost two thirds of the tumors occur in the right side, and they are usually located in the main-stem or lobar bronchi.1-3 Smoking and obesity are significant risk factors for EL development.4 Cough, hemoptysis, and shortness of breath are common symptoms,4,5 and patients are often misdiagnosed and treated for asthma for many years, as was our patient.

Owing to the endobronchial tumor growth, irreversible changes in distal pulmonary parenchyma often occur: atelectasis, overinfection, pneumonia, and fibrosis are frequently observed. Bronchoscopy is necessary to identify and locate the lesion and to differentiate it from a malignant tumor. Differential pathologic diagnosis between bronchial carcinoid and lipoma is often difficult because of the paucity of tissue sampling at biopsy. The tumor’s surface is usually paler in the lipoma and more friable in the carcinoid. EL sometimes coexists with lung cancer, especially in patients who have smoked heavily.1

Bronchoscopic resection should be considered the first choice of treatment for EL; Muraoka and associates2 report that no patient presented with bronchial tumor recurrence after EL bronchoscopic resection. Tumor surgical resection is preferred in some cases: (1) extrabronchial growth or subpleural lipomatous disease; (2) expected technical difficulties during the bronchoscopic resection; (3) possible malignant tumor coexisting with EL; and (4) peripheral irreversible lung parenchyma changes after long-term atelectasis or pneumonia.

In conclusion, EL is a rare benign endobronchial tumor that can be safely resected by a bronchoscopic procedure. Surgery is recommended when the distal lung parenchyma is damaged by an obstructive pneumonia or fibrosis.

References

A new type of diaphragmatic hernia: Anterolateral hernia

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Heterogeneous types of acquired hernias in the diaphragm have been described in the literature: the hiatal hernia through the esophageal foramen and the posterolateral hernia through the Henle costolumbar orifice or anterior foramen of Larrey-Morgagni hernia (retroxyphoid hernias).1,2 To our knowledge, no anterolateral hernia has been reported. We describe the case of a hernia between the eighth and ninth anterior costochondral insertion of the muscular fibers of the diaphragm.

Clinical Summary
A 76-year-old man had a medical history of diabetes mellitus, arterial hypertension, a bilateral inguinal hernia (operated), and a 2/3 gastrectomy for gastric ulcer (50 years ago). In 1999, a chest x-ray showed an air-filled opacity in the left thoracic base. No further investigations were done. In September of 2005, the patient experienced a blocking respiration symptom in the left hemithorax, especially after meals and occasionally associated with vomiting. The chest x-ray showed colic haustration at the left thoracic base. A gastroesophageal endoscopy showed no anomaly. The cardiac sphincter was 40 cm from the incisor teeth, and the gastric stump and duodenal-gastric anastomosis were healthy. The barium esophagography showed a normal cardiac region. The chest computed tomography scan demonstrated a colic ascent in the left thorax and the small intestine through an anterolateral hole of the diaphragm (Figure 1).

Surgery was performed via a low left posterolateral thoracotomy. We found the peritoneal sac containing the digestive loops (Figure 2). After clearly identifying the edges, we resected and closed the sac. These edges corresponded to the muscular fibers of the diaphragm inserted at the eighth and ninth chondrocostal cartilage. The hole measured 10 cm in the anteroposterior diameter, was located far from the retroxyphoid region, and ended at the central tendon. The closure was done with nonabsorbable sutures. The