This report, emphasizing a possible complication of en bloc resections involving the vertebral bodies, especially after chemotherapy, radiotherapy, or both, might prompt the thoracic surgeon to evaluate a possible solution of continuity of the dura mater during extended lung resections.

References

Rupture of a chronic expanding hematoma of the thorax into lung parenchyma

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C hronic expanding hematoma of the thorax is a rare entity and usually identified with mediastinal compression or chest wall protrusion in patients with a history of thoracotomy, tuberculous pleurisy, or thoracic trauma. We present a patient with chronic expanding hematoma of the thorax who had a massive hemoptysis caused by rupture into lung parenchyma.

Clinical Summary
A 59-year-old woman with a history of thoracoplasty, which included resection of the 9th to 11th ribs for tuberculous empyema 24 years earlier, presented with a massive hemoptysis and dyspnea. She had had a chest wall protrusion pointed out for 4 months. She was brought to our hospital by ambulance, and chest roentgenography showed bilateral lung opacity. The chest roentgenogram 2 months before showed a right intrathoracic mass-like shadow with pleural calcification (Figure 1). Because of the repetitive hemoptysis and the progression of cyanosis, emergency tracheal intubation was done to secure the airway. The bleeding was suspected from the right middle or lower lobe, and the right bronchus intermedius was blocked with a balloon within an endotracheal tube (Univent; Fuji System Corp, Tokyo, Japan). Computed tomography showed a heterogenous mass in the right thorax protruding into the chest wall and the right lung (Figure 2). The left lung was also opacified with infiltrate, indicating contralateral spread of the bleeding. Urgent bronchial arteriography was performed; the dilated bronchial artery and intercostal arteries were embolized with platinum coils and gelatin (Spongel; Yamanouchi Co, Tokyo, Japan). Despite the arterial embolization, hemostasis was not obtained immediately, and the balloon was kept inflated until the next day. Six days later, when the left infiltrate improved, the patient underwent a right pneumonectomy through a median sternotomy. Fresh hematoma was identified within the chronic organized hematoma in the thorax and penetrated into the right lung parenchyma. All the hematoma was extirpated along with pneumonectomy. Operating time was 315 minutes, and blood loss was 2077 mL.

Postoperatively, the patient required ventilatory support for respiratory failure. After 6 days of ventilatory support, she was weaned from the respirator. After respiratory rehabilitation, she was discharged from the hospital on postoperative day 43. She has been doing well and has been free from any symptoms of the disease for 2.5 years.

Discussion
Chronic expanding hematoma of the thorax is a rare entity. Affected patients typically have a history of medical or surgical therapy for tuberculosis. Since Iuchi and associates1 reported 5
cases of empyema without pus as organized empyema in 1988, most reports of chronic expanding hematoma of the thorax have been from Japan.2 Recent literature from the United States3,4 indicates a certain incidence of chronic expanding hematoma of the thorax in western countries.

Despite the rare entity arising from the thorax, chronic expanding hematomas occur in many locations. Although most hematomas resolve spontaneously, a few persist for long periods as slowly expanding space-occupying masses.5 The pathogenesis of chronic expanding hematoma is poorly understood. Labadie and Glover6 theorized that this self-perpetuating expanding process is due to the irritant effects of blood and its breakdown products, which cause repeated exudation or bleeding from capillaries in the granulation tissue. Clinically, intermittent episodes of bleeding, probably caused by respiratory motion and coughing, might cause a progressive increase in the volume of hematoma.

The presenting symptom is usually dyspnea related to lung compression or a slowly growing chest wall mass, both of which develop even 30 years or more after thoracotomy for tuberculosis or trauma. These lesions are frequently thought to represent soft tissue sarcoma or other malignant tumors. Surgical procedure has been controversial. Palliative procedures, such as curetting or removal of the inner substance without whole-capsule excision, are reported as a result of the difficulty of the procedure or for fear of

Figure 1. A, Chest roentgenogram on admission showing bilateral opacity of the lung. B, Chest roentgenogram 2 months earlier showing a mass-like lesion in the right thorax with pleural calcification and chest deformity.

Figure 2. Chest computed tomography on admission demonstrating a heterogeneous mass in the right thorax protruding into the chest wall and right lung.
possible complications. However, incomplete resection of the hematoma resulted in uncontrollable bleeding from the subcapsular lesion or recurrence of hematoma within several years in some publications. Complete extirpation, including the capsule, would be desired for cure.

In our patient acute massive hemoptysis was treated with emergency tracheal intubation, balloon occlusion of the bleeding bronchus, and bronchial artery embolization. In the literature neither massive hemoptysis nor emergency management for chronic expanding hematoma of the thorax has been reported. Because the bleeding spread to the contralateral lung, the operation was intentionally delayed until the remaining lung function recovered. Thereafter, the patient successfully underwent a pneumonectomy with extirpation of the hematoma. Division of the hilum first and sequential distal dissection of the lung and hematoma through a median sternotomy might have worked well for bleeding control.

**References**


**Thyroid metastasis after resection of atypical bronchial carcinoid**

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Bronchial carcinoids (BCs) are rare and constitute less than 2% of pulmonary tumors. They are characterized by slow, mainly endobronchial growth, with infrequent regional lymph node involvement or distant metastases.

Atypical carcinoids are part of the spectrum of neuroendocrine bronchopulmonary tumors, according to the 1999 World Health Organization–International Association for the Study of Lung Cancer (WHO-IASLC) Lung Tumor classification. They present 2 low-grade (typical and atypical carcinoids) and 2 high-grade malignant varieties (large cell neuroendocrine carcinoma and small cell lung carcinoma), the latter characterized by a high tendency toward mediastinal and distant metastatic spread.

Among low-grade malignant tumors, atypical carcinoids show a more aggressive biologic behavior than typical carcinoids: mediastinal lymph node metastases occur at presentation in about 15% of cases. Distant metastases are generally in the liver and in the bone.

We present a case of a woman in whom thyroid metastasis occurred 30 months after the resection of an atypical BC.

**Clinical Summary**

A 53-year-old white woman was referred to us in June 1999 because of the presence of a large pulmonary mass (7 × 3 cm in size) in the right lower lobe. The lesion had been detected with chest radiography performed in July 1998, and it was 5 × 2 cm in size, but at that time, the patient refused the intervention. She eventually agreed to be operated on because of the progression of the lesion and the associated cough and dyspnea. A preoperative transthoracic fine-needle aspiration biopsy was performed, and neoplastic cells with neuroendocrine features were observed. 

111-In-DTPA-Pentetreotide scintigraphy (Octreoscan) was performed for a correct preoperative assessment, resulting in an elective uptake in correspondence of the pulmonary lesion with no other abnormal uptakes. Chromogranin A and neuron-specific enolase serum levels were performed and increased: 197 ng/mL (normal values, 20-100 ng/mL) and 62 ng/mL (normal values, <12.5 ng/mL), respectively.

The patient underwent a right pneumonectomy with systemic lymphadenectomy: a more conservative intervention was not possible because of the tumor size. A residual tumor was left on the inferior pulmonary vein stump.

Grossly, an 8-cm mass with poorly demarcated borders infiltrating the mediastinal pleura and the adipose mediastinal tissue was found. The cut surface was tan-yellow, and a brown lymph node was adherent and directly infiltrated by the tumor. Microscopically, the tumor was not encapsulated, and it had a uniform organoid growth pattern composed of nests and sheets of cuboidal and eosinophilic cells. Rosette-like structures were also

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