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# A Case of Invasive Thymoma

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A 52 year-old man with a past medical history of an isolated seizure presented to the Veterans Affairs hospital complaining of weakness which was most prominent in the face and upper extremities. About 6 months prior to this admission he developed intermittent episodes of weakness when chewing and swallowing. He would often have to use his hands to close his jaw when eating or talking. He noticed that his voice had developed a nasal quality but he did not have slurred speech. He denied drooling, ptosis, cramping or muscle twitches. He was seen 6 months ago at another VA hospital where a neurologic workup was done. This included a cranial nerve EMG positive for denervation of cranial nerves, R arm and T- and L-spine. MRI and CT scans could not be done because the patient aspirated when supine. He was given a presumptive diagnosis of motor neuron disease, specifically bulbar ALS. At that time he was told he would need a PEG-feeding tube placed due to significant weight loss. The patient refused the feeding tube and did not follow up until again until 6 months later when the symptoms had worsened.

The patient's past medical history included an isolated seizure about 8 years ago. He had been taking phenytoin but it was discontinued several months ago after periodic negative EEG's. His family history was significant for a transient ischemic attack in his mother. There was no other family history of neurologic or systemic disorder. He had a 35 pack-year history of tobacco smoking. He also occasionally smoked marijuana. He quit drinking beer about a year ago and said he never drank heavily. The patient had a remote history of intravenous heroin use and was eager to undergo testing for HIV. He was employed as a roofer but quit recently because he was having to use his hands to hold his head up while working. There were no known occupational exposure to asbestos or chemicals.

The patient's heart, lung and abdominal exams were normal. On neurologic and musculoskeletal exam, the cranial nerves were intact. He had no tongue or facial fasciculations but did have bilateral temporal and masseter wasting. He had full strength throughout, but had slightly diminished vibratory sensation of the feet. There were no peripheral fasciculations or wasting. Deep tendon reflexes were +2/4 throughout. Cerebellar and Romberg exams were normal and there was no Babinski sign. The diagnosois of ALS was questioned and the patient underwent further neurologic workup.

His metabolic panel, complete blood count and thyroid function tests were within normal limits, as were the B12 and folate levels. His HIV test was negative, while Hepatitis C antibody was positive with an undetectable viral load. Albumin was low at 2.7. The patient was able to lie supine for a brain MRI. The MRI and a chest radiograph were unremarkable. He underwent a videofluoroscopic swallowing study which revealed moderate oropharyngeal dysphagia with increased vallecular residue and impaired elevation of the larynx. There was no esophageal dysfunction. Pulmonary function tests showed a decrease in forced inspiratory flow @ 39%.

He had an acetylcholine receptor antibody level was 3.62 nmol/L (normal up to 0.4). His SPEP was near-normal with a slight polyclonal elevation (IgM and IgG). The IgA was slightly low. A noncontrast chest CT showed a lobulated mass in the anterior mediastinum measuring 5 x 3.8 cm, extending from the sternum and abutting the SVC and aorta. (see Figure 1) The mass contained few calcifications. A lesion was seen in the anterior mediastinum just to the right of the sternum. It was pleural-based and measured 2 x 0.8cm. Another pleuralbased lesion was seen in the right posterolateral chest. Finally, there were several smaller, nodular soft tissue lesions of similar density within the right lung. A CTguided biopsy was done on the anterior mediastinal lesion. It revealed a WHO Type B3 epithelial thymoma. (See Table 2).

While reviewing the CT scan for a possible substernal thymectomy, surgery requested that interventional radiology biopsy of one of the pleural lesions. The biopsy of the pleural-based posterolateral chest lesion (see Figure 2) was also consistent with epithelial thymoma. CT scans of the neck, abdomen and pelvis showed no evidence of other metastases. Since thymectomy would not be curative for a metastatic lesion the patient was referred to oncology where he was started on chemotherapy with

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cisplatin, cyclophosphamide and adriamycin. He was also started on pyridostigmine for control of the myasthenia gravis. Chemotherapy was uncomplicated but a few weeks later the patient was admitted for community-acquired pneumonia and diarrhea. He was started on the appropriate antibiotic regimen and his pyridostigmine was withheld for diarrhea. During this acute illness the patient was so profoundly weak he had to constantly lean forward in bed to allow saliva to drain. Just as the decision was made to move the patient to an intensive care unit and electively intubate him for airway protection, his condition began to improve. He was once again able to swallow and within 3 days he had regained almost full strength. At this point he was a few weeks out from chemotherapy and a follow-up chest CT showed a marked decrease in the size of the patient's mediastinal mass (and pleural/lung masses).

### Discussion

This patient was initially diagnosed with motor neuron disease, specifically bulbar amyotrophic lateral sclerosis. It is difficult to tell why the diagnosis of myasthenia gravis was not made initially. The patient had less than optimal follow up and the clinical picture was further complicated because the patient's report of symptom onset at the time of a neck injury several years before. The physicians who first saw him must have wondered whether his fall from a ladder was a cause or result of his symptoms. Furthermore, the patient was taking his anticonvulsant medication eratically and his desire to taper off his anticonvulsant was a priority.

Thymomas are the most common primary anterior mediastinal tumor and account for 15-20% of all mediastinal masses1. Thymomas are derived from epithelial thymus cells with varying degrees of atypia, along with normal lymphocytes. These tumors should be distinguished from thymomas involving lymphomas, germ cell tumors, carcinoid tumors a small cell cancer. Thymomas are associated with many paraneoplastice syndromes, the most common of which is myasthenia gravis, which is seen in about 30% of patients with thymomas. Other paraneoplastic syndromes include pure red cell aplasia, acquired hypogamaglobulinemia and connective tissue disorders such as Sjogren's syndrome, myocarditis, dermatomyositis, SLE and rheumatoid arthritis.

About 15% of myasthenia gravis patients have thymomas. This has resulted in widespread screening for thymoma in MG patients. In other cases, thymomas are usually picked up by routine CXR or when patients present with symptoms secondary to mass effects, such as coughing, shortness of breath or chest pain.

There are several classification systems for thymomas. The Lattes/Bernatz, Rosai/Levine and Muller-Mermelink are based on microscopic appearance alone while the more commonly used Masaoka system is based on degree of invasion. A fourth classification system proposed by the World Health Organization was used with this patient. There is controversy regarding the most useful classification system in terms of predicting survival. Several of the studies used to develop these classification systems involved small numbers of patients. From a practical standpoint, the degree of invasiveness.

While it has often been reported that myasthenia gravis portends a worse prognosis in patients with thymoma, a more recent study patients with thymomas and myasthenia gravis appear to have a significantly better overall prognosis that those without myasthenia symptoms. It would appear that patients with myasthenia symptoms present earlier in the course of disease and the thymoma was caught at an earlier stage. However, one series has shown that this is not the case. Also, patients with myasthenia gravis are more likely to follow-up with their doctors and get better care for all of their medical problems. It has been shown that most patients with thymomas (invasive or noninvasive) tend to die of medical problems other than the thymoma. A third possible explanation was myasthenia patients seem to do better may be because some of them get treated with steroids, which can lead to involution of thymomas. Interestingly, while myasthenia appears to be a good independent predictor of survival, resection or "cure" of thymomas typically does not affect the myasthenia symptoms. The most significant independent predictor of survival is surgical resectability at initial presentation. Other independent risk factors for improved survival are (Continued from previous page)

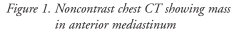
absence of advanced pathologic stage (per the Lattes/Bernatz classification) and age < 57 yrs.<sup>1</sup>

Treatment for thymoma is generally involves surgical resection, with a recurrence rate of about 2% for encapsulated tumors and about 20-40% for invasive lesions. Preoperative radiation therapy does not generally improve survival, but adjuvant postoperative therapy is considered the standard of care for completely or incompletely resected stage III or stage IV thymomas. For metastatic thymoma, this is generally not possible chemotherapy should be considered. Candidates for chemotherapy include about 1/3 of patients with invasive thymoma and all patients with stage IV disease. The usual regimen is cisplatin, doxorubicin and cyclophosphamide (PAC). Chemotherapy is followed by radiation and in one study this resulted in a 52% five year survival rate.<sup>2.3</sup>

In summary, thymoma should be considered in any patient with myasthenia gravis. The most important predictor of survival is surgical respectability at initial presentation. Finally, the presence myasthenia gravis is a predictor of improved survival in thymoma.

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#### Table 1. Masaoka Classification

Stage	Extent of disease
Ι	Totally encapsulated
II	Capsular invasion and/or invasion into surrounding fat or pleura
III	Invasion into organs (pericardium, lung, great vessels)
Iva	Pleural or pericardial implants
IVb	Hematogenous metastases

### Table 2. WHO Classification

Histological Types	Characterization
WHO Classification ofThymic Epithelial Tumors(1999)	This system is based upon the morphology of the epithelial cells as well as the lymphocyte to epithelial cell ratio
Туре А	Homogeneous population of neoplastic epithelial cells having spindle/oval shape, lacking nuclear atypia, and accompanied by few or no non-neoplastic lymphocytes
Type AB	Foci having the features of type A admixed with foci rich in lymphocytes; the segregation of the two patterns can be sharp and distinct
Type B1	Resembles the normal functional thymus by combininglarge expanses with an appearance practically indistinguishable from normal thymic cortex with areas resembling thymic medulla
Type B2	Neoplastic epithelial component appears as scattered plump cells with vesicular nuclei and distinct nucleoli among a heavy population of lymphocytes; perivascular spaces are common
Type B3	Predominately composed of epithelial cells having round or polygonal shape and exhibiting mild atypia admixed with a minor component of lymphocytes, foci of squamous metaplasia and perivascular spaces are common
Туре С	Thymic carcinoma

Figure 2. CT-guided biopsy of pleural-based mass



