Angiofollicular lymph node hyperplasia arising from the intercostal space

HIROYUKI MATSUDA, MASAKI MORI, KOSEI YASUMOTO, KEIZO SUGIMACHI

From the Department of Surgery II, Faculty of Medicine, Kyushu University, Fukuoka, Japan

Angiofollicular lymph node hyperplasia (Castleman’s disease) was first described in 1956 by Castleman et al. as asymptomatic benign, hyperplastic mediastinal lymph nodes that resembled thymoma. This disease has been reported in various sites and organs but a careful search of published reports showed no case arising from the intercostal space. We report a case that arose in the chest wall.

Case report

A 69 year old Japanese man was admitted to Kyushu University hospital for evaluation of a long standing abnormal shadow on his chest radiograph, first detected nine years previously on a routine chest radiograph. Initially it had been 1-3 cm in diameter, but three years later it had grown to 2-3 cm and by six years it was 3-4 cm. The patient had been in good health, without peripheral lymphadenopathy or abnormal palpable masses.

On examination there was dullness to percussion with decreased breath sounds and diminished vocal fremitus at the base of the left lung. The erythrocyte sedimentation rate was 18 mm in one hour, and other laboratory data were within normal limits. A chest radiograph showed a uniform shadow 5-7 x 4-0 cm in the left upper field, suggesting an extrapleural tumour and a large pleural effusion (fig 1). Thoracocentesis yielded serous fluid, which showed no specific features on microbiological and cytological examination. There was no evidence of tuberculosis or malignancy in a needle biopsy specimen of the pleura.

A thoracotomy was performed via a left posterolateral incision. When the left pleural cavity was entered about 800 ml of serous fluid was found. A mass, covered by parietal pleura, was situated on the posterolateral chest wall at the level of the 2nd-3rd rib. It appeared to originate in the second intercostal space and protruded into the pleural cavity. It was removed en bloc together with surrounding intercostal tissues. The postoperative course was uneventful and the patient was discharged 20 days after surgery. There has been no recurrence of either the mass or the effusion during the ensuing three years.

The resected specimen measured 5-5 x 4-5 x 4-5 cm and weighed 56 g. It was well encapsulated with fibrous tissues and had a hard elastic consistency. The cut surface was partly yellow grey and partly dark brown, with no necrosis. On microscopic examination there were numerous small lymphoid follicles, the centres of which were composed of concentrically layered large, pale cells with a squamoid appearance. In the interfollicular areas there was extensive capillary proliferation, and lymphocytes, plasma cells, and eosinophils were also present. The capillaries were lined by plump endothelial cells and ensheathed by a varying amount of hyalinised collagenous tissue (fig 2). All these findings were compatible with the hyaline vascular type of angiofollicular lymph node hyperplasia.

Discussion

Since Castleman et al. first described angiofollicular lymph node hyperplasia many investigators have reported the condition under various names, including large lymphphnodal hamartoma,1 follicular lymphoreticulosioma,2 angiofollicular mediastinal lymph node hyperplasia,4 and angiomatous lymphoid hamartoma.3 Keller and coworkers6 divided the disease histologically into the hyaline vascular type, characterised by small hyaline vascular follicles and interfollicular capillary proliferation, and the plasma cell type, consisting of large follicles with intervening sheets of plasma cells. The hyaline vascular type, which accounts for 90% of cases, does not give rise to systemic manifestations. The much rarer
plasma cell variant, on the other hand, is associated with anaemia, a raised erythrocyte sedimentation rate, hypoglobulinaemia, hypoproteinaemia, and fever.

These lesions are benign and grow slowly. Our patient had been followed for nine years, during which time the mass had gradually increased in size from 1.3 cm to 4-0 cm in diameter. The doubling times in the three year periods were calculated to be 386 days, 819 days, and 1551 days. Tanaka et al reported data on a patient followed for four years and here the doubling time was 501 days, the mass enlarging from 2.3 to 4.5 cm during four years.

Angiofollicular lymph node hyperplasia has a predilection for certain sites and organs. It most commonly arises in the mediastinum, followed by the neck, lung, retroperitoneum, intrapelvic cavity, axilla, and mesentry.6 A few cases of multicentric disease have been described, but these are generally considered to represent a separate clinical entity from the localised tumour.

The current case is, so far as we are aware, the first report of angiofollicular lymph node hyperplasia arising from the intercostal space and presenting as a chest wall tumour.

Another interesting finding was the accompanying pleural effusion. Although typically the hyaline vascular type tumour is usually without systemic manifestation the effusion in our patient seemed to be related to the disease as it disappeared rapidly after excision of the lesion.

An unusual reactive lymphoid response, a hamartomatous process, and true lymphoid tumour development have all been suggested for the pathogenesis. A viral causation, with decreased T4:T8 lymphocyte ratio, has been proposed in several recent studies.10 Immunocytochemical analysis was not, however, carried out in the present case.

Surgical removal seems the most appropriate treatment since there has been no recurrence of either the lesion or the effusion after three years of postoperative follow up.

References

2 Abell MR. Lymphnodal hamartoma versus thymic choristoma of the pulmonary hilum. Arch Pathol 1957;64:584–8.