Thymoma and systemic lupus erythematosus

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Thymoma and systemic lupus erythematosus. The simultaneous occurrence of a thymoma and systemic lupus erythematosus (SLE) is reported. Disease states associated with thymoma are reviewed and the possible immunological basis for this spectrum of diseases is discussed. The role of angiography in the diagnosis of thymic tumours is described.

This is the report of a case of thymoma associated with systemic lupus erythematosus. Although the association of these two pathological states is rare (Larsson, 1963; MacKechnie, Squires, and Platts, 1973), they have been found together in association with other diseases (Clinical Pathologic Conference, 1973; Funkhouser, 1961; Galbraith, Summerskill, and Murray, 1964; Kough and Barnes, 1964; Montes, Carter, and Moreland, 1968; Peterson and Lund, 1969; Singh, 1969).

In December 1972, the patient developed pains in the knees and metacarpophalangeal and proximal interphalangeal joints and also a Raynaud phenomenon. The ESR was elevated at 55 mm in 1 hour and the anti-nuclear factor was 1:1024. LE preparation showed one cell to be positive. A repeat ANF at a later time was 1:4096. The patient responded well to a brief course of prednisone but in September 1973 presented at another hospital with chest pain and a pericardial friction rub. A chest radiograph at that time revealed a right pleural effusion and a large mediastinal mass (Figs. 1 and 2). A mediastinoscopy was performed and a biopsy specimen showed normal lymphoid and involuted thymic tissue.

In November 1973, chest radiographs showed that the mediastinal mass was smaller. The ANF was positive to a titre of 1:256 and the LE preparation was positive. Arteriography via a right internal mammary artery in December 1973 demonstrated a vascular tumour (Fig. 3). Exploratory thoracotomy in January 1974 revealed a large, firm, lobular mass occupying the right lobe of the thymus which was enlarged to 8 x 7 x 5 cm. The tumour did not involve other structures. A total thymectomy was performed and histological examination showed an epithelioid and lymphoid thymoma (Fig. 4). Many cystic and necrotic areas were present and the tumour was observed to invade the capsule of the thymus as well as surrounding normal thymic tissue. Lymph nodes at some distance from the tumour were not involved microscopically. In January 1974, radiation therapy was given to a total dose of 4800 rads.

Removal of the thymoma temporarily reduced the clinical and laboratory manifestations of SLE. At the time of discharge, the patient was free of previously described symptoms, and no LE cells were found but
The patient fulfils the majority of the generally accepted criteria for systemic lupus erythematosus which include an ANF titre of more than 1:64, pleurisy, pericarditis, LE cells, arthralgia, and Raynaud's phenomenon (Wolf, Gokcen, and Good, 1963).

Within the past few years the number of different diseases found in association with thymoma has been steadily increasing (Rubin, 1964). The earliest and most firmly established connections include myasthenia gravis (Castleman, 1955), erythroid aplasia (Chalmers and Reheimer, 1954), and agammaglobulinaemia (MacLeary, Zak, and Varco, 1956). Other associations are numerous and include the

**DISCUSSION**

Masses occurring in the anterior mediastinum include thymoma, teratoma, dermoid cyst, aneurysm, substernal thyroid, lymph nodes, and metastatic neoplasms. Subclavian and internal mammary angiography can help to clarify and distinguish these mediastinal masses (Boijsen and Reuter, 1966). If the blood supply, as in this case, is derived primarily or exclusively from the thymic artery, the tumour is probably a thymoma. Although it is difficult to distinguish benign from malignant thymomas, the former is suggested by an absence of tumour vessels and a smooth margin to the mass. In contrast, a dermoid cyst is usually a totally avascular lesion. Substernal thyroids are very vascular masses which are supplied by an enlarged inferior thyroid artery. Many small tortuous tumour vessels arising from the internal mammary artery characterize metastatic deposits. Mediastinal angiography, in this case, was of value not only in establishing a diagnosis but also in determining the presence and extent of the tumour's blood supply. However, its obvious limitation in distinguishing benign from malignant thymomas should be noted.
following: hypoplasia of other marrow elements (Havard and Scott, 1960), carditis (Funkhouser, 1961), positive Coombs' test (Jahsman, Monto and Rebuck, 1962), Cushing's syndrome (Pimstone, Uys, and Vogelpoel, 1972), Hashimoto's thyroiditis (Dawson, 1972), pemphigus (Lundstrom, Pertschuk, and Zimmerman, 1973), candidiasis (Stillman and Baer, 1973), myositis and myopathy (Rubin, 1964), multiple myeloma (Lindstrom, Williams, and Brunning, 1968), leukaemia (Anderson and Henning, 1967), erythrocytosis (Lundstrom, 1970), and megaoesophagus (Demos, Yadasky, and Tjin, 1966).

The association of thymoma and systemic lupus erythematosus remains to be explained. Bach (1974) has presented evidence to support the existence of a circulating thymic hormone and has noted that the level of this hormone is decreased in some immune deficient states and in patients with SLE. Disease in NZB mice (East, 1970), similar to human lupus erythematosus, is characterized by abnormalities in T cells before autoimmune manifestations appear. Thymoma and SLE occurring in the same patient lends support to the hypothesis that the thymus gland is primarily responsible for the development and regulation of auto antibodies.

A consistent relation between thymoma and SLE has not been proven, but accumulating data would support more than a casual relationship between these two relatively rare disease states. Improvement in our patient's clinical condition following thymectomy might suggest that a thymic tumour should be excluded in patients with SLE refractory to the usual therapeutic measures.

REFERENCES


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