LYMPHO-EPITHELIOMA OF THE THYMUS
NERVOUS AND OTHER CLINICAL SYMPTOMS IN THE ADULT

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In this paper we are concerned, not with the simple tumours—fibroma, lipoma, lymphangioma, lymphosarcoma, round-cell sarcoma—which occasionally (though rarely) arise in the thymus, as in other parts of the body, but with those which arise from the essential epithelioid cells of the thymic medulla associated in greater or lesser proportion with lymphocyte-like cells (probably really lymphocytes), such as constitute the normal thymic cortex. These primary thymic tumours—true “thymomata” may be termed “lympho-epitheliomata” (Grandhomme, 1900; Schmincke, 1921; Regaud, 1921), and are much less often malignant than are the primary lympho-epitheliomata of the tonsils, pharynx, and naso-pharynx (Cappell, 1934; Harvey, Dawson and Innes, 1937). For recent historical accounts of thymic lympho-epitheliomata, see Wu (1935) and Obiditsch (1937).

The thymic lympho-epitheliomata seem in some cases to exert an endocrine effect, and are specially related to myasthenia gravis. Most important and interesting was the original observation by Weigert (1901) of a thymus tumour in a case of myasthenia gravis in which he wrongly interpreted the “lymphorrhages” in the muscles as tumour-metastases. To these questions we will return in the discussion further on, but for a summary of the literature of the subject we must refer to the recent papers by Gold (1935), Meister (1936), Norris (1936), Miller (1940), Blalock et al. (1941), and Poer (1942).

Case Record

The patient, Mrs. F. F., aged 58 years, was admitted on 5th November, 1941, with irregular pyrexia of uncertain origin. The history was that after a sore throat 6 weeks ago she had been suffering from pains in various parts. Blood-count: Hæmoglobin, 60 per cent.; erythrocyts, 3,610,000; colour-index, 0.8; leucocytes, 9,000 (polymorphs, 53 per cent.; lymphocytes, 40 per cent.; monocytes, 7 per cent.). Blood-sedimentation: first hour, 70; two hours, 120. Blood-culture: negative. Urine: nothing abnormal. After a bursting systolic murmur could be heard in the left intercostal space close to the sternal margin. X-ray examination of the heart (Dr. F. G. Wood) showed a rounded mass projecting to the left of, and apparently continuous with, the heart shadow and moving with it; right and left oblique views were thought to point to this projection being an enlargement of the pulmonary artery and conus (Fig. 1). Brachial blood-pressure: 135/80 mm. Hg. Blood-Wassermann reaction: negative. Ophthalmoscopic examinations (Dr. C. Markus) showed hyperaemia, especially left eye, apparently connected with hypermetropia. The patient at various times complained of a severe kind of acroparasthesia and of numbness (no real anaesthesia) of the right lower limb and afterwards of the right upper limb and both lower limbs. Severe cramps were sometimes complained of. The knee-jerks, which were equal and natural on admission, were unequal at the end of November, the left one being greatly exaggerated. In March 1942 the patellar and Achilles reflexes were absent on both sides; the triceps reflex was absent on the right side, present on the left side; the plantar reflex was of the normal flexor type on both sides; no superficial abdominal reflexes were obtained. Both pupils reacted normally. The speech was rather sluggish. Very little tactile responses in either lower limb; some hypersensitiveness to pressure over the sciatic nerve, both sides. The patient tended to keep her knee-joints flexed. Although general asthenia was very marked, no definite characteristic features of myasthenia gravis were noted.

Focal infection was thought of, but no focus could be discovered. There was moderate fever from admission to 25th November—then a little at the commencement of December and from 10th December to the commencement of January 1942. About 5th February there was again fever, and then from 22nd February to the patient’s death on 17th March, 1942, which was due to a final broncho-pneumonia with some decubitus. Frequent profuse night sweatings constituted also a clinical feature. The last blood-culture yielded a growth of staphylococcus albus. Amongst the drugs tried sulphapyridine on one or two occasions seemed to exert temporary beneficial effect.

Necropsy

Over the front upper part of the heart and slightly to the left, firmly connected with the fibrous layers of the parietal pericardium, was a bun-shaped mass, about the size of half a large orange, 14×8×6 cm. in measurement (weight after preparation as a museum specimen: 112 gm.). The multilocular cystic central portion of the mass had a honeycomb appearance on section; most of the cysts freely communicated with each other and were filled with a slimy, translucent mucoid fluid; their walls had a smooth inner surface. In the upper portion of the “tumour” was a slightly larger cavity, which was filled with what macroscopically appeared to be pus, but no microbes could be detected in a film stained with methylene blue, or by Gram’s method.

Thyroid gland: slightly enlarged, showing nodular adenomatous condition, one of the nodules in the left lobe having a calcified capsule. Bronchopneumonia of the lower lobe of the right lung; purulent bronchitis. Nothing specially noteworthy in the other organs: brain, heart, liver, spleen, pancreas, adrenals, kidneys, uterus, ovaries, gastrointestinal tract, larynx.
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Microscopical Examination

The solid part of the mass consisted of a dense collagenous fibrous framework, in which were embedded cellular islands consisting of delicate spindle and epithelioid cells with vesicular nuclei, associated with lymphocyte-like cells as in so-called thymic lympho-epitheliomata (Fig. 2). No Hassal’s corpuscles were seen, but at one spot several elongated, slightly curved ("banana-shaped") vesicular nuclei were noted, grouped together, so as to suggest a faint attempt at the formation of a Hassal’s corpuscle. In their shape some of the so-called "spindle cells" resemble rather the cells of "oat-cell" bronchial carcinoma. Microscopical examination of what macroscopically had appeared to be pus showed that it consisted of this cellular material breaking down ("softened") as a result of (ischemic?) necrosis. It may be mentioned that a film of this "pseudo-pus" stained with methylene blue showed the presence of a considerable number of tissue mast-cells, which were not clearly differentiated in sections stained in other ways. The honeycomb cystic portion of the tumour represented apparently a further stage of aseptic necrotic involution of the lympho-epitheliomatous tissue. The process was probably analogous to the occasional changes in the thymus gland of children leading to the formation of Dubois’s "abscesses" or pseudo-abscesses formerly supposed to be necessarily connected with congenital syphilis.

Is the mass to be regarded as a real lymphoepitheliomatous tumour of an ectopic thymus (or thymic "rest") with secondary cystic degenerative changes, as above explained, or is it an example of hypertrophy of an ectopic thymus (or thymic "rest") containing cysts resulting from involutionary focal ischemic necrotic processes (Kopac, 1939)? The spindle and epithelioid cells resemble those of normal thymic medulla and the lymphocyte-like cells resemble those of normal thymic cortex—which most authors now regard as true lymphocytes—but the grouping of these two constituent types

FIG. 1.—Radiogram of the thorax taken on 20th November, 1941.
of cells is not sufficiently clearly marked out into cortex and medulla to permit the mass to be regarded as the result of mere thymic hypertrophy; nor does the complete absence of Hassal's corpuscles favour the view of mere hypertrophy. The mass must therefore, we think, be regarded as a true primary thymic lympho-epitheliomatous tumour ("thymoma"). Whether the tumour is a primary one of an ectopic thymus or of a thymic "rest" makes no difference from the pathological point of view; we did not specially search for any thymic remnant in the normal position. By our term "ectopic" we do not imply that the position of the tumour, in front of the pericardium, is abnormal for thymic tissue in early childhood. In the present case there were no metastases and nothing in the microscopical examination pointed to malignancy.

**Discussion**

Clinical symptoms from thymic hypertrophy or tumour may of course be mechanical, from local pressure, and in the present case the systolic murmur heard to the left of the sternum was probably due to tumour pressure.

If the thymus is an endocrine organ one would expect that constitutional endocrine symptoms would often result—as they often do in primary tumours of other endocrine organs—from a primary neoplasm arising from the essential (endocrine) cells, that is to say, so long as the neoplastic cells still retain something of the original functional (endocrine) activity, in fact, so long as the neoplastic cells functionally still resemble the cells of simple hyperplasia.

In something like 55 per cent. of cases of myasthenia gravis hypertrophy or primary tumour (lympho-epithelioma) of the thymus has been present, but in reality the percentage is probably considerably higher, for in many cases thymus or thymic tumour has not been sufficiently looked for. On the other hand, myasthenic symptoms may have been overlooked in some cases of thymic tumour in surgical wards (Mann, 1934). Although in our present case characteristic symptoms of myasthenia gravis were absent, it is highly probable that the great general asthenia, the extreme acroparaesthesia, numbness and other nervous symptoms were in some way con-

![Fig. 2.—Photomicrograph of part of the tumour, showing the characteristic epithelioid and lymphocyte-like cells (× 400).](image)

The recurrent sweating and pyrexial periods may possibly, by analogy with what happens in some cases of lymphogranulatosis maligna, be attributed to the necrotic softening and formation of "pseudo-pus" in the lympho-epithelioma, with resulting toxic absorption. Patients with thymic lympho-epithelioma, according to the literature (Matras and Priesel, 1928; Zajewlo- 

schin, 1929 and 1933; Nemenow, 1932; Ercklentz, 1936), have died of pleuritis, empyema, pneumonia, furunculosis, and decubitus. The loss of resistance to toxic-infectious agents culminated in our case with the decubitus and broncho-pneumonia, the latter first manifest 4 days before the patient's death, the former a week earlier. It is possible that the absence of definite symptoms of myasthenia gravis might be due to the diminution of the (pre-
Leyton, O., Kopac, F., Gold, E. B., Harvey, M., Mann, L., Dudgeon, L., W. R. Brain, D., Cappell, H., latter occurrence occasional an ideal tumour, thought of. dilatation of the congenital pericardial tumour, diagnosis differential the on it shadow. The position of the tumour mediastinum overlapping the parietal of the latter "pseudo-pus" similar to those of myasthenia gravis.

It seems that in very rare cases malignant thymoma or primary carcinoma of the thymus may by some unknown process disturb the endocrine balance in such a way as to give rise to symptoms somewhat resembling Cushing's syndrome (Leyton, Turnbull and Bratton, 1931), but into this question of the "Leyton-Turnbull-Bratton syndrome" we cannot enter here.

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**REFERENCES**