Secondary neoplasm of the thyroid presenting as a goitre

J. N. HARcourt-Webster

From the Department of Pathology, University of Edinburgh

SYNOPSIS Secondary neoplasm is a very rare finding in a goitre despite the numerous reports of metastatic deposits in the thyroid gland at necropsy. Eleven cases are reported of such deposits presenting clinically as a thyroid nodule, the sites of origin including a meningioma and an unusual carcinoma of the skin. Some of the factors, including pre-existent degenerative processes, believed to determine the incidence of secondary neoplasm in the thyroid, are discussed together with the problems of establishing the diagnosis.

The thyroid gland has long been recognized, despite its great vascularity, as one of the less common sites for secondary neoplasm. There is, however, some variation in the incidence at necropsy. Abrams, Spiro, and Goldstein (1950) recorded 19 (0·19%) thyroid metastatic deposits in 1,000 consecutive cases of disseminated carcinoma whilst Willis (1931) described nine (5·2%) among 170 similar cases, and Shimoaka, Sokal, and Pickren (1962) reported another 188 (8·6%) such deposits in 1,980 examples of malignant disease. The deposits are derived from a wide variety of neoplasms, the lungs, melanomata, the breasts, and the kidneys being the most frequently recorded sites of origin.

In contrast to the numerous records of metastatic deposits in the thyroid at necropsy they are only rarely reported presenting clinically as a swelling. The true diagnosis may be suspected from the history but frequently the deposit simulates a primary carcinoma; occasionally they are an unexpected finding in an involutionary or degenerative goitre.

This paper records 11 cases of goitre in which there were secondary deposits, five mimicking a primary carcinoma, from a variety of different sources.

CASE REPORTS

CASE 1 A 56-year-old woman presented with a one-year history of a slowly enlarging, firm, slightly mobile nodule in the right lobe of the thyroid. It was regarded as a degeneration cyst and excised.

Some years previously the right breast had been removed for a grade I carcinoma; subsequently the patient remained well.

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case 4 A 40-year-old woman presented with a one-month history of a limp and pain in the left hip. On examination the principal finding was a moderately enlarged, firm thyroid with displacement of the trachea to the right. Radiography demonstrated an osteolytic lesion centred in the left ilium and roof of the acetabulum but there was no abnormality of the spine, long bones, or lungs. A diagnosis of carcinoma of the thyroid with a bone metastasis was made and a biopsy taken from the thyroid.

Histology Both lobes included several, variably sized microfollicular adenomata. Partially replacing some of the adenomata and the neighbouring tissue was a pleomorphic, clear cell, solid alveolar carcinoma (Fig. 1); there was an abrupt transition between it and the thyroid parenchyma but no significant stromal reaction.

The provisional histological diagnosis of a metastatic deposit from a renal carcinoma was confirmed after a retrograde pyelogram had demonstrated a still symptomless mass in the right kidney. Death occurred within eight months, there being multiple metastatic deposits in the bones and lungs.

case 5 A 66-year-old woman was first seen with a one-month history of a small, ulcerated nodule on the lateral aspect of the right thigh. Four months later a firm, nodular swelling appeared in the left lobe of the thyroid together with supraclavicular and inguinal lymphadenopathy; the skin nodule remained unchanged. The provisional diagnosis was carcinoma of the thyroid with metastatic deposits.

Histology Biopsies of the thyroid swelling, the supraclavicular lymph nodes, and the nodule on the thigh all showed a keratin-forming, squamous cell carcinoma. However, that from the skin and, to a lesser degree, that from the thyroid, also included large infiltrating foci of pleomorphic and actively mitotic spindle cells arranged, for the most part, in parallel rows (Fig. 2). Many of these cells contained small quantities of brown pigment some of which was demonstrated to be melanin (Fontana's method) and some to be haemosiderin (Perls's Prussian blue method). In several areas and particularly at the edges of the skin nodule, where an origin from the epidermis could be traced, there was a gradual transition between the squamous and the pigmented spindle cells (Fig. 3). There was no evidence of involutionary changes in the thyroid.

The melanosquamous carcinoma of the skin was regarded as the primary neoplasm with metastatic deposits in the thyroid and lymph nodes.

Over the ensuing 10 weeks there was increasing cervical lymphadenopathy followed by a respiratory infection from which she died: no necropsy was obtained.

case 6 A 52-year-old man presented with a firm, immobile swelling in the right lobe of the thyroid; there was no evidence of cervical lymphadenopathy. It was regarded as a carcinoma and a complete excision of the right lobe was attempted.

Nine years previously a tumour had been removed from the falk cerebri of the parietal area. This had recurred after seven years and removal was again attempted. On both occasions the histological interpretation had been difficult but the tumour was considered to be an active, vascular meningioma.1

Histology There was extensive infiltration of the right lobe of the thyroid by sheets of undifferentiated, pleomorphic polygonal cells (Fig. 4), amongst which were numerous vascular channels. The neoplastic cells abutting upon the endothelial lining of the sinusoids were frequently fusiform but there were few giant cells. There were numerous bizarre and hyperchromatic nuclei and moderate numbers of mitotic figures (Fig. 5). Only scattered, small foci of thyroid tissue remained and there was no stromal reaction.

A diagnosis of an undifferentiated malignant neoplasm of undetermined histogenesis was made; at this stage the detailed previous history and histological sections of the meningioma were not available.

Necropsy, eight months later, showed multiple small tumour deposits in the meninges close to the site of excision of the meningioma and in the lungs but there were none elsewhere. The histological features of both these deposits and the thyroid tumour closely resembled

![FIG. 1. Case 4: A large metastasis from a renal carcinoma which presented as a goitre. Haematoxylin and eosin × 140.](http://jcp.bmj.com/)

1A detailed account of the histological findings will be published by Drs. A. Gordon and A. F. J. Moloney. J. Neurol (in press).
FIG. 2. Case 5: Part of the skin nodule showing large numbers of spindle cells (pigment not shown) arranged in parallel rows amongst which are scattered clusters of pleomorphic squamous cells. Haematoxylin and eosin × 140.

FIG. 3. Case 5: Higher-power view of right half of Figure 2. Note the transition between, right, the pleomorphic and bizarre squamous cells and, left, the pleomorphic spindle cells in which fine granules of pigment can be seen. Haematoxylin and eosin × 350.

FIG. 4. Case 6: A deposit from an undifferentiated meningioma of the falx replacing much of the right lobe of the thyroid. Haematoxylin and eosin × 100.

FIG. 5. Case 6: Higher-power view of Figure 4 to show the pleomorphic polygonal cell structure of the deposit. Note the fusiform character of many of the cells adjacent to the endothelium of the vascular channels. There are numerous bizarre and hyperchromatic nuclei in addition to mitotic figures. Haematoxylin and eosin × 300.
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Histology The nodule from the buttock showed the features of the moderately pigmented malignant melanoma. The thyroid nodule was a macro-follicular adenoma; a small focus of melanin-pigmented (Fontana’s method) pleomorphic spindle cells, identical with those seen in the melanoma of the buttock, was however found amongst the colloid-laden follicles of the adenoma. It was regarded as a secondary deposit.

The patient remained well with no evidence of any further metastatic deposits one year later.

CASE 9 A 74-year-old woman had a slightly pigmented malignant melanoma excised from the left forearm. Three years later the right lobe of the thyroid became moderately enlarged but there was no evidence, at the site of excision, of any recurrence. A biopsy of the thyroid was taken.

Histology The thyroid parenchyma was infiltrated by non-pigmented fusiform and polyhedral cells, including small giant forms, arranged in interlacing bundles with a scanty but vascular fibrous stroma. Despite the absence of pigment the similar cytological features of the previously excised melanoma and the thyroid biopsy indicated that the latter was a metastatic deposit.

Over the next few months numerous enlarged lymph nodes appeared in the neck and axillae, death following from a respiratory infection. No necropsy was performed.

CASE 10 A 27-year-old woman presented with discomfort and dysphagia attributed to a nodular goitre. There were no symptoms or signs of abnormality in the other systems. Partial thyroidectomy was performed for the relief of the symptoms.

Histology The thyroid was extensively infiltrated by sheets of undifferentiated, small, dark-staining round cells similar to but slightly larger than lymphocytes. There was no stromal reaction nor was there any other abnormality of the thyroid. The tumour was provisionally regarded as a lymphosarcoma.

Necropsy, a few weeks later, showed an adrenal swelling together with a moderate sized mediastinal mass and a deposit in the cavity of the right auricular appendage of the heart. These all showed similar histological features to the thyroid tumour but also included small numbers of rosette formations; a diagnosis of an adrenal neuroblastoma with metastatic deposits was made.

CASE 11 An otherwise healthy 50-year-old woman was seen with a nodular colloid goitre which was treated by partial thyroidectomy.

Histology In many areas the thyroid showed the classical features of a nodular goitre; however, one area included a small focus of a poorly differentiated, cuboidal cell adenocarcinoma demarcated from the adjacent thyroid tissue by a thin capsule of connective tissue (Fig. 7). The structure of the tumour and the lack of any transition between it and the thyroid vesicles suggested secondary neoplasm.

A small nodule was eventually found in one of the breasts and excised; this proved to be a typical adenocarcinoma of the breast and was consequently regarded as the primary.

FIG. 6. Case 7: A solitary metastatic deposit from an epidermoid carcinoma of the cervix uteri in a thyroid also showing the features of Hashimoto’s disease. Haematoxylin and eosin × 300.

those of the previously removed meningioma which was concluded to be the site of origin.

CASE 7 A 52-year-old married woman presented with a history of vaginal bleeding and a firm, symmetrical swelling in the neck associated with discomfort and dysphagia. After a provisional diagnosis of carcinoma of the cervix uteri with a metastatic deposit in the thyroid gland, both biopsy of the cervix and a partial thyroidectomy were performed.

Histology The cervical specimen was a moderately well-differentiated epidermoid carcinoma. The thyroid showed the features of Hashimoto’s disease; however, a microscopic deposit of carcinoma similar to that in the cervix was seen in the isthmus (Fig. 6). Metastatic carcinoma also infiltrated a supraclavicular lymph node removed at thyroidectomy.

CASE 8 An otherwise fit 21-year-old man was seen with a small, cutaneous nodule on the right buttock thought to have been present since birth but recently enlarging. There was also a small, mobile lump in the left lobe of the thyroid believed to have gradually appeared over the previous nine months. Both were excised.
Bruce and Michie, 1954), with atrophic and degenerative conditions of the thyroid, in which the oxygen tension and iodine content are probably reduced, supports this belief.

The neoplasms recorded, in clinical cases, metastasizing to the thyroid are shown in the Table.

<table>
<thead>
<tr>
<th>Primary Site</th>
<th>No. of Cases in Literature</th>
<th>Author's Cases</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kidney</td>
<td>35</td>
<td>2</td>
<td>37</td>
</tr>
<tr>
<td>Alimentary tract</td>
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<td>0</td>
<td>9</td>
</tr>
<tr>
<td>Lung</td>
<td>8</td>
<td>1</td>
<td>9</td>
</tr>
<tr>
<td>Breast</td>
<td>6</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Skin</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Malignant melanoma</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Melano-squamous</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Adrenal</td>
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<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Adrenal neuroblastoma</td>
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<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Liver</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Meningioma</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Cervix uteri</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>62</td>
<td>11</td>
<td>73</td>
</tr>
</tbody>
</table>

1Carcinoma unless otherwise stated

The preponderance of renal cancer contrasts with the finding in most necropsy series in which the lungs, melanomata, the breasts, and the kidneys are almost equal in their frequency as the site of origin. This is of interest when considering the route of spread to the thyroid, for Willis (1952) considered entry via the lymphatics to be very rare in this organ; the predilection of renal cancer for haematogenous distribution is well known. The more equal distribution of origin in necropsy reports may be explained by the dissemination of malignant cells via the bloodstream as a near terminal event in many cancers.

The skin, with the exception of melanomata, is a very rare source of secondary neoplasm in the thyroid. The carcinoma with mixed squamous and melanotic features (case 5) is therefore of interest in that whilst its histogenesis remains in doubt, although it is tempting to regard it as an example of what has been termed 'melanogenen metaplasia' (Shivas and MacLennan, 1963), its spread to the thyroid is more in keeping with the behaviour of a melanoma than that of an epithelioma. The meningioma (case 6) is also of unusual interest; these neoplasms are occasionally recorded as metastasizing to various tissues, including the lungs (Jurav, 1941) and the liver (Hamblet, 1944), but there is no previous report of spread to the thyroid.

Secondary neoplasm in the thyroid is an indicator of grave prognosis; in seven of the nine patients being reported, whose subsequent history is known, metastatic deposits were identified at other sites, either before or a few months after biopsy of the thyroid. This is in accord with the observation that

![FIG. 7. Case 11: A deposit of poorly differentiated adenocarcinoma from the breast unexpectedly found in a nodular colloid goitre. Haematoxylin and eosin × 120.](image-url)
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spread to the thyroid at necropsy is virtually unknown in the absence of multiple deposits elsewhere (Bruce and Michie, 1954; Mortensen, Woolner, and Bennett, 1956).

The recognition of a metastatic deposit in the thyroid, when in association with a previously identified neoplasm, may present no particular difficulty to the histopathologist. In the absence of such a history or with a poorly differentiated deposit there may, however, be considerable difficulty but nevertheless, as in this and previous reports (H'Doubler, 1936; Elliott and Frantz, 1960), the possibility of a secondary deposit mimicking a primary carcinoma of the thyroid should not be overlooked. Particular care must be exercised in the diagnosis of a deposit from a renal carcinoma because of the histological similarity between the latter and the primary clear cell carcinoma occasionally seen in the thyroid (Chesky, Hellwig, and Barbosa, 1957). On the other hand, squamous cell carcinoma arising in the thyroid is very rare (I have seen one amongst 359 carcinomas of this organ) and the recognition of this type of neoplasm should arouse the suspicion of it being a metastatic deposit. In the final analysis the differentiation between a primary and a secondary neoplasm will often be made only by careful clinico-pathological consultation.

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REFERENCES


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