Langerhans Cell Histiocytosis Co-Existing with Papillary Carcinoma of Thyroid – A Rare Surgical Challenge

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Abstract: Langerhans cell histiocytosis (LCH) is a rare disorder of unknown origin characterized by infiltration of the involved tissues by large numbers of Langerhans cells, often organized into granulomas. Here we report a case of Langerhans cell histiocytosis associated with papillary thyroid carcinoma (PTC) in a 31 year old man who presented with an enlarged, diffusely firm, non tender, non mobile thyroid swelling. Fine needle aspiration cytology showed involvement of thyroid by langerhans cell histiocytosis. Patient was evaluated, and no evidence of systemic involvement was found. Patient received ten cycles of chemotherapy with vinblastine and etoposide, no decrease in size of swelling was noted. Subtotal thyroidectomy was performed. Histopathologic study revealed Langerhans cell histiocytosis along with papillary carcinoma in biopsy tissue. After histopathologic confirmation, patient was subjected to radioiodine ablation therapy.

Keywords: Langerhans cell histiocytosis, CD 1a, papillary carcinoma thyroid

1. INTRODUCTION

Langerhans cell histiocytosis (LCH) is a group of clinical conditions of unknown etiology characterized by clonal proliferation of bone marrow Langerhans like cells either in situ, leading to isolated bone lesions or, having spread in different remote tissues, potentially capable of inducing multisystem disease. Sometimes there are mutations (BRAF gene)¹ in Langerhans cells as they form. These changes may make the Langerhans cells grow and multiply quickly. This causes LCH cells to build up in certain parts of the body, where they can damage tissue or form lesions. Historically, the disease was confirmed by showing birbeck granules by electron microscopy, but currently the diagnosis is based on immunostaining showing presence of CD1a and / or langerin. Papillary thyroid carcinoma (PTC) is a well differentiated epithelial neoplasm. It is more common in women. Although it affects all age groups, the mean age at diagnosis is 40 years. Prognosis is good in patients aged ≤45 years (more than 90%). PTCs manifest themselves as nodules, and diagnosis relies on patient’s clinical parameters, ultrasonography, and fine needle aspiration cytology. Few cases have been reported of thyroid gland involvement coexistently by LCH and Papillary carcinoma. We do present a case of thyroid involvement by LCH and papillary carcinoma coexistently with diagnostic controversy between aspiration cytology and histopathology.

2. CASE REPORT

A 31 years old man presented in otolaryngology outdoor with complaint of swelling in anterior part of neck for past two years which was insidious in onset and gradually progressive. He also had complaints of polyyuria and polydyspsia for past six months and has been diagnosed with diabetes insipidus and this disorder was controlled by desmopressin nasal spray.

His other health problems included diabetes mellitus, hypothyroidism and thalassemia trait which were well controlled with medical treatment.

Physical examination of the patient revealed approximately 7cmx8cm firm swelling in right lower part of neck which moved with deglutition (Fig 1). The swelling was non tender, non pulsatile...
with restricted mobility. There were no abnormal findings in ear, nose and throat.

**Figure 1. Clinical picture showing thyroid swelling**

A fine needle aspiration cytology showed presence of langerhans cell histiocytosis which was CD1a positive on immunostain.

Contrast enhanced computed tomography scan of the neck (Fig 2) revealed heterogeneous enlargement of right thyroid lobe (10.4x6.4x8 cm) with retrosternal extension. Right common carotid artery was encased by the mass. Left lobe of thyroid was normal.

**Figure 2. CECT neck showing heterogeneous enlargement of right thyroid lobe with carotid vessel encasement**

CT Angiography neck (Fig 3) showed hypodense mass in right lobe of thyroid encasing distal brachiocephalic trunk and proximal segment of right common carotid artery circumferentially. Right IJV was not visualized along the course of the lesion with dominant drainage into the external and anterior jugular veins.

**Figure 3. CT Angio showing hypodense mass in right lobe of thyroid encasing distal brachiocephalic trunk and proximal segment of left common carotid artery circumferentially**

Magnetic resonance imaging of the brain, nuclear bone scan, and chest X ray were normal. He had received chemotherapy with vinblastin and etoposide but no reduction in size of swelling was noted. Patient developed neutropenia and drug induced colitis as a side effect which was controlled by medical treatment.

Surgical excision was planned under GA. Intraoperative findings (Fig 4) revealed 10x8cm tense cystic swelling involving right lobe of thyroid, isthmus and left lobe. The cyst was circumferentially encasing the right internal jugular vein and adhered to anterior wall of right common carotid artery. We performed subtotal thyroidectomy. Right internal jugular vein was ligated. Cyst wall adherent to anterior wall of common carotid was left in situ. Patient recovered from anesthesia without any complications. The post operative period was uneventful.

**Figure 4. Showing thyroid specimen after subtotal thyroidectomy**

Histopathology revealed papillary arrangement of tumor cells with nuclear crowding, overlapping, grooving and optical clearing. Few aggregates of langerhans cells with vesicular cleared nuclei were also noted which were highlighted on CD1a immunostain. Focal giant cell reaction and calcifications were also identified along with moderate lymphoplasmacytic infiltrate admixed with neutrophils and eosinophils.

After histopathologic confirmation of coexistent papillary thyroid carcinoma and langerhans cell histiocytosis, patient was subjected to radioiodine ablation therapy. Patient is disease free for 6 months after treatment.

**3. DISCUSSION**

Langerhans cell histiocytosis is a rare disease. In 1953 Lichtenstein coined the term ‘histiocytosis X’ to encompass three disease entities of langerhans cells including eosinophilic granuloma of bone, Hand Schüler Christian disease (the triad of exophthalmos, diabetes insipidus and multiple bony lesions) and Letterer Siwe disease (a
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syndrome of rapid, progressive multiple soft tissue involvement). In fact, there is a considerable overlap between these entities.

LCH usually presents as a multisystem granulomatous infiltrate (in two thirds of cases) in adults. The condition typically presents with the involvement of bone, lung, skin, lymph nodes, and multisites. Most common endocrinological manifestation of classical LCH is associated with posterior pituitary involvement presenting as diabetes insipidus. However, LCH involving the thyroid gland is extremely rare. Thyroid involvement of LCH can be indistinguishable from other thyroid disorder presenting with goiter. It is important to distinguish isolated thyroid LCH from multisystemic cases because single organ involvement is associated with an excellent survival of close to 100%. Thyroidal diseases secondary to LCH may present as thyroid enlargement because of infiltration of the thyroid by LCH.

They do demonstrate a characteristic nuclear groove in histopathology and positive immuno staining with CD68, S100, and CD 1a as in our case.

Under the recommendation of the Writing Group of the Histiocyte Society, a definitive diagnosis of LCH can only be rendered when either there is demonstration of Birbeck granules on electron microscopic study or there is the demonstration of CD1a expression with immunohistochemistry, the latter being present in our case.

Due to its rarity, diagnosing LCH of the thyroid can be challenging. It can be clinically confused with the far more common benign goiters; un differentiated carcinoma, lymphoma, lymphocytic thyroiditis, and chronic granulomatous thyroiditis.

Langerhans cell histiocytosis of the thyroid may occasionally be associated with a nor or with hypothyroism as in our patient.

The role of chemotherapy in histiocytosis X is controversial. There have been no trials large enough to confirm the benefit of a specific regime.

Multiple chemotherapeutic regimens have been attempted: glucocorticoids, vinblastine, etoposide, cyclophosphamide, methotrexate, and doxorubicin. Our patient had also received chemotherapy with vinblastine and etoposide but no reduction in size of swelling was noted.

Surgical treatment is the suitable option. For LCH localized to the thyroid, resection of LCH by subtotal, near total, or total thyroidectomy is the treatment of choice.

Papillary thyroid carcinoma is the most common thyroid malignancy with a high rate of BRAF V600E mutations. Most commonly found in younger (<45 years old) patients. Histologically, it consists of epithelial papillary projections between which calcified spherules (psammoma bodies) may be present. Nuclei are characteristically large with central clear areas (orphan Annie nuclei). Papillary carcinoma metastasizes via the lymphatics to the cervical lymph nodes. Diagnosis of papillary carcinoma relies on patient's clinical parameters, ultrasonography, and fine needle aspiration cytology.

LCH is known to be associated with other neoplasms, such as lymphoma, leukemia or lung cancer, but the association with PTC remains rare and unclear, as there have only been a few reported cases. However, a recent study showed an increased presence of CD1a positive dendritic cells in PTC. The association of LCH with PTC, which is rare, can be attributed to the fact that the same mutation may play a role in the underlying pathogenesis of both diseases.

Thyroid involvement in LCH is rare, as is its association with PTC. Surgical removal seems to be the gold standard treatment for isolated LCH, where as disseminated disease requires systemic therapy. Associated PTC may require further treatment, such as radioiodine therapy, but there are no standard recommendations.

In summary, in patients with thyromegaly, the possibility of Langerhans cell histiocytosis, although rare, should be kept in mind. A careful microscopic examination, immunohistochemical and ultrastructural studies are very important to make a correct diagnosis. In view of the increasing incidence of PTC, especially in women, one possible clinical implication of these findings is that patients with LCH characterized by activating BRAF mutations should be monitored for PTC.

**REFERENCES**


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