A rare incidental dual malignancy in Ectopic and Orthotopic Thyroids

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ABSTRACT

Ectopic thyroid tissue may be encountered anywhere from the foramen caecum to the lower neck. It is rarely seen in the mediastinum. True malignant transformation in ectopic thyroid tissue is rare. Cases of ectopic primary thyroid carcinoma with another primary malignancy in the orthotopic thyroid is extremely rare. We report on an extremely rare case of true mediastinal primary thyroid carcinoma with another malignancy in the orthotopic thyroid suggesting a de novo process in a 45 year old female. The clinicopathologic features and diagnosis of the lesion with regard to its mediastinal location are discussed.

KEYWORDS: Dual malignancy, Ectopic thyroid, papillary carcinoma, hurthle cell carcinoma, Orthotopic thyroid.

INTRODUCTION

Heterotopic thyroid tissue can be found not only as a component of thyroglossal duct cyst but anywhere along the course of the thyroglossal duct cyst, sometimes as the sole abnormality. The most frequent location is the base of the tongue. At a microscopic level lingual thyroid is found in 10% of normal individuals [1]. Other sites of ectopic thyroid tissue are the anterior tongue, submandibular region, larynx, trachea, mediastinum usually superior and heart. Heterotopic thyroid tissue in any of these locations is subject to the same diseases that can affect the main gland, including inflammation, hyperplasia and tumors. Thus several cases of follicular carcinoma arising from lingual thyroid have been reported [2].

CASE REPORT

A 45 Year old female patient was admitted to the hospital for incisional hernia repair. During her surgical work up incidentally she was found to have mediastinal mass on chest x-ray. On physical examination, thyroid was normal and no lymph node was palpable in the neck. Ultrasound examination showed enlarged left lobe of thyroid CT neck showed enlarged left lobe thyroid that appears continuous with above mass. Radioactive iodine scintigraphy revealed cold nodule in left lobe of thyroid. No iodine uptake for mediastinal mass. No association of mass with thyroid is established.

The patient was subjected to median sternotomy for excision of anterior mediastinal mass. Intraoperative evaluation of mediastinal mass showed no glandular continuity between the thyroid gland and the mass. Grossly mediastinal mass is measuring 10X8X3 cm. Cut section shows encapsulated tumor measuring 6X5 cm, grey white to grey brown in color. Microscopically sections show normal thyroid tissue along with tumor tissue arranged in papillary pattern.

Total orthotopic thyroidectomy was done one month after mediastinal mass resection. Totalthroidectomy specimen is measuring 7X3x1cm with nodular surface. CT neck showed enlarged left lobe thyroid that appears continuous with above mass. Radioactive iodine scintigraphy revealed cold nodule in left lobe of thyroid. No iodine uptake for mediastinal mass. No association of mass with thyroid is established. The patient was subjected to median sternotomy for excision of anterior mediastinal mass. Intraoperative evaluation of mediastinal mass showed no glandular continuity between the thyroid gland and the mass. Grossly mediastinal mass is measuring 10X8X3 cm. Cut section shows encapsulated tumor measuring 6X5 cm, grey white to grey brown in color.

Individual cells show vesicular nuclei with ground glass appearance. A diagnosis of papillary carcinoma was made. [Figure 2]. Microscopically sections show normal thyroid tissue along with tumor tissue arranged in papillary pattern. Individual cells show vesicular nuclei with ground glass appearance. A diagnosis of papillary carcinoma was made. [Figure 3].

Total orthotopic thyroidectomy was done one month after mediastinal mass resection. Total thyroidectomy specimen is measuring 7X3x1cm with nodular surface. Cut section shows 2.5X2 cm grey brown nodule. Sections show a tumor tissue composed of hurthle cells arranged in nest pattern. The tumor tissue is seen infiltrating into the surrounding normal thyroid in focal areas. A diagnosis of hurthle cell carcinoma was made. The post-operative period is uneventful. The patient is asymptomatic and disease free sixteen months after surgery.
DISCUSSION
Historically there have been two distinct categories of aberrant or ectopic thyroid tissue, one is lateral in its location and the other is in the midline. Median ectopic thyroid may be encountered from the tongue to the diaphragm. The development and descent of the thyroid is in anatomical juxtaposition with the heart, which lies just inferior to the tuberculum impar in the embryo. The mechanical effects of the descending heart on the thyroid gland contribute significantly to the development of the various anomalies of thyroid position anywhere from the foramen caecum to the diaphragm. Lingual thyroid is the most common type accounting for 90% of cases. The sublingual types may be suprahoid, infrathyroid or at the level of the hyoid bone[3].

In 1869, Hickman reported the first case of ectopic thyroid tumor of the base of the tongue, pressing down the epiglottis on the larynx and causing death by suffocation sixteen hours.
after birth[4]. The Ectopic thyroid may lie in the mediastinum, larynx, trachea and esophagus[5]. The presence of ectopic thyroid tissue in other places distant from the neck region include the heart, aorta, thymus, oesophagus, duodenum, gall bladder [6], stomach, pancreas, mesentry of the small intestine, porta hepatis, adrenal gland, ovary, fallopian tube, uterus, and vagina[7].

Genetic research has been shown that the gene transcription factors TITF-1(NKx2-1), Foxe 1 (TITF-2) and PAX-8 are essential for thyroid morphogenesis and differentiation. Mutation in these Genes may be involved in abnormal migration of the thyroid[8, 9]. Primary thyroid carcinomas arising from ectopic thyroid tissue are uncommon and have been reported to arise from thyroid tissue in the thyroglossal cysts, lateral aberrant thyroid tissue, lingual thyroid, mediastinal, and Struma ovarii. Most tumors in the ectopic locations have been papillary carcinomas, mixed follicular and papillary carcinomas or hurthle cell tumors [10].

Patients with intrathoracic goiter are usually asymptomatic with the tumor reported as an incidental finding on chest roentgenogram[11]. Similar to our case. Mediastinal ectopic thyroid carcinoma is extremely rare as most cases do not meet the criteria of ectopic thyroid tissue. Our case is a true ectopic mediastinal thyroid as it met the criteria of ectopic thyroid i.e. Intraoperative evaluation of mediastinal mass showed no glandular continuity between the thyroid gland and the mass. The cervical thyroid gland does not have a similar pathologic process as the ectopic tumor[12].

Surgical excision is the mainstay of treatment as these tumors usually give rise to compressive symptoms. Thoracotomy or sternotomy is usually required for mediastinal thyroid tumors. The primary treatment after mediastinal mass resection is total thyroidectomy. Complete thyroid resection, not only excludes the thyroid gland as the primary source of malignancy but also facilitates future patient management by serum thyroglobulin measurement and whole body iodine scintigraphy.

CONCLUSION

Mediastinal ectopic thyroid carcinoma is extremely rare. Synchronous presentation of carcinomas of two different histologic variants as primary malignancies in orthotopic and ectopic thyroids have not been reported in literature till date.

REFERENCES


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