Bulky Mediastinal Metastasis in Neglected Follicular Thyroid Carcinoma: A Case Report

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ABSTRACT
Follicular carcinoma of thyroid usually behaves in an indolent manner with low metastatic potential. Distant metastases as initial presentation are rare in follicular carcinoma, especially in young patients. Blood borne metastasis is common with spread to lung, bone and other solid organs. However, metastatic mediastinal tumors are rare.

Here, a neglected case of follicular carcinoma of thyroid (FTC) is reported. The patient is a 43 year old female who was referred to the Thyroid Division of NINMAS for radioiodine therapy for an inoperable metastatic FTC. She was presented with fever, shortness of breath and chest discomfort. $^{99m}$Tc scan showed concentration of most of the isotope in the big metastatic mediastinal mass. Because of the sheer volume of the mass, a plan was undertaken to first reduce her tumor burden with external beam radiation therapy (EBRT) and then considered adjuvant therapy with radiiodine after manageable regression in size of the metastatic tumor. She had a history of thyroidectomy for FTC 14 years back without radio iodine ablation.

Negligence about proper management for FTC may result in poor outcome like huge mediastinal mass as in this reported case. Radioiodine therapy is usually the first line approach for functional metastasis, but in exceptional cases when the tumor size is very big, EBRT may be considered first. Radiotherapy (RT) is effective in relieving compression symptoms and may improve the quality of life in these patients.

Key words: Follicular Thyroid Carcinoma, Mediastinal mass, Metastasis

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INTRODUCTION
Follicular Thyroid Carcinoma (FTC) is the second most prevalent of the thyroid carcinomas and it usually spreads by hematogenous route. The treatment protocol for FTC consists of surgery followed by ablation with radioactive iodine. Rarely FTC metastasis occurs in mediastinum and 3% have been reported as tumors within the chest. However, the incidence of malignancy in mediastinal masses reported ranged from 25 to 49% (1). Here, an interesting case of FTC is reported with unusual and extensive anterior mediastinal invasion.

CASE REPORT
A 43 year old female noticed a swelling in the neck and visited a physician 14 years back. She had no past medical history or risk factors for thyroid malignancy. Laboratory findings and thyroid function tests were normal. Thyroid ultrasonography revealed a predominantly solid, hypoechoic thyroid nodule in left lobe with sonographic features suggestive of malignancy. The right lobe was considered normal. Ultrasound guided FNAC prepared her for RAIT as a first line approach. Treatment plan was designed to reduce her tumor burden and pushed the trachea posteriorly. CT guided FNAC revealed metastatic FTC. There was no lung metastasis. Invasion of tumor to ascending aorta, superior vena cava and right atrium could not be excluded but showed compressions. Patient was referred to NINMAS for RAIT as surgeons found her inoperable. Laboratory findings revealed hyperthyroidism (TSH <0.01mIU/L) and her serum thyroglobulin (Tg) level was found to be more than 300 ng/ml. Thyroid scan (with $^{99m}$Tc) showed no radiotracer concentration in the thyroid bed but almost all radiotracer concentrated in the big mediastinal mass. High resolution ultrasound of neck (HRUS) detected a big (definite measurement was not possible),
predominantly solid, heterogeneous mass behind the sternum. Since the tumor size was huge, a customized treatment plan was designed to reduce her tumor burden with external beam radiation therapy (EBRT) rather than prepare her for RAIT as a first line approach. EBRT was chosen as the first line of treatment with two major expectations: 1) it would help to alleviate the compression symptoms that the patient was experiencing 2) it would be effective in regressing the tumor to a manageable size for subsequent radioiodine therapy.

Figure 1: CT image showing enhancing soft tissue density mass (11X09cm) located in anterior mediastinum having internal calcifications.
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The case reported here is remarkable in that she survived

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criteria. Minimally invasive follicular carcinoma is an

encapsulated tumor with microscopic penetration of the

tumor capsule without vascular invasion (6, 7). Patients

since mediastinal tumors are uncommon and represent

only 3% of the tumors seen within the chest (1).

In reports on mediastinal masses, the incidence of

malignant lesions ranged from 25 to 49% (4, 5). The

anterior mediastinal tumors include teratomas, lymphomas, thymic cancers, thymic carcinoids, thymic cysts and other metastatic tumors. Thyroid cancer rarely

metastasizes to the mediastinum and there have been few

reports of metastatic mediastinal tumors of thyroid origin. Follicular carcinoma is divided into minimally invasive and invasive variants based on morphologic

criteria. Minimally invasive follicular carcinoma is an

capsule without vascular invasion (6, 7). Patients
with minimally invasive follicular carcinoma tend to be younger than patients with invasive follicular carcinoma, and it has been suggested that minimally invasive follicular carcinoma may be a precursor to its invasive counterpart (8). Shaha reported an overall survival at 5, 10, and 20 years for patients with follicular thyroid cancer of 85%, 80% and 76%, respectively (9). When patients with follicular carcinoma were divided into low, intermediate, and high-risk groups based on age, T stage, distant metastases, histologic type, and grade, their survival rates were 98%, 88%, and 56% at 10 years and 97%, 87%, and 49% at 20 years, respectively. D’Avanzo and colleagues reported that patients with minimally invasive follicular carcinoma have a 98% 10-year survival, compared with 80% at 10 years for patients with angioinvasive follicular carcinoma with or without capsular invasion and 38% with extensive invasion of the tumor capsule and the thyroid parenchyma (10). The cause of death is most commonly from progression of distant metastases (11).

The case reported here is remarkable in that she survived more than 14 years without RAIT and/or hormone replacement. But it is evident that the cancer was progressing slowly and insidiously albeit sub-clinically until it became clinical with compression symptoms 12 year later. This slow progression is consistent with the indolent character of the follicular variety of the thyroid cancer.

The compression symptoms that the patient experienced were attributed to the slowly enlarging mediastinal metastasis. This was revealed on CT scan as a strongly enhancing soft tissue density mass having internal calcifications in anterior mediastinum. The mass compressed great vessels and trachea posteriorly. Core biopsy revealed metastatic follicular carcinoma in this patient.

The standard procedure for initial treatment of DTC is surgery followed by radiiodine therapy and hormone replacement. The management plan may differ from patient to patient and depends on several risk factors. However surgery followed by post-operative RAIT has been associated with improved prognosis (12). It is a well established fact that optimal management of patients with complex thyroid cancer requires an integrated team approach involving endocrinologists, NM physicians, medical oncologists, radiation oncologists, and surgeons (13). Radical surgery, remnant thyroid ablation and RAIT all improve survival, but this is less apparent for patients presenting with distant metastases as this subgroup shows worse prognosis. Accordingly surgery would have been the first choice in the management of the case presented here. However, due to the sheer volume of the mass and tracheal invasion surgical resection was not possible.

EBRT is recommended for palliative purposes to obtain local control for extensive diseases as first line therapy (14). The case reported here was referred to oncologist for EBRT with two goals. (1) To alleviate the compression symptoms (2) regression to a manageable size for subsequent RAIT.

Consequently the patient received a course of conventional radiation with 50 Gy in 25 fractions over 4 weeks targeting on the area volume encompassing the thyroid bed and the gross disease. She tolerated EBRT very well and her stridor subsided. There was tracheal inflammation and skin erythema which are expected side-effects of the radiotherapy, but these were transient. She also had complaints of dysphagia, but did not require a feeding tube. The patient is now recuperating and radiiodine will be considered as an adjuvant in the treatment plan. However, if RAIT fails patient may be considered for Tyrosine Kinase Inhibitors (TKIs) which is reported to be effective for RAI-refractory DTC (15).

**CONCLUSION**

Outcome of DTC is excellent with early diagnosis, meticulous surgery, proper staging, RAIT, lifelong thyroxine supplementation and regular follow-up but it may turn aggressive without appropriate management.

Individualized and an alternative therapeutic approach should be considered in patients of DTC with inoperable, huge metastatic mediastinal mass. Palliative use of EBRT for local control of extensive disease may be considered as first line of therapy; when other modality of treatment cannot be applied. Radiotherapy (RT) seems effective in relieving compression symptoms and may prolong survival and improve QOL.

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