

Simultaneous Occurrence of Papillary Carcinoma and Medullary Carcinoma

Lyza Camille Gadong and Thelma Crisostomo

Section of Endocrinology, Diabetes and Metabolism, Department of Internal Medicine, Makati Medical Center, Philippines

Abstract

The cell origin, histopathologic features, and prognosis of medullary and papillary thyroid carcinoma are different and to have them occur simultaneously in a single patient is a rare occurrence.

This is a case of a 38-year-old female who presented with an enlarging anterior neck mass whose fine needle aspiration biopsy could not rule out a papillary lesion. Thus, she was advised to undergo total thyroidectomy, and her final histopath showed a simultaneous medullary and papillary thyroid carcinoma. Her initial serum calcitonin was elevated at 252 pg/ ml, and it remained persistently elevated over the course of 7 months. A repeat ultrasound revealed solid nodules with coarse calcifications and enlarged lymph nodes at both submandibular regions. This warranted a repeat surgery with neck dissection with the finding of eight lymph nodes positive for metastatic carcinoma. On follow up after her second surgery, the calcitonin decreased to 42.70 pg/ml.

Knowledge of this simultaneous occurrence of medullary thyroid carcinoma and papillary cancer is important for its prognostic implications and therapeutic plan.

Key words: thyroid cancer, papillary thyroid cancer, medullary thyroid cancer

INTRODUCTION

The cell origin, histopathologic features, and prognosis of medullary thyroid carcinoma and papillary thyroid carcinoma are different and to have them occur simultaneously in a single patient is a rare event. Data about the simultaneous occurrence of these tumors are based mainly on case reports and series.

CASE

This is a case of a 38-year-old female who has simultaneous medullary thyroid cancer in the left thyroid lobe and papillary thyroid cancer in the right thyroid lobe. She was diagnosed to have thyrotoxicosis at 18 years old presenting with palpitations and an anterior neck mass. She was maintained on an unrecalled dose of carbimazole and propanolol for 1 year which was eventually discontinued since her thyroid function test normalized, with noted decrease in size of her anterior neck mass.

Five years prior, she began noticing an enlarging anterior neck mass. There was no associated hoarseness of voice, dysphagia, weight loss, palpitations or difficulty of breathing. She consulted an endocrinologist and work up was done. Thyroid function test showed an elevated TSH at 11.84 uIU/ml and a normal FT4 at 15 uIU/ml. Her ultrasound showed an enlarged left thyroid gland with a 2.38 x 1.4 x 1.69 solid nodule with calcifications

and minimal hypervascularity. Few coarse parenchymal calcifications are noted at the mid portion of the right lobe. She underwent fine needle aspiration biopsy (FNAB) of her left thyroid lobe with a result of an adenomatous goiter with concomitant lymphocytic thyroiditis. She was given Levothyroxine 50 mcg daily with serial monitoring of her thyroid function test and ultrasound.

Repeat ultrasound two years later showed interval increase in the size of the previously noted solid nodule with calcifications in the left lobe, with new nodules noted on the right lobe. This warranted a repeat biopsy. Repeat ultrasound guided fine needle aspiration biopsy of the nodule on the right thyroid gland showed nodular hyperplasia but cannot rule out papillary lesion. While the FNAB of the nodule on the left showed nodular hyperplasia of left thyroid lobe. Thus, she underwent total thyroidectomy.

Microscopic examination of the isthmus and left lobe were consistent with medullary carcinoma, with the tumor size 6 cm in its greatest dimension. The right lobe showed papillary thyroid carcinoma, with the tumor size in greatest dimension 1.1 cm, with no lymphovascular or perineural invasion (Figure 1). Immunohistochemistry report for the isthmus and left lobe were chromogranin positive, calcitonin positive supporting the diagnosis of medullary thyroid carcinoma.

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Figure 1. Histopath. **(A)** Areas of normal parenchyma, thyroiditis, and papillary carcinoma (H&E, 40x). **(B)** Papillary Thyroid carcinoma, Right lobe. Overlapping clear ground glass nuclei with eosinophilic inclusions (H&E, 400x). **(C)** Medullary carcinoma. Cells with eccentrically located round nuclei with salt and pepper nuclear chromatin pattern (H&E, 100x).



Figure 2. (A) CT scan of the neck with contrast. Calcification in the left infrahyoid carotid space possibly calcified lymph nodes. (B) Neck ultrasound. Nodes with calcifications in the left parajugular region (level 3).

After the surgery, serum thyroglobulin was 0.307 ng/ml with normal antithyroglobulin 90.859 IU/ml, while serum calcitonin was elevated at 252 pg/ ml. The elevated serum calcitonin warranted a work up for possible metastasis. Thus, a neck ultrasound was requested which showed post thyroidectomy. Solid nodules with coarse calcifications, left parajugular region. Enlarged lymph nodes, both submandibular regions (Figure 2).

Findings on CT scan of the neck, chest and whole abdomen were nonspecific. CT scan of the neck revealed no evidence of cervical lymphadenopathy (Figure 2).

Chest CT scan showed nonspecific nodules along the left lung fissure and along the posterior hemithoraces. Whole abdomen CT scan showed mild hepatosplenomegaly, uterine myoma and Nabothian cyst. For further workup, a whole-body fluorodeoxyglucose PET scan was requested. Results showed no hypermetabolic malignant looking disease in the neck and elsewhere. The mildly hypermetabolic calcified left submandibular and left supraclavicular lymph nodes look inflammatory/reactive.

However, serum calcitonin on serial monitoring was persistently elevated (362, 197 pg/ml) over the course of 7 months (Table 1). A repeat thyroid ultrasound showed solid nodules (4) with coarse calcifications at the left parajugular area (level 3).

Thus, she underwent neck exploratory dissection, with the finding of eight of twenty-five lymph nodes with metastatic carcinoma. On follow up three months after

Table 1. Laboratory Results			
	April 2017	May 2017	July 2017
Calcitonin (0 – 11.5 pg/ml)	252	362	197
Thyroglobulin (2 – 70 ng/ml)	0.307		
Antithyroglobulin (up to 100 IU/ml)	90.859		

the operation, the calcitonin decreased to 42.70 pg/ ml. Nature and prognosis of papillary and medullary thyroid carcinoma were explained to the patient, with plans of monitoring for recurrence. Medullary thyroid carcinoma can occur sporadically or it may be associated with hereditary syndromes such as Multiple Endocrine Neoplasia Type 2. Thus, molecular studies investigating RET germline mutations in exons 11, 15 and 16 were performed, and patient was found to be positive for a mutation in exon 15. She is now recommended for work up for pheochromocytoma and hyperparathyroidism to screen for multiple endocrine neoplasia 2 syndromes.

DISCUSSION

Papillary thyroid carcinoma (PTC) is the most common type of thyroid carcinoma (90%), and it originates from the follicular cells of the endoderm. Medullary thyroid carcinoma, on the other hand, arises from the parafollicular C cells of the ultimobranchial body of the neural crest, and it represents only 5% of all thyroid cancers.¹

The simultaneous occurrence of an independent medullary thyroid carcinoma, a tumor showing morphological features of medullary thyroid carcinoma (MTC) with positive immunoreactive calcitonin cells, along with papillary carcinomas in the same thyroid gland is uncommon, but it has been reported.

Although the simultaneous occurrence of papillary and medullary thyroid carcinoma in one patient may be entirely coincidental, several authors have tried to explain as to why it happens. According to Erhamamci (2014), the simultaneous occurrence of MTC and DTC in the same thyroid gland can occur in two ways: as a mixed tumor showing dual differentiation or as a collision tumor showing two separate different carcinomas.² They concluded that out of 1420 patients, 0.28% of patients with DTC displayed simultaneous MTC.² Machens (2011) followed 1019 patients diagnosed with papillary thyroid carcinoma, and found 26 patients had simultaneous MTC with a prevalence rate of 2.6.³

Previous studies have also mentioned several other theories about this concurrence. The first is the 'stem cell theory' which describes a common stem cell that transform into both follicular and C-cell lineages.⁴ Another is the 'field effect theory' which proposes that simultaneous transformation of both follicular and C cells is a result of common neoplastic stimuli. On the other hand, the 'collision theory' suggests that two independent tumours are located in the same thyroid by simple coincidence.⁴

Papillary carcinoma generally has a good prognosis since most patients are diagnosed during the early stage of the disease (stage I or II), with 5% having a lethal outcome.⁵ Medullary thyroid carcinoma, on the other hand, may have an indolent clinical course or an aggressive variant with a high mortality rate.5 Distant metastases are present in 13% of patients at initial diagnosis and portend a poor prognosis, with a 10-year survival rate of only 40%.5 MTC is difficult to cure with a high recurrence rate of 50%.5 According to WHO, the prognosis of mixed medullary and follicular thyroid carcinoma depends upon the medullary component. Thus, the presence of the medullary component makes the prognosis worse as compared to a pure papillary carcinoma. Patients who have clinically evident disease are best treated with a minimum of a total thyroidectomy and bilateral central neck dissection.5 The use of adjuvant radioiodine therapy for the papillary carcinoma component of this disease has been mentioned in several case reports with good results. In a study by Diogini et al. (2007), they presented a 65-year-old man with multicentric PTC and MTC associated with diffuse lymphocytic-type thyroiditis who underwent total thyroidectomy with neck dissection and subsequent radioiodine treatment.⁵ This patient had normal thyroglobulin and calcitonin levels on follow up.⁵

CONCLUSION

Knowledge of this simultaneous occurrence of medullary thyroid carcinoma and papillary cancer is important because of the poor prognosis associated with medullary thyroid carcinoma. This report also highlights the importance of immunohistochemical markers to make a correct diagnosis.

Ethical Considerations

Patient consent was obtained before submission of the manuscript.

Statement of Authorship

All authors certified fulfillment of ICMJE authorship criteria.

Author Disclosure

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