Multiple endocrine neoplasia type 2A with cutaneous lichen amyloidosis

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DESCRIPTION
A 33-year-old woman presented with a decade-long history of a painless, gradually increasing neck swelling not associated with any compressive symptoms. She also complained of intermittent unprovoked episodes of headache, palpitations, anxiety and sweating for the past 6 months. She had similar episodes 3 years back, and perusal of old documents revealed that she had undergone right adrenalectomy for pheochromocytoma. At that time, she was also documented to have elevated serum calcitonin (1271 (<6 pg/mL)), and multiple endocrine neoplasia type 2 (MEN 2) was suspected. However, further evaluation could not be done as the patient was lost to follow-up due to financial constraints. There was history of goitre in her maternal grandmother, mother and maternal uncle. Her mother and maternal uncle had also been operated for adrenal tumour and were advised neck surgery, which they had refused. On examination, she was normotensive. A firm and diffuse goitre was palpable in the neck associated with level II cervical lymphadenopathy on the left side. A hyperpigmented, pruritic, velvety plaque with fine scales was noted in the right scapular region (figure 1).

MEN 2A with cutaneous lichen amyloidosis (CLA) was suspected. Fine needle aspiration cytology from the thyroid revealed medullary thyroid carcinoma (MTC), Bethesda category VI. Serum calcitonin (6715 (<6 pg/mL)), 24 hours of urine metanephrines (963.53 (74–297 µg/day)) and normetanephrines (1063 (73–808 µg/day)) were elevated. She was normocalcemic with a normal intact parathyroid hormone level. Contrast-enhanced CT scan of neck, chest and abdomen revealed multiple thyroid nodules, with the largest nodule being 1.5×1.2 cm in size with calcification, nodule in the right upper lobe of the lung and 5×3.5×1.7 cm left...

Figure 1 Cutaneous lichen amyloidosis in multiple endocrine neoplasia type 2A. Hyperpigmented, velvety plaque with fine scales in the right scapular region.

Figure 2 (A) Maximum intensity projection image of 68Ga DOTANOC PET-CT showing focal areas of radiotracer uptake in the neck region and left suprarenal region, corresponding hypodense lesions in both the lobes of the thyroid (B,D) with the left one showing coarse calcifications. SSTR expression was noted in both the lesions in fused PET-CT images (C,E). Axial CT abdomen (F) showing enlarged left adrenal gland with SSTR expression in fused PET-CT image (G). Axial CT thorax (H), showing a nodule in the right lung upper lobe apical segment with SSTR expression in fused PET-CT image (I). PET, positron emission tomography; SSTR, somatostatin receptor.

Patient’s perspective
It has been explained by my doctors that I have a malignant cancer syndrome which runs in families and involves thyroid, adrenal and parathyroid glands. It is crucial to detect thyroid malignancy early to avoid involvement of other glands and spread of the disease to other organs. There is a family history of goitre in relatives on my maternal side, but I ignored my symptoms and now have advanced disease. I am now convinced that timely screening of at-risk family members and prophylactic thyroidectomy of family members who have the genetic mutation can be curative.
adrenal lesion with heterogeneous contrast enhancement. These lesions showed increased somatostatin receptor expression on $^{68}$Gallium-DOTANOC PET-CT scan (figure 2) consistent with MTC with lung metastasis and left adrenal pheochromocytoma. Genetic testing confirmed RET (REarranged during Transfection) codon 634 mutation confirming the diagnosis of MEN 2A.

She underwent laproscopic left adrenalectomy followed by total thyroidectomy with modified radical neck dissection and pulmonary metastasectomy and started on hydrocortisone, fludrocortisone and levothyroxine replacement. She is on outpatient follow-up. Her younger brother and daughter have tested negative for targeted RET codon 634 mutation.

MEN 2A with cutaneous lichen amyloidosis (CLA) is a rare MEN 2A variant almost exclusively associated with RET codon 634 mutation. About 35% of patients with this mutation have CLA in their lifetime, which manifests as an intensely pruritic, scaly and pigmented rash, typically located in the interscapular region.\(^1\)\(^2\) It is a precocious marker for the presence of a mutated RET allele and the impending development of MTC in a MEN 2A kindred.\(^3\)

Learning points

- There has been a paradigm shift in the diagnosis of multiple endocrine neoplasia type 2 (MEN 2) syndrome with the discovery of RET gene. The disease course can be predicted based on the RET point mutation. Targeted screening approach should be used for at-risk family members, and those positive for mutation should be convinced for thyroidectomy.
- Non-endocrine manifestations like CLA should be carefully looked for in a MEN 2A kindred as it may predate the onset of MTC by years.