A Carotid Body Tumor Mimicking a Thyroid Nodule

Husniye Basera, Baris Ayhanb, Meryem Ilkay Eren Karanisc, Salih Baserd, Deniz Karasoyd, Kemal Kalkand, Samil Ecirlid

Abstract

Paragangliomas (PGLs) are neuroendocrine tumors arising from the extra-adrenal chromaffin tissue of the autonomic nervous system. They are most frequently found in the head and neck, mainly associated with the carotid body, vagus nerve, jugulotympanic paraganglia, and occasionally, the superior-inferior paraganglia. Paragangliomas are rarely encountered in thyroid as well, and thyroid paragangliomas can be misinterpreted as medullary thyroid carcinomas. However, cases of paragangliomas mimicking thyroid nodules were also reported in literature rarely. In this article, we reported a case first giving an impression of a thyroid nodule, then suggesting the likelihood of a medullary thyroid carcinoma or intrathyroidal paraganglioma via core needle biopsy, but finally was diagnosed as a carotid body tumor.

Keywords: Paraganglioma; Carotid body tumor; Thyroid nodule

Introduction

Paragangliomas (PGLs) are uncommon tumors, incidence rate is 1 to 2 per 100,000, and the tumors are often given special designations, depending on their locations [1]. Only 3% of all PGLs occur within head and neck, the majority of which are located in carotid body (carotid body tumors), temporal-bone/middle-ear (glomus jugulare) and vagus nerve in neck (vagal PGLs) [2, 3]. Ninety percent of head and neck PGLs are sporadic, while only 10% are hereditary in nature [2]. In literature, cases of PGL mimicking thyroid nodules are rarely encountered [4, 5]. In the report, we will present a case of PGL (a carotid body tumor) that we initially evaluated as a thyroid nodule, and then diagnosed as a PGL.

Case Report

A 74-year-old woman was admitted due to a mass in the right side of neck growing rapidly within the last 1 month. The case reported no complaints of dysphagia or dysphonia. The family history of the case revealed no carcinoma and thyroid disease. While no history of flushing, hypertensive attacks, diarrhea, and other symptoms related to catecholamines hypersecretion was detected, hypertension was present in her personal medical history. She was lack of the history of neck irradiation. Physical examination revealed a hard and painless mass with smooth surface, extending from right thyroid lobe level to angulus mandibula and measuring 8 cm without palpable cervical lymph nodes. Indirect laryngoscopy showed normal mobile vocal cords. Thyroid stimulating hormone, free triiodothyronine and free tetraiodothyronine...
were within normal ranges, and anti-peroxidase and anti-thyro
globulin antibodies were found to be negative. Ultrasound
(US) examination showed two hypoechoic and non-homoge-
neous nodules displaced the trachea to left in the right thyroid
lobe, one in size of $46 \times 37$ mm and the other in size of $44 \times 38$ mm, and a isoechoic nodule in size of $23 \times 15$ mm in
the left thyroid lobe. For 3 nodules, a US-guided fine needle
aspiration biopsy (FNAB) was performed, and the nodules
were reported to be benign. On neck magnetic resonance
imaging (MRI) (Fig. 1), a mass of $7 \times 5.5$ cm with intense
pathologic contrast, extending from right thyroid lobe level
to angulus mandibula, and displaced larynx and trachea to
left was observed. Because of the existence of rapidly grow-
ing thyroid nodules in the case, core needle biopsy was per-
formed for the nodules in right thyroid lobe due to suspected
thyroid malignancy, and histopathologic findings were found
to be consistent with neuroendocrine tumor. Immunohisto-
chemical examination revealed that synaptophisin and chro-
mogranin A were positive, but calcitonin, thyroid transcrip-
tion factor-1 (TTF-1) and thyroglobulin (Tg) were negative.
The level of serum calcitonin was 2.8 pg/mL (normal range
0-11.5 pg/mL), fractionated urinary catecholamine levels
were normal, and on positron emission tomography (PET)-
computerized tomography (CT), increased 2-[fluorine-18]-
fluoro-2-deoxy-D-glucose (FDG) involvement was observed
in thyroid (SUVmax 7.8) and right paratracheal, aortopul-
monary, subcarinal and bilateral hilar regions (SUVmax
15.5). The excision of mediastinal lymphnodes and bilateral
total thyroidectomy were performed. During the surgery, a
mass of $4 \times 3$ cm was seen in the right carotid artery bifurca-
tion and excised. Macroscopically, the mass was encapsu-
lated, yellowish-white and soft. Microscopically, the tumor
cells were arranged in well-defined nests (Zellballen) and
encircled by a thin layer of S-100 protein and GFAP posi-
tive, spindle-shaped sustentacular cells (Fig 2, 3). Tumor
cells vary in size and shape, and have a finely granular cyto-
plasm. The nuclei were round to oval with coarsely granular
chromatin with a so-called salt-and-pepper appearance. No
definite pleomorphism, necrosis or mitosis was witnessed.
In tumor cells, immunohistochemical CD56, synaptophisin
(Fig. 4), neuron specific enolase (NSE) and chromogranin
A (Fig. 5) were positive, but calcitonin, TTF-1 and Tg were
negative. The proliferation index of Ki-67 was lower than
1%. In light of these findings, the case was diagnosed with
PGL, and the histopathologic findings of other thyroid nod-

Figure 2. Microscopic photograph of paraganglioma, HE ×200.

Figure 3. Immunstain for S-100 protein shows positive staining in sustentacular cells, S-100 ×400.

Figure 4. Immunstain for Synaptophysin shows positive staining in tumor cells, Synaptophysin ×200.

Figure 5. Immunstain for chromogranin A shows positive staining in tumor cells, Chromogranin A ×200.
ules were consistent with colloidal nodules. In addition, the histopathology of mediastinal lymph nodes was evaluated as reactive hyperplasia.

Discussion

PGLs are rare tumors derived from extra-adrenal paraganglionic system, which consist of cells from the neural crest associated with autonomous nervous system [6, 7]. PGLs are seen at many sites in head and neck areas, including carotid body, jugular-typanic region and vagus nerve [7, 8]. Unlike pheochromocytomas, PGLs in head and neck regions are usually non-functioning tumors.

The classic radiographic features are homogeneous or heterogeneous hyper-enhancing soft-tissue mass as shown by CT scan, multiple areas of signal void interspersed with hyperintense foci (salt-and-pepper appearance) within a tumor mass as shown by MRI, and as an intense tumor blush with enlarged feeding arteries by angiography [9].

Histopathologically, PGLs have a characteristic growth pattern often referred to as a “zellballen” growth pattern. Tumor cells are predominantly chief cells with round, hyperchromatic nuclei, a dispersed chromatin and abundant granular cytoplasm ranging from eosinophilic to basophilic in color. Neurromelanin pigment may occasionally be seen, and amyloid deposition has also been described in PGLs. Like the pheochromocytes of adrenal pheochromocytomas, tumor cells demonstrate reactivity to chromogranin and synaptophisin stains by immunohistochemical techniques, as well as other markers of neuroendocrine differentiation such as CD56 and NSE. The population of sustentacular cells can usually be identified at the periphery of nests and are thought to be modified Schwann cells and to be spindle-shaped, and can be highlighted with S-100 protein staining.

Among the disorders most easily mistaken for cervical PGLs, particularly those arising within larynx or thyroid, are neuroendocrine carcinomas, such as atypical carcinoids and medullary thyroid carcinoma (MTC) [10-15]. Another disorder that may be confused with PGLs is hyalinizing trabecular adenoma, a benign thyroid neoplasm showing a prominent nesting pattern resembling a PGL [14, 16, 17]. This rare entity is usually negative for chromogranin and positive for Tg and cytokeratins.

In literature, PGLs are generally reported as slowly growing neck masses without pain [7, 18]. Rarely, they may attain large size and infiltrative growth, and the local recurrence of PGLs may lead to death. Our case was admitted first as a thyroid nodule on US imaging, a carotid body tumor was observed in the right carotid artery bifurcation during the operation; on the other hand, thyroid histopathology was benign.

FNAB can be performed on numerous occasions to establish a diagnosis of PGLs [19-24]. However, the exact role of FNAB in the diagnosis of these tumors currently remains controversial. The main concern in performing a pre-operative FNAB on suspected PGLs is the risk of hemorrhage and hematoma [22, 24]. In our case, carotid body tumor suggested the impression of a thyroid nodule, so FNAB was performed, but no complication was seen as a result of biopsy.

Malignancy cannot be determined by cytology or even by histologic findings and can only be defined when the tumor metastasizes to regional lymph nodes or more distant sites. Nuclear pleomorphism, neurovascular invasion, high mitotic index and necrosis have been described in both benign and malignant types of tumors [2]. In literature, the ratio of proven cervical malignant PGLs is low ranging from 2 to 10% for carotid and vagal body tumors and is less than 2% for laryngeal PGLs [11, 12]. In our case, the determination of mediastinal lymphadenopathies on PET-CT suggested the likelihood of a malignant PGL; however, no malignancy was encountered in the histopathology of mediastinal lymph nodes.

In conclusion, cervical PGLs are uncommon tumors, so healthcare providers should take the likelihood of PGLs into account in the differential diagnosis of neck masses.

Conflicts of Interest

Authors declare that there is no conflict of interest.

References