Laryngeal Neuroendocrine Tumor: Atypical Presentation

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ABSTRACT

Neuroendocrine tumors of the larynx are the most common nonepidermoid tumors of the larynx and comprise less than 1% of the laryngeal tumors. Most of the symptoms and presentation mimic a usual laryngeal malignancy making the diagnosis difficult.


INTRODUCTION

Neuroendocrine tumors (NETs) of the larynx, which represent a heterogeneous group of neoplasm, are rare, but still are the second most common neoplastic lesion of the larynx. The first case of NET was reported by Goldman et al in 1969. Almost 500 cases of NETs have been reported in the literature so far.1 The origin of these tumors is obscure with many theories of origin advocated with the most commonly accepted being that they originate from the precursor cells of neuroendocrine system. This spectrum of condition forms the most common nonepidermoid carcinomas of the larynx.2 The most important risk factor associated is smoking. Here, we report a case of laryngeal neuroendocrine carcinoma that was managed with total laryngectomy.

CASE REPORT

A 50-year-old male presented to the emergency department with progressive difficulty in swallowing and change in voice since 3 months. He also complained of difficulty in breathing since the last 15 days, which had progressed to a frank stridor for which he was referred to the ENT department where he underwent an emergency tracheostomy. On general examination, he was conscious with pulse rate of 84 beats/minute and respiratory rate of 18/minute, and chest examination revealed conduct ed sounds over the chest. Indirect laryngoscopy showed an ulceroproliferative growth involving the left aryepiglottic (AE) fold and piriform sinus and extending to the midline causing edema in the left false vocal cord causing reduced glottis chink. On hematological investigation, he had hemoglobin of 105 gm/L with ESR of 55/hour. His liver function, kidney function, and other biochemical parameters were within the normal limits. Contrast-enhanced computer tomography (CECT) of neck showed soft tissue lesion with smooth outlines in left piriform fossa with involvement of left AE fold. The mass showed heterogeneous attenuation with no evidence of calcification or cavitation (Figs 1A and B). A CECT scan of chest, abdomen, and brain was done to know the staging and to rule out metastasis. The tumor was found to be limited to the laryngeal framework with no distant metastasis. The TNM staging of the tumor was deemed to be T3N0M0. He underwent a direct laryngoscopic biopsy under general anesthesia, which revealed features of NET. However, further grading of the tumor could not be done on the biopsy specimen. Thereafter, the patient was worked up and counseled for a total laryngectomy. The surgery was performed and no intraoperative complications were noted. The immediate postoperative period was uneventful.

Upon final histopathologic examination, there was an ulceroproliferative lesion involving the left AE fold and piriform sinus measuring 2 × 2 × 1.5 cm, which stained intensely with epithelial membrane antigen, neuron-specific enolase (NSE), and chromogranin (Fig. 2). Thus, a diagnosis of atypical carcinoid tumor was made. The patient, however, expired due to sudden cardiac arrest 10 days postoperatively, so radiotherapy and follow-up could not be done.

DISCUSSION

Neuroendocrine tumors of the larynx are the most common nonepidermoid tumors of the larynx and comprise less than 1% of the laryngeal tumors.2 The male to female ratio is 3:1, and the most common risk factor associated with this condition is smoking.3

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The spectrum of NETs can only be differentiated from one another histopathologically and they constitute paragangliomas, atypical carcinoids, small and large cell neuroendocrine carcinoma. The most frequent site involved is the supraglottis. Atypical carcinoid tumor is the most commonly reported variant in the literature, with the typical carcinoid being the rarest.

Most of the symptoms and presentation mimic a usual laryngeal malignancy and they can be distinguished only upon histopathologic examination. Features suggestive of NETs are neurosecretory granules, cells with variable degrees of pleomorphism and organization into nests, cords and trabecules with submucosal origin of the neoplasm, evidence of mucin (Periodic acid–Schiff+) and immunohistochemical positivity for neuroendocrine markers, such as chromogranin, NSE, synaptophysin, and neurofilaments. Positive staining for calcitonin in neuroendocrine carcinoma is usually the norm; however, in our case, this was not so. Negative staining is suggestive for paragangliomas; however, in our case, the other histopathological features suggested atypical carcinoid tumor.

Other nonspecific markers for paragangliomas like urinary levels of catecholamines could also be done to confirm the diagnosis. Urinary levels of catecholamines and some metabolites (vanillylmandelic and homovanillic acid) are often raised in paraganglioma, whereas increased levels of 5-HIAA, NSE, and CgA can be diagnostic for carcinoids. The most frequent sites of metastasis are lymph nodes, skin, liver, and lung.

The therapeutic approaches to NETs of the larynx vary according to the biological behavior of the various histological types and disease stage. Conservative surgery is the standard treatment for local typical carcinoid and paraganglioma but locoregional lymph node dissection is not indicated because metastatic lymph node involvement is rare and the long-term prognosis is good. In NETs and atypical carcinoids, depending upon the extent of disease, total or subtotal laryngectomy with lymph node dissection is the most appropriate treatment for carcinoma. The overall 5-year survival rate is 48%. The role of radiotherapy and/or chemotherapy is still doubtful, as these tumors are usually resistant.

We would like to highlight this case as laryngeal NETs form a relatively rare entity when compared with the commoner epidermal origin tumors. Our case was also unique in the sense that even though all histopathologic features pointed toward atypical carcinoid tumor, calcitonin negativity usually indicates paragangliomas. Thus, in cases of atypical presentations, optimal management of this subgroup of tumors can be difficult.
REFERENCES