Primary neuroendocrine tumors of the ear, nose and throat: A report of three cases and a review of the literature

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Abstract. The aim of the present study was to review all cases of neuroendocrine tumors of the ear, nose and throat in a tertiary care center, as well as the data published in the literature. The study presents all the cases of neuroendocrine tumors (NETs) in the Hotel Dieu De France Hospital (Beirut, Lebanon) between January 2004 and January 2014. The data reported in the English and French literature is also reviewed with regard to the typical clinical presentation and management of these tumors. Three cases of NETs presented to the Department of Otolaryngology-Head and Neck Surgery during the study period. One case was of an atypical carcinoid (AC) tumor of the larynx, one case was of a typical carcinoid tumor in the middle ear and the third case was, to the best of our knowledge, the first reported case of an AC tumor of the nasopharynx. Overall, NETs are rare in the head and neck. The clinical presentation can mimic any other tumor in the same localization in the absence of a carcinoid syndrome. Management of these tumors remains controversial, but a complete excision of the tumor is crucial, followed by possible adjuvant treatment.

Introduction

Neuroendocrine tumors (NETs) are rare tumors arising from enterochromaffin (Kulchitsky) cells, which are part of the amine precursor uptake and decarboxylation system. The tumors are thus found in locations with a wide distribution of these cells, such as in the gastrointestinal tract, lungs and bronchi (1).

The World Health Organization (WHO) classification of NETs groups the tumors into those of epithelial origin, namely typical (well-differentiated) carcinoid (TC) tumors, atypical (moderately-differentiated) carcinoid (AC) tumors and small cell (undifferentiated) neuroendocrine carcinoma (SmCC), and those of neural origin, namely paraganglioma (2).

The clinical and pathological features of NETs are characteristic of the organ of origin, however, these tumors can share other attributes irrespective of their anatomical site (3). Although NETs may have a similar presentation to other head and neck tumors in the same locations, their behavior and management are not clearly established and vary according to their histological type (4). Functioning tumors, particularly carcinoid NETs, can present with symptoms caused by hormone secretion; for example, patients may present with carcinoid syndrome, which is characterized by flushing, diarrhea and abdominal pain (3,5). A diagnosis of carcinoid NET is suspected when features of carcinoid syndrome are present and high urinary 5-hydroxyindoleacetic acid levels are identified, with histological analyses used to confirm the diagnosis (3,5). Therapeutic strategy selection depends on the site of origin of the tumor and its extent. Treatment usually consists of surgery and/or radiotherapy with or without chemotherapy (6). Rarely, NETs occur in the head and neck region, predominantly in the larynx and the middle ear, however, few cases have been described in the nasopharynx (2).

In the present study, three cases of head and neck NETs in the three different aforementioned locations are described along with their management and follow-up.

Case report

Data collection. All head and neck NET cases treated in Hotel Dieu de France Hospital (Beirut, Lebanon), between January 2004 and January 2014, were retrospectively reviewed. Data regarding clinical presentation, management, pathology results and follow-up were retrieved from hospital charts. A total of three cases were found. Written informed consent was obtained from all patients prior to surgery and prior to writing this study. A review of the English and French literature regarding NETs of the head and neck region was performed. This review was based on a search of the US National Library of Medicine (PubMed) between 1990 and 2014.

Case 1. A 64-year-old Caucasian male presented in June 2007 with hoarseness and dysphagia that had persisted for 2 years.
The patient had smoked 2 packs of cigarettes per day for 20 years until quitting 20 years ago. The patient denied a history of alcohol consumption. The patient also experienced moderate dyspnea, particularly when in the left lateral decubitus position. A total of 20 kg of weight loss had been noted over the last 2 years. At 1 month prior to the current visit, the patient developed right-sided otalgia. A laryngoscopy revealed a mass in the right aryepiglottic fold (Fig. 1A). An endoscopic excisional biopsy of the lesion was performed. The tumor was composed of tubular structures and compact cords. Moderate anisonucleosis was present and the cells exhibited powder-like chromatin. Few mitotic and apoptotic cells were noted. Stroma was mildly abundant and endocrinoid in aspect. Periodic acid-Schiff coloration showed no intracytoplasmic mucosecretions. Immunohistochemistry was positive for chromogranin A, synaptophysin and cytokeratin 7 (CK7). The final diagnosis was of an AC tumor of the larynx in the right aryepiglottic fold.

Radiotherapy was suggested, but the patient refused this therapeutic option. After 3 years, recurrence of the disease was noted on the right arytenoid cartilage, which was treated by endoscopic resection. The pathology report showed recurrence of the AC tumor, with negative resection margins. The patient was then followed up regularly. At the 5-month follow-up visit, a mass was observed in the right arytenoid and aryepiglottic fold on direct laryngoscopy (Fig. 1B). Endoscopic excision of the lesion was performed using a CO₂ laser. The pathology report once again showed recurrence of the AC tumor, with clean resection margins. The patient then underwent 28 sessions of adjuvant radiotherapy (56 Gy) for a total of 4 weeks and is currently free of disease 9 months after this treatment.

Case 2. A 43-year-old Caucasian female presented in July 2006 with the sensation of ear fullness and mild hearing loss that had persisted for 5 months, but with no otorrhea, vertigo or tinnitus. A physical examination showed a white retro-tympanic mass in the right ear and conductive hearing loss in the same ear, confirmed by an audiogram. A computed tomography (CT) scan showed a well-defined mass in the middle ear, with no erosion of the ear ossicles. The patient subsequently underwent a radical mastoidectomy with excision of the middle ear mass.

The tumor was composed of a tubular and trabecular proliferation with no patent signs of cytological malignancy. Immunohistochemistry was positive for chromogranin A, synaptophysin and cytokeratin 7 (CK7). The final diagnosis was of a TC tumor of the middle ear. No clinical evidence of disease recurrence was found after 8 years.

Case 3. A 51-year-old Caucasian male presented in February 2013 with atypical vertigo that had first occurred 2 months previously. The patient had smoked 2 packs of cigarettes per day for 30 years and denied a history of alcohol consumption. No additional complaints, other than an old history of chronic nasal obstruction, were noted. Otoscopy, anterior rhinoscopy and an oral cavity examination were normal, as was the neck palpation. Brain magnetic resonance imaging was performed for the workup of the vertigo, which showed no abnormal findings in the brain parenchyma, but
Table I. Diagnostic criteria of neuroendocrine tumors in the head and neck region.

<table>
<thead>
<tr>
<th>Criteria</th>
<th>TC</th>
<th>AC</th>
<th>SmCC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neuroendocrine differentiation</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Mitotic count</td>
<td>0-1/10 HPFs</td>
<td>2-10/HPFs</td>
<td>&gt;10/10 HPFs</td>
</tr>
<tr>
<td>Ki-67</td>
<td>&lt;2%</td>
<td>3-20%</td>
<td>&gt;20%</td>
</tr>
<tr>
<td>Cytoplasm amount</td>
<td>Moderate</td>
<td>Moderate</td>
<td>Little/scanty</td>
</tr>
<tr>
<td>Nucleoli</td>
<td>Inconspicuous</td>
<td>Inconspicuous</td>
<td>Inconspicuous</td>
</tr>
<tr>
<td>Nuclear polymorphism</td>
<td>Little</td>
<td>Moderate</td>
<td>Moderate</td>
</tr>
<tr>
<td>Necrosis</td>
<td>No</td>
<td>Focal punctate or mild</td>
<td>Marked</td>
</tr>
</tbody>
</table>

TC, typical carcinoid tumor; AC, atypical carcinoid tumor; SmCC, small cell neuroendocrine carcinoma; HPFs, high-power fields.

Discussion

Neuroendocrine neoplasms are rare tumors that are mainly found in the GI tract, pancreas and lungs (5).

Numerous proposals have been put forward with regard to the classification and nomenclature of NETs, and a number of these differ in their use of specific terminology and the criteria for grading and staging (3). In order to predict the patient outcome and therapy, the use of a single system of nomenclature, grading and staging is required for NETs of all anatomical sites, as there are a number of similarities among NETs throughout the body. However, certain systems that have arisen independently have become firmly established and are recognized by organizations charged with standardizing terminology, such as the World Health Organization (WHO). No data favors one system over another (3). The basic data that underlie the systems are similar, even if the criteria differ.

NETs are rarely found in the head and neck, particularly the larynx, the middle ear and the nasopharynx. The WHO classification of laryngeal and middle ear NETs is generally consistent with that for pulmonary neuroendocrine carcinoma. The tumors can be subdivided into TC, AC, SmCC, combined SmCC with non-small cell carcinoma, and those with neuroendocrine cells have been considered as a separate subtype by one previous study (6). However, NETs of the nasopharynx are not included in any of the WHO classifications of nasopharyngeal tumors.

Histopathologically, the tumor was composed of an endocrine proliferation containing sheets and nests separated by a thin vascularized stroma. The tumor cells exhibited round nuclei containing salt and pepper-like chromatin. Cytoplasm was abundant and amphophilic. Mild nuclear pleomorphism was noted and punctuate areas of necrosis were present. Mitotic activity was low with a rate of 1/10 high-power fields (Fig. 2A). Immunohistochemistry was diffusely positive for chromogranin A (Fig. 2B), synaptophysin and pancytokeratin. There was no S100 staining of the tumor cells, nor of the sustentacular cells. Calcitonin, thyroid transcription factor-1 and caudal-type homebox-2 staining was also negative. The proliferation index was 3%. The final diagnosis was of an AC tumor of the nasopharynx.

Positron emission tomography-CT scanning with gallium-68-DOTA-NOC was performed 1 month post-operatively to search for other localizations and revealed positive uptake in the nasopharynx only. The urinary 5-hydroxyindoleacetic acid level was 9.1 mg/24 h (normal range, 2-6 mg/24 h). At 3 months post-surgery, the patient was administered 28 sessions of adjuvant radiotherapy (56 Gy) for a total of 4 weeks, and is currently free of disease at 14 months of follow-up.

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The study found that the more benign end of the spectrum is represented by TC tumors, and that local excision alone is curative. Radiotherapy does not induce a good response in patients with AC tumors, which are therefore best managed by radical surgical excision in combination with elective bilateral neck dissection due to the high propensity for regional metastasis. The most benefit for cases of SmCC or large cell neuroendocrine carcinoma appears to be gained from a combination of radiotherapy and chemotherapy, although patient survival remains poor (14).

Laryngeal paragangliomas are almost always benign and should be treated accordingly. Partial laryngectomy is preferable to radiation as a cure is usually achieved without loss of laryngeal function (15). Laser surgery is not widely used due to the vascular nature of these tumors (13).

Carcinoid tumors of the middle ear are primarily treated by surgical excision. There is no known established surgical approach, perhaps owing to the limited incidence of the condition, but complete removal by tympano-mastoidectomy or radical mastoidectomy is the most commonly used technique (16). In the literature, middle ear carcinoid tumors were successfully removed by tympanotomy alone in certain studies, while in other cases, a subtotal petrosectomy was performed (17). No sufficient data exist regarding the role of chemotherapy or radiation therapy in middle ear carcinoid tumors (17). Radiation therapy was used as adjuvant therapy in certain studies, but the number of cases was insufficient to draw a conclusion concerning better local control and recurrence rates (18).

In view of the small number of cases, no clear treatment policy has been established for neuroendocrine carcinomas of the nasal cavity, paranasal sinuses and nasopharynx. In almost all the cases reported in the English literature, the treatment consisted of surgery (7,19-25). In certain studies, conventional radiotherapy was used alone or in combination with either surgery or chemotherapy, but the efficacy of such regimens is yet to be proven (22). In a review by Furuta et al., it was proposed that radiation therapy and chemotherapy should be performed as post-operative adjuvant therapy when a complete resection is difficult and when surgery is ineffective (24).

NETs are rarely diagnosed in the head and neck region. The tumors are found mainly in the larynx, less frequently in the middle ear and scarcely in the nasal cavity or nasopharynx. While a well-defined classification exists for NETs of the larynx and the middle ear, these tumors are not mentioned in the most recent WHO classification of nasopharyngeal tumors. Clinicians should be aware of this tumor in the nasopharynx and should consider it in the differential diagnosis of tumors located there. Following the recent meta-analysis by van der Laan et al (14), treatment schemes for laryngeal NETs have been well established considering the low level of evidence on this subject. However further studies are required to establish treatment protocols for the middle ear and nasopharyngeal locations.

References