Sinonasal Carcinomas with Neuroendocrine Features: Histopathological Differentiation and Treatment Outcomes

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Keywords
► olfactory neuroblastoma
► esthesioneuroblastoma
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► sinonasal neuroendocrine carcinoma
► sinonasal small cell cancer
► sinonasal carcinoid
► sinonasal large cell cancer
► sinonasal cancers with neuroendocrine features
► sinonasal cancers of neuroectodermal origin

Abstract

Objectives Sinonasal cancers with neuroendocrine features share similar clinical, radiological, and histopathological features; however, these tumors often exhibit varying degrees of aggressive behavior presenting significant treatment challenges. The objective of this study was to report our experience with these rare malignancies and to present a review of current literature.

Methods Following institutional review board approval, the records of all patients with biopsy-proven sinonasal malignancies over a 5-year period were reviewed.

Results The study included 14 patients with olfactory neuroblastomas (ONBs), 7 patients with sinonasal undifferentiated carcinomas (SNUC), and 2 patients with sinonasal neuroendocrine carcinomas (SNEC). Histopathologic markers aided in final diagnosis, but showed variable specificity. In patients with sufficient follow-up, the 2-year disease-free survival rate was 81% (9/11) for ONB and 75% (3/4) for SNUC. Three patients developed a regional or distant recurrence (two with ONBs and one with SNUC). Two patients, one with SNEC and one with ONB, succumbed to brain radionecrosis related to proton radiation therapy.

Conclusions Overlapping clinical and histopathological features in poorly differentiated sinonasal cancers with neuroendocrine features continue to present a diagnostic challenge. Individualized assessment and treatment strategies can improve the accuracy of the initial assessment and the treatment outcomes.
Introduction

Sinonasal malignancies with neuroendocrine features are rare head and neck tumors that can be divided into three major histological phenotypes: olfactory neuroblastoma (ONB), sinonasal undifferentiated carcinoma (SNUC), and sinonasal neuroendocrine carcinoma (SNEC). SNEC meets all of the diagnostic criteria that define neuroendocrine tumors, while ONB and SNUC exhibit some neuroendocrine features.

The defining features for neuroendocrine neoplasms continue to be a source of ongoing debate due to the fact that some neuroendocrine markers are nonspecific and can be expressed in nonneuroendocrine malignancies. Bell et al. reported the following features as diagnostic for neuroendocrine malignancies:

- Histological features of neuroendocrine differentiation (stippled chromatin, rosette formation, organoid, and trabecular growth)
- Focal or widespread expression of immunohistochemical neuroendocrine markers with synaptophysin and chromogranin being more specific
- Some degree of epithelial differentiation as evidenced by the expression of cytokeratin

SNEC is the only true “neuroendocrine” tumor as it meets all of the criteria required for diagnosis of neuroendocrine malignancies. ONBs are considered to be neuroectodermal in origin and are classified as such in the World Health Organization (WHO) classification of head and neck tumors; however, these tumors share some of the features present in neuroendocrine neoplasms and are often difficult to distinguish from SNECs. SNUCs can present with weak focal positivity for neuroendocrine markers, but in most cases completely lack any evidence of neuroendocrine markers. SNUCs are considered here due to overlapping morphology with poorly differentiated ONBs and SNECs.

SNEC is an umbrella term that encompasses both typical and atypical carcinoids, as well as small cell (SmCC) and large cell neuroendocrine carcinomas. The most commonly used WHO classification published in 2005 does not recognize large cell-type SNEC; however, the existence of these malignancies has been established in recent reviews suggesting future revisions of the existing WHO classification. Sinonasal malignancies with neuroendocrine features can be classified based on the cells of origin (Fig. 1). ONBs arise from neuroectodermal cells of the olfactory epithelium. Although, the cells of origin for SNUCs have not been identified unquestionably, SNECs and SNUCs are thought to arise from the respiratory epithelium of the sinonasal cavity.

SNECs can be further subdivided based on the cell size and differentiation. Well-differentiated SNECs are also known as carcinoid tumors, while moderately differentiated SNECs are classified as atypical carcinoid tumors (Fig. 1). SmCC and large cell neuroendocrine carcinomas are poorly differentiated tumors of epithelial origin.

Sinonasal carcinomas with neuroendocrine features share similar clinical and radiological attributes, and their histopathology often shows poorly differentiated morphology with several overlapping features, which creates diagnostic challenges (Fig. 2). A correct diagnosis is imperative for appropriate treatment and prognosis as these malignancies exhibit varying degrees of aggressive behavior. This study presents a case series of the sinonasal malignancies with neuroendocrine features and a review of the current literature with an emphasis on diagnosis, treatment strategies, and outcomes.

![Fig. 1](Sinonasal Carcinomas with Neuroendocrine Features: Classification)
Patients and Methods

Following approval by the institutional review board, medical records of the patients treated by the skull base team at the Ohio State University James Cancer Hospital were reviewed over a 5-year period. Records of histopathologically confirmed sinonasal carcinomas with neuroendocrine features were selected. All three types of carcinomas were staged using Kadish, Dulguerov/University of California, Los Angeles (UCLA) staging systems and the 7th American Joint Committee on Cancer (AJCC) TNM staging classification of the nasal cavity and paranasal sinuses. Length of follow-up was calculated from the last treatment date to the date of last follow-up visit.

Results

We identified 14 patients with ONBs, 7 patients with SNUCs, and 2 patients with SNECs in 7 women and 16 men. Records were reviewed with emphasis on diagnosis, tumor stage, treatment, and outcomes. One ONB, one SNEC, and three SNUC specimens required expert pathological consultation at another institution due to diagnostic challenges.

ONB patients presented with Kadish stages ranging from A to D (2/14 Kadish A, 4/14 Kadish B, 5/14 Kadish C, and 2/14 Kadish D), Dulguerov UCLA staging T1–T3, or AJCC 7th TNM staging T1–T4b. One ONB patient was initially treated at another institution with no information about the initial tumor stage. Nine of the 14 patients with ONB were found to have low-grade tumors (Hyams grade I–II), and 5/14 had high-grade (Hyams grade III) lesions (►Table 1).

All patients with SNUC presented at advanced stages T4a–T4b by AJCC 7th TNM staging system, with assigned Dulguerov UCLA T2–T4 stages and Kadish C-D stage (5/7 Kadish C stage and 2/7 Kadish D). The SNEC cases were poorly differentiated malignancies, one T4a large cell neuroendocrine type carcinoma, and the other a T4b SmCC (►Table 2).

At the time of diagnosis, two patients with ONB and two patients with SNUC had cervical node metastases. No distant metastases were noted at the time of diagnosis in any of the patients included in this study.

In patients diagnosed before April 2013 (11 of ONBs, 4 of SNUCs), the 2-year disease-free survival rates were 81% (9/11) for those with ONB and 75% (3/4) for those with SNUC. One of the patients with SNUC presented with a T4bN2cM0, an AJCC Stage IVB tumor that was treated with surgery followed by chemoradiation. This patient developed distant metastases to the brain and the bone 5.5 months after the initial treatment. These lesions were treated with radiation therapy; nonetheless, the patient passed away 26 months after treatment due to extensive distant metastases to the brain and bones. One patient with large cell type SNEC succumbed to treatment complications 21 months after the treatment due to brain radionecrosis. Of patients treated at our institution, two patients with ONB and one with SNUC suffered a regional or distant recurrence (i.e., no local recurrence was encountered).

Discussion

Histopathological Diagnosis

Sinonasal malignancies with neuroendocrine features cannot be distinguished based on clinical presentation or radiological studies. These malignancies always require histopathological assessment of the specimen by an expert pathologist. Under light microscopy, well-differentiated ONBs are composed of homogenous small-to-medium size cells with round to oval nuclei, sparse cytoplasm, punctate chromatin, small or absent nucleoli and fibrillar cytoplasmic material.7,8

Initially, ONBs were described as tumors with rosette or pseudorosette cell arrangement.9 Presence of rosettes or pseudorosettes and ganglion cells is one of the considerations for the most commonly used ONB grading system introduced...
<table>
<thead>
<tr>
<th>Age (y)</th>
<th>Sex</th>
<th>Kadish stage</th>
<th>Dulguerov UCLA/AJCC TNM stage</th>
<th>Hyams grade</th>
<th>Treatment</th>
<th>Recurrence</th>
<th>TTR (mo)</th>
<th>Tx</th>
<th>F/u</th>
<th>Status</th>
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</thead>
<tbody>
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<td>M</td>
<td>C</td>
<td>T3N0M0/T4aN0M0 stage IVA</td>
<td>2</td>
<td>EEA + post-op RT</td>
<td>Regional</td>
<td>76</td>
<td>EEA</td>
<td>77</td>
<td>NED</td>
</tr>
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<td>M</td>
<td>C</td>
<td>T3N0M0/T3N0M0 stage III</td>
<td>1</td>
<td>EEA + post-op RT</td>
<td>Regional</td>
<td>33</td>
<td>Chemo + EEA</td>
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<td>D</td>
<td>T2N1M0/T4bN1M0 stage IVB</td>
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<td>EEA + LND + post-op RT</td>
<td>Regional, distant</td>
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<td>Chemo + EEA</td>
<td>42.5</td>
<td>AWD</td>
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<td>B</td>
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<td>B</td>
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<td>EEA + post-op RT</td>
<td>Regional, distant</td>
<td>19</td>
<td>Chemo + EEA</td>
<td>41</td>
<td>NED</td>
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<tr>
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<td>M</td>
<td>C</td>
<td>T3N0M0/T4bN0M0 stage IVB</td>
<td>2</td>
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<td></td>
<td></td>
</tr>
<tr>
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<td>M</td>
<td>A</td>
<td>T1N0M0/T1N0M0 stage I</td>
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<td>EEA</td>
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<td></td>
</tr>
<tr>
<td>72</td>
<td>F</td>
<td>C</td>
<td>T3N0M0/T4bN0M0 stage IVB</td>
<td>3</td>
<td>EEA + post-op RT</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>65</td>
<td>M</td>
<td>C</td>
<td>T3N0M0/T4bN0M0 stage I</td>
<td>3</td>
<td>EEA + post-op RT</td>
<td></td>
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<td></td>
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<td></td>
</tr>
<tr>
<td>53</td>
<td>M</td>
<td>A</td>
<td>T1N0M0/T1N0M0 stage I</td>
<td>2</td>
<td>EEA + post-op RT</td>
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<tr>
<td>77</td>
<td>M</td>
<td>D</td>
<td>T3N1M0/T4bN1M0 stage VIB</td>
<td>2</td>
<td>Chemo + EEA + BND + post-op chemo/RT</td>
<td></td>
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<td>24</td>
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<td>B</td>
<td>T1N0M0/T1N0M0 stage I</td>
<td>2</td>
<td>EEA</td>
<td></td>
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</table>

Abbreviations: AJCC, American Joint Committee on Cancer; AWD, alive with disease; BND, bilateral neck dissection; chemo, chemotherapy; DOC, dead of other causes; DOD, dead of disease; EEA, endoscopic endonasal approach resection; F, female; F/u, follow-up from the last day of treatment in months; LND, L neck dissection; M, male; NED, no evidence of disease; post-op, postoperative; RT, radiotherapy; TNM, TNM classification of malignant tumours; TTR, time to recurrence in months; Tx, treatment of recurrence; UCLA, University of California, Los Angeles.
Table 2  SNUC and SNEC outcomes at the Ohio State University James Cancer Center

<table>
<thead>
<tr>
<th>Age (y)</th>
<th>Sex</th>
<th>Tumor type</th>
<th>Kadish stage</th>
<th>Dulguerov UCLA/AJCC TNM stage</th>
<th>Treatment</th>
<th>Recurrence</th>
<th>TTR (mo)</th>
<th>Tx</th>
<th>F/u</th>
<th>Status</th>
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</thead>
<tbody>
<tr>
<td>81</td>
<td>F</td>
<td>SNUC</td>
<td>C</td>
<td>T3N0M0/T4bN0M0 stage IVB</td>
<td>EEA (debulk) + chemo/RT</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>59</td>
<td>F</td>
<td>SNUC</td>
<td>D</td>
<td>T3N2cM0/T4bN2cM0 stage IVB</td>
<td>EEA + Caldwell–Luc (debulk) + chemo/RT</td>
<td>Regional, distant</td>
<td>5.5</td>
<td>RT</td>
<td>26</td>
<td>DOD</td>
</tr>
<tr>
<td>43</td>
<td>M</td>
<td>SNUC</td>
<td>C</td>
<td>T3N0M0/T4bN0M0 stage IVB</td>
<td>Chemo + RT</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>60</td>
<td>M</td>
<td>SNUC</td>
<td>D</td>
<td>T2N1bM0/T4aN2bM0 stage IVA</td>
<td>Chemo + RT</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>45</td>
<td>F</td>
<td>SNUC</td>
<td>C</td>
<td>T4N0M0/T4bN0M0 stage IVB</td>
<td>Chemo + EEA (debulk) + RT</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>46</td>
<td>M</td>
<td>SNUC</td>
<td>C</td>
<td>T3N0M0/T4bN0M0 stage IVB</td>
<td>Chemo + RT</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>83</td>
<td>M</td>
<td>SNUC</td>
<td>C</td>
<td>T3N0M0/T4bN0M0 stage IVB</td>
<td>Chemo</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>66</td>
<td>M</td>
<td>SNEC (large cell)</td>
<td>C</td>
<td>T3N0M0/T4bN0M0 stage IVB</td>
<td>EEA (debulk) + RT + chemo/RT</td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>55</td>
<td>M</td>
<td>SNEC (SmCC)</td>
<td>C</td>
<td>T3N0M0/T4bN0M0 stage IVB</td>
<td>Chemo + chemo/RT</td>
<td></td>
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Abbreviations: AJCC, American Joint Committee on Cancer; AWD, alive with disease; BND, bilateral neck dissection; chemo, chemotherapy; DOC, dead of other causes; DOD, dead of disease; EEA, endoscopic endonasal approach resection; F, female; F/u, follow-up from the last day of treatment in months; LND, L neck dissection; M, male; NED, no evidence of disease; post-op, postoperative; RT, radiotherapy; SNEC, sinonasal neuroendocrine carcinomas; SNUC, sinonasal undifferentiated carcinomas; TNM, TNM classification of malignant tumours; TTR, time to recurrence in months; Tx, treatment of recurrence; UCLA, University of California, Los Angeles.
by Hyams. However, these features are absent in high-grade ONBs. Fibrillary cytoplasmic background is a diagnostically helpful microscopic feature seen in 85% of ONB cases.8

A distinction between poorly differentiated variants of ONBs, SNECs, or SNUCs can be particularly difficult. Some authors consider Hyams grade III/IV ONBs and SNUC to be the same entity, although this notion is controversial as the first grading system for ONB, the Hyams grading system, preceded the first mention of SNUC by 10 years.10

SNUCs are comprised of pleomorphic cells organized into sheets, large nests, trabeculae, and ribbons with frequent necrosis and no differentiation on light microscopy. High-mitotic activity, atypia, and lymph-vascular invasion can be observed.7,11,12

Immunohistochemical markers aid in the exclusion of other diagnoses, but are rarely diagnostic for SNUCs. Strong positivity for cytokeratin with negative synaptophysin and chromogranin staining suggests the diagnosis of SNUC. Although SNUCs often lack neuroendocrine features, neuron-specific enolase-positive staining can be seen. Neuron-specific enolase is a nonspecific marker and is not considered to be diagnostic for true neuroendocrine neoplasms. However, when expressed in SNUC tumors, it can differentiate them from ONBs based on the pattern of staining. Unlike ONB, which stains diffusely positive for neuron-specific enolase, SNUC usually exhibits focal neuron-specific enolase positivity.7,13,14

A recent study by Stelow et al reported that some SNUCs could be categorized as a nuclear protein in testis (NUT) midline carcinomas.15 These tumors are undifferentiated but may have focal squamous differentiation, which has been previously noted in SNUC tumors and was later suggested as a potential addition to the histologic diagnostic criteria for SNUC.12,15 Nonetheless, SNUCs and NUT midline carcinomas are classified as two separate malignancies, and clinically most SNUCs are highly responsive to chemotherapy, whereas NUT midline carcinomas are not.

SNEC is often used as an umbrella term, which includes both typical and atypical carcinoid tumors of sinonasal space as well as poorly differentiated SmCC and large cell neuroendocrine carcinomas. The latter malignancy is extremely rare and can be distinguished from moderately differentiated atypical carcinoid tumors by the presence of high-mitotic activity.3 SmCC is morphologically identical to the small cell carcinoma of the lung. In some cases SmCCs have a characteristic appearance on light microscopy in addition to the distinct immunohistochemical profile, which can help distinguish them from other tumors with neuroendocrine features.14 In the past, some authors argued that SmCC should be classified as SNUC; however, it is widely accepted that SNUCs and SNECs are two separate entities.8,14

In contrast to well-differentiated ONBs, SNEC tumors lack rosette formation. With the exception of SmCC, SNUC tumor cells are larger than ONB cells, have bigger nucleoli, abundant cytoplasm, and neurosecretory granules, while lacking neurofibrillary background.16 These distinguishing features are not as helpful when the differential diagnosis includes poorly differentiated ONB as they may lack rosettes or neurofibrillary background. In contrast to SNUCs, SNECs are often positive for chromogranin and synaptophysin. However, there are cases in which a distinction between SNEC and SNUC cannot be made.7,12,14

Poorly differentiated ONBs are difficult to distinguish from other sinonasal malignancies. The differential diagnosis is broad and includes malignant melanoma, rhabdomyosarcoma, Ewing sarcoma (ES, primitive neuroectodermal tumor or PNET), malignant lymphoma, and extramedullary plasmacytoma among others. Tumor cell positivity for neuronal markers (chromogranin, synaptophysin), S-100 in sustentacular cells, diffuse positivity for neuron-specific enolase, with absence of epithelial, mesenchymal, myogenic and leukocytic markers is characteristic of ONB (Table 3). However, if there is a coexpression of neuroendocrine markers and cytokeratin, typical for simple epithelia, neuroendocrine carcinoma is more likely.

Sinonasal malignant melanomas are often amelanotic and can be composed of small blue cells resembling ONBs. Immunohistochemical markers for melanoma, such as

<table>
<thead>
<tr>
<th>Malignancy</th>
<th>SYN</th>
<th>CHR</th>
<th>NSE</th>
<th>CK</th>
<th>S-100</th>
<th>HMB45</th>
<th>DES</th>
<th>MSA</th>
<th>VIM</th>
<th>CD99</th>
<th>LCA</th>
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<tbody>
<tr>
<td>ONB</td>
<td>+</td>
<td>+</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SNUC</td>
<td>−†</td>
<td>−†</td>
<td>+‡</td>
<td>+</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SNEC</td>
<td>+†</td>
<td>+†</td>
<td>+‡</td>
<td>+</td>
<td>−−−−−−</td>
<td></td>
<td></td>
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<td>−</td>
<td>+‡</td>
<td>+</td>
<td>−</td>
<td>−</td>
<td>V</td>
<td>−</td>
<td></td>
</tr>
<tr>
<td>Rhabdomyosarcoma</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>V</td>
<td>V</td>
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<td>+</td>
<td>−</td>
<td>+</td>
<td>V</td>
<td></td>
</tr>
<tr>
<td>ES/PNET</td>
<td>+‡</td>
<td>+‡</td>
<td>−−−</td>
<td>+</td>
<td>V</td>
<td>−</td>
<td>V</td>
<td>V</td>
<td>+</td>
<td>−</td>
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</tr>
<tr>
<td>Lymphoma</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
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<td>−</td>
<td>−</td>
<td>−</td>
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</tr>
</tbody>
</table>

Abbreviations: CD99, product of MIC-2 gene; CHR, chromogranin; CK, cytokeratin; DES, desmin; HMB45, melanoma-associated marker; LCA, leukocyte antigen; MSA, muscle-specific actin; NSE, neuron-specific enolase; S-100, S-100 protein; SYN, synaptophysin; VIM, vimentin.

Note: Based in part on data from Haas and Ganzer.63

†+, positive; −, negative; V, variable; +‡, positive in most cases; −−, negative in most cases.

‡NSE staining is diffuse in ONBs and focal in SNUCs.

§S-100 is positive in sustentacular cells only in ONBs and stains diffusely in melanoma.

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Sox100, S-100, HMB-45, and melanin A are helpful in diagnosis of these tumors. Sox100 is a highly sensitive and specific immunohistochemical marker for melanoma. It is nearly always expressed in sinonasal melanomas, and is often positive when other markers characteristic of melanoma are negative. In low-grade ONBs with the classic rosette architecture and the presence of sustentacular cells, the location of S-100 positive cells can distinguish these tumors from sinonasal malignant melanoma. The S-100 positive cells in ONBs are limited to the sustentacular cells at the periphery of the cell nests, and melanomas are diffusely positive for S-100 protein. Additionally, tumor location might be helpful in these situations with malignant melanomas limited to the lower part of the nasal cavity beyond the distribution of the olfactory epithelium that gives rise to ONBs.

ES/PNET is another mimic of ONB, with the similar microscopic appearance of small, undifferentiated cell sheets, Homer–Wright rosettes and centrally located fibrils. However, unlike ES/PNET tumors that are CD99 positive in more than 95% cases and FLI1 immunopositive in 71% of cases, virtually all ONB tumors are CD99 and FLI1 negative. Approximately 85% of the ES/PNET tumors show characteristic (t [11; 22] [q24; q12]) translocation with EWS and FLI1 gene fusion, which can be confirmed by fluorescence in situ hybridization.

**Treatment and Outcomes**

**Olfactory Neuroblastoma**

Rosenthal et al first described the classification of sinonasal malignancies with neuroendocrine features into ONB and non-ONB subtypes reporting significant differences in outcomes between the two categories. Although the treatment of SNUC and carcinoid type SNEC tumors is similar, the management approach to ONBs and poorly differentiated SmCC and large cell type SNECs is vastly different. Therefore, a thorough assessment and accurate diagnosis are important for the treatment selection and prognosis.

ONBs have a better prognosis and survival compared with other sinonasal neuroendocrine malignancies. In general, a combination of surgery (craniofacial resection) and radiotherapy has shown superior results to radiation alone and has become a standard of care for all resectable cases of ONBs.

Traditionally, ONB resections involved an open craniofacial approach. Dulguerov et al reported a meta-analysis of 390 patients treated with an anterior craniofacial approach yielding a 5-year disease-free survival of 45% for surgery and radiation. Conversely, the median overall survival for a case series of 70 patients treated at MD Anderson was 126.3 months (10.5 years) with a median disease-specific survival of 139 months (11.6 years). Recent advances in endoscopic techniques have led to an increased use of endoscopic resection for ONBs, in an effort to avoid brain manipulation and external facial incisions. Multiple case series from around the world have yielded promising results with similar outcomes to that of open approach.

Several small case series reported 3- and 5-year survival range of 89 to 100%. A meta-analysis of 23 publications comparing endoscopic surgery to open approach showed a 25% improvement in 10-year overall survival with endoscopic approach. However, these results have to be interpreted with caution due to a strong selection bias; a larger number of patients with advanced disease (Kadish C–D) were treated by open craniofacial approach than by endoscopic approach.

In this study, all ONB cases, including Kadish C and D tumors, were treated with endoscopic resection. With the exception of two T1 Kadish B tumors, all ONB patients underwent postoperative radiation therapy. A neck dissection was only performed for clinically positive nodes. One patient, who was initially treated with an open approach at another institution, underwent an endoscopic resection for regional tumor recurrence at our institution. Two patients with regional and regional-distant recurrence underwent endoscopic resection combined with chemotherapy.

The 2-year disease-free survival for 11 ONB patients with a minimum of 2-year follow-up was 81% (9/11). One patient with Dulguerov/UCLA T2N0M0 stage and AJCC T2N0M0 stage II ONB was found to have regional and distant metastases 19 months after the initial treatment. This patient was still alive 23 months after the treatment of the recurrence and 42 months after the initial treatment. One patient with Dulguerov/UCLA T3N0M0 stage and AJCC T3N0M0 stage III ONB died of brain radionecrosis following postoperative proton radiation therapy. Despite these results and the seemingly superior outcomes reported in the literature following endoscopic resection, long-term follow-up is needed to further assess the outcomes of endoscopic approach. ONB is known for its propensity to recur later and requires long-term surveillance with previously reported disease-free survival reduction of 34% on 15-year follow-up.

Chemotherapy as primary treatment for ONBs has shown inferior outcomes. It is primarily reserved for the treatment of nonresectable disease, as neoadjuvant therapy to decrease the size of the tumor and the extent of the brain or orbit manipulation, and as a radiosensitizer. A series of 50 patients treated with neoadjuvant chemoradiation at the University of Virginia showed outcomes comparable to traditional open resection and postoperative radiotherapy. In this series, patients with Kadish A and B ONBs were treated with preoperative radiation, while Kadish C cases received sequential preoperative chemotherapy and radiation 4 to 6 weeks before open approach resection. The 5- and 15-year disease-free survival rates were 86.5 and 82.6%, respectively. The local-regional recurrence was 24% and distant recurrence was 10%.

In our case series, one patient with Kadish C, Dulguerov/UCLA T3N1M0 disease received neoadjuvant chemotherapy and postoperative chemoradiation following endoscopic resection with bilateral neck dissection; however, follow-up was limited to 6 months.

Some authors suggested combining an elective neck dissection with radiation therapy. Most surgeons agree that a neck dissection is only indicated in patients with cervical metastases, arguing that prophylactic treatment...
adds considerable morbidity and should be reserved for salvage therapy. Noh et al reported a reduction in the rate of relapse in patients undergoing elective neck irradiation, especially radiation to level 1.

Sinonasal Undifferentiated Carcinoma
SNUCs are known for their aggressive nature and often present with locally advanced disease, orbital and periorbital invasion, and intracranial extension. Surgical management of SNUCs is challenging and the indications for resection vary from institution to institution.

Despite poor outcomes reported in early studies, recent reports have shown 5-year overall survival rates between 40 and 75%. Due to the rarity of these tumors, case series lack uniform treatment paradigms, which complicates the comparison of outcome. Multimodality treatment with or without craniofacial resection (in cases of resectable disease) has yielded improved outcomes. Most series reported utilizing radiotherapy or chemotherapy and radiation in addition to surgery, but no paradigm has demonstrated clear therapeutic advantage.

Currently, the trend is to treat patients presenting with advanced SNUC with combined chemotherapy and radiation leaving surgery as a salvage option. Tanzler et al reported on 15 patients, 9 of whom underwent open resection with subsequent radiation or chemotherapy with 3-year overall survival of 67%. In a case series of 16 patients half of whom received neoadjuvant chemotherapy and the other half received surgery, the overall survival was 62.5%. Similarly, a 2-year overall survival rate of 64% was reported by the University of Virginia in a series of 10 patients treated with neoadjuvant chemotherapy followed by radiotherapy.

Conversely, Al-Mamgani at al showed superior outcomes with a 5-year overall survival rate of 74% when a tailored approach was utilized with surgery and postoperative radiation or chemoradiation used for resectable disease and neoadjuvant chemotherapy or chemoradiation alone used for nonresectable cases. Multimodality treatment for SNUC has been proven to improve survival regardless of the primary treatment.

More recently, small case series treated with endoscopic resection reported comparable results to the open approach with local and regional control of up to 78% for endoscopic resection followed by radiation. Surgery as primary treatment in combination with neoadjuvant therapy showed improved survival and local-regional control.

In this study, three SNUC cases underwent endoscopic gross total removal followed by chemotherapy and radiation therapy (two patients had an initial diagnosis of ONB). All of our cases were treated with chemotherapy and radiation. The 2-year disease-free survival for the four patients was 75%.

Sinonasal Neuroendocrine Carcinomas
Typical and atypical carcinoids are generally considered separately from SmCC and large cell neuroendocrine carcinoma as their treatment and outcomes differ significantly. A limited number of prior studies with the lack of uniform treatment protocols for these malignancies make outcome comparison difficult.

Some centers favor treating patients with these malignancies similar to those with SNUCs and others use treatment paradigms similar to those used for patients with ONBs.

A recent SNEC case series showed a 5-year overall survival of 66.9%, although the authors did not distinguish between the histopathological types of SNEC. This study included patients with various treatment strategies that were based on the interdisciplinary consensus with nearly half the patients undergoing surgery for definitive management of their disease. As with SNUCs, neoadjuvant chemotherapy followed by surgery or chemoradiation is a preferred strategy due to high chemoresponse of most SNEC cases. A complete response to neoadjuvant chemotherapy correlated with improved survival in prior studies.

SmCC is a rare malignancy with outcomes that are inferior to those of other sinonasal neuroendocrine cancers, but comparable to the outcomes of small cell carcinoma at other sites. It is difficult to recommend a single treatment strategy since prior studies lacked a uniform treatment approach. Chemoresponsive cases showed better 3-year survival. Neoadjuvant chemotherapy with subsequent chemoradiation or surgery and postoperative radiation have been utilized in previous studies.

One SNEC patient at our institution had a diagnostically challenging histopathological profile with variable reactivity for cytokeratin, synaptophysin, chromogranin, peripheral sustentacular S-100 protein, and neuron-specific enolase. Although the pattern of S-100 staining was similar to that of ONB and the overall findings did not show classic features of poorly differentiated neuroendocrine carcinoma (SmCC or large cell type), an outside pathological consultation suggested to manage this tumor as a sinonasal SmCC. Therefore, this lesion was staged as T4bN0M0, AJCC stage IVB neuroendocrine carcinoma and was treated with neoadjuvant chemotherapy followed by the chemoradiotherapy and radiation. At the time of this report the follow-up was insufficient.

Our study included only one patient with T4aN0M0, AJCC stage IVA large cell neuroendocrine carcinoma. The patient underwent neoadjuvant chemotherapy and radiation before endoscopic gross total removal with postoperative radiation treatment. Unfortunately, this patient died due to brain radionecrosis secondary to proton radiation therapy.

Large cell neuroendocrine cancer is extremely rare; thus, data regarding this tumor are sparse in the literature. Based on the existing reports, these malignancies have poor outcomes comparable to that of SmCC. Aggressive multimodality therapy is usually applied, but the outcomes remain poor.

Conclusions
Sinonasal malignancies with neuroendocrine features cannot be distinguished without a thorough histopathological evaluation. Individualized assessment and treatment strategies that take into account tumor stage, grade, and histopathological characteristics can improve the accuracy of the initial assessment and treatment outcomes of sinonasal carcinomas with neuroendocrine features. Multicenter studies and
collaboration with uniform treatment protocols are essential to improve the reporting of treatment outcomes. Endoscopic resection of these malignancies has shown outcomes similar to previously reported open resections, but long-term follow-up is needed to further assess the outcomes.

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