Pearls & Oy-sters: Bilateral cavernous sinus syndrome as presenting manifestation of nasopharyngeal carcinoma

PEARLS

- Nasopharyngeal carcinoma commonly presents with trismus, pain, otitis media, nasal regurgitation (due to paresis of the soft palate), hearing loss, and cranial nerve palsies. Incidence of cranial nerve involvement varies from 12% to 35%.
- In order of prevalence, the most commonly affected cranial nerves include the trigeminal, abducens, oculomotor, and hypoglossal.

OY-STER

- Cranial nerve palsy at initial presentation is observed in 12%–35% of patients, but it is not clear what percentage of patients present only with cranial nerve involvement.

CASE REPORT

A 48-year-old man presented to our hospital with an acute-onset, severe, persistent holocranial headache of 3 months’ duration. He also noted binocular horizontal diplopia, drooping of the left eyelid, and inward deviation of the left eye that began at the same time as the headache. The symptoms worsened gradually and within a few weeks he developed drooping of the right eyelid as well. He also complained of gradually progressive hearing loss with tinnitus, beginning in the left ear and progressing to the right ear. There were no other systemic or neurologic symptoms. There was no history of smoking or drug/alcohol abuse. He was a security guard by profession. General physical examination revealed a firm 1 × 1 cm painless cervical lymph node. Ophthalmologic examination revealed bilateral sixth, bilateral partial third, and right fourth cranial nerves palsy (figure 1). Fundus examination did not reveal any abnormality. Bilateral conductive hearing loss was also present. The remainder of the neurologic examination was normal. He was diagnosed with bilateral cavernous sinus syndrome and bilateral sequential conductive deafness. The etiologic possibilities considered at this stage included noninfectious granulomatous conditions (such as sarcoidosis and Tolosa-Hunt syndrome), infectious etiologies (fungal, tuberculosis, or other chronic infection), or malignancies (a primary sellar mass with parasellar extension, an extrasellar mass with bilateral extension into parasellar space, or metastasis). His hematologic profile, blood glucose, renal and liver function tests, serum angiotensin-converting enzyme levels, and CSF examination results were within normal limits. HIV serology was negative. He underwent a noncontrast CT scan of the head (figure 2), which revealed an ill-defined mildly hyperattenuating mass causing destruction of the sphenoid bone including the sella and clivus. The mass extended to the parasellar region, sphenoid sinuses, and prepontine region bilaterally. On MRI (figure 2), a large mass was seen involving the posterouserior aspect of the nasopharynx, as well as the sphenoid bone with destruction of the sella and clivus, sparing the basilar artery. The mass encased both internal carotid arteries without any luminal compromise. It appeared isointense with central hypointensity on T1-weighted imaging and hyperintense on T2-weighted images with heterogeneous postcontrast enhancement.

Histopathologic examination of the mass was consistent with an undifferentiated squamous cell nasopharyngeal carcinoma (NPC) (figure 3). Radiotherapy with a midplane dose of 30 Gy in 15 fractions weekly was started, and chemotherapy was planned. Since initiating therapy, the patient has noted some improvement in the double vision and ptosis. His headache persists, though the intensity decreased significantly.

DISCUSSION

NPC is a tumor arising from the epithelial cells that line the nasopharynx. NPC was first described as a separate entity by Regaud in 1921. Currently, 3 subtypes of NPC are recognized in the World Health Organization classification: type 1, squamous cell carcinoma, typically found in the older adult population; type 2, nonkeratinizing carcinoma; and type 3, undifferentiated carcinoma. Most cases seen in childhood and adolescence are of the undifferentiated type, and approximately one-third of all undifferentiated NPCs are diagnosed in adolescents or young adults. The annual incidence of NPC varies by geographic region. Incidence is high in Tunisia.
and moderately high in Southeast Asia, China, the Mediterranean basin, and Alaska. It is low in the United Kingdom and India (0.9 per million). The Epstein-Barr virus, ethnic background of the patient, and environmental carcinogens (volatile nitrosamines) all seem to play an important role in the pathogenesis of NPC.5

NPC usually originates in the lateral wall of the nasopharynx in the nasopharyngeal recess. It can then extend within or out of the nasopharynx to the contralateral wall or posterosuperiorly to the base of the skull or the palate, nasal cavity, or oropharynx.1 It then can metastasize to cervical lymph nodes. Distant metastases may occur in bone, lung, mediastinum, and rarely in liver. Cervical lymphadenopathy is a common initial presentation and in many patients the diagnosis is made by lymph node biopsy. Symptoms related to the primary tumor include trismus, pain, otitis media, nasal regurgitation, hearing loss, and cranial nerve palsies. Larger growths may produce nasal obstruction or bleeding and a “nasal twang.”1

Most of these neoplasms infiltrate aggressively. They grow within the nasopharynx or can spread anteriorly, laterally, posteriorly, superiorly, and inferi- orly to the contralateral wall. NPC can also infiltrate other structures toward the base of the skull and invade the palate, nasal cavity, or the oropharynx. NPC can spread intracranially via the foramen lacerum or foramen ovale or through both foramina. It can also spread by direct erosion, which results in multiple cranial deficits. Perineural infiltration through the foramen ovale is an important route of spread resulting in invasion of the cavernous sinuses as was seen in our patient. Cranial nerve palsies, if present, are associated with a worse prognosis as compared to invasion of the base of the skull.5 Patients with cranial nerve palsies may need more aggressive treatment. A favorable response to definitive radiation
therapy has been documented in such patients. Local growth of the tumor can involve the Eustachian tube (by direct involvement, invasion of the tensor palatini muscle, or because of tube displacement) leading to middle ear effusion and hearing disturbances.1,5,7–9

Ophthalmic-neurologic manifestations are common at time of first presentation in patients with NPC. In a series of 124 patients, signs of cranial nerve involvement were seen in 48 (38.7%) patients at presentation.2 The most commonly affected cranial nerves were the abducens in 11 (22.9%), trigeminal in 10 (20.8%), oculomotor in 6 (12.5%), and hypoglossal in 5 (10.4%). However, multiple cranial nerve dysfunction was seen in only 3/124 (2.4%) patients. This suggests that though cranial nerve involvement is common, finding multiple cranial nerve palsies is relatively rare at presentation in NPC. With regards to cavernous sinus syndrome, Agrawal et al.7 reported unilateral cavernous sinus syndrome as a presenting manifestation of NPC. However, bilateral cavernous sinus syndrome as a presenting manifestation of NPC, as seen in our patient, has not been reported in literature to our knowledge. This case expands the spectrum of neurologic findings as the presenting manifestation of NPC. Our case further expands the clinical spectrum of this tumor and illustrates the importance of keeping a high index of suspicion for NPC in patients presenting with cavernous sinus syndrome. In this regard, it is important to emphasize that though NPC is commonly not amenable to surgery by virtue of its location, it is an exquisitely radiosensitive tumor.10

AUTHOR CONTRIBUTIONS
Bhawana Kumari: acquisition of data, data analysis and interpretation. Manoj Goyal: critical revision of the manuscript for important intellectual content. Vivek Lal: study concept and design, study supervision.

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REFERENCES


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