Nonvestibular Schwannoma Tumors in the Cerebellopontine Angle: A Single-Surgeon Experience

Abstract

**Background:** The most common cerebellopontine angle (CPA) tumor is a vestibular schwannoma. Schwannomas account for 8.5% of all intracranial tumors and more than 90% of the tumors originate from the eighth cranial nerve, but one in five CPA tumors are not vestibular schwannoma. These tumors may have different manifestations and require different management strategies.

**Methods:** We report 224 consecutive NVCPAT operated in the Department of Neurosurgery, P.D. Hinduja National Hospital and Research Centre, Mumbai, by the senior author between 2001 and 2014 and discuss the different approaches and outcomes in NVCPAT.

**Results:** The age range was 20–60 years and there were 129 females and 95 male patients. The clinical material consisted of 81 cases of meningioma (36.1%), 44 cases of epidermoid (19.64%), 34 cases of trigeminal schwannoma (15.17%), 26 cases of jugular foramen schwannoma (11.60%), and 39 cases of other tumors (17.41%). In nonvestibular schwannoma (NVS), symptoms and signs from cranial nerve VIII are less frequent and other cranial nerves and cerebellar symptoms and signs predominate.

**Conclusion:** Symptoms and signs are different in NVCPAT from those found in patients with vestibular schwannoma. Hearing loss is not the predominant symptoms. Cerebellar signs and trigeminal dysfunction are more common. The most common approach used in the current series was retrosigmoid craniotomy. Gamma knife radiosurgery was a useful adjunct in a subset of these patients.

**Keywords:** Cerebellopontine angle, endolymphatic sac tumors, epidermoid, meningioma, nonvestibular schwannoma

Introduction

About 6%–10% of all intracranial tumors arise in or involve the cerebellopontine angle (CPA) and the vast majority of these (80%) are vestibular schwannomas.[1-5] Meningioma and epidermoid account for 10% and 6%, respectively, and the remainder consists of an extremely heterogeneous group of tumors that affect the region.[1-5] These tumors often resemble vestibular schwannomas in their clinical presentation and otologists must be aware that approximately one in five CPA tumors is not a vestibular schwannoma. A thorough history and examination will often provide clues leading to a different diagnosis and management strategy. In recent years, advances in neuroradiology have facilitated the preoperative differentiation of CPA lesions and have aided the surgical planning. Magnetic resonance imaging (MRI) is the investigation of choice.

Vestibular schwannomas are usually the cause for the typical symptoms of a CPA syndrome. Hearing and vestibular disturbances are by far the most common symptoms. The usual natural history of vestibular schwannoma is an insidious hearing loss that develops over several years, a pattern quite different from the less common CPA tumors.[6] Larger CPA tumors can inflict functional deficits on any of the cranial nerves that traverse the angle or neural structures that form part of its boundaries, the pons, and cerebellum. The progression and sequence of symptoms depend on individual regional anatomy and compliance, growth rate, and invasive nature of the tumor. The sequence of these symptoms may suggest a nonvestibular schwannoma (NVS) lesion.[1,3-7] Surprisingly, 30% of patients who present with a CPA syndrome have no diagnosable tumor at all but suffer from cerebrovascular disease, migraine, or other neurological disorder.[8] This original article provides approaches to diagnosis of NVS cerebellopontine tumors (NVCPAT) and management guidelines.
Materials and Methods

Between 2001 and 2014, 224 consecutive NVCPAT were operated in the Department of Neurosurgery, P.D. Hinduja National Hospital and Medical Research Centre, Mumbai, by a single surgeon.

Recorded documents were retrospectively studied for age, sex, clinical profile, investigations, microneurosurgical management, complications, and outcomes. Postoperatively, the patients were followed up clinically and radiologically at regular intervals. Postoperative computed tomography (CT) scan was performed in the immediate postoperative period. Follow-up MRI scan of the brain was done 3 months after the operation, 1 year after the operation, and at regular intervals.

Results

The age range was 20–60 years and there were 129 females and 95 male patients. The clinical material consisted of 81 cases of meningioma (36.1%), 44 cases of epidermoid (19.64%), 34 cases of trigeminal schwannoma (15.17%), 26 cases of jugular foramen schwannoma (11.60%), and 39 cases of other tumors (17.41) [Table 1]. Trigeminal neuralgia was found as a significantly frequent sign of epidermoid and meningioma [Table 2]. Epidermoid was more common among males (24 out of 44), whereas meningioma was more common in females (61 out of 81 patients). Meningioma was found more commonly in the age group of fifth-to-sixth decade (46 out of 81 patients), whereas epidermoid was found more commonly in third-to-fourth decades (27 out of 44) [Table 3]. Other less common tumors operated were malignant skull base tumor (9), glomus jugulare (5), ependymoma (4), hemangioblastoma (4), endolymphatic sac tumor (ELST) (4), and facial nerve schwanna (4). The other rare diagnoses were choroid plexus papilloma (2), pilocytic astrocytoma (2), cholesteatomas (1), cavernous hemangioma (1), solitary fibrous tumor (1), and melanoma (1). Many approaches were used for resection. Retrosigmoid was the most common approach; other common approaches used were Kawase’s, fronto-temporo-orbito-zygomatic approach, far lateral, intralabyrinthine, endonasal, and petrosal [Graph 1]. Complete resection of tumor depended on type of NVCPAT and were as follows: meningioma, 55%; epidermoid, 95%; trigeminal schwannoma, 88%; and jugular foramen schwannoma, 62% [Table 3].

<table>
<thead>
<tr>
<th>NVSCPA tumor</th>
<th>n (%)</th>
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<tbody>
<tr>
<td>Meningioma</td>
<td>81 (36.1)</td>
</tr>
<tr>
<td>Epidermoid</td>
<td>44 (19.64)</td>
</tr>
<tr>
<td>Trigeminal schwannoma</td>
<td>34 (15.17)</td>
</tr>
<tr>
<td>Juglar foramen schwannoma</td>
<td>26 (17.41)</td>
</tr>
<tr>
<td>Others</td>
<td>39 (17.41)</td>
</tr>
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</table>

NVSCPA: Nonvestibular schwanna cerebellopontine angle

<table>
<thead>
<tr>
<th>NVSCPA tumor</th>
<th>Hearing loss (%)</th>
<th>V nerve dysfunction (%)</th>
<th>Cerebellar sign (%)</th>
<th>VIII nerve preservation (%)</th>
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</thead>
<tbody>
<tr>
<td>Meningioma</td>
<td>32</td>
<td>36</td>
<td>22</td>
<td>81</td>
</tr>
<tr>
<td>Epidermoid</td>
<td>16</td>
<td>45</td>
<td>20</td>
<td>95</td>
</tr>
<tr>
<td>Juglar foramen schwannoma</td>
<td>58</td>
<td>4</td>
<td>38</td>
<td>73</td>
</tr>
<tr>
<td>Vth nerve schwannoma</td>
<td>20</td>
<td>65</td>
<td>18</td>
<td>85</td>
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</table>

NVSCPA: Nonvestibular schwanna cerebellopontine angle

<table>
<thead>
<tr>
<th>NVCPAT</th>
<th>Number of patient</th>
<th>Male:female</th>
<th>Median age (years)</th>
<th>Most common approach (%)</th>
<th>Complete excision (%)</th>
<th>Mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meningioma</td>
<td>81</td>
<td>1:4</td>
<td>43.5</td>
<td>Retrosigmoid (76)</td>
<td>55</td>
<td>0</td>
</tr>
<tr>
<td>Epidermoid</td>
<td>44</td>
<td>1.4:1</td>
<td>34.5</td>
<td>Retrosigmoid (82)</td>
<td>95</td>
<td>1</td>
</tr>
<tr>
<td>V nerve schwannoma</td>
<td>34</td>
<td>1.1:1</td>
<td>41</td>
<td>Subtemporal Petrous Apex</td>
<td>88</td>
<td>0</td>
</tr>
<tr>
<td>Juglar foramen schwannoma</td>
<td>26</td>
<td>1.1:1</td>
<td>45.5</td>
<td>Retrosigmoid (77)</td>
<td>62</td>
<td>0</td>
</tr>
</tbody>
</table>

NVSCPA: Nonvestibular schwanna cerebellopontine angle
Two patients died postoperatively in this series, one had epidermoid and the other malignant skull base tumor.

Discussions

Schwannomas account for 8.5% of all intracranial tumors and more than 90% of the tumors originate from the eighth cranial nerve. The CPA is covered or lined by the meninges and in addition to cerebrospinal fluid, contains nerves, vessels, and possibly embryologic remnants. Each of these structures can be the tissue of origin of an NVS CPA lesion

Meningioma

Meningioma accounts for 10%–20% of all intracranial neoplasms. The incidence increases with age and the average age at the time of diagnosis of posterior fossa meningioma is 43.5 years. About 5% to 10% of all meningioma are found in the CPA, predominately in middle-aged women. Second to vestibular schwannomas, meningioma is the second most common tumor in the CPA and constitutes approximately 10% of these tumors. Meningioma is generally a benign tumor but is locally aggressive invading bone along the Haversian canals. This feature may produce radiologically demonstrable hyperostosis. Tumors displace or surround the cranial nerves and vessels rather than invade them and can become strongly adherent to these structures. The tumor is best classified according to where the bulk of its volume is located and by its relationship to major neurovascular structure, information generally more useful to the surgeon. Hormonal influence of these tumors has been investigated. Meningioma tumor cells contain a high concentration of progesterone receptors, moderate numbers of androgen receptors, and a low level of estrogen receptors. The potential of hormone therapy is being investigated. Hearing loss is found at presentation in 50%–80% of patients with meningioma compared with almost all patients with vestibular schwannomas. Tinnitus is experienced by anywhere from 15% to 60% of patients with meningioma compared with 80% with vestibular schwannoma. Disequilibrium troubles 30%–60% of patients with meningioma and is a presenting symptom in 80% of patients with vestibular schwannoma. Facial pain is rarely encountered in patients with vestibular schwannomas but is a presenting symptom in 5%–30% of patients with meningioma. The outcome from surgical resection of meningioma varies depending on tumor location and size. In recent series, gross total resection was achieved in 45%–86% and mortality ranged between 0% and 5%. Permanent postoperative facial weakness occurred in 6%–11% with as many as 30% having postoperative facial paresis. Swallowing problems occurred in 2%–12%. Hearing declined in 17% of patients in hearing preservation surgeries, although one large study found 91% of functional hearing preservation in their series.

In our series of 81 patients of meningioma (51 patients of petroclival, 27 patients CPA, and 3 patients of jugular foramen meningioma), the male-to-female ratio was 1:4 and the median age was 49 years. Trigeminal nerve dysfunction (36%) was the predominant symptoms followed by hearing loss (32%) and cerebellar signs (22%). The most common surgical approach used was retrosigmoid (77%). Total excision was achieved in 55% of patients and postoperative facial function was preserved in 81% of patients. There was no mortality [Figure 1a and b]. Excision may be accomplished through several standard approaches that are also employed for vestibular schwannomas. The choice depends on the patient’s hearing status, the size and location of the tumor, and its involvement with neurovascular structures. The retrosigmoid approach has the advantage of offering the possibility of hearing preservation. Other approaches such as middle fossa approach, translabyrinthine approach, transcoclear and transtentorial approach, and sometimes combined approach are used depending on tumor size, location, and severity of seventh and eight nerve dysfunctions.

Radiosurgery may be used for CPA and petroclival meningioma, either primarily or as an adjunct to microsurgery. Good long-term control and a low side-effect profile have been demonstrated. By the time patients present to the surgeon, most petroclival meningiomas have reached a large size with a wide attachment and the tumor often invades the exit foramina of multiple cranial nerves. Total excision of the tumor with its dural and bony attachment is not possible in such cases without significant risks and unacceptable morbidity. In several cases, the difficulty of excision is further compounded by arterial and brain stem involvement. A review of the literature clearly demonstrates the trend toward less aggressive surgery and an emphasis on the functional outcome, as reported in various series [Table 4]. Facial pain was the most common new symptom after radiosurgery. In one study examining CPA meningioma, facial pain tended to persist after radiosurgery in petroclival meningioma despite effective tumor control.

![Figure 1: (a) Right side petroclival meningioma. Preoperative contrast magnetic resonance imaging axial image. (b) Postoperative contrast computed tomography scan of the brain demonstrates no residual tumor](image-url)
Epidermoid

Epidermoid account for 0.2%–1.8% of all intracranial neoplasm and an equal or slightly male-dominated incidence has been reported.[35,29,30] About 30% to 40% of epidermoid are found in the CPA, where they account for 5%–9% of all tumors.[3,26] Epidermoid is the third most common CPA lesion and represent approximately 6% of such lesion and 1% or all intracranial tumors.[31] Epidermoid is thought to develop from sequestered epithelial cells rests from the laterally migrating secondary optic and otic capsule or from developing embryonic neurovasculature but is not associated with other congenital abnormalities.[34] These grow slowly through accumulation of keratin and cholesterol from their squamous epithelial lining.[32] Their peak age of occurrence is 40 without gender predilection. It tends to spread along normal cleavage planes and surround, not displace, cranial nerves and blood vessels.[33] These are benign lesions, although malignant transformation has been reported.[2] The gross total resection of such lesions ranges from 33% to 88%.[34] In several larger studies, rates of tumor recurrence have ranged from 7% to 45%. Clinical improvements have been seen in 50%–100% of the patients. As in other surgeries of CPA, removal of these tumors has risks of facial weakness (0%–23%), worsening of hearing (8%–10%), and swallowing problems (0%–10%). The resection of epidermoid carries a 3%–8% risk of aseptic meningitis and 0%–4% of hydrocephalus requiring shunt.[35]

In our series of 44 patients of epidermoid, the male-to-female ratio was 1.4:1 and the median age was 34.5 year. Trigeminal nerve dysfunction (45%) was the predominant symptom followed by cerebellar sign (20%) and hearing loss (16%). The most common approach used was retrosigmoid (82%). Complete excision was achieved in 95% of patients and postoperative facial function was preserved in 95% of patients. There was one mortality. As with meningioma, hearing loss was less frequent (50%–80%) compared to vestibular schwannoma and early progressive facial nerve symptoms predominated.[1,4,7,29,35–37] Other less common symptoms were trigeminal neuralgia, facial numbness/spasm, cerebellar signs, and signs of elevated intracranial pressure. The lesions have a characteristic heterogeneous, low-signal-intensity appearance with no gadolinium enhancement on T1 images. Special fluid-attenuated inversion recovery and diffusion sequence MRI scans have proven useful in differentiating between arachnoid cysts and epidermoid.[38] Microsurgical removal of epidermoid was the treatment of choice and the retrosigmoid approach was the preferred technique. As with meningioma, alternative approaches such as the translabyrinthine or middle fossa can be employed if required.[1,3,4,5,7,35,36,39] The goal of surgery is decompression of the cyst and removal of the capsule. The interior of the tumor is soft and caseous and can be easily removed with suction or curettage. The capsule, however, is more difficult to remove. Because of the pattern of growth, neurovascular structures are often engulfed in the tumor and total excision can be difficult without increased mortality and morbidity.[1,3,4,5,7,35,36,39]

### Table 4: Petroclival meningiomas: Rate of total excision

<table>
<thead>
<tr>
<th>Authors (year)</th>
<th>Number of patients</th>
<th>Gross total resection (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Samii et al., 1989[37]</td>
<td>24</td>
<td>71</td>
</tr>
<tr>
<td>Sekhar et al., 1990[38]</td>
<td>41</td>
<td>78</td>
</tr>
<tr>
<td>Al-Mefty and Smith 1991[19]</td>
<td>18</td>
<td>83</td>
</tr>
<tr>
<td>Misra 1991[16]</td>
<td>11</td>
<td>82</td>
</tr>
<tr>
<td>Kawase et al., 1991[30]</td>
<td>10</td>
<td>70</td>
</tr>
<tr>
<td>Bricolo et al., 1992[21]</td>
<td>33</td>
<td>79</td>
</tr>
<tr>
<td>Spetzler et al., 1992[22]</td>
<td>46</td>
<td>91</td>
</tr>
<tr>
<td>Jung et al., 2000[33]</td>
<td>49</td>
<td>20</td>
</tr>
<tr>
<td>Little et al., 2005[24]</td>
<td>137</td>
<td>40</td>
</tr>
<tr>
<td>Mathiesen et al., 2007[25]</td>
<td>29</td>
<td>41</td>
</tr>
<tr>
<td>Natarajan et al., 2007[26]</td>
<td>150</td>
<td>32</td>
</tr>
<tr>
<td>Bambakidis et al., 2007[27]</td>
<td>46</td>
<td>43</td>
</tr>
</tbody>
</table>

### Trigeminal nerve schwannoma and other cranial nerve schwannoma

Trigeminal nerve schwannomas tend to involve the ganglion, nerve root, or both. Symptoms of trigeminal dysfunction tend to dominate over dysfunction of nerve VII and the patients may present with facial pain or numbness. Other symptoms of an expanding CPA tumor may also be present.[1,10] CT scans demonstrate enlargement of Meckel’s cave or foramen lacerum and the tumors tend to be hypo- or isodense and show contrast enhancement. On MRI, the tumors appear iso- or hypointense on T1-weighted images and isointense or hyperintense on T2-weighted images. They enhance as brightly as other schwannomas. Various approaches have been employed and large tumor removal is often possible through combined posterior and middle fossa approaches with lateral opening of Meckel’s cave[1,39] [Figure 2a and b].

In our series of 34 patients of trigeminal nerve schwannomas, the male-to-female ratio was 1:1 and median age was 41 years. Trigeminal nerve dysfunction (65%) was the predominant symptoms followed by hearing loss (20%) and cerebellar signs (18%). The most common approach...
used was subtemporal petrous apex approach. Complete excision was achieved in 88% of patients and postoperative facial function was preserved in 85% of patient. There was no mortality.

Lower cranial nerve schwannomas account for <1% of all CPA lesions. Symptoms reflect the nerve involved. Treatment, when necessary, is usually surgical. The future role of stereotactic radiotherapy or gamma knife for these schwannomas remains to be seen. In the future, more cases are likely to be treated this way and will probably have much the same outcome as with vestibular schwannoma [30] [Figures 3a, b and 4a, b].

In our series of 26 patients of jugular foramen schwannomas, the male-to-female ratio 1.1:1 and the median age was 45.5 years. The hearing loss (58%) was the predominant symptoms followed by cerebellar sign (38%) and trigeminal nerve dysfunction (4%). The most common approach used was retrosigmoid approach. Total excision was achieved in 62% of patients and postoperative facial function was preserved in 73% of patient. There was no mortality. Treatment strategy of jugular foramen tumor depends on the age of patients, size of tumors, and lower cranial nerve involvement [40] [Flow chart 1].

Paragangliomas (glomus tumors)

Paraganglioma is a benign but locally aggressive tumor and destroys the petrous bone to enlarge to the CPA. Symptoms include pulsatile tinnitus, headache, or hearing loss. On otoscopy, a red pulsatile mass can be seen behind the tympanic membrane with increased vascularization of the floor of the external auditory canal. Most patients have conductive hearing loss but with tumors that occupy the CPA either a mixed loss or profound sensorineural loss is not uncommon. Arteriography demonstrates the extent of the tumor and source of its blood supply. [1,2] On CT, tumors appear well defined with adjacent bone erosion and marked enhancement after contrast injection. Paraganglioma has a characteristic “salt and pepper” appearance on both T1- and T2-weighted MRI images because of intratumoral blood vessels and hemorrhages. The tumors enhance intensely after gadolinium administration. [1,2] Treatment options include radiosurgery or surgery.

Many centers have moved away from microsurgical resection in favor of stereotactic radiosurgery for such lesions because of the risk of lower cranial nerve dysfunction. Stereotactic radiation therapy is also a primary option for lesions with more limited extension. A newer paradigm for these lesions is dictated by patient symptomatology. [42] When pulsatile tinnitus and conductive hearing loss are bothersome to a patient, a targeted debulking of the middle ear and mastoid component may be undertaken. If indicated, based on the growth of the remaining lesion, adjuvant radiotherapy may be pursued. Unlike when complete resection is planned, the authors have found preoperative embolization is not needed when only a limited debulking is planned. In our series, the five cases of glomus tumors were operated by far lateral intralabyrinthine approach [Figure 5a and b].

Endolymphatic sac tumors

ELSTs are uncommon tumors and most often associated with von Hippel–Lindau (VHL) disease, although approximately 20% of cases result from sporadic mutations. Nevoux et al. have posited that sporadic tumors behave quite differently from those associated with

![Figure 3: (a) Left side hypoglossal schwannoma. Contrast magnetic resonance imaging axial image demonstrates enhancing lesion with central necrosis arising from left hypoglossal nerve extending to left cerebellopontine angle cistern. (b) Clinical photograph of left hypoglossal schwannoma demonstrating ipsilateral tongue atrophy and deviation](image)

![Figure 4: (a) Left side jugular foramen schwannoma. Contrast magnetic resonance imaging axial image demonstrates enhancing lesion with central necrosis arising from the left jugular foramen. (b) Postoperative contrast magnetic resonance imaging axial image demonstrates no residual tumor](image)
VHL. VHL may affect other organ systems, including cysts throughout the urogenital system, hemangioblastomas of the central nervous system (that may also present in the CPA), and certain malignancies. A majority of patients with VHL inherit this through an autosomal dominant pattern from mutations on chromosome 3. Pathologically, ELSTs are benign low-grade tumors but may be locally aggressive including erosion into the otic capsule. They are present in 10% of patients with VHL. They may cause progressive hearing loss as well as tinnitus and vertigo. Characteristic imaging findings include T1 hyperintensity with heterogeneous enhancement after gadolinium administration. CT may also be helpful in demonstrating a destructive lesion centered at the posterior petrous temporal bone in the region of the vestibular aqueduct. Calcifications may be present as well [Figure 6]. Histologically, these lesions are characterized as papillary adenomatous lesions, and in some cases, are classified as adenocarcinomas. Early gross total resection is advocated because of the high risk of recurrence when a subtotal resection is undertaken. In addition, larger lesions pose increased risk to postoperative facial nerve function. With larger lesions, preoperative embolization may be prudent because these tumors may be supplied by branches of the external carotid or vertebral arteries. In cases where complete surgical resection is not possible or in poor surgical candidates, data suggest a role for adjuvant radiation therapy or primary stereotactic radiotherapy. Small, less-extensive lesions may be approached from a retrolabyrinthine approach, allowing access to the endolymphatic sac for complete resection of these lesions. This approach also allows for preservation of hearing, as Kim et al. demonstrated in maintaining stable pure-tone average in 30 of 31 ears operated through this approach. Depending on a patient’s preoperative hearing status, other approaches, including the translabyrinthine route, may be used. In our series, all four patients of ELST were operated by intralabyrinthine approach [Figure 6a and b].

**Choroid plexus papilloma**

Choroid plexus papilloma is a rare tumor. They represent <1% of intracranial neoplasms. Derived from the

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**Flow Chart 1: Treatment strategy in jugular foramen tumors**

For Young Patients:
- Lower Cranial Nerve Affected
  - Microsurgical Total Excision
  - Observation
- Observation
- Gamma Knife Radiosurgery

For >60 Years:
- >2.5 cm
  - Excision
  - Observation
- Gamma Knife Radiosurgery
- Gamma Knife Radiosurgery

For <2.5 cm:
- Observation
- Gamma Knife Radiosurgery
- Gamma Knife Radiosurgery

*Figure 5: (a) Right side glomus tumor. Contrast magnetic resonance imaging axial image demonstrates enhancing lesion with central necrosis arising from right jugular foramen “salt and paper appearance.” (b) Postoperative contrast magnetic resonance imaging axial image demonstrates no residual tumor

*Figure 6: (a) Right side endolymphatic sac tumor. Contrast magnetic resonance imaging coronal image demonstrates heterogeneous enhancement with lobulated lesion. (b) Postoperative contrast computed tomography scan of the brain demonstrates no residual tumor
epithelial cells of the choroid plexus, they have the same microstructure of normal choroid plexus when benign. Most often affecting children, they arise typically in the lateral ventricles. In adults, the most common site is the fourth ventricle. Malignant forms are very rare. Patients with choroid plexus papilloma may present with signs of a CPA tumor and if of significant size, raised intracranial pressure is almost always present. CPA choroid plexus papilloma should be evaluated using MRI. Their prognosis is excellent when total surgical excision is possible. Radiotherapy has been employed when surgery is not possible, but the results are variable.[2,48,49]

Choroid plexus papilloma has been reported both extending from the 4th ventricle through the foramen of Luschka and primarily in the CPA. Fourth ventricular tumors and tumors primarily in the CPA tend to present in adults.[50] These are benign lesions and can be managed solely with surgery. Less common are more aggressive lesions – so-called atypical papilloma and carcinomas.[31] Overall, 6% of papilloma recur and need repeated surgical intervention.[51] In the pre-MRI era, a series of 12 choroid plexus papilloma of the CPA were resected either through midline approach through the cerebellomedullary fissure or a retrosigmoid approach. Seven of those patients improved but two patients recurred.

Conclusions
Symptoms and signs are different in NVCPAT from those found in patients with vestibular schwannoma. Hearing loss is not the predominant symptom. Cerebellar signs and trigeminal dysfunction are more common. The most common approach used in the current series was retrosigmoid craniotomy. Majority was benign tumors and had an excellent outcome. Early diagnosis, proper investigations, and evaluation, along with appropriate decision-making and surgical planning with microsurgical techniques are the essential factors that can result in an optimum outcome with complete resection and no new neurological deficit. Some of these tumors, especially petroclival meningioma and jugular foramen tumors are better managed by safe subtotal excision and radiosurgery.

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Nil.

Conflicts of interest
There are no conflicts of interest.

References
23. Jung HW, Yoo H, Paek SH, Choi KS. Long-term outcome and