REVIEW

Trends in the Management of Non-Vestibular Skull Base and Intracranial Schwannomas

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Abstract: The aim of this review is to analyze the latest trends in the management of non-vestibular skull base and intracranial schwannomas in order to optimize tumor control and quality of life. Non-vestibular cranial nerve schwannomas are rare lesions, representing 5–10% of cranial nerve schwannomas. Management decisions should be individualized depending on tumor size, location and associated functional deficits. Generally, large sized schwannomas exerting significant mass effect with increased intracranial pressure are treated surgically. In some cases, even after optimal skull base resection, it is not possible to achieve a gross total resection because tumor location and extent and/or to reduce morbidity. Thus, subtotal resection followed by stereotactic radiosurgery or fractioned radiotherapy offers an alternative approach. In certain cases, stereotactic radiosurgery or radiotherapy alone achieves good tumor control rates and less morbidity to gross total resection. Finally, given the slow growth rate of most of these tumors, observation with periodic radiographic follow-up approach is also a reasonable alternative for small tumors with few, if any, symptoms.

Keywords: schwannoma, head and neck, cranial nerves, non-vestibular, skull base

Introduction

Schwannomas are primary peripheral nervous system tumors arising from Schwann cells. Schwann cells function to myelinate peripheral nerves beginning at the Obersteiner-Redlich zone, the transition from the central to the peripheral nervous system.¹ They may originate from any peripheral, cranial or autonomic nerve. Schwannomas are usually solitary neoplasms; multifocal presentation should raise the suspicion for neurofibromatosis type 1 and type 2 or schwannomatosis.²

Sporadic non-vestibular cranial nerve schwannomas (NVCNS) are rare lesions, representing 5–10% of cranial nerve schwannomas.³ Nerve sheath tumors account for 8% of primary intracranial tumors and more than 90% of intracranial nerve sheath tumors are vestibular schwannomas (VS).^{4,5} In descending frequency, NVCNS involve the trigeminal nerve, lower CNs, the facial nerve and lastly the oculomotor nerves.

NVCNS are characterized by slow growth. It has been estimated that over 50% of the NVCNS exhibit an annual growth rate of more than 5% of the initial volume. This ratio is higher than in sporadic vestibular schwannomas, but lower than in patients with neurofibromatosis type 2 (NF2).⁶ Schwannomas usually remain benign, however there is a possibility of malignant transformation mainly in the setting of neurofibromatosis type 1 (NF1).^{5,7,8}

The morbidity associated with skull base and intracranial schwannomas and their treatment has decreased with the improvement of open surgical approaches to

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the skull base and with the development of endoscopic skull base surgery. However, the morbidity of resection is still relatively high depending on the size and location of the tumor.

The purpose of this review is to discuss the latest trends in the management of these lesions in order to optimize disease control and quality of life.

Trigeminal Schwannomas (TS) Epidemiology

TS are a rare entity, accounting for 0.07-1% of all intracranial tumors. However, these tumors are the second most common intracranial schwannomas, after VS, and account for 3–9% of intracranial schwannomas.^{9–11} While most TS develop in the Gasserian ganglion within the middle fossa, TS can occur anywhere along the course of the trigeminal root, ganglion, and peripheral branches. Malignant TS are rare. Only 22 cases of histologically malignant TS have been reported in the literature.¹² A review of 514 patients surgically treated for TS reported malignancy in 1.5% of them.⁹

Classification Systems

Several classification systems of TS have been employed, some of them are similar. The original classification of these tumors by Jefferson includes type A (mainly in the middle fossa), type B (mainly in the posterior fossa) and type C (dumbbell type, in both the middle and posterior fossae).¹³ Type D was added later and includes those arising from one of the three divisions of the trigeminal nerve.^{12,14,15} In a review of 455 cases the following incidence was observed: 40% type A, 22% type B, 32% type C and 6% type D.⁹ A more complex classification into 6 different types has been proposed by Yoshida and Kawase:¹⁶ type P (posterior fossa tumor); type M (middle fossa tumor); type E (extracranial tumor). The other 3 types refer to dumbbell type tumors in multiple fossae: type MP (middle and posterior fossae); type ME (middle fossa and extracranial space); type MPE (posterior fossa, middle fossa, and extracranial space). Employing this system, Yoshida and Kawase reported the location of 429 tumors: type M 38.5%; type P 23.5%; type E 5.4%; type MP 28.3%; type ME 3.5%; and type MPE 0.7%.¹⁶ In summary, TS arising in the middle fossa are the most common.

Symptoms and Signs

Trigeminal nerve dysfunction is seen in most patients. The most common trigeminal nerve symptoms are pain, paresthesia, numbness and decreased motor function. Symptoms and signs are reflected in Table $1.9^{-12,15-20}$ However, it has been reported that up to 18% of tumors can be found incidentally with magnetic resonance (MR) imaging performed for other reasons.¹⁶

Management

Surgical resection is the mainstay of management, although the therapeutic modality used should be tailored individually according to tumor location and size, histology, patient comorbidities and symptoms.²¹

Table 1 describes the main open approaches.^{11,12,18,19,22,23} Lateral approaches via temporal craniotomy provide the shortest corridor to the Meckel's cave.¹² Tumors originating intracranially from the second (V2) or third (V3) division of the trigeminal nerve are better approached through a subtemporal preauricular infratemporal fossa approach (Figure 1).^{22,23} In tumors with a large extracranial component and anterior and inferior growth into the maxillary sinus, a facial translocation approach may be indicated (Figure 2).^{24,25} Transnasal endoscopic access to type A TS offers advantages similar to open skull base approaches in terms that there is no need for temporal lobe exposure or retraction with either technique, and both provide direct access to the tumor (Figure 2). Endoscopic approaches may provide a minimally invasive and safe approach to radically resect selected tumors.

Although the use of transpetrosal approaches for type B TS is under debate, the retrosigmoid approach is usually employed for posterior fossa tumors (Figure 1).^{11,12,18} In dumbbell shaped tumors (type C) with both posterior and middle fossa extension, in addition to a subtemporal approach a pre- or retrosigmoid craniectomy may provide additional exposure to resect.^{11,18}

Schwannomas frequently do not invade nerve fibers but compress them. However, the nerve fibers are frequently adherent to the tumor.¹⁵ MRI sequences will soon allow preoperative determination of whether the trigeminal nerve fibers are medial or lateral to the tumor. Transcranial approaches might then be chosen for tumors that displace the trigeminal nerve medially while endonasal techniques may be selected for tumors displacing the nerve laterally.²⁶

The treatment aims of TS focus on improvement in neurologic symptoms, relief of mass effect, and

Location	Frequency	Classification	Main Symptoms	Other Symptoms/ Signs	Surgical Approach
TS	3–9% of intracranial schwannomas	-Type A: middle fossa (40%) -Type B: posterior fossa (22%) -Type C: middle and posterior fossae (32%)	Pain (23–95%) Paresthesia (36–46%) Numbness (10–76%) Decreased motor function (24–60%) Headache (14–43%) Abnormal corneal reflex (36–72%)	Diplopia (10–52%) Oculomotor nerve paresis (5–40%) Exophthalmos (4–14%) Visual loss (5–24%) FN weakness (0–20%) Hearing loss (5–30%) Lower cranial nerves deficit (5–18%) Cerebellar/pyramidal system symptoms (8–25%)	-Type A: transzygomatic extradural temporopolar/EEA -Type B: retrosigmoid -Type C: transzygomatic frontotemporal and retrosigmoid
LCNS	-2.9-4% of all intracranial schwannomas -10-30% of tumors of the JF	-Type A: primarily intracranial (25–32%) -Type B: primarily within the JF (15–22%) -Type C: primarily extracranial (12–15%) -Type D (dumbbell-shaped): intra- and extracranial (40–68%)	Hearing loss (45–84%) Headache (15–27%) Dizziness (15–23%) Dysphagia (9–23%) Dysphonia (9–21%) Tongue atrophy (80–92% of the HS)	CN IX palsy (30–40%) CN X palsy (11–27%) CN XI palsy (9–17%) CN XII palsy (15–18%) CN VII palsy (8–18%)	-Type A: retrosigmoid -Type B: modified infratemporal fossa approach type A -Type D: Infratemporal fossa/transcoclea HS: far lateral/transcondylar
FNS	Less than 1% of all intrapetrous mass lesions	Cerebellopontine angle (18.7%) Canalicular segment (24.9%) Labyrinthine/geniculate segment (43.6%) Tympanic segment (42.2%) Mastoid segment (35.8%) Peripheral segment (14.6%) Chorda tympani (1.1%) Stapedial nerve (0.9%)	Facial weakness (62.5%) Hearing loss (50.3%) Tinnitus (20.6%), Vestibular symptoms (14.0%) Ear canal mass (10.8%)	Pain (8.0%) Otorrhea (3.4%) Loss of taste (2.8%) Parotid mass (2.8%) Facial spasm (2.1%)	-Retrosigmoid approach in CPA tumors -Translabyrinthine approach in tumors from the brainstem to the mastoid -Middle cranial fossa approach for tumo between IAC and the tympanic segment -Transmastoid approach for tumors of th tympanic and segments
OMNS		-Cisternal 42% -Cavernous 30% -Cisterno-cavernous 20% -Orbito-cavernous 6%	Diplopia (42%) Headache (32%) Ptosis (27%) Vision loss (13%)	Hemiparesis (7%) Facial hypoesthesia (4%)	-Cisternal: subtemporal transtentorial approach -Cisterno-cavernous: subtemporal transtentorial approach -Orbito-cavernous: orbito-zygomatic approach
TNS		Cisternal type >50% Cisterno-cavernous type Cavernous type	Diplopia (76%) Paresis Sensory changes Ataxia Other cranial nerve palsies		Subtemporal transtentorial approach
ANS		Cisternal (33%) Cisternocavernous (24%) Cavernous sinus (27%) Caverno-orbital (3%) Intraorbital (12%)	Diplopia in horizontal gaze Headache Proptosis Other oculomotor signs	Trigeminal signs Signs of intracranial pressure	-Lesions involving cavernous sinus: frontotemporal transcavernous approac -Cisternal lesions: retrosigmoid approac

Table I Main Features of Non-Vestibular Intracranial Schwannomas

(Continued)

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Table I (Continued).

Location	Frequency	Classification	Main Symptoms	Other Symptoms/ Signs	Surgical Approach
OGS			Headache (53%) Seizures (41%) Anosmia (46%)	Diplopia Signs of intracranial pressure	Transfrontal approach EEA
ONS		Cisternal Optic canal	Proptosis Visual acuity disturbance Visual field defect	Retro-orbital pain Headache	Transfrontal approach EEA

Abbreviations: TS, trigeminal nerve schwannoma; LCNS, lower cranial nerves schwannoma; FNS, facial nerve schwannoma; OMNS, oculomotor nerve schwannoma; TNS, trochlear nerve schwannoma; ANS, abducens nerve schwannoma; OGS, olfactory groove schwannoma; ONS, optic nerve schwannoma; EEA, endoscopic endonasal approach; CN, cranial nerves; JF, jugular fossa; HS, hypoglossal schwannoma; SRS, stereotactic radiosurgery; CPA, cerebellopontine angle; IAC, internal auditory canal; HB, House-Brackman; CN, cranial nerves.

preservation of cranial nerve function. After the application of skull base techniques, gross total resection (GTR) has been achieved in 69–87%. Near GTR (4–26% of patients) or subtotal tumor resection (STR) (7–21% of patients) are performed when there are difficulties dissecting tumor from cranial nerves, blood vessels, or the brainstem (Table 2).^{11,12,15,18,19,27} An improvement in facial pain post-operatively is seen between 45% and 100% of cases;^{12,15,18,19} trigeminal hypesthesia improved in 16% to 62%.^{12,15,18,19} Temporalis and pterygoid function will rarely, if ever, fully improve when present pre-operatively.^{18,19} Other cranial nerve deficits, such as

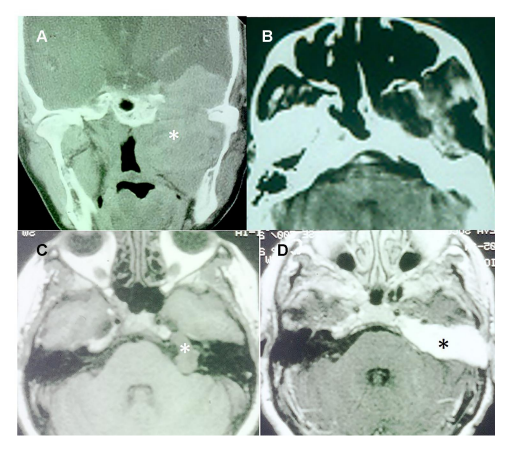


Figure I Surgical approaches in trigeminal schwannomas: subtemporal-preauricular and transcochlear. (A) Type ME2 TS with massive involvement of the middle cranial fossa and the infratemporal fossa (*). (B) Postoperative CT showing the resection of the tumor through a subtemporal-preauricular approach. (C) TI-weighted MRI of a Type P TS in the posterior cranial fossa (*). (D) TI-weighted MRI with contrast showing total resection of the tumor through a transcochlear approach. The operative cavity is filled by fat (*).

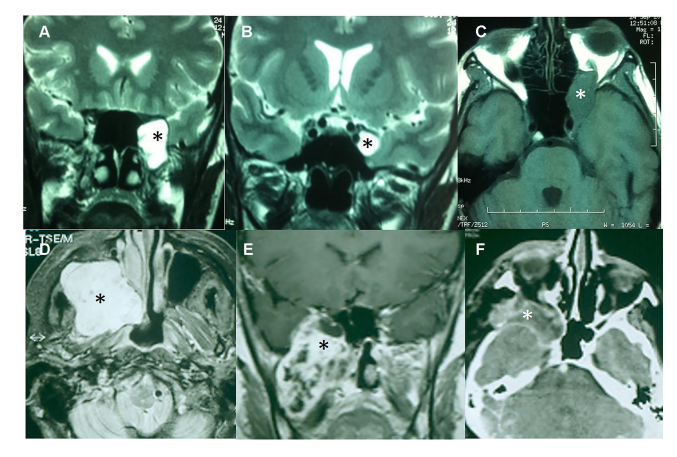


Figure 2 Surgical approaches in trigeminal schwannomas: EEA and facial translocation approach. (A) T2-weighted MRI of a Type MEI TS. The tumor is in contact with the transition between ethmoidal cells and the sphenoid sinus (*). (B) Tumor into the Meckel's cave (*). (C) Orbital progression in a T1-weighted MRI (*). This tumor was excised through an EEA. (D) T1-weighted MRI with contrast of an extensive dumbbell-shaped tumor (*) Type ME2 involving the middle and infratemporal fossae as well as the maxillary sinus. (E) T1-weighted MRI with contrast of a TS with a similar pattern of extension (*). (F) Postoperative CT scan of the anterior case showing the removal of the tumor. The operative cavity was obliterated with a temporalis muscle flap (*).

diplopia caused by palsy of the abducens nerve, improved in 44-75% of patients,^{12,17-19} However surgical injury to the abducens nerve (8%), trochlear nerve (3.7%), and oculomotor nerve (1.9%) has been reported.²⁸ Long tract and cerebellar signs disappeared in 78-100%. 15,17-19 Severe surgical complications are rare (<2% of cases) and include meningitis, cerebrospinal fluid (CSF) leak, arterial vasospasm, hematoma, and hydrocephalus.^{9,11,18,19} The overall mortality rates of surgical series reported since 1990 range from 0 to 3%.11,12,19,27,29,30 The sensory and motor outcomes of endoscopic approaches compare favorably with open surgical approaches. However, some patients may develop dry eye, as a result of vidian nerve sacrifice.³¹

Stereotactic radiosurgery (SRS) is an alternative to surgery. Most of series report a tumor control rate greater than 80%, which is somewhat lower than that achieved with vestibular schwannomas.^{3,26,32–34} The incidence of tumor regression ranges from 34% to 87% and tumor stability from 8% to 50%.^{32–37} A significant symptomatic improvement following SRS varies from 31% to 72% across the series, whereas the incidence of symptomatic deterioration varies from 0% to 27%.9 The factors affecting outcomes include tumor size and site, radiation dose and the radiological appearance of the mass (solid or cystic). The optimal tumor marginal dose appears to be 13-16 Gy. SRS is associated with a very low risk of a new neurologic deficit or other type of complications (less than 16% of patients) (Table 3).^{32–37} There has been speculation regarding the risk of malignancy after treatment with SRS. With a median follow-up of 8.1 years, 2 of 3251 (0.0006%) patients with benign tumors treated with SRS were diagnosed with suspected malignant transformation and 1 of 4905 (0.0002%) patients was considered a case of radiosurgeryassociated intracranial malignancy, resulting in an incidence of 6.87 per 100,000 patient-years for malignant transformation and 2.26 per 100,000 patient-years for radiosurgery-associated intracranial malignancy, similar to the risk of developing a malignant CNS tumor in the general population.³⁸

Author	Location	No. Cases	Total Removal (%)	Near Total or Subtotal Removal (%)	Increased or Newly Developed Morbidity (%)	Mortality (%)
Konovalov et al ²⁷	TN	111	77	23	13	3
Goel et al ¹⁹	TN	73	70	30	31	2,7
Wanibuchi et al ¹²	TN	105	75	25	9	0
Chen et al ¹⁸	TN	55	69	31	5	0
Liu et al ¹¹	TN	84	75	25	12	0
Yoshida & Kawase ¹⁶	TN	27	67	33	52	0
Sharma et al ³⁰	TN	68	76	24	15	1,5
Day & Fukushima ¹⁴	TN	39	77	23	16	0
Jeong et al ²⁹	TN	49	92	8	41	0
Cornelius et al ⁵⁵	FN	10	70	30	60	0
Sherman et al ⁵⁶	FN	10	90	10	20	0
Bakar ⁴⁰	LCN*	199	79	21	38	0,5
Bulsara et al ⁵³	LCN	53	90	10	62	0

Table 2 Main Surgical Series of Intracranial Schwannomas

Abbreviations: TN, trigeminal nerve; FN, facial nerve; LCN, lower cranial nerves. *Systematic review of 19 series.

Author	Location	No.	Decrease	Unchanged	Increase	Deterioration of	
		Cases	Tumor Size (%)	Tumor Size (%)	Tumor Size (%)	Symptoms (%)	
Elsharkawy et al ³²	All locations NVCNS	36	56	25	19	12	
D'Astous et al ³	All locations NVCNS	63	57	35	8	6	
Peker et al ³³	TN	15	87	13	0	7	
Ryu et al ³⁴	TN	32	34	50	16	16	
Snyder et al ³⁵	TN	22	54	23	23	16	
Pan et al ³⁶	TN	56	85	8	7	7	
Sun et al ³⁷	TN	52	77	10	13	4	
Kida et al ⁶³	FN	14	57	43	0	7	
Hasegawa et al ⁶²	FN	42	55	45	3	12	
Hasegawa et al ⁸⁰	LCN	117	53	36	11	7	
Kano et al ⁷⁹	LCN	92	51	36	13	15	
Martin et al ⁸¹	LCN	34	48	46	6	3	
Peciu-Florianu et al ⁶⁵	OMNS/TNS/ANS	30	63	36	3	0	

Abbreviations: NVCNS, non-vestibular cranial nerves schwannomas; TN, trigeminal nerve; FN, facial nerve; LCN, lower cranial nerves; OMNS, oculomotor nerve schwannoma; TNS, trochlear nerve schwannoma; ANS, abducens nerve schwannoma.

Small lesions with few, if any, symptoms may be managed with watchful waiting. Makarenko et al observed 8 patients with TS; an 11.1% progressed over a median of 7.1 years.³⁹

Lower Cranial Nerves Schwannomas (LCNS)

Epidemiology

Among NVCNS, LCNScomprise the second most common site after TS. LCNS can originate intracranially or in the cervical region. Intracranial LCNS usually extend to the jugular foramen or the hypoglossal canal. Jugular foramen schwannomas arising from cranial nerves IX, X or XI are rare and comprise 2.9% to 4% of all intracranial schwannomas, and represent 10–30% of all tumors observed around the jugular foramen.^{40,41} In a review of 204 patients, the tumor originated from the glossopharyngeal nerve (IX) in 47 cases (23.6%), in 26 cases (13%) from the vagal nerve (X) and in 11 cases (5.5%) from the accessory nerve (XI). In the remaining 58% of the cases, the origin of the tumor remained unknown.⁴⁰ Hypoglossal schwannomas (HS) constitute only about 1% of all intracranial schwannomas.⁴²

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Classification

Pellet et al classified jugular foramen schwannomas into 4 classes.⁴³ Type A (primarily intracranial): minimal extension into the jugular foramen; type B (primarily within the bone): with or without an intracranial component; type C (primarily extracranial): only a minor extension into the jugular foramen or into the posterior fossa; and type D (dumbbell-shaped): intra- and extracranial extension. According to Pellet's classification, type Dschwannomas are the most common (40–68%), followed by type A (25–32%), type B (15–22%) and type C (12–15%).^{40,44}

HS have been classified by Kaye et al⁴⁵ in type A: intradural tumor; type B: dumbbell-shaped tumor; and type C: extracranial tumor. In 2 reviews of the literature consisting of 94 and 160 cases, the following distribution has been reported: type A, 29–32%; type B, 39–50%; and type C, 21-29%.^{42,46}

Symptoms and Signs

Hearing impairment is the most frequent presenting symptom in intracranial and jugular foramen LCNS.^{40,41,43,47} Most of patients show a mid-frequency hearing loss, in contrast to the high-frequency hearing loss observed with VS.⁴⁷ Type A tumors may cause symptoms related to increased intracranial pressure, without or with minimal deficits of the lower cranial nerves, whereas a tumor expanding into the jugular foramen or upper cervical region might cause earlier damage of the nerves.⁴¹ Other presenting symptoms are shown in Table 1.

HS are characterized by tongue atrophy. Lower cranial nerves are affected in up to 50% of the cases, mainly in tumors with extension to the jugular foramen.^{45,46}

Management

Schwannomas generally arise in nerves with a sensory component and are associated with sensory ganglia. In the case of the vagal nerve, tumors are likely to involve sensory nerves and it is possible to preserve their function with an intracapsular enucleation, which can increase the neural function preservation by more than 30%.^{7,48} Continuous vagal nerve monitoring during surgery is a new strategy developed with the aim of reducing post-operative morbidity.⁴⁹

Table 1 summarizes the main surgical approaches used in LCNS resection according to their location.^{40,50–52} For tumors confined intracranially (type A), a retrosigmoid approach may provide adequate exposure (Figure 3). In tumors centered in the jugular foramen (type B) or with extracranial extension the modified infratemporal fossa approach type A (ITFA) provides the greatest exposure of the jugular foramen, carotid artery, and anterior temporal bone and it allows direct access to the posterior fossa (Figures 3 and 4).^{51,52} Other skull base techniques such as far lateral, transcondylar, and supracondylar approaches have been advocated for the surgical excision of HS (Figure 4).^{41,45,46}

Bakar⁴⁰ reviewed the outcomes obtained after the surgical treatment of 204 patients. He reported a GTR in 86.9% of the patients, near GTR in 3.3% of the patients and STR in 9.8% of the patients (Table 2).40 Park et al44 analyzed the results of surgery in 275 cases collected in large series of jugular foramen schwannomas and found postsurgical lower cranial nerve palsies in 34.9% of the patients. New neurological deficits are common, especially involving the facial nerve, which was reported in 6-34% of cases^{40,43,47} Other reported complications include CSF leak (3-7%), aspiration pneumonia (1.5-6%), venous (2%). thrombosis (3%), and meningitis sinus Postoperative mortality ranges between 0.5% and 5%.40,46,53

Near GTR followed by SRS or SRS alone offers an alternative approach in some patients (Tables 3 and 4). There are limited data based largely on extracranial schwannomas that fractionated external beam radiotherapy (RT) using doses of approximately 50 Gy at 1.8 Gy per fraction results in a high likelihood of local control.⁵⁴ RT usually induces cessation of lesion growth or reduction in size, rather than complete disappearance of the tumor, which is the goal of the surgical intervention. Side effects are negligible when doses are 50 Gy or less and modern precision RT techniques are used. This is additional option for patients requiring treatment who are unsuitable for surgery.

Observation is another option in asymptomatic patients, particularly for those who are elderly and/or infirm.

Facial Nerve Schwannomas (FNS) Classification

About 650 cases of FNS have been reported in the literature.⁵⁵ The site of origin can be anywhere along the course of the facial nerve from the glial-Schwann cell transition site at the cerebellopontine angle to an extracranial peripheral nerve

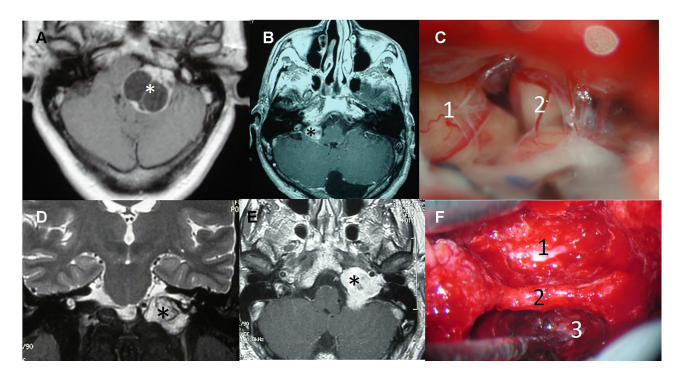


Figure 3 Surgical approaches in lower cranial nerve schwannomas. (A) TI-weighted MRI with contrast of a Type A cystic vagal schwannoma (*). (B) TI-weighted MRI with contrast of a Type A vagal schwannoma (*). (C) Retrosigmoid approach of the last case showing the tumor (1) and the cochleo-vestibular nerve (2). (D) Coronal T2-weighted MRI delimiting a Type B lower cranial nerve schwannoma in the jugular fossa (*). (E) Axial TI-weighted MRI with contrast of the same tumor (*). (F) Modified infratemporal fossa approach type (A) internal carotid artery (1), third portion of the facial nerve (2), tumor in the jugular fossa (3).

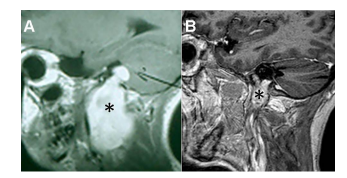


Figure 4 Lower cranial nerve schwannomas with intra- and extracranial extension. (**A**) TI-weighted MRI with contrast of a Type D vagal schwannoma (*). (**B**) TI-weighted MRI with contrast of a Type B hypoglossal schwannoma (*).

location. They have been classified based on their site of origin into cerebellopontine angle (CPA), labyrinthine, geniculate, tympanic, mastoid, and parotid segments (Figure 5).⁵⁶ The percentage involving each segment in a review of 438 cases was reported by Cornelius et al (Table 1).⁵⁵

It is imperative to include FNS in the differential diagnosis of VS during preoperative planning and counseling. However, this is often difficult to achieve preoperatively, even with a careful evaluation of the MRI and computed tomography (CT). FNS involving the CPA and/or the IAC segment of the FN cannot be differentiated from a VS on imaging unless the tumor extends to the labyrinthine segment of the FN. Erosion of superior part of the internal auditory canal and eccentricity of tumor in relation to porus acousticus are not reliable signs. However, facial paralysis is a sign that would support the origin in the facial nerve since VS presenting with facial paralysis is rare.

Symptoms and Signs

The growth rate of FNS is slow. Thus, in a series of 13 patients treated expectantly with a mean follow-up period of 5 years, only 4 demonstrated tumor growth based on serial MRs. The average annual growth rate was 1.4 mm/ year.⁵⁷

Patients frequently present with facial nerve weakness [House-Brackmann (HB) III or IV], as well as hearing loss.^{58–60} Tumors located at CPA usually present with sensorineural hearing loss (SNHL) and tinnitus instead of facial paresis. Patients with tumors originating from the labyrinthine segment tend to present with slowly progressive facial paresis and SNHL, whereas patients with tumors of the tympanic and mastoid segments show progressive facial paresis and conductive hearing loss. The

Author	Location	No. Cases	Total Removal	Subtotal Removal + SRS	Newly Developed Cranial Nerve Palsies	Mortality
Park et al ⁴⁴	LCN LCN	22 22	13	9	23 4	0 0
Author	Location	No. Cases	Total Removal %	Near Total or Subtotal Removal %	Increased or Newly Developed Morbidity %	Mortality
Sedney et al ⁷⁷	LCN LCN	53 28	90 24	10 76	62 25	0 0
Author	Location	No. Cases	Total/Subtotal Removal	Conservative Management	Worsened Facial Function %	Mortality
Liu & Fagan ⁸²	FN FN	22 22	12	10	50 20	0 0
McMonagle et al ⁸³	FN	53	33		33	0
	FN	53		20	15	0

Table 4 Radical Resection versus More Conservative Management

Abbreviations: FN, facial nerve; LCN, lower cranial nerves.

distribution of presenting symptoms in 438 patients with FNS is depicted in Table 1.55

Management

If the patient is asymptomatic and the tumor is small, conservative management with careful observation and followup with serial MRI are recommended. SRS may be an option for small and symptomatic tumors with good facial function (HB I or II). If the tumor is larger (more than 3 cm) or the patient has facial palsy (HB III or worse), surgical resection should be considered.⁶¹ Alternatively, fractionated RT is an option if the anticipated morbidity of surgery is significant. The surgical approach depends on the tumor location and extent. The retrosigmoid approach permits access to the nerve from the brainstem to the IAC and allows resection of large tumors in the CPA. The translabyrinthine approach gives access to the entire nerve course from the brainstem to the stylomastoid foramen. This approach may only be chosen if hearing is not serviceable or if tumor volume or inner ear erosion makes hearing preservation impossible. A middle cranial fossa approach allows access to the nerve between IAC and the proximal tympanic segment. Combining this approach with a transmastoid approach the entire nerve from the IAC to the stylomastoid foramen can be exposed. Hearing preservation is possible by employing this route that make if possible, to repair ossicular chain defects.55

There are different options to preserve the facial nerve function: nerve conservation, nerve resection with immediate grafting, or hypoglossal-facial nerve anastomosis.⁵⁹ Nerve preservation has been reported in 58–71% of the patients.^{55–57} However, facial nerve function was, in the majority of cases, a HB grade III, depending on surgical strategy (Table 2).^{58,60} Regardless of the type of facial nerve repair, patients can expect no better than an eventual HB grade III palsy.^{58,59}

SRS achieves a good tumor control rates.^{62,63} Thus, a multi-institutional study reported 42 patients treated with SRS, with partial regression in 23 patients (54.7%) and stability in 19 patients (45.2%) (Table 3). The actuarial 5-year progression-free survival rate was 92%.⁶³

Oculomotor Nerve Schwannomas (OMNS)

Classification

There are only 54 cases of isolated OMNS reported in indexed literature.⁶⁴ The transition of the oculomotor nerve from the central to the peripheral nervous system occurs 0.6 mm distal to the brainstem. OMNS develop distal to this transition zone. OMNS have been divided into 4 types based on their location: cisternal, cavernous, cisterno-cavernous, and orbito-cavernous (Table 1).^{65,66}

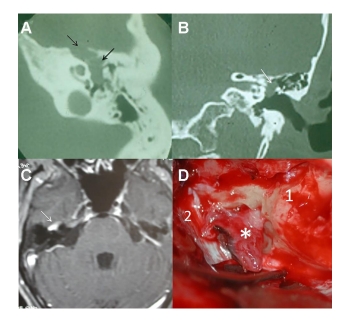


Figure 5 Facial nerve schwannomas. (A) The CT scan shows a tumor involving the tympanic and geniculate segments of the facial nerve (arrows), with displacement of the ossicular chain. (B) Small tumor limited to the tympanic segment (arrow). (C) T1-weighted MRI with contrast of a facial nerve schwannoma of the geniculate ganglion (arrow). (D) Transtemporal approach. Geniculate ganglion schwannoma (*), petrous pyramid (1), dura (2).

Symptoms and Signs

The clinical features depend on the location of the tumor with cisternal schwannomas showing isolated deficits of the nerve. When these tumors originate into the cavernous sinus, they can present with either a cavernous sinus syndrome or an orbital apex syndrome. Symptoms can vary from incomplete oculomotor palsy and exophthalmos to brainstem compression signs.⁶⁵ A systematic review has summarized the most frequent symptoms (Table 1).⁶⁴

Management

Treatment for OMNS remains multimodal, depending on clinical presentation, tumor size, histology, and location. A careful preoperative evaluation including MR guides in distinguishing different histologies. Although most tumors arising from the oculomotor nerve are well encapsulated schwannomas and, in a few cases, nerve sparing surgery may be possible, neurofibromas have also been described. Neurofibromas tend to invade the entire nerve fibers and always require the complete sacrifice of the nerve.⁶⁷

Often microsurgery remains the treatment of choice for the GTR of cisternal schwannomas after identification of the nerve of origin. However, when tumors are located in the cisternocavernous area, total removal of tumor and identification of the parent nerve is barely achieved in half of the cases and a combined approach with STR and SRS should be employed.⁶⁸ Tumors with small-to-medium sized volumes are preferably treated by SRS with a high rate of tumor control and stabilization and/or improvement of symptoms.⁶⁵

Functional recovery of the oculomotor nerve after STR is successful in less than 10% of the patients. Since the oculomotor nerve innervates multiple ocular muscles, the functional result after nerve anastomosis may be limited, and aberrant regeneration may interfere with coordinated binocular movements.⁶⁴

Trochlear Nerve Schwannomas (TNS)

Classification

A total of 85 cases reporting on the management of TNS have been published.⁶⁶ Three types of TNS have been described: cisternal type, cisterno-cavernous type, and the cavernous type. In more than half of the cases reported so far, the cisternal type was diagnosed.^{65,66}

Symptoms and Signs

According to a systematic review from Torun et al⁶⁶ 76% of the entire cohort presented with diplopia, which was the solitary symptom in over half of the cases, and 29% of cases presented with more severe symptoms, such as paresis, sensory changes, ataxia, and other cranial nerve palsies.⁶⁶

Management

All symptomatic patients with a lesion exerting significant mass effect should undergo decompression. The most widely employed microsurgical approach has been the subtemporal transtentorial approach. Nevertheless, nearly 80% of those with surgical resection had persistent trochlear nerve palsy at the last follow-up.

SRS is a good option for patients presented with diplopia only, and 75% of the patients show improved symptoms after therapy.⁶⁶

Patients who only have stable ocular misalignment due to trochlear nerve palsy can also be managed with prism glasses to alleviate diplopia or may choose to undergo strabismus surgery, combined with observation and serial MR imaging.⁶⁶

Abducens Nerve Schwannomas (ANS)

Classification

ANS form an extremely rare entity with only 33 surgical cases published to date⁶⁹ and 16 cases treated by SRS.⁶⁵ The most common locations are shown in Table 1.^{69,70}

Symptoms and Signs

The cardinal neurological sign of ANS is diplopia in the horizontal gaze. However, patients may be asymptomatic or present with headache, proptosis, other oculomotor and/ or trigeminal signs, and signs of increased intracranial pressure.⁶⁵ Involvement of other cranial nerves has been noted in over half the patients.⁶⁹

Management

Surgical resection is indicated in symptomatic patients with neurological deficits, and most importantly to relieve any mass compression of surrounding structures.⁶⁹ Extracavernous extension involving the orbital apex, superior orbital fissure and petroclival junction limits complete resectability. GTR with graft reconstruction of the abducens nerve has been recommended,⁶⁹ but gross involvement of the cavernous sinus make more convenient to achieve a STR in order to avoid additional neurological damage. A frontotemporal transcavernous approach has been suggested for cavernous sinus and parasellar lesions, whereas the retrosigmoid approach is indicated for tumors involving the preportine region.⁷¹ In a systematic analysis, Sun et al⁶⁹ have reported abducens nerve recovery in less than half of the patients (45%). Tumor extension to the cavernous sinus was significantly associated with a lower likelihood of postsurgical recovery.

SRS (single dose of 12.0–12.5 Gy) is recommended for small and moderate size tumors. Tumor volume was reported to remain stable in 18%, reduced in 37%, and increased in 12% of the patients, respectively (not reported in 31%), treated with gamma knife SRS.⁶⁵

Olfactory Groove Schwannomas (OGS)

Histogenesis

Thus far, only 46 cases with an OGS have been reported.⁸ Schwannomas of the olfactory bulb are uncommon as the olfactory nerve is unsheathed by Schwann cells. There have been various hypotheses proposed to explain the origin of OGS: the developmental hypothesis suggesting transformation of mesenchymal pial cells into ectodermal Schwann cells, and the non-developmental theory stating that these schwannomas may arise from Schwann cells of adjacent normal structures, such as Schwann cell hyperplasia occurring within the perivascular nerve plexuses.^{59,72} In contrast, neural crest cells have the ability

to transform to other cells in the line or acquire abilities from some of these.⁷³

Symptoms and management

In most reported cases of OGS, the chief complaints were headache, vomiting, seizure, diplopia, hyposmia or anosmia (Table 1).^{8,59}

Surgery, either transfrontal or endoscopic, is the preferred treatment for olfactory schwannoma. GTR done by careful extra-arachnoidal dissection is possible as the tumor is well encapsulated and a plane of cleavage from olfactory tract is often present thus allowing preservation.⁵⁹

Optic Nerve Schwannomas (ONS) Histogenesis

Schwannomas partially located in the optic canal with an orbital and/or cisternal component are rare with only few cases reported, but those confined entirely within the optic canal are much rarer. As the optic nerve is in fact a part of the central nervous system, it is myelinated by central nervous system oligodendrocytes rather than Schwann cells.^{1,74} ONS originate from ectopic Schwann cells around the optic nerve or from the sympathetic nerves around the optic nerve in the orbit, probably originating from peripheral sympathetic nerves.^{1,74}

The microscopic appearance of schwannomas of the optic nerve is the same as for all schwannomas of peripheral nerve origin, thus distinguishing them from similar appearing lesions, such as optic nerve gliomas or meningiomas. Optic nerve sheath schwannomas probably cannot be diagnosed on clinical grounds alone, and their neuroimaging appearance mimics that of the more common optic nerve gliomas and the diagnosis is usually made at surgery.⁷⁵

Symptoms and Management

Clinical presentation of lesions arising in the optic nerve is nonspecific and ranges from insidious proptosis, severe visual acuity disturbance and/or visual field defect, retroorbital pain and headaches, to in rare cases blindness. Since the optic canal is a narrow bony structure with a mean diameter of 4 mm, the optic nerve can be compressed by even a tiny tumor.

The tumor can be approached extradurally by drilling the anterior clinoid and unroofing the optic canal. Orbital

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tumors may be safely and effectively resected via a various number of approaches. Currently, a transorbital endoscopic approach alone or combined with an expanded endoscopic endonasal approach is proposed as a method to improve visualization and maximize instruments' maneuverability. Nevertheless, traditional external approaches, with different skin or trans-conjunctival incisions, represent an effective and sound surgical option for the management of these lesions.^{74,76} SRS would be not be indicated because of the risk of optic neuropathy associated with a single large fraction of irradiation. Patients unsuitable for surgery may be considered for fractionated proton irradiation to provide a high likelihood of tumor control while minimizing the dose to the retina.

Discussion

In the era of personalized medicine, trends in NVCNS management are giving way to a more individualized treatment plan based on the combination of surgery, radiation therapy and/or observation.

Because NVCNS are benign tumors, complete surgical excision while avoiding functional and aesthetic sequelae, is a primary objective. Despite the progress made in surgical approaches, resection is still associated with some morbidity in certain circumstances. The treatment goals for NVCNS are heavily dependent on the tumor presentation, location, histology and size.⁶⁷ In addition, patient characteristics such as age and comorbidities must be taken into account. Not all patients should be operated, and surgery would not be an option if the risks are greater than those of a non-surgical treatment or observation. When dealing with tumors involving critical structures (i.e.: cavernous sinus, optic nerve, jugular fossa, etc.), whose damage has a significant impact on quality of life, management decisions must be individualized. Although some authors have reported improvement of symptoms with GTR, in most cases the presenting symptoms persisted or worsened after surgical resection.31

In older and in asymptomatic patients with small tumors, when imaging studies and examination strongly suggest a benign tumor, close surveillance may be considered even without histological evidence, to avoid the risks inherent in biopsy or surgical resection. Likewise, asymptomatic patients with slow-growing small tumors should be managed with caution because they are at high risk of cranial nerve damage if operated on. In these cases, surgery should be avoided if it is expected to cause high morbidity. Gradual loss of nerve function is usually better tolerated than sudden loss as a result of surgery, so observation or deferral of surgery is recommended in patients without neurological alterations.

Safer treatment options have been proposed in some cases. RT has been used as either a primary or an adjunct treatment for NVCNS and offers the advantages of a high tumor control rate and a low rate of radiation associated complications. In contrast to surgical resection, injuries to the brain, vascular, or other cranial nerves are unlikely. SRS may offer therapeutic advantages over conventional RT, particularly logistically because it employs one fraction as opposed to 5 to 6 weeks of daily fractionated RT. Additionally, a tighter margin is employed with SRS so that less normal tissue is irradiated. Disadvantages of SRS are that the risk of a neurologic deficit may be higher due to the large dose per fraction and the risk of a marginal miss may be increased.

Patients with small- to moderate-sized tumors with intact cranial nerve function are optimal candidates for SRS. Furthermore, symptoms related to cranial nerve dysfunction may improve after SRS. In addition, patients with previous contralateral neurological deficits of the lower cranial nerves may not be suitable for surgery because of the risk that bilateral deficits will cause problems with swallowing and laryngeal mobility. In these cases, SRS may be useful. However, it is important to note that sometimes small-sized and slow-growing tumors are very symptomatic, and the symptoms are refractory to medical treatment (i.e., TNS refractory to carbamazepine). In these cases, surgical treatment is safer than follow-up or SRS. RT has been also used for patients who present large tumors or tumors affecting critical areas with high surgical morbidity. Residual tumor after surgery should be treated and SRS or RT are good options with acceptable morbidity.

When it is not possible to achieve a GTR because of technical operative difficulties or when surgery causes significant morbidity, STR or near GTR followed by SRS is gaining acceptance. Concerning TS, Pan et al³⁶ reported that tumor growth control in 56 patients treated by using SRS, was obtained in 93% of the patients (including patients who had previously undergone STR). Cranial nerve preservation in patients with LCNS extending to the jugular foramen is challenging. Lower cranial neuropathies can severely affect quality of life requiring tracheostomy or gastrostomy. Park et al⁴⁴ reported on 13 patients with LCNS treated with GTR and 9 patients

treated by STR followed by SRS. In the latter group, 4 patients (44%) showed a reduction in tumor size and 5 patients (56%) showed no evidence of tumor progression. Postoperative CN deficits were higher in the GTR group (23 vs. 4 CNs affected, respectively). In addition, patients treated by STR and SRS showed a statistically significant improvement in the level of dysphagia. Sedney et al⁷⁷ compared the outcomes of 53 LCNS patients who underwent a GTR with 28 patients receiving a more conservative resection. There was a statistically significant decrease in permanent deficits of CN IX/X with a conservative technique, whereas the recurrence rate was similar in both series (Table 4). The satisfactory clinical results obtained support that an appropriate strategy in the management of some LCNS is surgical resection to reduce tumor size followed by SRS in situations where GTR would likely cause significant morbidity.78-81 We believe that this alternative is potentially safe and effective and should be considered.

The best timing for surgery in FNS is controversial: it depends on facial nerve function, hearing, size, intracranial mass effect and patient choice. If hearing remains good and there is no threat from intracranial mass effect, observation until HB III is possible and radical tumor excision may be performed then. The reason is that even for good preoperative facial function (HB I–II) and regardless of the grafting technique, no better postoperative result than HB-III can be expected, although it is possible to preserve the nerve in approximately 25% of cases if the tumor is located eccentrically on the nerve.⁶¹ Liu and Fagan⁸² analyzed a series of 22 patients with FNS, of whom 12 underwent definitive excision and 10 were managed more conservatively. The best postoperative facial function in the group where the tumor was removal was a HB-III, while in the group treated conservatively, 8 had normal facial function. The role of "wait and see" approach has been evaluated by McMonagle et al⁸³ in a series of 53 patients with FNS where 20 patients were managed conservatively and 33 underwent surgery. There was a GTR in 21 cases and near GTR was obtained in 12 cases. HB grade remained stable in all patients in whom no removal or STR was undertaken, while it worsened in 55% of patients receiving GTR. They conclude that observation is preferred until facial function deteriorates to a HB-III (Table 4). Another alternative is RT to avoid facial nerve deterioration and have a high likelihood of tumor control. Wilkinson et al⁸⁴ reported a series of patients with FNS; through 1995, 85% of cases had surgical resection and none had observation only. Of the patients seen after 1995, 27% had surgical resection and grafting, 33% had bony decompression, 29% were managed with observation alone, and 11% had RT. Facial nerve grade was maintained or improved over the follow-up period in 78.9% of the decompression group and 100% of the observation and RT groups, compared to 54.8% of the resection group (P < 0.012). The trend is towards conservative management of these tumors.

Non-vestibular skull base and intracranial schwannomas

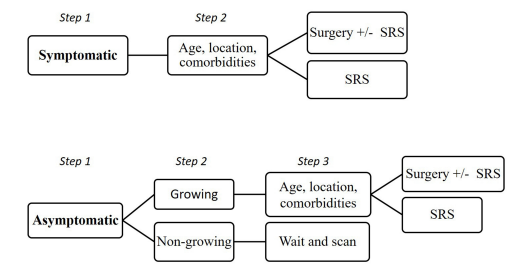


Figure 6 Steps in the management of non-vestibular skull base and intracranial schwannomas.

Conclusion

NVCNS are rare tumors with a diversity of clinical manifestations, depending on the location and cranial nerve of origin. Management decisions are based on tumor size and functional deficits. The treatment of choice is total surgical resection, but SRS or RT are alternatives in those cases where surgical resection would be associated with substantial morbidity. Given the slow growth rate of most of these tumors, observation is also an alternative in asymptomatic patients (Figure 6).

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