Vestibular Schwannoma
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Vestibular schwannoma (VS) is a benign Schwann cell-derived tumour arising from the vestibulocochlear nerve. Although benign, it represents a risk to intracranial structures due to mass effect and carries a small risk of malignant transformation. VS therefore represents an important healthcare burden. We review the literature regarding pathogenesis, risk factors, and diagnosis of VS. The current and future potential management strategies are also discussed. A narrative review of all relevant papers known to the authors was conducted. The majority of VS remain clinically stable and do not require interventional procedures. Nevertheless, various surgical techniques exist for removing VS, the most common of which are translabyrinthine and retrosigmoid approaches. Due to surgical risks such as hearing loss, facial nerve dysfunction, post-operative headache, and cerebrospinal fluid leakage, a “watch and rescan” approach is adopted for most patients. Radiotherapy is a useful alternative and has been shown to have a similar response for growth restriction. Due to the heterogeneous nature of VS, there is a lack of consensus regarding management of tumours that are too large for conservative management but too small to indicate surgery. Emerging biologic therapies, such as Bevacizumab, Everolimus, and Lapatinib, as well as anti-inflammatories like aspirin are promising potential treatments; however, long-term evidence of their efficacy is required. The knowledge base regarding VS continues to improve. With increased understanding of the pathogenesis of these tumors, we believe future work should focus on pharmacologic intervention. Biologic therapies aimed toward improved patient outcomes are particularly promising.

Keywords Vestibular · Schwannoma · Review · Pathology · Diagnosis · Management

Introduction

Vestibular schwannoma (VS), also called acoustic neuroma, is a benign Schwann cell-derived tumour arising from the vestibulocochlear nerve. These tumors represent 85% of intracranial growths arising at the cerebellopontine angle [1]. The Koos grading scale is commonly used to classify tumour size with respect to extrameatal extension and brainstem compression (Table 1) [2]. There is an ongoing debate as to the terminology of VS versus acoustic neuroma. As the majority of tumours arise from the vestibular aspect of the vestibulocochlear nerve and the tumour cells are Schwann cells rather than neuronal, we prefer “vestibular schwannoma” for this article [3].

Although benign, VS represents a risk to various intracranial structures due to mass effect. The most common symptoms include progressive hearing loss and tinnitus which are reported in over 60% of patients. Larger tumours can cause hydrocephalus and brainstem compression leading to symptoms such as facial paraesthesia, vertigo, and headache [4].

VS accounts for approximately 8% of all intracranial tumours with an incidence of 10.4 per million per year [5]. The majority of tumors are unilateral and sporadic. Bilateral disease accounts for less than 5% of cases and is a hallmark of a hereditary disease related to neurofibromatosis type 2 (NF2). Patients typically present between the ages of 20–40; however, those associated with NF2 often manifest earlier [6, 7]. The documented incidence of VS is rising, but there is a general consensus that this is a reflection of increased reporting. The now common use of magnetic resonance imaging (MRI) for symptoms of tinnitus and earlier care
seeking patient behaviours are contributing factors to these higher numbers [8].

This article aims to provide an up to date review of our understanding of the pathogenesis and diagnosis of VS with a focus on current management strategies. Emerging treatment options are also discussed and explored.

**Histologic Features**

The majority of VS grow from the inferior vestibular nerve with rare cases arising from the superior vestibular, or cochlear portion of the nerve [9]. The histologic features of bipolar spindle cells arranged in distinctive Antoni A and Antoni B tissue types are characteristic (Fig. 1).

| Koos grade | Description |
|------------|-------------|
| I          | Intracanicular |
| II         | Extension into cerebellopontine angle, < 2 cm |
| III        | Occupies cerebellopontine angle, no brainstem displacement, < 3 cm |
| IV         | Brainstem displacement, > 3 cm |

**Pathogenesis**

**Molecular Pathogenesis**

Mutations to \( NF2 \), a tumour suppressor gene on chromosome 22, play a vital role in the development of both sporadic and NF2-related disease [11]. Inactivation of the \( NF2 \) protein product, Merlin (schwannomin), leads to deregulation of various intracellular signalling pathways such as Rac1, Ras, PAK1, and mTORC1. Inactivation of other tumour suppressor genes including \( LZTR1 \), \( SMARCB1 \), and \( COQ6 \) are also linked to schwannoma development (Fig. 2) [12, 13].

Although the role of \( NF2 \) mutations was reinforced by recent large-scale sequencing studies, there are data to suggest that NF2-associated VS has a different, polyclonal mutation pattern [17]. This has been postulated to account for variance in treatment outcomes as compared to sporadic VS [18].

**Risk Factors**

Presently, much of the research on risk factors for development of VS is focused on radiation exposure and mobile phone use. High dose ionizing radiation exposure (mean dose 4.6 ± 1.9 Gy) to the cerebellopontine angle in children has been linked to a higher incidence of VS after a latency period of 20–55 years [19]. Other studies from atomic bomb survivors in Japan revealed similar trends [20, 21]. The contrary has been observed with lower dose radiation where no significant link was found between the use of ionizing laden with foamy macrophages and surrounding foci of necrosis, cystic changes, and haemorrhage. In some tumor cells, degenerative nuclear changes can be seen (200×, hematoxylin and eosin). Reproduced with permission from ‘Frequency of loss of heterozygosity of the \( NF2 \) gene in schwannomas from Croatian patients’. Reproduced with permission from Pećina-Slaus et al. [10]
medical imaging and VS development (odds ratio (OR) 0.97, 95% confidence interval (CI) 0.54–1.75) [22].

Multiple epidemiological studies failed to establish a significant link between mobile phone exposure and VS [23–25]. However, a 2009 systematic review and meta-analysis evaluating long-term mobile phone use found a 60% increase in risk for VS following ten years of ipsilateral mobile phone usage (OR 1.6, 95% CI 1.1–2.4) [26]. These findings led to a call for awareness of governments to this risk and even for the classification of mobile phone radiation as a probable human carcinogen [27, 28]. A potential flaw to these data is that most studies relied on patient recall of mobile phone usage introducing recall bias. Furthermore, the majority of the data relied on MRI to confirm the diagnosis without the analysis of tumour histopathology leaving the potential for inaccuracies in the reported findings. Other potential risk factors studied include smoking, occupation, allergic rhinitis, noise exposure, asthma, and eczema [29].

**Diagnosis**

VS is often diagnosed due to otological or neurological symptoms. Otological symptoms include progressive sensorineural hearing loss, unilateral tinnitus, and vertigo. These are more common than the neurological symptoms such as trigeminal and facial nerve impairment, headache, and hydrocephalus amongst the general population [4, 30].

Up to 20% of patients presenting to Ear, Nose, and Throat (ENT) clinics have symptoms suggesting a lesion at the cerebellopontine angle [31]. Therefore, patients presenting with the aforementioned symptoms are generally investigated with otoscopy, pure tone audiometry, and MRI of the internal acoustic meatus. As hearing loss typically presents first, the use of brainstem-evoked response audiometry has proven to be a useful screening tool for a suspected diagnosis. Despite this, it is no longer used as a first-line investigation due to high false-negatives, up to 30%, for small schwannomas and a false positive rate of 10% [32].

All patients presenting with unilateral tinnitus or sensorineural hearing loss require MRI as the gold standard investigation [30, 33]. Fortnum et al. conducted a systematic review and cost-effectiveness study on the use of MRI in diagnosis of VS. Although the use of gadolinium-enhanced T1 weighted MRI is considered the gold standard, there was little difference in sensitivity and specificity compared to non-contrast T2 weighted scans. Furthermore, the use of non-contrast T2 weighted scans was deemed more cost-effective for clinical practice [34].

**Current Treatment Options**

Various approaches exist to treat patients with VS. These include observation, termed a ‘watch and rescan’ approach, surgical removal, and radiotherapy. The main aim of interventional treatment is to remove or debulk the tumour to prevent mass effect [30].

**Conservative**

Observation is an accepted treatment option due to the slow growth rate of VS. A systematic review of 41 papers showed a mean tumour growth rate of 1–2 mm/year with up to 75% of tumours showing no further growth [35]. An observational study of 436 patients showed similar findings where 68% of tumours did not grow during the follow-up period (mean 3.6 years). A third study demonstrated that patients...
with NF2-related disease had higher mean growth rates (1.7 mm/year) as compared to sporadic tumours (1.1 mm/year) [36]. Interestingly, one study observed that sporadic tumours reaching two centimetres in intracranial diameter are likely to continue growing. Despite this, there is no accepted cut-off to dictate when an observational approach transitions to an interventional one. Other cited risk factors for increased growth rates include the presence of an extracanalicular component and diagnosis at a young age; however, these observations are based on limited sample sizes from their respective studies [30, 37].

Although observation can increase the risk of tumour progression and mass effect, it seems a safe approach due to the minimal growth rate. Furthermore, delaying surgical intervention appears to have no increased risk in mortality [38]. Since growth usually manifests within the first 3 years after presentation, a recommended protocol for observation is serial MRI scans at 6 month intervals for 2 years and then another scan 2 years later. Following this, patients should be scanned every 5 years with lifelong follow-up [39].

Surgery

A range of surgical methods for VS removal are discussed in the literature and mainly pertain to large tumours. The approaches include: middle fossa (MF), extended middle fossa, transtotic, and endoscopic resection. The translabyrinthine (TL) and retrosigmoid (RS) approaches are the most commonly used in the United Kingdom (UK) and are further discussed below [30]. The benefit of the MF approach for small tumours of greater hearing preservation versus the RS approach is worth mentioning. However, its use is now limited in the UK due to high risks of damage to the facial nerve and seizures caused by temporal lobe manipulation [40, 41].

The TL approach refers to a retroauricular incision with posterior extension toward the mastoid tip. This provides quick access to the internal acoustic canal, facial nerve, and tumour. A major disadvantage is poor preservation of hearing, limiting the use of the TL approach to those with minimal or absent preoperative hearing.

With the RS approach, a suboccipital incision exposes the transverse and sigmoidal sinuses, and subsequent cerebellar manipulation exposes the internal acoustic canal. Although this method is superior in preserving hearing, there is an increased risk of facial and cochlear nerve damage [30]. Of course, surgical intervention within the cranium is not without other risks such as meningitis, cerebellar injury, epilepsy, persistent headache, and cerebrospinal fluid leakage [42–45].

In a systematic review of 35 studies (n = 5064), it was reported that facial nerve dysfunction was less likely to occur with the RS versus TL approach for tumours greater than 1.5 cm [40]. Evidence exists to suggest the size of the tumour preoperatively is an important prognostic factor for facial nerve dysfunction postoperatively [46]. The study compared surgical interventions based on tumour size which represents a potential confounder to the data. They also reported postoperative headaches and CSF leak rate to be significantly greater following the RS approach versus TL. It is postulated that postoperative headaches are more common with the RS approach due to the incision size and remnant intracranial bone dust [30]. Conflicting data exists regarding CSF leak; Magnus et al. reported no difference between RS and TL approaches in 1,922 patients while Sughrue et al. found the TL approach was riskier [47, 48].

Mortality and tumour recurrence did not differ between surgical approaches with mortality rates as low as 0.2% [48]. Non-specific neurological complications such as cerebellar dysfunction, stroke, and epilepsy were rare and occurred in less than 3% of cases [40]. The impact of surgery on patient well-being was notable as the procedure potentially delays a return to work for up to four months. This may lead to financial hardship and adverse affects on mental health [49].

Radiotherapy

There are three forms of radiotherapy treatment for VS management: stereotactic radiosurgery (SRS), fractionated stereotactic radiotherapy (FSRT), and proton beam therapy. SRS and FRST are most commonly used, as there is limited availability of proton beam therapy and inadequate evidence of its efficacy [50]. The main aim of radiotherapy is to prevent tumour growth, therefore this is not considered a suitable approach for large tumours with mass effect.

Various advantages and disadvantages apply to SRS and FRST. SRS uses Gamma Knife technology to expose the tumour to a singular dose of radiation and is less applicable to large lesions (> 2.5 cm extracanalicular diameter) [51]. FRST requires numerous sessions of radiotherapy in an attempt to target the tumour at the most radiation-sensitive phase of the cell cycle for, theoretically, greater efficacy [52]. In addition, FRST systems are more readily available in hospitals and can be used on larger lesions [51].

A recent systematic review comparing FRST and SRS reported similar rates of tumour control with an average of 4.8% and 5% of patients undergoing treatment requiring rescue therapy, respectively. Facial and trigeminal nerve deterioration was less for SRS. These comparisons were based on limited evidence; however, and without randomised controlled trials (RCTs). Furthermore, only two studies on FRST were evaluated limiting the validity of estimates regarding this treatment. More studies are required to confidently compare these two therapies [53].

Ranges of controlled studies demonstrate comparable progression-free survival rates and side effects, such as nerve palsies and hearing deterioration, between radiotherapy
Emerging Treatments

As knowledge surrounding the molecular pathology of VS improves, targeted biologic therapies are emerging at the forefront of treatment. Bevacizumab, Everolimus, and Lapatinib are potential options for treating VS. Furthermore, a link between tumour growth and non-steroidal anti-inflammatory medications such as aspirin has been recognised.

Bevacizumab is a monoclonal antibody and vascular endothelial growth factor (VEGF) inhibitor. VEGF is a key mediator of angiogenesis and subsequently aids tumour growth [66]. Plotkin et al. conducted the earliest research on the use of Bevacizumab in NF2 patients with progressive disease. Although these studies reported improvement in hearing and restriction of tumour growth in over 50% of patients, they were based on limited sample sizes (n = 10 and n = 31, respectively). The later 2012 study was also based on retrospective data with a median treatment duration of only 14 months [67, 68].

Since then, more promising data has arisen. In a systematic review on the safety and efficacy of Bevacizumab, 41% of patients receiving treatment had tumour regression, 20% experienced hearing improvement, and 69% had no further hearing deterioration [69]. This promising evidence supports the funding of Bevacizumab by NHS England for VS treatment; however, these data were only based on NF2 patients. Therefore is difficult to know whether the results will translate to the majority of the sporadic VS cases [70, 71].

Furthermore, various side effects have been reported in a dose-dependent relationship with Bevacizumab including hypertension, proteinuria, and infection [69, 72]. Although the most recent systematic review by the Congress of National Neurological Surgeons (CNS) supports the use of Bevacizumab, the recommendations are based on level 3 evidence [73]. It seems clear more long-term evidence pertaining to safety and appropriate dosing is required.

Everolimus is an mTOR complex 1 (mTORC1) inhibitor. mTORC1 activation has been implicated in tumour growth as a result of merlin deficiency. Furthermore, inhibition of mTORC1 has demonstrated antiangiogenic properties [74, 75]. In theory, this seems a promising treatment; however, there is limited evidence of its clinical use. Phase II trials in NF2 patients have shown mixed results. Karajannis et al. found no response to Everolimus on tumour growth or hearing improvements [76]. Although Goutagny et al. found a 66.5% reduction in tumour growth during Everolimus use in ten NF2 patients, growth resumed following the discontinuation of treatment [77]. At the time of writing, the CNS does not support use of Everolimus for VS management [73].

Lapatinib is an EGFR/ErbB2 inhibitor with demonstrated promise for tumor growth inhibition during in vitro trials [78]. The CNS recognises Lapatinib as a potential agent for growth management and hearing improvement due to encouraging early clinical evidence [73]. A phase II clinical trial involving 21 NF2 patients found a ≥ 15% decrease in tumour volume in 23.5% of participants following serial MRI scans, and 30.8% of participants showed improved hearing. Only 14 of the 21 participants were eligible for evaluation of audiological response, however. The lack of a control group also limits these findings. Encouragingly, there were low levels of reported toxicity with Lapatinib which may prove important when comparing to Bevacizumab and its propensity for adverse side effects [79]. Again, long-term controlled studies are required to provide more robust evidence.

Studies also suggest that aspirin may be beneficial to delay tumour growth through its anti-cyclooxygenase 2 (COX-2) effect. A proliferative effect of COX-2 expression in 30 tumours was observed [80]. Another study demonstrated the application of acetylsalicylic acid decreased COX-2 expression and in vitro cell growth [81]. Furthermore, a retrospective analysis followed 347 patients with VS, taking aspirin for unrelated reasons, to inspect the effect on volumetric growth versus non-aspirin users. They found a significant inverse association with aspirin users and tumour growth (OR 0.32, 95% CI 0.11–0.91) [82]. Although two similar retrospective analyses did not find an inverse association, they did not observe increased tumour growth with aspirin or other NSAID users [83, 84]. Therefore, a level 3 recommendation exists in the CNS stating aspirin may be considered for patients undergoing observation [73].
More recent evidence is not as promising. In a 2019 study, tissue samples of 1048 tumours were analysed. It was discovered that while COX-2 expression increased with tumour proliferation, the use of acetylsalicylic acid did not alter COX-2 expression [85]. This study used tumour tissue retrospectively which can lend itself to selection bias since large tumours are more likely to be managed surgically. Most of the tissue samples were likely obtained from clinically stable tumours that may not have grown independent of the presence of the acetylsalicylic acid. In any case, the conflicting research on the use of aspirin necessitates further investigated through large-scale trials. Currently, there is one prospective phase II randomised double blind trial recruiting patients to assess the effect of acetylsalicylic acid on tumour growth and hearing preservation (NCT03079999).

Conclusions

VS is a clinically important disease with an evolving knowledge base. There is now a vast scope of literature pertaining to risk factors, diagnosis, and treatment. With improved understanding of the pathogenesis, future work should focus on biologic interventions due to the current risks and lack of high quality RCTs of different surgical and radiologic treatment options.

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Compliance with Ethical Standards

Conflict of interest  On behalf of all authors, the corresponding author states that there is no conflict of interest.

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References

1. Singh K, Singh MP, Thukral C, Rao K, Singh K, Singh A. Role of magnetic resonance imaging in evaluation of cerebellopontine angle schwannomas. Indian J Otolaryngol Head Neck Surg. 2015;67(1):21–7. https://doi.org/10.1007/s12070-014-0756-0.
2. Koos WT, Day JD, Matula C, Levy DJ. Neurotographic considerations in the microsurgical treatment of small acoustic neurenomas. J Neurol Surg. 1998;58(3):506–12. https://doi.org/10.3171/jns.1998.88.3.0506.
3. Dublin A. Acoustic neuroma or vestibular schwannoma? Skull Base. 2009;19(5):375. https://doi.org/10.1055/s-0029-1220205.
4. Matthies C, Samii M. Management of 1000 vestibular schwannomas (acoustic neuromas): clinical presentation. Neurosurgery. 1997;40(1):1–9. https://doi.org/10.1097/00006123-199701000-00001 (discussion 10).
5. Evans DG, Moran A, King A, Saeed S, Gurusinghe N, Ramsden R. Incidence of vestibular schwannoma and neurofibromatosis 2 in the North West of England over a 10-year period: higher incidence than previously thought. Otol Neurotol. 2005;26(1):93–7. https://doi.org/10.1017/S00129492-200501000-00016.
6. Miller AB, Morgan LL, Udassin I, Davis DL. Cancer epidemiology update, following the 2011 IARC evaluation of radiofrequency electromagnetic fields (Monograph 102). Environ Res. 2018;167:673–83. https://doi.org/10.1016/j.envres.2018.06.043.
7. Koo M, Lai JT, Yang EY, Liu TC, Hwang JH. Incidence of vestibular schwannoma in Taiwan from 2001 to 2012: a population-based national health insurance study. Ann Otol Rhinol Laryngol. 2018;127(10):694–7. https://doi.org/10.1177/0003489418788385.
8. Stangerup SE, Tos M, Thomsen J, Caye-Thomasen P. True incidence of vestibular schwannoma? Neurosurgery. 2010;67(5):1335–400. https://doi.org/10.1227/NEU.0b013e318f22660 (discussion 40).
9. Khrais T, Romano G, Sanna M. Nerve origin of vestibular schwannoma: a prospective study. J Laryngol Otol. 2008;122(2):128–31. https://doi.org/10.1017/S0022215107001028.
10. Pecina-Slaus N, Zeljko M, Pecina HI, Nikuseva Martic T, Bacic N, Tomas D, et al. Frequency of loss of heterozygosity of the NF2 gene in schwannomas from Croatian patients. Croat Med J. 2012;53(4):321–7. https://doi.org/10.3325/cmj.2012.53.321.
11. Warren C, James LA, Ramsden RT, Wallace A, Baser ME, Varley JM, et al. Identification of recurrent regions of chromosome loss and gain in vestibular schwannomas using comparative genomic hybridisation. J Med Genet. 2003;40(11):802–6. https://doi.org/10.1136/jmg.40.11.802.
12. Brodhun M, Stahn V, Harder A. Pathogenesis and molecular pathology of vestibular schwannoma. HNO. 2017;65(5):362–72. https://doi.org/10.1007/s00106-016-0201-3.
13. Mehta GU, Feldman MJ, Wang H, Ding D, Chittiboina P. Unilateral vestibular schwannoma in a patient with schwannomatosis in the absence of LZTR1 mutation. J Neurol Surg. 2016;125(6):1469–71. https://doi.org/10.3171/2015.11.JNS151766.
14. Thaxton C, Lopera J, Bott M, Fernandez-Valle C. Neuregulin and laminin stimulate phosphorylation of the NF2 tumor suppressor in Schwann cells by distinct protein kinase A and p21-activated kinase-dependent pathways. Oncogene. 2008;27(19):2705–15. https://doi.org/10.1038/sj.onc.1210923.
15. Fiala C, Utermark T, Parkinson DB, Poetsch A, Hanemann CO. Impaired intercellular adhesion and immature adherens junctions in merlin-deficient human primary schwannoma cells. Glia. 2008;56(5):506–15. https://doi.org/10.1002/glia.20629.
16. Jacoby LB, Jones D, Davis K, Kronn D, Short MP, Gusella J, et al. Molecular analysis of the NF2 tumor-suppressor gene in
schwannomatosis. Am J Hum Genet. 1997;61(6):1293–302. https://doi.org/10.1086/301633.

17. Havik AL, Brunland O, Myrseth E, Miletic H, Aarhus M, Knappskog PM, et al. Genetic landscape of sporadic vestibular schwannoma. J Neurosurg. 2018;128(3):911–22. https://doi.org/10.3171/2016.10.JNS161384.

18. Dewan R, Pemov A, Kim HJ, Morgan KL, Vasquez RA, Chit-tiboina P, et al. Evidence of polyclonality in neurofibromatosis type 2-associated multilobulated vestibular schwannomas. Neuro Oncol. 2015;17(4):566–73. https://doi.org/10.1093/neuonc/nou317.

19. Schneider AB, Ron E, Lubin J, Stovall M, Shore-Freedman E, Tolentino J, et al. Acoustic neuromas following childhood radiation treatment for benign conditions of the head and neck. Neuro Oncol. 2008;10(1):73–8. https://doi.org/10.1215/15227517-2007-0047.

20. Preston DL, Ron E, Yonehara S, Kobuke T, Fujii H, Kishikawa M, et al. Tumors of the nervous system and pituitary gland associated with atomic bomb radiation exposure. J Natl Cancer Inst. 2002;94(20):1555–63. https://doi.org/10.1093/jnci/94.20.1555.

21. Yonehara S, Brenner AV, Kishikawa M, Inskip PD, Preston DL, Ron E, et al. Clinical and epidemiologic characteristics of first primary tumors of the central nervous system and related organs among atomic bomb survivors in Hiroshima and Nagasaki, 1958–1995. Cancer. 2004;101(17):1644–54. https://doi.org/10.1002/cncr.20543.

22. Blettner M, Schlehofer B, Samkange-Zebs F, Berg G, Schlaefer K, Schurz J. Medical exposure to ionizing radiation and the risk of brain tumours: interphone study group. Germany Eur J Cancer. 2007;43(13):1990–8. https://doi.org/10.1016/j.ejca.2007.06.020.

23. Christensen HC, Schuz J, Kosteljanczetz M, Poulsen HS, Thomsen J, Johansen C. Cellular telephone use and risk of acoustic neuroma. Am J Epidemiol. 2004;159(3):277–83. https://doi.org/10.1093/aje/kwh032.

24. Ahlbom A, Feychtling M, Green A, Kheifets L, Savitz DA, Swedlow AJ, et al. Epidemiologic evidence on mobile phones and tumor risk: a review. Epidemiology. 2009;20(5):639–52. https://doi.org/10.1097/EDE.0b013e3181b0927d.

25. Takebayashi T, Aki sa B, Kikuchi Y, Taki M, Wake K, Watanabe S, et al. Mobile phone use and acoustic neuroma risk in Japan. Occup Environ Med. 2006;63(12):802–7. https://doi.org/10.1136/oem.2006.028308.

26. Hardell L, Carlberg M, Hansson MK. Epidemiological evidence for an association between use of wireless phones and tume. Pathophysiology. 2009;16(2–3):113–22. https://doi.org/10.1016/j.pathophys.2009.01.003.

27. Hardell L, Carlberg M, Hansson MK. Use of mobile phones and cordless phones is associated with increased risk for glomi and acoustic neuroma. Pathophysiology. 2013;20(2):85–110. https://doi.org/10.1016/j.pathophys.2012.11.001.

28. Morgan LL, Miller AB, Casco A, Davis DL. Mobile phone radiation causes brain tumors and should be classed as a probable human carcinogen (2A) (review). Int J Oncol. 2015;46(5):1865–71. https://doi.org/10.3892/ijo.2015.2908.

29. Berkowitz O, Iyer AK, Kano H, Talbott EO, Lunsford LD. Epidemiology and environmental risk factors associated with vestibular Schwannoma. World Neurosurg. 2015;84(6):1674–80. https://doi.org/10.1016/j.wneu.2015.07.007.

30. Halliday J, Rutherford SA, McCabe MG, Evans DG. An update on the diagnosis and treatment of vestibular schwannoma. Expert Rev Neurother. 2018;18(1):29–39. https://doi.org/10.1080/14737715.2018.1399795.

31. Harcourt JP, Vijaya-Sekaran S, Loney E, Lennox P. The incidence of symptoms consistent with cerebellopontine angle lesions in a general ENT out-patient clinic. J Laryngol Otol. 1999;113(6):518–22. https://doi.org/10.1017/s0022215100144391.

32. Doyle KJ. Is there still a role for auditory brainstem response audiometry in the diagnosis of acoustic neuroma? Arch Otolaryngol Head Neck Surg. 1999;125(2):232–4. https://doi.org/10.1001/archotol.125.2.232.

33. NICE. Tinnitus. 2017. https://cks.nice.org.uk/tinnitus—!topicSummary.

34. Fortnum H, O’Neill C, Taylor R, Lenthall R, Nikolopoulos T, Lightfoot G, et al. The role of magnetic resonance imaging in the identification of suspected acoustic neuroma: a systematic review of clinical and cost effectiveness and natural history. Health Technol Assess. 2009;13(18):iii-iv, ix-xi, 1–154https://doi.org/10.3310/hta3180.

35. Nikolopoulos TP, Fortnum H, O’Donoghue G, Baguley D. Acoustic neuroma growth: a systematic review of the evidence. Otol Neurotol. 2010;31(3):478–85. https://doi.org/10.1097/MAO.0b013e3181d279a3.

36. Suryanarayanan R, Ramsden RT, Saeed SR, Aggarwal R, King AT, Rutherford SA, et al. Vestibular schwannoma: role of conservative management. J Laryngol Otol. 2010;124(3):251–7. https://doi.org/10.1017/S0022215109992362.

37. Ogawa K, Kanzaki J, Ogawa S, Yamamoto M, Ikeda S, Shibara R. The growth rate of acoustic neuromas. J Neurosurg. 1995. Cancer. 2004;101(7):1644–54. https://doi.org/10.1017/s0022215109992362.

38. Martin TP, Senthil L, Chavda SV, Walsh R, Irving RM. A protocol for the conservative management of vestibular schwannomas. Otol Neurotol. 2009;30(3):381–5. https://doi.org/10.1097/mao.0b013e31819a8d16.

39. Ansari SF, Terry C, Cohen-Gadol AA. Surgery for vestibular schwannomas: a systematic review of complications by approach. Neurosurg Focus. 2012;33(3):E14. https://doi.org/10.3171/2012.6.FOCUS12163.

40. Irving RM, Jackler RK, Pitts LH. Hearing preservation in patients undergoing vestibular schwannoma surgery: comparison of middle fossa and retrosigmoid approaches. J Neurosurg. 1998;88(5):840–5. https://doi.org/10.3171/jns.1998.88.5.0840.

41. Irving RM, Jackler RK, Pitts LH. Hearing preservation in patients undergoing vestibular schwannoma surgery: comparison of middle fossa and retrosigmoid approaches. J Neurosurg. 1998;88(5):840–5. https://doi.org/10.3171/jns.1998.88.5.0840.

42. Douglas SA, Yeung P, Daudia A, Gatehouse S, O’Donoghue GM. Spatial hearing disability after acoustic neuroma removal. Laryngoscope. 2007;117(9):1648–51. https://doi.org/10.1097/MLG.0b013e3180ca162.

43. Schick B, Dlugaczky J. Surgery of the ear and the lateral skull base: pitfalls and complications. GMS Curr Top Otorhinolaryngol Head Neck Surg. 2013;12:Doc05. https://doi.org/10.3205/cto00097.

44. Tufarelli D, Meli A, Labini FS, Badaracco C, De Angelis E, Alesii A, et al. Balance impairment after acoustic neuroma surgery. Otol Neurotol. 2011;32(9):1525–9. https://doi.org/10.1097/MAO.0b013e31823e4a4.
Pawlik TM, Keyomarsi K. Role of cell cycle in mediating sensitivity to radiotherapy. Int J Radiat Oncol Biol Phys. 2004;59(4):928–36. https://doi.org/10.1016/j.ijrobp.2004.03.005.

Webber DC, Chan AW, Bussiere MR, Harsh GRT, Ancukiewicz M, Barker FG 2nd, et al. Proton beam radiosurgery for vestibular schwannoma: tumor control and cranial nerve toxicity. Neurosurgery. 2003;53(3):577–86. https://doi.org/10.1227/01.NEU.0000093699.59231.e0 (discussion 86–8).

Weber DC, Chan AW, Bussiere MR, Harsh GRT, Ancukiewicz M, Barker FG 2nd, et al. Proton beam radiosurgery for vestibular schwannoma: tumor control and cranial nerve toxicity. Neurosurgery. 2003;53(3):577–86. https://doi.org/10.1227/01.NEU.0000093699.59231.e0 (discussion 86–8).

Chen AW, Bussiere MR, Harsh GRT, Ancukiewicz M, Barker FG 2nd, et al. Proton beam radiosurgery for vestibular schwannoma: tumor control and cranial nerve toxicity. Neurosurgery. 2003;53(3):577–86. https://doi.org/10.1227/01.NEU.0000093699.59231.e0 (discussion 86–8).

Combs SE, Welzel T, Schulz-Ertner D, Huber PE, Debus J. Differences in clinical results after LINAC-based single-dose radiosurgery versus fractionated stereotactic radiotherapy for patients with vestibular schwannomas. Int J Radiat Oncol Biol Phys. 2010;76(1):193–200. https://doi.org/10.1016/j.ijrobp.2009.01.064.

Pawlik TM, Keyomarsi K. Role of cell cycle in mediating sensitivity to radiotherapy. Int J Radiat Oncol Biol Phys. 2004;59(4):928–42. https://doi.org/10.1016/j.ijrobp.2004.03.005.

Persson O, Barket J Jr, Shalom NB, Wangerd T, Jakola AS, Forander P. Stereotactic radiosurgery vs. fractionated radiotherapy for tumor control in vestibular schwannoma patients: a systematic review. Acta Neurochir (Wien). 2017;159(6):1013–21. https://doi.org/10.1007/s00701-017-3164-6.

Chen AW, Black P, Ojemann RG, Barker FG 2nd, Kooy HM, Lopes VV, et al. Stereotactic radiotherapy for vestibular schwannomas: favorable outcome with minimal toxicity. Neurosurgery. 2005;57(1):60–70. https://doi.org/10.1227/01.NEU.0000163091.12329.bb (discussion 60–70).

Combs SE, Engelhard C, Kopp C, Wiedenmann N, Schramm O, Proikic V, et al. Long-term outcome after highly advanced single-dose or fractionated radiotherapy in patients with vestibular schwannomas—pooled results from 3 large German centers. Radiother Oncol. 2015;114(3):378–83. https://doi.org/10.1016/j.radonc.2015.01.011.

Hasegawa T, Kida Y, Kato T, Iizuka H, Kuramitsu S, Yamamoto T. Long-term safety and efficacy of stereotactic radiosurgery for vestibular schwannomas: evaluation of 440 patients more than 10 years after treatment with Gamma Knife surgery. J Neurol Surg. 2013;118(3):357–65. https://doi.org/10.1055/s-0032-132523.

Myrseth E, Moller P, Pedersen PH, Vassbolth FS, Wenzel-Larsen T, Lund-Johansen M. Vestibular schwannomas: clinical results and quality of life after microsurgery or gamma knife radiosurgery. Neurosurgery. 2005;56(5):927–35. https://doi.org/10.1093/neo.56.5.927 (discussion 35).

Pollock BE, Driscoll CL, Foote RL, Link MJ, Gorman DA, Bauch CD, et al. Patient outcomes after vestibular schwannoma management: a prospective comparison of microsurgical resection and stereotactic radiosurgery. Neurosurgery. 2006;59(1):77–85. https://doi.org/10.1227/01.NEU.0000192177.14930.14 (discussion 77–85).

Sun S, Liu A. Long-term follow-up studies of Gamma Knife surgery with a low margin dose for vestibular schwannoma. J Neurol Surg. 2012;71(1):57–62. https://doi.org/10.1055/s-0031-12783.

Myrseth E, Moller P, Pedersen PH, Lund-Johansen M. Vestibular schwannoma: surgery or gamma knife radiotherapy? A prospective, nonrandomized study. Neurosurgery. 2009;64(4):654–61. https://doi.org/10.1227/01.NEU.0000340684.60443.55 (discussion 61–3).

Wolbers JG, Dellangia AH, Mendez Romero A, van Linge A. What intervention is best practice for vestibular schwannomas? A systematic review of controlled studies. BMJ Open. 2013. https://doi.org/10.1136/bmjopen-2012-001345.

Muzevic D, Legcevic J, Splavski B, Caje-Thomasen P. Stereotactic radiotherapy for vestibular schwannoma. Cochrane Database Syst Rev. 2014;12:CD009897. https://doi.org/10.1002/14651858.CD009897.pub2.

Yang I, Aranda D, Han SJ, Chennupati S, Sughrue ME, Cheung SW, et al. Hearing preservation after stereotactic radiosurgery for vestibular schwannoma: a systematic review. J Clin Neurosci. 2009;16(6):742–7.

Frischer JM, Gruber E, Schoffmann V, Ertl A, Hoftberger R, Mallouhi A, et al. Long-term outcome after Gamma Knife radiotherapy for acoustic neuroma of all Koos grades: a single-center study. J Neurosurg. 2018. https://doi.org/10.3171/2017.8.JNS171281.

Hentschel M, Rovers M, Markodimitraki L, Steens S, Kunst H. An international comparison of diagnostic and management strategies for vestibular schwannoma. Eur Arch Otorhinolaryngol. 2019;276(1):71–8. https://doi.org/10.1007/s00405-018-5199-6.

Shih T, Lindley C. Bevacizumab: an angiogenesis inhibitor for the treatment of solid malignancies. Clin Ther. 2006;28(11):1779–802. https://doi.org/10.1016/j.clinthera.2006.11.015.

Plotkin SR, Merker VL, Halpin C, Jennings D, McKenna MJ, Harris GJ, et al. Bevacizumab for progressive vestibular schwannoma in neurofibromatosis type 2: a retrospective review of 31 patients. Otol Neurotol. 2012;33(6):1046–52. https://doi.org/10.1097/MAO.0b013e3182573f5f.

Lu VM, Ravindran K, Grafeo CS, Perry A, Van Gompel JJ, Daniels DJ, et al. Efficacy and safety of bevacizumab for vestibular schwannoma in neurofibromatosis type 2: a systematic review and meta-analysis of treatment outcomes. J Neurooncol. 2019;144(2):239–48. https://doi.org/10.1007/s11060-019-03234-8.

Morris KA, Golding JF, Axon PR, Afridi S, Blessing C, Ferner RE, et al. Bevacizumab in neurofibromatosis type 2 (NF2) related vestibular schwannomas: a nationally coordinated approach to delivery and prospective evaluation. Neurooncol Pract. 2016;3(4):281–9. https://doi.org/10.1093/npwp/wpw065.

England N, 2013/14 NHS Standard Contract for Neurofibromatosis Type 2 Service (All ages) 2013. https://www.england.nhs.uk/wp-content/uploads/2013/06/6b13_neurofib2.pdf.

Morris KA, Golding JF, Blessing C, Evans DG, Ferner RE, Foweraker K, et al. Toxicity profile of bevacizumab in the UK Neurofibromatosis type 2 cohort. J Neurooncol. 2017;131(1):117–24. https://doi.org/10.1007/s11060-016-2276-9.

Van Gompel JJ, Agazzi S, Carlson ML, Adewumi DA, Hadjipanayis CG, Uhm JH, et al. Congress of neurological surgeons systematic review and evidence-based guidelines on emerging therapies for the treatment of patients with vestibular Schwannomas. Neurosurgery. 2018;82(2):E52–E545. https://doi.org/10.1093/neuros/nyx516.

James MF, Han S, Poliziano C, Plotkin SR, Manning BD, Stemmer-Rachamimov AO, et al. NF2/merlin is a novel negative regulator of mTOR complex 1, and activation of mTORC1 is associated with meningioma and schwannoma growth. Mol Cell Biol. 2009;29(15):4250–61. https://doi.org/10.1128/MCB.01581-08.

Lane HA, Wood JM, McSheehy PM, Allegri PNG, Pollock CA, Brueggen J, et al. mTOR inhibitor RAD001 (everolimus) has antiangiogenic/vascular properties distinct from a VEGFR tyrosine kinase inhibitor. Clin Cancer Res. 2009;15(5):1612–22. https://doi.org/10.1158/1078-0432.CCR-08-2057.
76. Karajannis MA, Legault G, Hagiwara M, Giancotti FG, Filatov A, Derman A, et al. Phase II study of everolimus in children and adults with neurofibromatosis type 2 and progressive vestibular schwannomas. Neuro Oncol. 2014;16(2):292–7. https://doi.org/10.1093/neuonc/not150.

77. Goutagny S, Raymond E, Esposito-Farese M, Trunet S, Mawrin C, Bernardeschi D, et al. Phase II study of mTORC1 inhibition by everolimus in neurofibromatosis type 2 patients with growing vestibular schwannomas. J Neurooncol. 2015;122(2):313–20. https://doi.org/10.1007/s11060-014-1710-0.

78. Ammoun S, Cunliffe CH, Allen JC, Chiriboga L, Giancotti FG, Zagzag D, et al. ErbB/HER receptor activation and preclinical efficacy of lapatinib in vestibular schwannoma. Neuro Oncol. 2010;12(8):834–43. https://doi.org/10.1093/neuonc/noq012.

79. Karajannis MA, Legault G, Hagiwara M, Ballas MS, Brown K, Nusbaum AO, et al. Phase II trial of lapatinib in adult and pediatric patients with neurofibromatosis type 2 and progressive vestibular schwannomas. Neuro Oncol. 2012;14(9):1163–70. https://doi.org/10.1093/neuonc/nos146.

80. Hong B, Krusche CA, Schwabe K, Friedrich S, Klein R, Krauss JK, et al. Cyclooxygenase-2 supports tumor proliferation in vestibular schwannomas. Neurosurgery. 2011;68(4):1112–7. https://doi.org/10.1227/NEU.0b013e318208f5c7.

81. Dilwali S, Kao SY, Fujita T, Landegger LD, Stankovic KM. Nonsteroidal anti-inflammatory medications are cytostatic against human vestibular schwannomas. Transl Res. 2015;166(1):1–11. https://doi.org/10.1016/j.trsl.2014.12.007.

82. Kandathil CK, Cunnane ME, McKenna MJ, Curtin HD, Stankovic KM. Correlation between aspirin intake and reduced growth of human vestibular schwannoma: volumetric analysis. Otol Neurotol. 2016;37(9):1428–34. https://doi.org/10.1097/MAO.0000000000001180.

83. Hunter JB, O’Connell BP, Wanna GB, Bennett ML, Rivas A, Thompson RC, et al. Vestibular Schwannoma growth with aspirin and other nonsteroidal anti-inflammatory drugs. Otol Neurotol. 2017;38(8):1158–64. https://doi.org/10.1097/MAO.0000000000001506.

84. MacKeith S, Wasson J, Baker C, Guilloyle M, John D, Donnelly N, et al. Aspirin does not prevent growth of vestibular Schwannomas: a case-control study. Laryngoscope. 2018;128(9):2139–44. https://doi.org/10.1002/lary.27114.

85. Behling F, Ries V, Skardelly M, Gepfner-Tuma I, Schuhmann M, Ebner FH, et al. COX2 expression is associated with proliferation and tumor extension in vestibular schwannoma but is not influenced by acetylsalicylic acid intake. Acta Neuropathol Commun. 2019;7(1):105. https://doi.org/10.1186/s40478-019-0760-0.

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