Cystic Vestibular Schwannomas and Post–Gamma Knife Radiosurgery-Induced Necrosis of Vestibular Schwannomas: Report of Two Cases and Review of Literature

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Abstract

Cystic vestibular schwannomas (VS) form a rare subgroup that differs from the solid variant clinically and radiologically. Certain unique features reported in cystic VS are larger size, presentation with atypical initial symptoms such as dysgeusia, vertigo, facial pain and gait unsteadiness, short duration of symptoms, rapid progression of symptoms or sudden deterioration and frequent involvement of facial nerve. Gamma Knife radiosurgery (GKRS) is a safe and effective treatment for VS, achieving tumor control and resulting in a very low morbidity rate. Here, we described two cases of cystic vestibular schwannomas prior to GKRS and their radiological appearances post-GKRS.

Case Report

A 76-year-old female presented with gradual left-sided hearing loss and impaired cerebellar function tests. Magnetic resonance imaging (MRI) revealed a cystic left cerebellopontine angle lesion causing significant mass effect on adjacent cerebellum, pons, and cerebellar peduncle with vasogenic edema in the ipsilateral cerebellum. Left vestibulocochlear, facial nerve complex root entry zone (origin), is masked by the lesion (►Fig. 1). A 58-year-old female presented with gradual hearing loss and left-sided tinnitus. MRI revealed marked peripheral enhancement of the left cerebellopontine angle lesion with central nonenhancing areas (►Fig. 2). A 51-year-old female presented with vertigo, facial weakness, and hearing loss emerging 6 weeks following Gamma Knife radiosurgery (GKRS) of a right-sided vestibular schwannoma (VS). Brain MRI demonstrated central nonenhancement within the schwannoma, likely related to radiation necrosis (►Figs. 3 and 4).

Discussion

VSs account for approximately 8 to 10% of primary intracranial tumors and 80% of cerebellopontine angle tumors. These tumors may undergo cystic changes, frequently as microcysts, caused by degenerative changes within the tumors. Occasionally, these areas may coalesce to form larger cysts. Piccirillo et al2 classified cystic VS as follows: type A—central and thick-walled cysts which can be further subdivided as (1) polycystic (multiple small intratumoral cysts with thick-walled cyst), (2) polycystic (multiple moderate-sized intratumoral cysts with thick-walled cyst), and (3) monocystic (single-large cyst); and type B—peripheral and thin-walled cysts which can be further classified as (1) anterior, (2) medial, (3) posterior, and (4) combined. It has also been found that a cystic VS is less responsive to Gamma Knife surgery than a solid counterpart.3 In conclusion, cystic VS are a heterogeneous group presenting with short duration and rapid progression of symptoms with frequent facial nerve involvement due to sudden expansion of the cystic elements. VSs are the most common tumors of the cerebellopontine angle (CPA), accounting for more than 90% of all such tumors.4 VS are typically benign, slow-growing tumors. Symptoms at diagnosis commonly include hearing deficiency or loss, tinnitus, loss of balance, and, more rarely, change in facial sensation and headache. Diagnosis is performed by MRI showing a typical gadolinium-enhanced lesion. The principal management options for VS are watchful waiting,

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Clinical studies by Yang et al, which utilized GKRS for the treatment of large VS (diameter greater than 3 cm), revealed worse tumor control rates and higher patient morbidity as compared with small tumors (diameter ≤3 cm). Based on the aforementioned studies, neurosurgeons and radiation
oncologists must take tumor size, growth rate, and location, as well as the patient's neurological function and age, into account when determining the ideal course of action for a VS. Large tumors causing significant brainstem and cranial nerve compression, or those that have already failed radiosurgery require surgical intervention. GKRS is a safe and effective treatment for VS, achieving tumor control and resulting in a very low morbidity rate. Few complications reported with GKRS are cysts with multiple septae developing medial to the tumor causing compression of the brain stem and fourth ventricle, thus necessitating post-GKRS surgery.

Multiple cranial neuropathies of the cerebellopontine angle after GKRS should raise suspicion of necrosis within the schwanna and urgent MRI should be considered.

Conflict of Interest
None declared.

References