Role of Susceptibility Weighted Imaging in Cerebellopontine Angle Schwannoma Vs Meningioma

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ABSTRACT

Introduction: Vestibular schwannomas may sometimes be difficult to differentiate from cerebellopontine angle (CPA) meningiomas. Identifying foci of blooming on Susceptibility Weighted Imaging (SWI) has the potential to differentiate Vestibular schwannomas from CPA meningiomas. Hence; we planned the present study to assess the role of susceptibility weighted imaging in cerebellopontine angle schwannomas vs meningiomas cases.

Material and methods: In this prospective study of 32 consecutive patients of CPA tumors (schwannomas and meningiomas), SWI sequence is done along with all other routine basic sequences in conventional MRI. For all cases histopathological diagnosis was compared after surgery.

Result: In our prospective study 32 patients were included, out of which 23 patients are diagnosed as Vestibular schwannoma and 9 patients as meningiomas. Microhemorrhage was noted in T2* sequence (SWI) as blooming foci in 22 cases out of 32 which gives a potential diagnosis of vestibular schwannomas and rest of 9 cases shows no blooming and they are diagnosed as meningiomas. All patients underwent surgery and Histopathological diagnosis was compared with our study. Statistical analysis shows sensitivity of 95.6%, specificity of 100%, positive predictive value of 100% and negative predictive value of 90% and diagnosing accuracy of 96.8% for T2* weighted SWI sequence for microhemorrhages in vestibular schwannomas.

Conclusion: MR imaging demonstrates microhemorrhages in most of VS in T2* -susceptibility weighted imaging. This helps to differentiate Vestibular schwannomas from CPA meningiomas. T2* - susceptibility weighted imaging should be used as a routine sequence to evaluate CPA tumors.

Key words: Vestibular Schwannomas, Susceptibility Weighted Imaging, Cerebellopontine Angle

INTRODUCTION

Cerebellopontine angle (CPA) tumors accounts for 5 –10% of all intracranial tumors in adults and 1% of all paediatric intracranial tumors. In which 80 to 90% constitute Vestibular schwannomas (VS) and 10 to 15% meningiomas.¹ Vestibular schwannomas may sometimes be difficult to be differentiated from cerebellopontine angle meningiomas. Magnetic resonance imaging is a vital tool to assess cerebellopontine angle tumors and to distinguish vestibular schwannomas from meningiomas. Demonstration of intratumoral microhemorrhages with T2* - susceptibility weighted imaging shows characteristic feature of identifying vestibular schwannomas from that of meningiomas.² In most cases, MR imaging and computed tomography (CT) display characteristic features of meningioma’s. Acoustic neuromas are routinely round masses in the cerebellopontine cistern that emerge from the internal auditory canal, widen the porus, and grow posteriorly be- cause of the anterior limit represented by the cisternal segment of the facial nerve. They can be heterogeneous due to cystic components. Conversely, meningiomas are usually hemispheric, semilunar masses with a broad petrous base to which they are attached and are usually asymmet- ric to the internal auditory canal.³ The aim of present study was to assess the role of susceptibility weighted imaging in cerebellopontine angle schwannomas vs meningiomas cases.

MATERIAL AND METHODS

In our prospective study from Jun 2015 to Dec 2017, conducted in the Department of Radiology in Chettinad hospital, we included 32 patients. After obtaining oral and written informed consent and ethical clearance, the patients were subjected to MR imaging with 1.5 Tesla SIGNA GE HDxt MRI scanner with an 8 channel NV (NAVIGATOR) radiofrequency coil. The bore size of the machine is 60cm. Image acquisition parameter are shown in Table 1.
Histopathologic confirmation of the diagnosis was done in all cases. Approximately 2 and half minutes was the acquisition time. We performed the post processing, followed by generation of thick minimum intensity projection (mIP) slabs. We, then, reviewed the images obtained on a DS3000 Impax workstation. For image analysis purpose, we used an image matrix of 374 512 and FOV of 220 mm 240 mm. The presence of intratumoral microhemorrhages was identified by hypointense dots on T2* MRI similar to that of cerebral microhemorrhages.²⁴

**STATISTICAL ANALYSIS**

All the results were analyzed by SPSS software version 17.0. Multivariate regression curve were used for assessment of level of significance.

**RESULTS**

In our prospective study 32 cases were included, out of which 23 patients were diagnosed as Vestibular schwannoma and 9 patients were diagnosed as meningiomas (fig.1). Out of 23 patients of vestibular schwannoma, 8 patients were women and 15 were men. In 9 cases of meningiomas 6 patient were women and 3 patients were men (fig.2). The age range of vestibular schwannomas are 21 to 55 years were as for meningiomas 40 to 60 years. According to the location of the CPA tumors in vestibular schwannomas 17 cases presented on right and 6 cases on left side whereas in meningiomas 6 cases on right side and 3 on left side (fig.3). Distribution of type of CPA tumor lesions, 22 cases were solid and 1 case is cystic in vestibular schwannomas whereas in meningiomas all the 9 cases were solid. Among 23 cases of vestibular schwannomas, IAC extension is noted in all cases. In T1 weighted imaging sequence tumor shows hypointense in 22 cases and isointense in 1 case. The matrix of tumor shows heterogeneously hyperintense on T2 weighted imaging in all the vestibular schwannoma cases. Intense enhancement is noted in 17 cases and moderate enhancement in 6 cases. Microhemorrhage were noted in T2* sequence (SWI) as blooming in 22 cases out of 32 which gives a potential

![Figure-1: PIE diagram showing distribution of sample](image1)

![Figure-2: PIE diagram showing sex distribution of patients](image2)

![Figure-3: Bar diagram showing distribution of sample based on location](image3)

<table>
<thead>
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<th>T1 SE</th>
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*Table-1: Conventional MR image acquisition parameters*
The first most common CPA (hearing loss, dysequilibrum, and tinnitus) are more common and account for about 6 to 10% of all intracranial tumors arising in or involving the cerebellopontine angle. CPA lesions are categorized into three major groups: 1) Lesions originating in CPA, 2) lesion primarily located in close anatomical sites and extending to the CPA and 3) Intraventricular and brain stem pathologies showing exophytic expansion to the CPA.

About 6 to 10% of all intracranial tumors arise in or involve the cerebellopontine angle. The term Cerebellopontine angle (CPA) was first described by Hennieberg and Koch in 1902, they reported two individuals with bilateral acoustic neuromas occurring in the location they described (in German) as kleinhirnbruchen-winkel (kleinhirn= cerebellum; bruchen= pons or bridge; and winkel= angle). Cerebellopontine angle (CPA) is lined or covered by the meningeal layers and added to that of CSF, nerves, vessels, and possibly embryologic remnants. CPA lesions can be arising from any of these structures. CPA lesions are categorized into three major groups: 1) Lesions originating in CPA, 2) lesion primarily located in close anatomical sites and extending to the CPA and 3) Intraventricular and brain stem pathologies showing exophytic expansion to the CPA. About 6 to 10% of all intracranial tumors arise in or involve the cerebellopontine angle. The first most common CPA tumors are vestibular schwannomas, which constitute 80 to 90%. The next most common CPA tumor is meningiomas. Clinically audiovestibular disturbance (hearing loss, dysequilibrium, and tinnitus) are more common symptom of vestibular schwannomas as compared with meningiomas. Hearing loss is found in 50 to 80% of patient with meningiomas as compared with almost all patient with vestibular schwannomas.

The globular shape of the tumor with dilatation of the internal auditory canal in association of canalicular component and formation of acute angle by anterolateral or posterolateral tumor border with the adjacent petrous bone is highly suggestive of vestibular schwannomas. Meningiomas are large, sessile and oval in shape and usually have a broad base against petrous bone. Hyperostosis and calcifications are more frequently seen in meningiomas and involvement of the internal auditory canal (IAC) is less frequent in meningiomas than vestibular schwannomas.

Vestibular schwannomas are isointense to hypointense to brain in T1W images and hyperintense on T2W images. They may be heterogenous due to cystic degeneration and haemorrhage. They are usually round or oval in shape and taper along the axis of the internal acoustic meatus like ‘ice cream cone appearance’. Meningiomas are usually isointense to brain in both T1 and T2 weighted images and with contrast, intense and homogeneous enhancement is seen. Even dural tail of enhancement may also be present. Mostly they do not show extension into the internal acoustic meatus and usually have a broad base against petrous bone.

In our study all patients underwent MRI brain are subjected to the T2* sequence (SWI) in which blooming are noted in 22 cases out of 32 cases which gives a potential diagnosis of vestibular schwannomas due to presence of microhemorrhages and rest of 9 cases shows no blooming and they are considered to be meningiomas. Hence Intratumoral microhemorrhages noted on T2* weighted SWI sequence holds a potential diagnosis for vestibular schwannomas and helps to differentiate from that of the meningiomas.
in cerebellopontine tumors. CT is done routinely to all preoperative patient to exclude calcification in all vestibular schwannomas cases. The treatment and operative approach varies for both vestibular schwannomas and meningiomas, so accurate preoperative diagnosis is evident to differentiate this two CPA tumors because the outcome with preservation of hearing is better in meningiomas. Similar studies have been done by other authors with T2* Weighted Gradient-Echo Imaging to differentiate vestibular schwannomas and meningiomas, but we have used susceptibility weighted imaging to differentiate these two lesions. Meningiomas are known to be richly vascular tumors and may occasionally present with intratumoral, subarachnoid, or subdural hemorrhage, particularly with angioblastic and meningotheliomatous meningiomas. Cystic changes in meningiomas have been associated with hemorrhage and aggressive behavior. Focal concentrations of thin-walled vessels and endothelial channels have been recognized close to the gross bleeding sites in meningiomas. Arteriolar hyalinization has been also recognized in meningiomas presenting with hemorrhage. Mishra A et al in 73 patients, conducted MRI with susceptibility weighted imaging sequence along with other routine used techniques. All the subjects were analyzed for the occurrence of foci of blooming inside the tumor on susceptibility weighted imaging. In all the cases, diagnosis was confirmed by assessment of histopathology reports. In the vestibular schwannoma group, there were overall of 59 patients, while in the meningioma group, there were overall of 14 patients. On assessing the Susceptibility weighted imaging results; we observed the presence of blooming (due to microhemorrhages) in 100 percent cases of cerebellopontine angle schwannomas. 13 cases showed no blooming within the lesion in the meningioma group, while single case demonstrated blooming which was established to be due to calcification on CT study. Under the light of above results, the authors concluded that in making accurate pre-operative differentiation of vestibular schwannoma from meningioma, susceptibility weighted imaging is a significant useful tool.

**CONCLUSION**

Conventional MR imaging demonstrates microhemorrhages in most of VS in T2* - susceptibility weighted imaging. This helps to differentiate Vestibular schwannomas from CPA meningiomas. T2* - susceptibility weighted imaging should be used as a routine sequence to evaluate CPA tumors.

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