Case Report

Otosclerosis associated with type B-1 inner ear malformation

Otosclerosi associata a malformazione dell’orecchio interno tipo B-1

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SUMMARY

Malformations of bony inner ear are rare anomalies occurring in approximately 20% of patients with congenital sensorineural hearing loss. Conductive hearing loss is usually associated with abnormalities of the external and middle ear. Recent reports of patients with lateral semicircular canal malformations indicate inner ear malformations to be associated with sensorineural or conductive hearing loss. Differential diagnosis of conductive hearing loss should include otosclerosis, isolated ossicular deformities, inner ear anomalies or a combination of these. In this report, a case is described with right vestibule-lateral semicircular canal dysplasia presenting at our centre with bilateral otosclerosis.

KEY WORDS: Inner ear malformation • Otosclerosis • Conductive hearing loss • Lateral semicircular canal dysplasia • Stapedotomy

INTRODUCTION

Malformations of bony inner ear are rare anomalies occurring in approximately 20% of patients with congenital sensorineural hearing loss (SNHL). Conductive hearing loss (CHL) is usually associated with abnormalities of the external and middle ear as both bear a common embryonic origin from the first and second branchial arches. Malformations of the inner ear result from an arrest of maturation during one of the stages of inner ear embryogenesis. Developmental malformations of the bony inner ear are best evaluated on computerized tomography (CT). Several Authors have classified the anomalies based on the radiological anatomy and embryogenesis of the inner ear. Recent reports of patients with lateral semicircular canal (LSC) malformations indicate inner ear malformations to be associated with SNHL and/or CHL. Other inner ear malformations, such as large vestibular aqueduct syndrome and X linked mixed deafness with perilymph gusher, may also present with SNHL or CHL.

Clinically, a normal otoscopic examination with conductive loss should include a differential diagnosis of otosclerosis, isolated ossicular deformities, inner ear anomalies or a combination of these. LSC dysplasia is the most common inner ear malformation. This may occur in isolation or be associated with middle ear, cochlear, or vestibular malformations.

In this report, the case is described of a patient with right vestibule-LSC dysplasia presenting at our centre with bilateral CHL.

Case report

A 39-year-old female presented with a 2-year history of progressively worsening right-sided hearing loss. There were no associated complaints of otorrhoea, tinnitus, ver-
tigo/imbalance, facial weakness or trauma. A positive family history of otosclerosis (father and brother) existed. Clinically otoscopic findings were normal. On audiometric assessment, Pure Tone Audiometry (PTA) showed a right CHL (air-bone gap 35 dB) and an initial air-bone gap of 10 dB for lower frequencies (250 Hz, 500 Hz), on left side. Speech discrimination score was 100% on left side and 80% on right side with bilaterally absent stapedial reflex (ipsi - contra). A provisional diagnosis of otosclerosis was made. However, a high resolution CT scan of temporal bone, performed 6 months prior to her visit at our Centre, incidentally showed a malformation of the right inner ear. Both axial and coronal sections, on the right side, showed the vestibule to be enlarged and forming a common cavity with a dysplastic LSC (Fig. 1). The superior vestibular nerve canal was also enlarged (Fig. 2). The superior and posterior semicircular canals, cochlea and the internal auditory canal were normal. There was no radiological evidence of any malformation on the left side. Consequently, the radiological findings on CT contraindicated any surgery for otosclerosis because of a severe risk of hearing loss or gusher.

**Discussion**

Malformations of the inner ear are present in about 20% of the cases with congenital SNHL. In the remaining 80%, the condition exists at a cellular level with membranous anomalies and normal bony structures of the inner ear. Usually, patients with CHL require appropriate investigations to exclude external or middle ear disease. However, recent studies have indicated that CHL is associated with inner ear anomalies. Johnson and Lalwani demonstrated that CHL is often present in LSC malformations. Karlberg et al. suggested Mondini-like cochlear dysplasia to be a cause of inner ear CHL. Superior semicircular canal dehiscence may present as CHL with or without vertigo and normal stapedial reflex. Air-bone gap has been reported in the enlarged vestibular aqueduct syndrome. Dysplasia of the inner ear may be inherited, sporadic or the result of chromosomal aberrations and results from an arrest of maturation during one of the stages of inner ear embryogenesis. Labyrinthine malformation presumably occurs due to an interruption in differentiation between the fifth and sixth weeks of foetal development. The LSC, which is the inner ear structure most frequently found to be malformed, may be particularly vulnerable due to its late stage of formation. Inner ear malformations may occur isolated or as a part of a syndrome. This variability in presentation makes diagnosis and treatment difficult.

This case report highlights an isolated unilateral malformation of the vestibule and LSC forming a common dysplastic cavity associated with CHL. This is in agreement with the classification of Jackler et al. of inner ear malformations, Vestibule-LSC dysplasia type B1. In type B1, the cochlea appears normal but the vestibule is enlarged with a short and dilated LSC. The remaining semicircular canals are normal. On audiometric assessment, these anomalies show an air-bone gap.

In the present case report, a family history of otosclerosis, normal otoscopic findings, air-bone gap on audiometry and bilaterally absent stapedial reflex, were typical of otosclerosis and thus stapedotomy was initially hypothesised as the primary treatment. However, as CT findings of the temporal bone showed malformation of the right inner ear, it was decided not to perform any surgical intervention as it carried a substantial risk of an intra-operative gusher and post-operative hearing loss. In inner ear anomalies, surgery for otosclerosis carries a severe risk of gusher as it is well recognized that congenital dysplasias provide a potential communication between the subarachnoid space and middle ear cavity. Gusher is a very rare phenomenon associated with stapes
fixation in children or otosclerosis in adults. This atypical anomaly, at times, presents as rhinorrhoea (cerebrospinal fluid) associated with recurrent meningitis or otorrhoea in the presence of a perforated tympanic membrane. However, it may often remain unobserved and asymptomatic for a long period. Thus the absence of definitive signs, prior to surgery for otosclerosis, makes it difficult to conclusively diagnose an impending gusher.

Intra-operatively, following stapedotomy, perilymphatic fluid gushes out under high pressure resulting in incomplete surgical treatment. Consequences are severe, as it exposes the patient to risk of recurrent meningitis and hearing loss.

**Conclusions**

This case report highlights the presence of bilateral otosclerosis with unilateral inner ear malformation. Increased conductive hearing loss, unilaterally, may reflect an enhanced otosclerotic process on the affected side or a combination of a hearing deficit due both to otosclerosis and inner ear malformation. It is possible that onset of otosclerosis, in the present case, led to worsening of an already existing hearing loss on the right side that had remained unnoticed because of good hearing on the other side. Radiological evidence of the inner ear malformation was revealed on a chanced CT, as, in the normal course of events, we do not advocate radiological evaluation for patients suspected of having otosclerosis. These patients are at high risk of gusher and/or hearing loss, if surgery is performed. We do not advocate surgery for these patients on account of the associated risks. Pre-operative CT of otosclerosis, is still debatable, in all patients, considering the rarity of inner ear malformations and the elevated cost of the procedure.

**References**


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