CASE REPORT

Bilateral posterior semicircular canal aplasia and atypical paroxysmal positional vertigo: a case report

Aplasia bilaterale del canale semicircolare posteriore e vertigine parossistica posizionale atipica: caso clinico

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SUMMARY

Isolated congenital malformations of semicircular canals are rare abnormalities. Most inner ear abnormalities occur in syndromes and are associated with hearing loss. Unilateral or bilateral single aplasia of one semicircular canal does not usually result in vertigo, but these become clinically important if there are clinical complaints of vertigo. Computed tomography imaging and high resolution magnetic resonance imaging may reveal inner ear abnormalities. The case is presented here of a 46-year-old male with a 10-year history of recurrent positional vertigo with strong onset when changing position to the left side. Magnetic resonance imaging of the inner ear showed a bilateral posterior semicircular canal aplasia as well as an enlarged vestibule on both sides. Dix-Hallpike positional manoeuvre revealed a positional nystagmus in the left head-hanging position of short duration and latency of a few seconds. When rising, vertigo occurred, but no nystagmus was visible. The fast phase of the nystagmus was mainly vertical down-beating with a slight torsional component to the uppermost ear. Although benign paroxysmal vertigo of the anterior canal was suspected, physical therapy was not effective using a modified liberatory manoeuvre. Brandt-Daroff therapy was effective permanently.

KEY WORDS: Vertigo • Semicircular canal • Bilateral aplasia • Diagnosis • Brandt-Daroff therapy

INTRODUCTION

Malformations of the semicircular canals are the most common congenital abnormalities of the inner ear. Depending on induction time during embryological development, they may involve simultaneous several structures such as the cochlea, vestibule, vestibular aqueduct or semicircular canals (SCC). Anterior semicircular canal (ASC) and posterior semicircular canal (PSCC) dysplasia are commonly associated with a dysplastic lateral semicircular canal and severe hearing loss. The horizontal semicircular canal (HSCC) is the last single structure to be formed during embryogenesis of the inner ear. Malformation can be found as an isolated anomaly. Absence of three semicircular canals has been reported in Goldenhaar syndrome and CHARGE association. Congenital isolated abnormalities may not become clinically significant if not associated with hearing loss or acute acquired vertigo symptoms. Computed tomography (CT)
and high resolution magnetic resonance imaging (HR-MRI) can identify peripheral vestibular lesions especially anomalies such as deformities of the bony labyrinth. T2 weighted turbo-spin-echo-sequences may show pathological changes of the membranous labyrinth. Detection of inner ear abnormalities is important in the diagnosis of vertigo symptoms. Furthermore, CT and MRI imaging are necessary for pre-operative planning of temporal bone surgery especially in preparation for cochlear implant surgery. However, there are still several vestibular diseases that cannot be visualized, such as vestibular neuritis, Meniere’s disease and benign paroxysmal positional vertigo (BPPV). BPPV of one PSCC is the most common. Involvement of the HSCC is rare. ASCC BPPV has been described, but opinions are controversial. Besides the typical forms, there are some idiopathic forms of paroxysmal positional vertigo that cannot be classified (atypical BPPV).

A case of bilateral posterior canal aplasia and enlarged vestibule on both sides, associated with an atypical positional nystagmus is described.

**Case report**

A 46-year-old male presented with a history of vertigo episodes for 10 years. Until the onset of vertigo no symptoms were present. There was no trauma and no history of inner ear diseases in the past. Clinical examination, including otomicroscopy, showed no pathological findings. Pure tone audiogram showed a bilateral symmetrical sensorineural hearing loss (high tone reduction) with a linear increase of hearing loss above 3,000 Hz.

There was no spontaneous nystagmus using Frenzel glasses. Since the patient reported dizziness when lying on his left side, vertigo and nystagmus were provoked using the Dix-Hallpike manoeuvre. Vertigo occurred after a latency of 3 seconds with head rotation, of approximately 45 degrees, to the left. A predominantly down-beating nystagmus with a slight torsional nystagmus component and a quick phase toward the uppermost ear was observed using Frenzel glasses and two dimensional video-oculography (Fig. 1). The duration of nystagmus and vertigo was approximately 20 seconds and showed a crescendo-decrescendo pattern. Repetition resulted in decreasing of symptoms.

In a sitting position, there was no positional nystagmus, but vertigo, that was reproducible on several days. Testing BPPV of the HSCC and in other positions of the head, no vertigo or nystagmus was observed. Caloric irrigation using water at 44 degrees (50 ml in 30 s) showed less excitation of the HSCC in comparison to the right side (Fig. 2) but...
was within the normal range. Testing the thermal stimulus with water at 20 °C when the patient was turned from the prone position in the face-down position after 40 s showed a change in direction of induced nystagmus. In conclusion, utriculus function was assessed as normal. Otolithic function (sacculus), also tested using vestibular evoked myogenic potentials (vEMG), off vertical axis rotation (oVAR) and eccentric rotation, static lateral tilting, showed no pathological findings. On account of the unusual nystagmus constellation and the fact that the liberatory manoeuvre for ASCC BPPV of the right side did not improve complaints, CT and MRI imaging were performed. Inner ear structures showed aplasia of both PSCC. No central nervous diseases were revealed at CT or MRI of the head. Brandt-Daroff exercises were performed successfully (Figs. 3, 4, 5a-d).

Discussion

Inner ear development is a complex process involving growth of labyrinth structures in three dimensions. The embryological origin of the inner ear is the so-called ear vesicle. This divides into a ventral part, composed of sacculus and ductus cochlearis and a dorsal component, the utriculus. Development of the inner ear begins after 6 weeks. Three SCC arise at the same time from three flattened protuberances of the utriculus. Further SCC development is a process of growth and fusion of the protuberances. Single malformation of the posterior semicircular canal has been described in several diseases. Higashi et al., in 1992, found in 26% of all cases, aplasia of the PSCC, as a characteristic symptom in Waardenburg syndrome. Irie et al., in 1990, reported on two cases of unilateral aplasia of PSCC in Waardenburg syndrome. They also described an association with enlarged vestibules. Furthermore, abnormalities of the SCC can be found in CHARGE association and trisomie 18. Among the SCC, HSCC malformations are the most frequent. PSCC aplasia seems to be comparatively rare. Probably, the SCC malformation is the result of an abnormal fusion during embryological development. Kiernan et al., in 2002, described a model for canal truncation in mutants. They demonstrated that SCC development is encoded in the proximal portion of chromosome 4 near the centromere.

To our knowledge, bilateral loss of PSCC has, until now, not been described in the present literature. Isolated uni- or bi-lateral SCC malformations may be clinically undetectable because of the complete compensation. Therefore, CT and MRI are indispensable diagnostic tools in some selected cases. Sennaroglu et al. recently presented a new classification for cochleo-vestibular malformations. Bilateral aplasia of the posterior SCC can be reassigned to the vestibular subgroup of this classification system with single semicircular malformations.

Since vertigo and nystagmus occurred strictly, in this case, at the left head hanging position and in the plane of the right anterior SCC during Dix-Hallpike manoeuvre, BPPV was a possible differential diagnosis in the absence of both PSCC. From the geometric orientation of the SCC, taking up the Dix-Hallpike position, either the right anterior and the left posterior SCC or the left anterior and the right posterior SCC are situated approximately in the same direction. Downbeat nystagmus, during Hallpike manoeuvre, with a slight counter-clockwise torsional component, in the present case, gave rise to the assumption of anterior canal BPPV as a differential diagnosis. The presence of anterior canal BPPV is rare. Some Authors doubt its existence. The anterior canal is normally the highest part of the inner ear.
ear. The otoconial debris in BPPV would naturally tend to fall into the posterior canal and, therefore, the PSCC BPPV is the most common variant of BPPV. In rare cases, the horizontal canal may be involved in BPPV.

Bertholon et al. described 50 cases with down beating nystagmus and assumed that some patients with idiopathic paroxysmal down beat nystagmus have paroxysmal positional vertigo with lithiasis of the anterior canal. The torsional component may be weak on account of the predominantly saggital orientation of the anterior canal and, clinically, may not be readily detected.

Rapoport and Sadeh also observed patients with an atypical paroxysmal positioning vertigo. They showed apogeotropic nystagmus when the Dix-Hallpike test was performed. Nystagmus was torsional and vertical with the vertical component beating downwards. The torsional component was beating away from the lowermost ear. The Authors assumed ASCC BPPV in these cases. The absence of central nervous system signs and symptoms as well as normal imaging suggested that this predominantly vertical nystagmus may be another variant of BPPV.

Typical forms of paroxysmal positional vertigo are the variants of BPPV (PSCC and HSCC BPPV) and the central paroxysmal positional vertigo (CPPV). The duration of CPPV is usually longer than BPPV and the direction of nystagmus does not correspond with the plane of the SCC.

On account of the absence of nystagmus in the sitting position during Halfpikke positioning and the fact that the laboratory manoeuvre for the right anterior SCC were not successful, in this case, other forms of paroxysmal positional vertigo seem to be likely. Some atypical forms, without any central nervous involvement, may, therefore, derive from peripheral vestibular origin but the mechanism is still not completely understood.

In summary, congenital bilateral aplasia of the semicircular canals may be clinically silent if there is no additional vertigo. Patients with atypical BPPV should be submitted to CT and MRI imaging.

References