New Imaging Findings of Incomplete Partition Type III Inner Ear Malformation and Literature Review

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ABSTRACT

SUMMARY: Incomplete partition type III, also referred to as X-linked deafness, is a rare genetic inner ear malformation. Its characteristic CT findings, including bulbous dilation of the internal auditory canal and absence of the modiolus with the interscalar septa present, have been well-recognized. In this series of 19 cases, we report the abnormalities of the vestibule and semicircular canals and provide a comprehensive description of their CT and MR imaging findings. The inner ear malformations in incomplete partition type III were bilateral and basically symmetric, with involvement of the internal auditory canal, nerve canals in the fundus, cochlea, vestibule, semicircular canals, vestibular aqueduct, otic capsule, round window, oval window, and stapes. An irregular vestibule with a cystic appearance is also a distinctive imaging feature, which could be seen in about 90% of our patients, with a cystic appearance of the semicircular canals present in nearly half of the cases.

ABBREVIATIONS: CN = cochlear nerve; DFNX2 = X-linked deafness type 2; IAC = internal auditory canal; IP-III = incomplete partition type III; SCC = semicircular canal; VA = vestibular aqueduct; VN = vestibular nerve

In 1971, Nance et al described X-linked mixed deafness with congenital fixation of the stapedial footplate and perilymphatic gusher. Based on the classification of sex-linked loci and genes implicated in nonsyndromic deafness, this type of hearing loss has been classified as X-linked deafness type 2 (DFNX2), and the responsible gene is POU class 3 homeobox 4 (POU3F4). Its distinctive CT features were first described by Phelps et al in 1991, and it was named incomplete partition type III (IP-III) by Sennaroglu et al in 2006.

The characteristic CT findings of the cochlea and internal auditory canal (IAC) in IP-III have been described in detail, including absence of the modiolus with interscalar septa present and a bulbous dilated IAC at the lateral end. However, there are fewer descriptions of abnormal vestibules and semicircular canals (SCC). By analyzing a case series of 19 patients with IP-III, we aimed to present the newly described imaging findings of the vestibular and SCC abnormalities in IP-III and provide a comprehensive description of its CT and MR imaging features with a literature review.

MATERIALS AND METHODS

This retrospective case series was performed with the approval of the institutional review board and exemption from informed consent. Imaging records of 2075 patients with inner ear malformation (based on the classification of Sennaroglu and Bajin) between August 2014 and December 2018 were reviewed, and 19 patients (18 males and 1 female, 6 months to 47 years of age) with the IP-III anomaly were finally included. Genetic analyses were not performed.

All 19 patients underwent CT examinations of the temporal bone, and 14 patients underwent MR imaging examinations. The CT scans were performed with a 16- or a 128-section multidetector CT scanner (Somatom Sensation 16 or Definition Edge; Siemens). Image acquisition and reconstruction parameters were as follows: helical acquisition, 120 kV, 240 effective mAs, 0.75 pitch, 0.75- or 0.6-mm thickness, 0.75- or 0.6-mm collimation, 220-mm FOV, and B70 reconstruction kernels. Multiplanar reformatted images were processed on a separate workstation (Carestream RIS 3.1; https://www.carestream.com/en/us/ris-software). The temporal bone MR images were obtained with a 3T MR imaging unit (Magnetom Verio; Siemens) using matched 12-channel phased array coils. The MR imaging protocol included an axial sampling perfection with application-optimized contrasts by using different flip angle evolution (SPACE; Siemens) scan plane (3D-SPACE).
sequence with 0.5-mm thickness, oblique sagittal 3D-CISS images perpendicular to IAC with 1-mm thickness, and routine an axial and coronal T1WI and T2WI. Two experienced neuroradiologists and 1 otologist evaluated these CT and MR imaging findings.

The cochlear dimensions, including basal turn length and lumen diameter and upper turn width and height, were measured according to the method of Purcell et al9 (On-line Figure). Fifteen cases without inner ear malformation were selected as a healthy control group. IP-III and healthy groups were compared using a t test.

RESULTS

The prevalence of IP-III among cases of total inner ear anomaly was calculated to be 0.9% (19/2075). The inner ear malformations in IP-III were bilateral and basically symmetric with involvement of the IAC, nerve canals in the IAC fundus, cochlea, vestibule, SCC, vestibular aqueduct (VA), otic capsule, oval and round windows, and stapes. The imaging findings are shown in the Table.

All 19 patients had characteristic cochlear and IAC appearances as follows: bulbous dilation of the lateral end of the IAC, absence of the bony modiolus with the interscalar septa present, hypoplasia at the cochlear base (or enlargement of the cochlear nerve [CN] canal), and thinning of the otic capsule (Fig 1A–B). The outline of the cochlea was similar to normal, but the dimensions of the cochlea were slightly smaller than normal (On-line Tables 1 and 2), including basal turn length (7.91-mm versus 8.66-mm, P = .000), basal turn lumen diameter (1.78-mm versus 1.93-mm, P = .004), and upper turn width (4.59-mm versus 5.87-mm, P = .000). However, the upper turn height showed no significant difference between the IP-III and healthy groups (3.45-mm versus 3.51-mm, P = .258). The CN and vestibular nerve (VN) were visible in all 14 patients who underwent MR imaging, and there was a hypointense spiral structure extending from the fundus of the IAC to the cochlea in 13 patients (Fig 1C–D).

In addition to the markedly widened CN canal, the superior VN canal was enlarged in 16 patients (Fig 2A), and the canals of the small branches of the superior VN from the ampulla of the lateral SCC and the utricle were also visible. The labyrinthine facial nerve canal was slightly enlarged in 11 patients (Fig 2A), and the singular nerve canal, in 14 ears of 8 patients (Fig 2B). The inferior CN canal showed no enlargement. On MR imaging, 14 patients with enlarged superior VN canals and 6 patients with enlarged labyrinthine facial nerve canals demonstrated fluid signal (Fig 2C). Twelve patients showed fluid signal in the singular nerve canal (Fig 2D), and 6 of them did not show enlargement of the singular nerve canal.

An irregularly shaped vestibule with small cystic bulges in the margin was observed in 17 patients (Fig 3). The small saccs were usually located in the superior margin, between the feet of the superior SCC. A multicystic appearance of SCC was also seen in 9 patients (Fig 4), with a widened lumen of the SCC in 1 patient. Enlargement of the VA at the part close to the vestibule was observed in 13 ears of 7 patients, with a cystic appearance present in 8 ears and a normal VA orifice (Fig 2A). No patient showed an enlarged endolymphatic sac on MR imaging.

The oval window was dysplastic in 12 patients, and the round window, in 11 patients, including 1 patient with bilateral oval and round window atresia (Fig 5). A thickened stapled footplate was observed in 5 patients, with malformed stapes in 1 patient.

DISCUSSION

X-linked nonsyndromic deafness has been estimated to contribute only 1%–
5% of genetic nonsyndromic deafness. DFNX2 is the most common form of X-linked nonsyndromic deafness, which accounts for approximately 50% of all families with X-linked nonsyndromic deafness. Most interestingly, DFNX2 is the only form of X-linked deafness to show anatomic anomalies on CT of the temporal bone. Sennaroglu et al considered these anomalies as a type between incomplete partition type I (cystic cochleovestibular malformation) and incomplete partition type II (classic Mondini deformity) and called it IP-III. The term IP-III could help radiologists better understand this rare and characteristic type of deformity in the comprehensive classification of inner ear malformations. Choi et al identified IP-III in 10 (4.8%) of 206 patients with inner ear abnormalities in their research. Sennaroglu and Bajin reported that IP-III constituted 2% of inner ear malformations in their database. In our case series, it only accounted for 0.9%. IP-III mainly occurs in males, but female carriers may have milder forms of some characteristic findings.

Phelps et al first reported the following characteristic CT findings: bulbous IACs, incomplete separation of the basal turn of the cochlea from the fundi of the IAC, and wide first and second parts of the intratemporal facial nerve canals. Talbot and Wilson later added absence of the bony modiolus, an abnormal VA, and a dilated singular nerve canal. Sennaroglu noted that the otic capsule around the membranous labyrinth was thinner than normal. Saylisoy et al observed an enlarged VA with a cystic appearance and middle ear anomalies, including dysplasia at the oval and/or round window and stapes abnormalities such as a thickened footplate and single crus. Gong et al reported the stapes abnormalities,
enlarged superior VN canal, and the cystic bulge in the superior vestibular margin. Anderson et al.\textsuperscript{15} described the cystic changes in the SCC and vestibule, and they also reported a potential association between IP-III and hypothalamic hamartoma.

This case series demonstrates that the entire bony labyrinth, otic capsule, and IAC are all involved in IP-III. A dilated IAC at the lateral end and absence of modiolus with intercalar septa present are the most typical imaging findings. The size of the cochlea is slightly smaller, especially in the middle turn width, but the upper turn height is within the normal range. Although the cochlear cavity appears to be empty on CT, a hypointense spiral structure can be observed on MR imaging. This structure probably represents the membranous labyrinth, which extends into the dilated IAC due to the incomplete separation of the cochlear base from the IAC, and correspondingly, the branches of the CN can also be observed in the IAC. The enlarged superior VN canal has some noteworthy details as well. Because the superior VN has several branches from the ampullae of superior and lateral SCC and the utricle, the enlarged bony canals of some of these small branches can also be seen in some cases. An irregular vestibule with a cystic appearance is also a very common imaging finding, which could be seen in about 90% of our cases, and a cystic appearance of the SCC was present in nearly half of our patients. A slightly enlarged VA with a cystic appearance can also be observed in some cases. These cystic bulges show the same signal as the inner ear fluid on MR imaging. Dysplasia of the oval and round windows is also common, and the stapes footplate is sometimes involved.

DFNX2 hearing loss is due to loss of function of the POU3F4 protein.\textsuperscript{2} The POU3F4 gene is expressed in early embryos in the otic capsule, which is involved in mesenchymal-mesenchymal signaling for the development of the inner ear.\textsuperscript{2} During the inner ear development of the mouse, the Brn4/Pou3f4 gene product is initially detected in the ventral aspect of the otic capsule and later throughout the otic capsule in the mesenchyme of both the cochlear and vestibular aspects.\textsuperscript{15} The subcellular localization of the Brn4/Pou3f4 gene product also changes during the differentiation of the otic capsule. It remains nuclear in those regions of the otic capsule that eventually give rise to the mature bony labyrinth. However, the subcellular localization of the product shifts from strictly nuclear to perinuclear in those regions of the otic capsule that will cavitate to form acellular regions in the temporal bone, such as the scala tympani, scala vestibuli, and the internal auditory meatus.\textsuperscript{15} Brn4/Pou3f4-deficient mice also show very similar anomalies to those in human DFNX2.\textsuperscript{16} Most of the phenotypic features of these mutant animals result from the reduction or thinning of the bony compartment of the inner ear, including malformations of the stapes, cochlea, IAC, and superior SCC.\textsuperscript{15}

The abnormal development of the otic capsule should be the key factor in these malformations in IP-III. It may lead to a thin otic capsule, absence of the bony modiolus, enlarged nerve canals in the IAC fundus, dysplastic oval and round windows, and an abnormal stapes footplate as well. Sennaroglu\textsuperscript{13} reported that the thin otic capsule may be formed by a thick endosteal layer, and probably the second and third layers are either absent or very thin instead of the usual 3 layers. It may not be possible to observe the normal endosteal layer or differentiate these 3 layers on CT with today’s level of radiologic precision. However, we still observed hypodense areas in the region of the fissula ante fenestram in the thin otic capsule in most of our young patients, which are prevalent among healthy children.\textsuperscript{17} This pericochlear hypodensity suggests that the thin otic capsule may have >1 layer. With respect to the cystic changes of the vestibule, SCC, and VA, we speculate that they might be caused by the abnormal development of the perilymphatic space.

The disadvantage of this case series is that we did not perform genetic analyses. However, the diagnosis of IP-III can be established with typical imaging features. Radiologists should be familiar to these imaging features, thereby preventing harmful interventions and providing proper genetic counseling.

CONCLUSIONS

IP-III is a rare genetic inner ear malformation with distinctive imaging features. It involves not only the cochlea and IAC, but also the whole otic capsule, including the vestibule and SCC.

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