Audiologic Features of Norrie Disease

Chris Halpin, PhD; Grace Owen, ScD; Gustavo A. Gutiérrez-Espeleta, PhD; Katherine Sims, MD; Heidi L. Rehm, PhD

Objectives: Norrie disease is an X-linked recessive disorder in which patients are born blind and develop sensory hearing loss in adolescence. The hearing loss associated with Norrie disease has been shown in a genetically altered knockout mouse to involve dysfunction of the stria vascularis; most other structures are preserved until the later stages of the disease. The objective of this study was to characterize the audiologic phenotype of Norrie disease for comparison with the pathophysiology of the disease.

Methods: The design combined two series of clinical audiologic evaluations, with special attention to speech intelligibility.

Results: The audiologic results for 12 affected individuals and 10 carriers show that patients with Norrie disease retain high speech intelligibility scores even when the threshold loss is severe.

Conclusions: The cochlear mechanism — failure of the stria vascularis — accounts for some of the higher values in the wide distribution of speech scores in cases with similar pure tone audiograms.

Key Words: audiometry, NDP gene, Norrie disease, speech intelligibility, stria vascularis.

INTRODUCTION

Norrie disease is an X-linked recessive disorder caused by point mutations in the NDP gene at Xp11.4. Affected individuals are born blind, with pseudogliomas of the eyes, and exhibit various degrees of mental and emotional dysfunction. The audiologic features of Norrie disease were described previously by Parving and Warburg. These authors found normal hearing at birth, with the onset of hearing loss in early adolescence and progression to anosmia late in life. They also found good word recognition scores (80% correct or better for monosyllables). Further study by this same group indicated a cochlear site of lesion. Molecular genetic studies have identified the Norrie disease gene (NDP), characterized the protein (norrin), and generated a knockout mouse. Rehm et al. used light and confocal microscopy to identify the specific sites of cochlear disease in the mouse model of Norrie disease. These authors showed gross developmental abnormalities of the stria vascularis in the knockout mice by 3 months of age compared with normal littermates. Most other structures in the cochlea appeared normal in affected mice at this age. Longitudinal analysis showed that the initial effects in the cochlea occur in the stria; neurons and hair cells are affected in the later stages of life. This article presents the audiologic results of two series of patients with Norrie disease.

METHODS

Two clinical investigations are presented. The first reflects a series of patients in the United States. After informed consent procedures were completed, 6 affected individuals (male), 2 carriers (mothers), and 4 unaffected family members (2 fathers and 2 female siblings) were tested. Diagnosis of Norrie disease had previously been made in these families and was confirmed by molecular analysis showing point mutations in the NDP gene. Most patients were evaluated by standard audiometry for pure tones. Speech intelligibility was evaluated with 50-item CID W22 word lists on compact disc. One affected adult, institutionalized for severe mental disability, was evaluated by threshold auditory brain stem response audiometry. One other affected adult, also severely mentally disabled, was evaluated behaviorally by a conditioning procedure. (For an overview of audiologic methods in genetic investigations, see the 1995 article by Halpin.)

The second investigation took place in Costa Rica. After giving informed consent, subjects were evalu-
<table>
<thead>
<tr>
<th>Age</th>
<th>Speech Level (db HL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>14</td>
<td>[Graph A]</td>
</tr>
<tr>
<td>40</td>
<td>[Graph B]</td>
</tr>
<tr>
<td>58</td>
<td>[Graph C]</td>
</tr>
</tbody>
</table>

Fig. 1. Output of Harvard Audiometer Operating System software. The Audiology case screen consists of standard pure tone audiogram (on right) and speech intelligibility space (on left). This space consists of speech signal intensity on horizontal axis from 0 to 110 dB hearing level, and percent correct on vertical axis. Clinical test results (CID W22) are plotted by level and score in this space such that horizontal tick on data point bars (red, right; blue, left) represent score, and extent of bars represents 95% critical difference from another score. Vertical axis is arc sine transformed such that critical difference bars will remain same size even though variance of speech scores changes, given range in which score is found. Ogive (“S”) curves represent performance function predicted by Articulation Index given right and left ear thresholds.

In Costa Rica, speech intelligibility was evaluated in adults with 50-item Spanish bisyllable lists delivered by compact disc. The lists used were selected...
because their published performance/intensity function had a practically identical slope and intercept to that of the CID W22 lists used in the United States. The expected values and variability of these lists are equivalent, and they are plotted interchangeably within the speech intelligibility space on the audiograms in Figs 1\textsuperscript{13-15} and 2.\textsuperscript{15}

RESULTS

Figure 1 shows 3 cases that illustrate the general progression of the hearing loss. Figure 1A reflects a 14-year-old individual with an intragenic deletion in exon 2. (Mutation identifications were performed in the laboratory of author K. Sims.) His documented hearing was normal up to 7 years of age. This audiogram shows the high-frequency emphasis and asymmetry often reported in the earlier stages. Speech intelligibility was also affected, but his scores did not fall below 78% correct. The sensory hearing loss in these two series (US and Costa Rica) progressed to a relatively stable state during the third and fourth decades of life, with features illustrated by Fig 1B. In this 40-year-old individual (mutation R109X), the threshold loss had become severe (approximately 70 dB hearing level), flat, and symmetric, with preservation of word recognition near 80% correct. Figure
Aging down. Although the right ear still exhibits the severe flat hearing loss with speech discrimination at 74%, the left ear shows a further threshold loss and an extreme decrease in speech intelligibility (18%).

Figure 2 shows 3 cases that reflect some findings of individual interest. The results in Fig 2A (38 years of age, the brother of patient in Fig 1B; R109X) should be noted for unusual interpretation in light of speech score analysis using the Articulation Index. Although the absolute scores (60% right and 52% left) could be considered poor, these scores are not significantly different from the predicted values for the Articulation Index calculation. (In Fig 2A, the variance bars touch the ogive curves.) This means that a normal listener would get a similar score with the same limited amount of audible speech. Therefore, this case, with its low score, does not detract from the general trend for very good speech intelligibility in the presence of severe cochlear disease.

Figure 2B shows the individual with the least hearing loss. At this age (44 years), he would be expected to exhibit a severe flat loss on both sides (as in Fig 1B). Although his sensory loss resembles a "4-kHz notch," this patient had no history of any exposure to gunfire or industrial noise. Interestingly, this patient has a mutation in the NDP gene (R121Q), which has been observed to be milder in terms of both the eye phenotype and the lack of hearing loss reported in one family. Further studies will be needed to investigate specific mutations that may be indicative of normal or near-normal hearing.

Finally, Fig 2C depicts the data of the mother of 2 affected individuals (Figs 1B and 2A), who is therefore an obligate carrier. This was the only 1 of the 10 carriers in these two series who had a severe, symmetric sensorineural hearing loss similar to that seen in Norrie disease. This is a very unusual and possibly coincidental finding, as hearing loss in carriers has not been seen in previous reports. It is also possible that the hearing loss is a manifestation of nonrandom X-inactivation (or chance segregation of such cells) in the ear tissue. Such a case involving eye disease in a female NDP gene mutation carrier has been described.

A summary of the combined clinical test results for all affected individuals can be found in Fig 3. These results show complete penetrance of sensory hearing loss in all affected individuals 14 years of age and older. (One US individual, shown in Fig 2B, is affected at frequencies not graphed.) The graph includes threshold data from 2 mentally disabled individuals. One of these data points (at 41 years of age) was interpolated from a 65 dB normal hearing level auditory brain stem response to a broadband click including 2 kHz within its bandwidth, and another (at 44 years of age) contributes a single 2-kHz threshold found in the sound field. The youngest individual (2 years of age) contributes a single 2-kHz value in the normal range, found with a conditioning procedure. None of these 3 individuals contribute to the speech data (Fig 3B). Although the progressive loss shows a high-frequency emphasis and notable asymmetry in the earlier stages, the progression seems to result in a symmetric, bilateral, flat sensory loss for patients above 35 years of age. This loss seems generally stable for 2 to 3 decades by both these data and clinical history. The speech intelligibility scores (Fig 3B) are fairly high (mean, 78%), given the severity of the hearing loss. These are also fairly stable up to 58 years of age.

**DISCUSSION**

The results of this investigation confirm and extend those reported by Parving and Warburg. In terms of threshold, the hearing losses are sensorineural, mild, asymmetric, and high-frequency in adolescence, and progress to a severe, symmetric, flat loss around 35 years of age. The previous audiometric and medical reports include several individuals around 25 years old who have been described.
of age and older than 50 years of age.\textsuperscript{3,21} This study provides a number of audiograms in the intervening age range, and these results indicate that the disease does not progress steadily, but generally achieves a steady state of severe loss from about 35 years of age to nearly 60 years of age. There is some evidence from this study, that of Parving and Warburg,\textsuperscript{3} and the reported temporal bone from a 77-year-old individual\textsuperscript{22} that suggests that a very severe degeneration follows at about 60 years of age. Whereas Warburg\textsuperscript{23} initially estimated the penetrance of sensorineural hearing loss at 33\%, several other reports showed loss in all tested individuals above the age of puberty.\textsuperscript{3,21,24} In this study as well, all men over 13 years of age showed clear sensorineural loss, indicating 100\% penetrance of this feature.

Norrie disease serves as a rare example in which audiologic speech intelligibility can be studied within a group whose cochlear pathophysiology arises from the same mechanism, and that mechanism is indicated by a supporting mouse model. Speech intelligibility can be a more informative challenge to a dysfunctional cochlea than the binary detection task of the pure tone audiogram, and therefore can be more informative about the underlying state of the organ of Corti.\textsuperscript{25} Although the audiograms of cochleas with different pathophysiologic mechanisms may be quite similar,\textsuperscript{26} the differences in clinical speech scores are more likely to be informative regarding different underlying disorders (and prognoses for hearing aids).\textsuperscript{27} This is especially true if at least one such mechanism allows for remarkably high scores, as is the case in Norrie disease. Clinicians routinely observe similar audiograms with much lower scores, and possibly the different mechanisms in such cases may be further quantified as similar models become available.

Some authors have suggested that all sensory hearing loss is the result of outer hair cell loss plus inner hair cell loss.\textsuperscript{28} Although not specifically stated, such an approach would suggest that speech intelligibility would be predictable from threshold audiogram values alone and that the range actually seen in the clinic could be attributed to poor test reliability. The results of this study clearly do not support this assertion, as they indicate the stria vascularis as at least one additional mechanism. With the clear possibility of complex interactions among the mechanisms resulting in both high and low scores, speech scores remain valuable in parsing individual cochlear dysfunction and should be measured and applied within clinical cases, rather than assumed to be predictable from pure tone threshold values.

In a more general sense, it is important to remember that “sensory hearing loss” is a symptom of a variety of cochlear dysfunctions and is not expected to give coherent results when different pathophysiologic mechanisms are grouped together. In this study, molecular genetics, pathophysiology, and speech audiometry provide insight into a specific cochlear mechanism that limits patient performance. Similar collaborations in the future may allow separate study of other specific cochlear disorders.

Acknowledgments: The authors thank Dr Agneta Parving for providing information regarding the speech testing from previous studies, and Dr Joseph Pillion for providing data on one of the affected individuals. David Corey, PhD, and colleagues also provided valuable advice and support.

REFERENCES


Para los efectos pertinentes, adjunto esta publicación.

Gustavo