Sensory–neural hearing loss in pseudoexfoliation syndrome

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Abstract

- AIM: To evaluate relationship between ocular pseudoexfoliation syndrome (PXF) and sensory-neural hearing loss (SNHL).
- METHODS: This prospective case-control study was designed on patients who referred to a general ophthalmic clinic at Imam Khomeini Medical Center, Urmia, Iran (March 2010 through November 2010). On routine ophthalmic examination, patients diagnosed with ocular PXF were referred to the ENT department and, selected cases (after evaluating inclusion and exclusion criteria) were referred to Audiometric Department. Pure tone hearing threshold level (HTL) was measured at 1, 2, 3 kHz for each ear and was compared with International Standard (ISO 7029) median age associated hearing loss at 1, 2, 3 kHz (AAHL).
- RESULTS: Overall 21 of 50 patients (42.0%) had a higher HTL than the ISO 7029 median AAHL at 1, 2 and 3kHz, which included 14 ears of 23 patients in the male group (30.4%) and 21 ears of 27 patients in the female group (38.8%). Approximately 12.0% of patients had glaucoma at the same time, however; no significant correlation was found in SNHL prevalence and severity between PXF patient and patients with simultaneous glaucoma. SNHL was more common in patients with ocular PXF compared to their age-sex matched controls (P<0.05).
- CONCLUSION: Most of patients with ocular PXF had SNHL compared to their age-sex matched controls, which could be due to PXF fibrils in the inner ear. These findings suggest PXF could be a systemic disease.
- KEYWORDS: pseudoexfoliation; sensory-neural hearing loss, glaucoma

INTRODUCTION

Pseudoexfoliation syndrome (PXF) is characterized by grey-white fibrillogranular extra cellular deposits in ocular structure such as anterior capsule, iris, anterior chamber angle, zonules, ciliary body, anterior vitreous face, and conjunctiva. It is an accidental finding most of the times, and seems to be unilateral 2/3 of cases [1]. In addition to intra ocular deposits, PXF material found in the heart, lung, liver and gallbladder and arteries, which cause systemic vascular disorders including systemic hypertension, abdominal aortic aneurysms, ischemic heart disease, Alzheimer's disease, retinal vascular disorders and age-related macular degeneration [2]. Also there are some reports about finding PXF fibrils in the basement membrane of extra ocular and orbital tissues, which are documented pathologically [1,4]. In immunologic studies, aggregates stained positively for elastin and human amyloid P protein [3]. In recent studies, there are evidences and traces of PXF fibrils in multiple organs (such as lung, heart, liver, gallbladder and etc) of autopsy tissues which suggest systemic process [4]. The organ of corti is a complex structure in the inner ear which contains hair cells that are located on the basilar membrane and are overlaid by the tectorial membrane. There are 2 types of hair cells in the cochlea: the inner hair cells and outer hair cells. One row of inner cells spirals up the cochlea near the central axis, while 3-4 rows of adjacent outer hair cells spiral up the cochlea further from the central axis. The tectorial and basilar membranes are connected centrally. Sound moves these structures differentially and produces a shear force that bends the steroielpia. The tectorial membrane covers the steroielpia. The conversion of the mechanical energy produced by a sound wave to electrical energy (resulting in hearing) requires deflection of steroielpia, induced by a shearing motion between the reticular lamina and the tectorial membrane. This shearing motion is produced by the basilar membrane moving as a result of the traveling sound wave [6-8]. Any dysfunction in this process will cause SNHL [9].
Sensory-neural hearing loss (SNHL) and presbycusis may be attributed to various etiologies such as toxic agents, acoustic trauma, or the aging process; however, the exact mechanism is unknown \cite{10, 11}. SNHL in PXF syndrome may be caused by deposits of its material in inner ear structures (organ of corti), it may cause slight alterations in fine vibrations induced by sound analogously \cite{2} and also inhibits the conversion of the vibration energy to bioelectric energy by depositing into tectorial and basilar membranes \cite{9}.

Glucoma is a condition characterized by raised intraocular pressure, typical glaucomatous optic nerve head damage and presence of glaucomatous visual field defects. PXF syndrome is relatively common but an easily overlooked cause of chronic open angle glaucoma. They are found at the same eye most of the time, but few studies demonstrated the relationship between PXF and SNHL \cite{1}. The goal of our study is to evaluate correlation between PXF and SNHL and to assess the possible systemic nature of this syndrome.

**MATERIALS AND METHODS**

**Subjects** Patients attending a general ophthalmic clinic in Imam Khomeini medical center (Urmia, Iran, March to November 2010) were interviewed and underwent a complete ophthalmologic examination including determination of best-corrected Snellen visual acuity, slit-lamp examination, Goldmann applanation tonometry, gonioscopy, and dilated ophthalmoscopy using a +90 diopter noncontact lens.

**Methods** The presence of pseudoexfoliation substance on the iris, lens capsule, angle, or corneal endothelium was observed and confirmed by one of the investigators. Patients were classified into groups according to their age and gender and matched controls were selected and screened for absence of pseudoexfoliation substance by the same investigator. Glaucoma was diagnosed based on presence of at least 2 of the 3 following criteria: (1) intraocular pressure >22 mmHg without anti-glaucoma medications; (2) typical glaucomatous damage of optic nerve’s head; (3) presence of glaucomatous visual field defects. Visual field defects were defined on the basis of Anderson’s criteria \cite{12}. In order to prevent any misinterpretation related to glaucoma diagnosis and terminology, only patients with at least one definite glaucomatous eye and those who were non-glaucomatous were enrolled in this study and glaucoma suspects (subjects with only one of the above diagnostic criteria) were not entered in the study.

Any subject with manifestations of pseudoexfoliation or with evidence of glaucoma in either eye was considered a case of PXF or glaucoma, respectively. Patients were excluded if there was a history of acute or chronic ear disease, head trauma, long-term exposure to heavy noise or gunfire, and intake of ototoxic agents such as gentamicin or streptomycin. All subjects were referred to an otolaryngologist who examined them and excluded cases with evidence of upper respiratory tract infection and external or middle ear abnormalities. One masked operator performed standard bilateral pure-tone Audiology using the same device for all subjects. Hearing thresholds were determined using pure-tone Audiology using air and bone conduction at 1-, 2-, and 3-kilohertz (kHz) frequencies, which are thought to be important for speech comprehension. The sum of these thresholds was compared with the ISO7029 standard \cite{11} which is the result of a meta-analysis of large community-based studies to determine the normal distribution of hearing thresholds at different frequencies in otologically normal white subjects. Hearing loss was diagnosed if HTLs 1, 2, and 3 were higher than the sum of corresponding normal median thresholds as defined by the ISO7029 standard. The level of hearing loss in one or both ears was compared between cases and controls; furthermore, average hearing thresholds at each frequency were compared between cases and controls.

According to ISO 7029 data, males aged (in terms of years old) from 18-30, 31-40, 41-50, 51-60, 61-70 and 71-80 had median AAHL1,2,3 of 0-5kHz, 5-10kHz, 10-25kHz, 25-40kHz, 40-60kHz and 60-85kHz, respectively. Similarly, females aged (in terms of years old) from 18-30, 31-40, 41-50, 51-60, 61-70 and 71-80 had median AAHL1,2,3 of 0kHz, 5-10kHz, 10-25kHz, 25-30kHz, 30-45kHz and 50-65kHz, respectively \cite{14}.

**RESULTS**

Overall 50 patients (23 males with mean age of 61 years, range 52-70 years and 27 females with mean age of 60.5 years, range 52-69 years) were included in the study group. 3 males and 3 females had unilateral glaucoma at the same time; 41 of them had bilateral PXF. After evaluating inclusion and exclusion criteria, cases of both genders were allocated into 5 year age strata (Table 1). Control group consisted of 50 patients (21 males with mean age of 59.2 years, range 51-69 years and 29 females with mean age of 61.5 years, range 53-69 years). Similar to the case group, they were classified into 5 year age strata and none of them were glaucomatous (Table 1).

In audiometric studies, mean hearing thresholds in PXF patients were obtained 40.6 db and 41.2 db, in the right and left ear, respectively. In patients with simultaneous glaucoma, mean hearing thresholds were 57.5 db and 60.8
db: decibel; 1 Sum of hearing thresholds at 1, 2, and 3 kHz.

Table 2 Mean hearing thresholds in study and control groups

<table>
<thead>
<tr>
<th>Frequency</th>
<th>Control group (db)</th>
<th>Study group (db)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Right ear</td>
<td>Left ear</td>
</tr>
<tr>
<td>1kHz</td>
<td>5.66</td>
<td>5.41</td>
</tr>
<tr>
<td>2kHz</td>
<td>8.65</td>
<td>8.89</td>
</tr>
<tr>
<td>3kHz</td>
<td>14.32</td>
<td>15.52</td>
</tr>
<tr>
<td>1 HTL 1,2,3 kHzs</td>
<td>28.58</td>
<td>30</td>
</tr>
</tbody>
</table>

Table 3 Number of patients in control and study group with and without hearing loss classified by their gender

<table>
<thead>
<tr>
<th>Groups</th>
<th>Control group</th>
<th>Study group</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Male</td>
<td>Female</td>
</tr>
<tr>
<td>With SNHL</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Without SNHL</td>
<td>19</td>
<td>25</td>
</tr>
<tr>
<td>Total</td>
<td>21</td>
<td>29</td>
</tr>
</tbody>
</table>

Table 4 SNHL in glaucomatous and non glaucomatous patients in study group

<table>
<thead>
<tr>
<th>Male</th>
<th>Female</th>
</tr>
</thead>
<tbody>
<tr>
<td>With SNHL</td>
<td>Without SNHL</td>
</tr>
<tr>
<td>PXF</td>
<td>8</td>
</tr>
<tr>
<td>PXF and Glaucoma</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>9</td>
</tr>
</tbody>
</table>

DISCUSSION

This study was conducted to evaluate correlation between PXF and sensorineural hearing loss and to find the possible systemic nature of this syndrome. The results of this study showed that sensorineural hearing loss is more common in study group with PXF than age and sex matched control group (P=0.001). This finding is based on comparison to ISO 7029 standard, which is consistent with recent studies and confirms the systemic nature of the PXF disease. In this study Severity and prevalence of SNHL were also studied between PXF patients and patients with concomitant PXF and Glaucoma. The results demonstrated that difference was not statistically significant (P=0.118) and (P=0.193) respectively (in both sides ears at HTL 1, 2, 3 kHzs).

In the study by Cahill et al. [1] sum of pure-tone hearing thresholds was measured at 1, 2 and 3 kHz (HTL 1,2,3) in each ear and compared with the ISO 7029 standard. Sixty-nine patients were studied. 101 ears (73.7%) had a higher HTL 1,2,3 than the ISO 7029 median AAHL 1,2,3. There was no significant difference between the proportion of ears with SNHL on the same side as eyes without PXF, with PXF but not glaucoma and with PXF and glaucoma, in either male or female groups. A large proportion of patients with PXF had sensorineural hearing loss in comparison to age-matched controls, regardless of whether or not there is associated glaucoma.

Similarly in our study there was no significant difference between the proportion of sensorineural hearing loss, in either male or female groups. SNHL prevalence was more common in patients with PXF and concomitant Glaucoma contrary to PXF patients but it was not statistically significant in either ear at HTL 1,2,3 kH(zs (P=0.193).

The significant difference between this study and previous studies is that the mean age of under study patients in this article were lower than previous studies [1,2,4,9,13,16]. We selected PXF patients at younger ages therefore it was possible for us to reduce the Presbycusis effect in patients. Also it is obvious that prevalence of SNHL in PXF patients is lower in our study in comparison to other articles.

In the study by Yazdani et al. [3] on 166 subjects which included 83 patients with pseudo exfoliation syndrome and 83 age and gender-matched controls, equal numbers of male and female subjects were allocated into each of the study groups. Prevalence of SNHL (in one or both ears) was
94.0% in case group and 69.5% in control group which demonstrated a significant difference.

Detorakis et al.\textsuperscript{[10]} evaluated the acoustic function in pseudoexfoliation syndrome and exfoliation glaucoma. Tympanometric peak values were significantly lower in study compared with control group (\(P=0.04\)). The reduced tympanometric peak values in cases imply impairment in the elastic properties of the middle ear in PXF syndrome. Finally it is concluded that SNHL is more common in frequencies important for speech in patients affected by ocular PXF than in single frequency in comparison to sex and age matched controls, which is irrelevant to simultaneous glaucoma. Similar to Detorakis's study, in our study also, hearing thresholds were evaluated at 1, 2 and 3 kHz; but results did not show a statistically significant difference between study and control ears at both sides (\(P=0.333\) for all frequencies).

Correlation and association of SNHL with PXF syndrome have been demonstrated in several studies. We also choose studying SNHL associated with PXF like, because it was easy and cheap way to evaluate the systemic character of this syndrome. More studies like biopsy and specific staining are needed to evaluate other organs such as ears in order to understand the necessity of additional tests in PXF patients. The findings of this study support presence of pseudoexfoliative material in ear tissues and imply that this disorder may truly a systemic condition with envolvement of multiple organs. More research on visceral complications of PXF is required to pronounce it as a systemic disease, and to prevent more complications.

REFERENCES

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