Unilateral Sensorineural Hearing Loss in the Southern Region of Saudi Arabia

Tawfiq Khurayzi*, Abdullah AlSkaini2 and Ibrahim Sumaily1
1Otolaryngology Department, King Fahd Central Hospital, Jizan, KSA
2Otolaryngology Department, Asir Central Hospital, Abha, KSA
Submission: May 08, 2018; Published: May 17, 2018
*Corresponding author: Tawfiq Khurayzi, Department of otolaryngology, King Fahd Central Hospital Jizan, KSA, Email: tawfiqak@gmail.com

Abstract
Background: Unilateral sensorineural hearing loss (USNHL) is important to be identified and diagnosed early. It is common and can be a symptom of a wide range of diseases. In our region, southern region of Saudi Arabia, the USNHL has not been studied before. In this study we report the incidence and common presentation and causes of this entity.

Method and Aim: A retrospective chart review of 209 patient with USNHL whom underwent full investigations in adult age group. We aim to find the risk factors, causes, common presentation, diagnostic work up and prognosis of USNHL in the southern region of Saudi Arabia and comparing the result with a literature review.

Result: Out of 209 patients, 183 met our criteria. Most of our sample had chronic mode hearing loss (74.8%). Tinnitus, vertigo, aural fullness and ear pain were the most common associated symptoms respectively. On audiological assessment: PTA at time of investigation, half of them at high frequency (54%) and severe hearing loss (31.1%). Speech discrimination was very poor in all patients with retrocochlear pathology. ABR done in all cases with suspected to have retrocochlear lesion with positive finding affected on wave V. All patients underwent magnetic resonance imaging and is was normal in most of them (75.4%). While in 3.8% there was retrocochlear lesion, (5%) there was cochlear lesion, and in (13.6%) there was central nervous system pathology. Common diagnoses found to be idiopathic SSNHL (14.7%), NIHL (13.3%), Meniere’s disease (14.7%), presbycusis (14.5%), central nerves system (11.4%), head trauma (8.2%), ototoxic medication (6.5%), infection (3.8%), cerebellopontine angle (2.7%), metabolic/autoimmune (1%) and congenital (0.5%).

Conclusion: All patients with USNHL need full audiological screening and early diagnosis. Regarding our study, we recommend MRI as part of their diagnostic workup. The choice of diagnostic laboratory tests should be directed by patient risk factors and exposures.

Keywords: Unilateral sensorineural hearing loss; Single side hearing loss; Sensorineural hearing loss

Introduction
Hearing disorders it can divide to sensorineural hearing loss (SNHL) and conductive hearing loss (CHL). SNHL is a common disorder that results from damage to the inner ear in over 95% of all cases. Therefore, retrocochlear hearing disorders are rare and cannot be differentiated from sensory losses by clinical symptoms alone. Unilateral hearing loss affect the quality of life in the affected patients. Conductive deafness has a readily determined etiology in most cases. In contrast, USNHL (USNHL) requires more refined and extensive investigation [1]. There are very few studies discussing the incidence of USNHL. SNHL, which accounts for 90% of the cases of Americans people those presenting with hearing loss and it can be caused by damage to the cochlea or the vestibule-cochlear nerve. The vast majority of patients with SNHL have bilateral hearing loss and on the other hand, approximately 60,000 new cases of USNHL occur annually in the United States 2. According to the World Health Organization (WHO), Hearing defects are the most frequent sensory deficit in human population, affecting more than 250 million people in the world with an average of 7 out of 100 of the population in Saudi Arabia and still need more and more searching about prevalence of USNHL. It is important to identify and diagnose it early. It occurs when the hearing in one ear is within normal hearing capacity and the other ear has a hearing loss. The normal audiometric function is defined as hearing thresholds that are not poorer than 20–25 dB hearing level (HL) for pure-tone audiometry (PTA).

The diseased ear can be range above that, above 25 dB, from mild to profound level hearing loss. Because a wide variety of pathologic processes may be responsible for the USNHL, numerous diagnostic tests are usually used during initial
evaluation, including pure-tone audiometry, acoustic reflex testing, imaging, serologic testing, and auditory brainstem response testing 1,2. USNHL can be a symptom of a wide range of diseases. Also, can be defined by site of disease, severity, audiometric configuration, and method of onset. The most frequent mechanism cause of USNHL or single-sided deafness (SSD) are sudden deafness; Meniere’s disease; cerebellopontine angle tumors / vestibular schwannoma (acoustic neuroma); neoplasms; demyelinating pathologies such as multiple sclerosis, vertebrobasilar arterial occlusion, stroke; acoustic trauma; head injury; perilymphatic fistula; ototoxic drugs; labyrinthitis; autoimmune disease as Cogan disease, Wegener’s granulomatosis, lupus, Takayasu arteritis, systemic sclerosis, and other rheumatological disorders and idiopathic SNHL 3,4,14.

**Aim:** Our aim is to find the risk factors and causes of USNHL, presenting symptoms, diagnostic work up and prognosis in southern region of Saudi Arabia and comparing the result with literatures.

**Method:** A retrospective chart review for last 10 years (2006–2016 G) in Asir Central hospital, a tertiary hospital in the southern region of Saudi Arabia. Medical record collected and reviewed for 209 patients with USNHL whom underwent full investigations at time of presentation and with closed audiological follow up in clinic for 1 years post first presentation of hearing loss in different adult age groups. We are excluded pediatric age group less than 12 years, patients with contralateral hearing loss, those with ipsilateral mixed and conductive hearing loss and who has history of ear surgery. Data analysis done with SPSS v.22.

**Result:** 183 patients met our criteria, 108 males (59%) and 75 females (41%). Onset and Duration; Most of them had gradual onset of hearing loss, (74.8%). while (25.2%) reported sudden onset. At the time of presentation, in 100 cases (54.6%) the hearing loss was prolonged for more than six months duration, 45 (24.6%) cases from 30 days to 6 months, and 38 cases (20.7%) presented within one month of the inset.

History of Risk factors; 12 cases (6.5%) of the 183 total patients had mechanical head trauma, 16 patients (8.7%) received ototoxic medications, and 16 patients (8.7 %) with positive criteria for exposed to risky noise. Only (35.5%) had chronic medical illness. The association symptoms were tinnitus in 135 cases (73.7 %), vertigo in 121 cases (66.1%), aural fullness in 36 cases (19.6%), pain in 28 cases (15.3%).

**Audiological assessment;** pure tone audiometry (PTA) at time of investigation and during audiological follow up, most of them, 99 cases (54%) at high frequency 4 kHz and above, 20 cases (11%) at low frequency less than 1 kHz, 18 cases (9.8%) at middle frequency 1kHz-2kHz and 46 cases (25%) at all frequency. On the other hand, 57 cases (31.1%) in severe level, 42 cases (23%) in moderate level, 33 cases (18%) in moderate-severe level, 33 cases (18%) at mild level, 13 cases (7.1%) at profound and 5 cases (2.8%) as dead ear. See Figure 1.

### Figure 1: Sex distribution on PTA.

<table>
<thead>
<tr>
<th></th>
<th>Mild</th>
<th>Moderate</th>
<th>Moderately-Sever</th>
<th>Sever</th>
<th>Profound</th>
<th>Dead ear</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>15 (8.1%)</td>
<td>24 (13.1%)</td>
<td>14 (7.6%)</td>
<td>45 (24.5%)</td>
<td>8 (4.37%)</td>
<td>2 (1%)</td>
<td>108 (59%)</td>
</tr>
<tr>
<td>Female</td>
<td>18 (9.8%)</td>
<td>18 (9.8%)</td>
<td>19 (10.4%)</td>
<td>12 (6.5%)</td>
<td>5 (2.7%)</td>
<td>3 (1.6%)</td>
<td>75 (41%)</td>
</tr>
<tr>
<td>Total</td>
<td>33 (18%)</td>
<td>42 (22.9%)</td>
<td>33 (18%)</td>
<td>57 (31.1%)</td>
<td>13 (7.1%)</td>
<td>5 (2.8%)</td>
<td>183 (100%)</td>
</tr>
</tbody>
</table>

Noise induced hearing loss notch at 4000 Hz found in 20 cases (11%) only. Speech discrimination which is ability to understand speech with a list of phonetically balanced words was done only in 141 patients as the following; normal in (10 % of 141), slight difficulty in (15.5% of 141), moderate difficulty (28.4% of 141), poor in (25.7% of 141), and very poor in (20.4% of 141). In addition, Speech discrimination was very poor result was in all patients with retrocochlear pathology. Auditory brainstem response (ABR) done in all cases suspected to have retrocochlear lesion and there were positive findings: 11 cases (6%) retrocochlear lesion and 32 cases (17.5%) was cochlear SNHL. MRI done for all of the patients and it was normal in 138 cases (75.4%), while in 9 cases (5%) it shows cochlear lesion, 7 cases (3.8 %) it shows retrocochlear lesion including cerebellopontine angle lesion 5 cases (2.7%). Also, 4 cases (2.2%) with middle ear pathology and (13.6%) have central nerves system pathology Figure 2.

### Figure 2: MRI finding.
We are categorized the Diagnosis as following: noise induced hearing loss in 25 cases (13.6%), Meniere’s disease in 26 cases (14.7%), presbycusis in 27 cases (14.5%), central nerves system pathology in 21 cases (11.4%), cochlear otosclerosis 15 cases (8.2%), idiopathic sudden sensorineural hearing loss (SSNHL) 27 cases (14.7%), head trauma 15 cases (8.2%), ototoxic medication SNHL 12 cases (6.5%), infection 7 cases (3.8%), cerebellopontine angle lesion 5 cases (2.7%), metabolic/autoimmune 2 cases (1%) and congenital 1 cases (0.5%) Figure 3.

**Figure 3:** Common causes of USNHL.

**Discussion**

Hearing loss is an extremely common disorder. Most of patients with SNHL have bilateral hearing loss and on the other hand, approximately 60,000 new cases of USNHL occur annually in the United States. According to the World Health Organization (WHO), Hearing defects are the most frequent sensory deficit in human populations, affecting more than 250 million people in the world with an average of 7 out of 100 of the population in Saudi Arabia. Few studies on USNHL are available in the literature and database. We are concern in our study on adult patients group. Also, in both sex, in our result the USNHL is more common in male than female, which is running in same configuration with the WHO Deafness and HL Fact Sheet No. 300 (2015). In our study, most of the patients presented with severe level hearing loss (31.1%) and (23%) in moderate level severity. Moreover, some patients can present with moderate-severe level (18%), mild level (18%) and (7.1%) at profound as in our study. When we comparing with other studies, common unilateral hearing loss ranged from mild (34.2%) to profound (31.96%) [16].

Unilateral hearing losses in children have traditionally been underappreciated, but health care practitioners are now beginning to understand their effect on development and the underlying pathophysiologic mechanisms. Average age of diagnosis unilateral SNHL is 8.7 years. 3/1000 Unilateral SNHL greater than 50dB, and Genetic cause in 50% of individuals 38. Universal infant screening has led to earlier diagnosis. On the other hand, congenital hearing loss may be delayed. In criteria of our study we are exclude all children. Moreover, we find 1 case (0.5%) diagnosed with unilateral congenital familial hearing loss.

In our study of 183 cases, 137 cases (74.8%) had chronic mode hearing loss and 46 cases (25.2%) in sudden mode of onset. 27 cases (14.7%) met the criteria of idiopathic sudden SNHL and we diagnosed him as idiopathic SSNHL. Divyank Bansal et al, he did similar as our study on 155 patients and he find (7.74%) had sudden onset and (92.26%) had gradual onset 16. In addition to that, another study published in 2015 on 820 patients with SNHL they find that 5% presented with sudden HL and 95% with gradual HL 18. But still there is variation in as in Hughes et al, in the United States; they find that only 1% of cases are of sudden hearing loss. Sudden SNHL (SSHL) have a strictest definition is a 30 dB loss over 3 contiguous frequencies occurring within 3 days and it is considered an otological emergency 19.

A number of factors have been identified that contribute to the presentation of unilateral sudden sensorineural hearing loss. Routinely, patients presenting undergo a battery of full investigation to determine a cause. The utility of the
Comprehensive evaluation should be reconsidered. The choice of diagnostic evaluation should be directed by patient risk factors and exposures. SSNHL is most commonly and differential for SSNHL is vast, for the majority of patients an etiologic factor is not identified.

In addition to above risk factor, ototoxic medications is considered strong risk, as we find, 16 cases (8.7%) received ototoxic medications before time of presentation with SNHL and 12 cases (6.5%) of those 16 cases developed SNHL within and early after receiving the course of this medication and we diagnose him as SNHL secondary to ototoxicity and all of them present with severe high-frequency SNHL. As we know most ototoxic medications cause bilateral damage and SNHL, but it can cause unilateral hearing loss also. Ototoxic drugs almost cause high-frequency SNHL. The most common drugs with ototoxic effects include aminoglycosides, loop diuretics, salicylates, and chemotherapy agents and the ototoxic effect increased in elderly patients and patients with poor renal function.

25 cases (13.6%) diagnosed as Noise-induced hearing loss (NIHL) and positive finding of Noise induced hearing loss notch 20 cases. Loud sound is risky to ear; this risk is increased by repeated exposure. Usually due to totally or partial damage to the organ of Corti and/or rupture of the cochlear membranes. NIHL it can be a temporary threshold shift followed by a permanent threshold shift after repeated exposure. In contrast, acoustic trauma is caused by a single high-intensity noise exposure that results in a permanent threshold shift without a temporary threshold shift and usually causes a sudden permanent unilateral or asymmetric. An audiogram of a patient with NIHL and acoustic trauma will show a typical noise notch, which is a hearing loss from 3000 to 6000 Hz, with the poorest hearing often centered at 4000Hz.

Meniere’s disease, known as idiopathic endolymphatic hydrops, presents with a combination of hearing loss, aural fullness, tinnitus, and/or vertigo. Meniere’s disease accounts for around 5% of SNHL cases. Comparing to 26 cases (14.7%) in our study and all of them collect the definite or probable criteria of meniere’s disease. The diagnosis is usually made by clinical history and audiological hearing tests as we did. However, electocochleography and cochlear hydrops masking procedure can be used to supplement the clinical history for diagnosis.

Autoimmune/metabolic can also affected on inner ear and cause SNHL bilaterally in most cases but still there are some patients present with unilateral SNHL as in our study: 2 cases (1%), one known case of systemic lupus erythematosus and other with severe rheumatoid arthritis. Systemic autoimmune disorders associated with SNHL include Cogan syndrome, polyarteritis nodosa, relapsing polychondritis, Wegener’s granulomatosis, systemic lupus erythematosus, Takayasu arteritis, systemic sclerosis, and other rheumatological disorders.

Cochlear otosclerosis can be classified as a mixed type or a sensorineural type according to the clinical appearance. Cochlear otosclerosis is defined as a focus of otosclerosis located in the otic capsule involving the cochlear endosperm and causing sensorineural hearing loss without any stapes fixation or any conductive component. If the definition of cochlear otosclerosis is accepted as the involvement of cochlear endosperm without associated stapes fixation, then the incidence among ears with pure progressive sensorineural hearing loss is about 1% 37. We collect only 15 (8.2%) cases meting criteria of cochlear otosclerosis.

Infection of inner ear still play important role in unilateral SNHL and almost patient present with vestibular symptoms. Labyrinthitis infection 7 cases (3.8%), as in our findings. Labyrinthitis can be suppurative or serous. Suppurative Infection can extend into the inner ear either by otogenic routes (acute or chronic otitis media) or meningogenic routes (the IAC or the cochlear/vestibular aqueduct. Infection also may cause bony erosion of the otic capsule. An infection causing meningitis can spread through the cochlear aqueduct or the IAC on the other hand; serous infection is defined as an abnormal process within the labyrinth caused by bacterial toxins, blood contamination of perilymph, or viruses 21. Common viruses related to unilateral hearing loss include measles, mumps, rubella, varicella zoster and cytomegalovirus. Other infections include syphilis, Rocky Mountain spotted fever, Lyme disease 24,25,26,27.

Presbyacusis is a complex and multifactorial disorder usually characterized by symmetrical progressive loss of hearing over many years and usually affects high frequencies of hearing. Also, can be asymmetry in PTA and speech levels, between the two ears. It is not uncommon to find the sensory loss more pronounced in one ear. We find (14.5%) 27 patients, all of them above 70 years, have mild to moderate high frequency hearing loss with good speech discrimination in 7 patients and 12 patients mild loss and 8 patients poor discrimination. all of them running with criteria of Presbyacusis after we exclude other causes. Diagnosis is frequently influenced by concurrent aging in other tissues and is frequently established by exclusion. Unilateral tinnitus or a gross asymmetry of response between the two sides frequently suggests pathology other than presbyacusis and warrants further investigation.

Central nervous system pathology can cause unilateral hearing loss. 21 cases (11.4%) of patients in our study carried with this diagnosis ether as stroke or demyelinating process according to MRI and neurologist assessment. Multiple sclerosis it common example of demyelination central pathology and can present almost with neurological symptom and between 4%-10% with SNHL 28,29. Hearing loss in multiple sclerosis patients usually present in various patterns, frequently, it can be sudden and unilateral hearing loss 31, 32.

An ABR and MRI play important role to rash to diagnosis 32, 33, 34, 35. Stroke (CVA) can affect cochlear and vestibular brain.
Loss can occur with cerebellopontine angle (CPA) tumors, but sensitivity and screening rate 12. Different pattern of hearing schwannoma screening protocols, the best compromise between by comparing published protocols. As regards vestibular other causes; and to assess proposed screening criteria other retro cochlear diseases. The pathologic correlates of many good screening test for cerebellopontine angle (CPA) tumors and reasonable cost-containment guidelines for the evaluation of health care resources challenges otolaryngologists to develop expensive, screening with audiometric brain stem response head is done on every patient, the cost is high. Although less tumor size 5. Magnetic resonance imaging is recognized as the gold standard for the evaluation of these problems, but if a complete examination of the internal auditory canals and head is done on every patient, the cost is high. Although less expensive, screening with audiometric brain stem response risks missing up to 33% of small tumors 14. The limitation of health care resources challenges otolaryngologists to develop reasonable cost-containment guidelines for the evaluation of patients with SNHL for the presence of retro cochlear pathology. On the other hand, auditory brainstem response (ABR) is a good screening test for cerebellopontine angle (CPA) tumors and other retro cochlear diseases. The pathologic correlates of many cases with positive ABR remain unknown 10.

To compare audiometric parameters in patients with vestibular schwannoma and in those with hearing loss from other causes; and to assess proposed screening criteria by comparing published protocols. As regards vestibular schwannoma screening protocols, the best compromise between sensitivity and screening rate 12. Different pattern of hearing loss can occur with cerebellopontine angle (CPA) tumors, but it typically causes high-frequency SNHL with word recognition is significantly reduced and is typically not in agreement with the pure-tone thresholds 20. In our study, 7 cases retrocochlear pathology on MRI report, 5 cases of them have cerebellopontine angle (CPA) tumors; two cases were Vestibular schwannoma, one case was arachnoid cyst and remaining two cases was without clear radiological diagnosis. 2 of those partially absent wave-V and 3 of those have delay in latency of wave-V with interaural delay of wave-V more than 0.2ms. Jack D. et al, he finds the False-negative responses are few and false-positive responses are common. Brain stem electric response audiometry BERAs is the most efficient audiometric test available today in our search for tumors affecting the auditory nerve (13).

Conclusion

After define a patient with USNHL (USNHL) or single-sided hearing loss need to identify the possible risk and cause of this hearing loss and to make early clear diagnosis and management plan. All patients with USNHL need full audiological screening. Regarding our study, we recommend MRI as part of their diagnostic workup to early rule out cerebellopontine angle lesion. The choice of diagnostic laboratory tests should be directed by patient risk factors and exposures. Few studies focused on adult unilateral SNHL and most of researches are advised further studies required in this field.

References


Global Journal of Otolaryngology

Your next submission with Juniper Publishers will reach you the below assets

- Quality Editorial service
- Swift Peer Review
- Reprints availability
- E-prints Service
- Manuscript Podcast for convenient understanding
- Global attainment for your research
- Manuscript accessibility in different formats (Pdf, E-pub, Full Text, Audio)
- Unceasing customer service

Track the below URL for one-step submission

https://juniperpublishers.com/online-submission.php

This work is licensed under Creative Commons Attribution 4.0 License
DOI: 10.19080/GJO.2018.15.555919