Update - Noise Induced Hearing Loss and the Military Environment

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SUMMARY: Although Noise Induced Hearing Loss (NIHL) is well recognised there are an increasing number of associations that are important in its study and to a lesser extent in clinical practice. For many there is controversy over whether there is a causal relationship or a confounding effect but of these presbyacusis is the most important. The PULHHEEMS assessment H3 is currently the first level of abnormal hearing. Between H2 and H3 there is a performance decrement and compensation is awarded at levels of NIHL less than H3. If the measurement of NIHL is to be used as a tool in the Army Hearing Conservation Program (AHCP) then the emphasis must be placed on the change from H1 to H2 and not from H2 to H3.

Introduction
NIHL may appear a spent subject but without a sound understanding of all its associations it is difficult to focus on the more meaningful issue of hearing conservation. In the Army NIHL is chiefly seen as being related to exposure to weapon noise and research tends to be in that context (1,2). There are however other sources of exposure in both military and civilian environments that potentially affect hearing loss and confound the study of NIHL. These may either cause hearing loss in their own right or modify the way the human auditory system reacts to noise. The last 10-15 years have seen advances in the study of NIHL and a degree of change in the perception of what constitutes significant loss. The AHCP has however remained essentially unchanged since its inception. This paper explores current issues in NIHL with particular regard for the practice and study of hearing conservation in the military.

Historical Perspective
In the nineteenth century Fosbroke noted that deafness in the military was caused by the explosions of cannons (3) and by the turn of the century standard Ear, Nose and Throat texts listed occupations that exposed workers to excessive noise and led to deafness; these included artillerymen, riflemen and sailors.

The First World War saw the first use of magazine and belt fed weapons using metal cartridges filled with smokeless-powder which greatly increased the energy and noise of the gases exiting the muzzle. Additionally high explosive replaced black-powder and there was a dramatic increase in the numbers with hearing damage (4). In the French Army the war-time prevalence of pensionable ear disabilities was 10-20% (5), the reporting author implying that almost all were due to NIHL. In England, however, the Oxford War Primers dismissed hearing loss as malingering and exaggeration and after the war Major T Jefferson Faulder RAMC reported hearing deficit from gunfire to be temporary.

In the Second World War there was a revival in military interest in NIHL as the operational consequences were recognised; Daggett stated “men with a high pitched loss were undoubtedly handicapped in night operations when the snap of a twig might be of significance”. Although by the end of the war all three Services were making some effort to protect hearing and it was not until 1974 that the AHCP was introduced against the background of the Health and Safety at Work Act of 1974.

Noise Induced Hearing Loss
The American Occupational Medicine Association Noise and Hearing Conservation Committee succinctly defined NIHL loss as a sensorineural loss, usually bilateral but rarely producing a profound deafness. The frequencies most commonly affected are 3-6 kHz with the greatest loss usually at 4 kHz; maximal levels of loss are reached in five to ten years (6). Unless there is a history of work related exposure this should not be labelled as occupational. Exposure to noise above 90 dB(A) L_{eq,d} causes damage in the unprotected ear and levels below 75-80 dB(A) L_{eq,d} do not cause damage (7,8). Between these levels there is debate over the size and cause (confounded by non-occupational exposure) (9) of the effect and although both UK and USA legislation are based on the upper limit they are said to protect only 85% of exposed workers (10).

Low level noise causes changes in the organelles and nuclei of the hair cells and an increase in the lysosome numbers leading to membrane damage, impaired regulation of ionic contents and ultimately cell loss. The mechanism of damage at higher levels of exposure (140-150 dB(A)) is mechanical. Shearing forces cause damage to the stereocilia of the hair cells leading to a loss of function. Damage also occurs at the synapses with the eighth cranial nerve. Other investigators have identified microvascular changes during noise exposure that induce ischaemia which may result in reduced auditory sensitivity (11). Although thought to be irreversible recent research has demonstrated regeneration of the sensory cells is possible at least in some animals (12).

Personal Factors
The most important confounding factor in the study of NIHL is presbyacusis. In contrast to NIHL the histopathological changes seen include degeneration of the stria vascularis and a loss of spiral ganglion cells and only
limited loss of hair cells. Presbyacusis has a similar pattern on audiometry to NIHL, although the characteristic 4 kHz notch is not seen, in practice however this is not useful for distinguishing the two conditions as the notch may not be present in NIHL. The fact that hearing diminishes with age or exposure to noise is well recognised, the relationship between the two is however much less clear. There are two distinct areas, the interaction between age related and noise related damage and the susceptibility of the ageing ear to NIHL. The different sites of damage mitigate against a simple additive interaction between noise and ageing. Rather the permanent damage caused to different structures, by both factors individually, leads to a cumulative interaction that does not reach the sum of the damage at the two sites (14). Finally there appears to be no clear association between previous hearing loss and future susceptibility to hearing loss, arguably those that have been affected are most sensitive to loss, but a simple alternative explanation is “the less there is to lose, the less will be lost” (15,16).

There is strong evidence that smokers have worse hearing than non-smokers (17). However different workers have interpreted this as smoking leading to an increased susceptibility to NIHL (18) or as a non-causal association with perhaps smoking itself causing the hearing loss (15).

Attempts to tie race and eye colour with melanin levels and NIHL have met with limited success. Blue eyes (relative lack of melanin) are weakly linked with NIHL (19) whereas black skin appears to have a protective effect. There is less hearing loss in a black population than white but before this difference is attributed to the occupational noise further research is required into differential ageing effects and socio-economic behaviour (20).

Gender has been linked to hearing levels and authors using ISO 1999 and ISO 7029 have necessarily assumed females to have less age related loss than males. This has been confirmed in a large longitudinal study in the USA (21) but there is controversy about how the sexes respond to noise. In the presence of noise males appear to fare worse than females but more rigorous analysis of the exposure history is required before this can be attributed to biological susceptibility.

Other factors that have been associated with NIHL include a raised cholesterol level, diabetes, high blood pressure and cardiovascular risk factors (22,23). A common thread running through these associations, including smoking, is cochlea blood supply, but research has so far failed to prove a conclusive link. In general with the exception of age these factors tend to be small when compared to the effects of noise but despite all the research there still appears to be considerable individual variation in the susceptibility of NIHL. Areas for future research include the protective effects of the acoustic reflex, toughening of the auditory system by exposure to noise and the role of the efferent auditory nervous system in mediating cochlea function (24).

**External Factors**

Ototoxic drugs can act synergistically with noise to produce hearing loss. However this has only been shown for aminoglycosides and cisplatin where a common mechanism of action exists. The evidence for the threshold shift seen with salicylate use interacting with that produced by noise is inconclusive and most likely does not exist (25). Unlike noise, chemical agents may cause damage to any part of the auditory system. Some common neurotoxic agents such as lead, mercury, toluene, styrene trichloroethylene, xylene and n-hexane are also ototoxic yet their interaction with noise is unclear but seems to be related to the site and mechanism of action (26). Of this group organic solvents have caused the most interest recently with toluene and styrene having been shown to cause damage to the hair cells (27). There may be a direct interaction with noise as in the case of toluene (28) but even if one does not exist for other solvents they clearly constitute a confounding factor in the study of hearing conservation.

Possible physical associations with NIHL include vibration, especially vibration white finger and the protective effect of a reduced body temperature (29). Exercise has been shown to increase hearing loss, possible as a result of increased core temperature. This may simply be due to the reduced efficiency of hearing protection that moves during physical activity thus breaking the seal (30).

**Noise Exposure**

Sources of noise exposure in the Army may be divided into continuous noise, similar to that found in industry, and impulse noise. Workshop and maintenance areas may house intense noise sources such as main battle tank engines which, when run up for servicing, produce 125 dB(A). Helicopter crews and passengers are exposed to cockpit noise of 102 dB(A) (Gazelle) whilst groundcrew who approach the aircraft are exposed to considerably more. Armoured fighting vehicles are a potent source of noise with their enclosed crew compartment having internal noise levels of 115 dB(A).

The most conspicuous cause of exposure to impulse noise in the military environment is small arms fire giving a higher peak level, shorter duration and lower repetition rate than industrial noise. Mechanical damage to the hair cells progresses quicker than metabolic damage and Table 1

<table>
<thead>
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<th>Table 1</th>
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<tr>
<td><strong>Typical military weapon noise (31,32)</strong></td>
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<tr>
<td>Weapon</td>
</tr>
<tr>
<td>--------</td>
</tr>
<tr>
<td>Thunderflash</td>
</tr>
<tr>
<td>84mm Anti-armour weapon</td>
</tr>
<tr>
<td>Medium mortar</td>
</tr>
<tr>
<td>Medium artillery</td>
</tr>
<tr>
<td>Service rifle at firer's ear</td>
</tr>
<tr>
<td>Tank gun inside closed down tank</td>
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<tr>
<td>Multiple launch rocket system crew position</td>
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reveals that most weaponry is in the category that causes this type of damage (31,32). Both industrial and weapon impulse noise are thought to produce greater hearing loss than continuous noise (33).

There is a clear association between shooting and hearing loss based on the asymmetry of the loss with right handed firers having greatest loss in the ear closest to the muzzle, the left (34). Even after matching for age, occupational exposure, salicylate consumption, blood pressure, cholesterol, smoking and vibration-induced white finger the effect still exists (35).

Impairment and Disability

Impairment of hearing is an abnormality of function of the auditory system usually assessed objectively by audiometry. Disability implies a state worse than impairment and in particular a difficulty in perceiving speech. The speech frequencies are 0.5-4 kHz with 1.5 kHz being the dividing line between the transmission of vowels and consonants. Difficulty in discriminating consonants is an early problem but it is only in more advanced disease that true disability occurs. Speech discrimination in noisy environments causes the most problems (33) and NIHL has been associated with poor word recognition due to a reduced ability to temporally resolve auditory information between gaps in the environmental noise (36). Other research has indicated the presence of a secondary distortion in the hearing impaired ear which makes speech perception worse than that implied by audiology alone (37). Mission success in the armoured battle has been directly linked to speech intelligibility (38).

In the American Army a striking difference in performance was found between soldiers with normal hearing level that fell between the British Army PULHEEMS grade H2 and H3 thresholds (39). The distances at which two military sounds can be heard is summarised in Table 2.

Table 2

<table>
<thead>
<tr>
<th>Sound</th>
<th>Normal Hearing</th>
<th>Impaired Hearing (H2-H3)</th>
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</thead>
<tbody>
<tr>
<td>Rifle bolt operating</td>
<td>1000m</td>
<td>210m</td>
</tr>
<tr>
<td>Approaching footsteps</td>
<td>100m</td>
<td>5.5m</td>
</tr>
<tr>
<td>Footsteps and temporary threshold shift</td>
<td>46m</td>
<td>0.5m</td>
</tr>
</tbody>
</table>

Statutory And Occupational Compensation

Although compensation is not paid by the Army it is useful to examine the levels at which other organisations award payment as a guide to the point beyond which meaningful loss exists. NIHL compensation schemes tend to be based around the frequencies of 1, 2 and 3 kHz, with some including adjacent frequencies. Table 3 gives the basic details of five schemes in use and one proposal for a replacement UK scheme. The average loss represents the point at which compensation is first paid and is commonly described as the "low fence".

Comparison with the AHCP reveals the lower limit for entry to the first band considered abnormal (H3) is less conservative than that of most schemes excepting the Industrial Injuries Compensation Scheme in the UK. A simple comparison can mislead as the frequencies used are different and in the Army hearing loss needs only to exceed the low or high frequency limit. A soldier could therefore have NIHL making them H3 and yet still have relatively intact speech frequencies. Even so the AHCP appears only to take account of hearing loss when it reaches levels that others would compensate and it cannot therefore function effectively. Arguably the system is intended as one documenting fitness for work and not compensation however communication needs must be represented when assessing fitness for work as well as compensation.

When the high frequencies are predominantly affected little day to day difficulty should occur. Even so in 1989 a court awarded compensation of between £2200 and £12500 for modest NIHL to 23 Ford Motor Company workers. Of these 10 equated to the H2 category with one "not troubled by hearing loss every day".

Trade Unions have negotiated no fault compensation deals with insurance companies and management that bypass statutory schemes. These rely only on documented exposure to 90dB(A) for one year and payment begins at levels as low as an average loss of 10dB, inside even the limit for H2 (40).

Conclusion

Noise as a cause of hearing loss has been recognised for at least two centuries but as research progresses an increasingly complex picture emerges (41). Although exposure levels exist they do not guarantee protection against either continuous or impulse noise. Race, eye colour, smoking and gender have been associated with NIHL but
not causally. Industrial chemicals, vibration and temperature may all play a part in mediating damage but perhaps the most rewarding lines for future research are those which link hearing loss with cochlea blood flow and the auditory system's own protective responses. Despite all these presbyacusis remains the main confounding factor, although the exact relationship has yet to be elucidated.

The value of good hearing to a soldier is apparent and yet investigation has shown a performance decrement even before the currently accepted limit for abnormality, H3, is reached. Compensation schemes, notably in the USA, make payments at levels similar to the threshold for entering the H3 bracket and UK civil courts at the H2 level. No fault schemes have taken this further and pay at levels around the H1-H2 border.

It is patently wrong that the action level in the AHCP should be set at a similar point to that which others use for compensation. In its current form the AHCP documents established hearing loss rather than providing a basis for monitoring and acting on early loss. It is felt, therefore, that greater emphasis should be placed on the change in hearing from H1 to H2 rather than from H2 to H3.

REFERENCES


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