Consensus practice parameter: audiological assessment and management of unilateral hearing loss in children

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Objective: Provide recommendations to audiologists for the management of children with unilateral hearing loss (UHL) and for needed research that can lend further insight into important unanswered questions.

Design: An international panel of experts on children with UHL was convened following a day and a half of presentations on the same. The evidence reviewed for this parameter was gathered through web-based literature searches specifically designed for academic and health care resources, recent systematic reviews of literature, and new research presented at the conference that underwent peer review for publication by the time of this writing.

Study sample: Expert opinions and electronic databases including Cumulative Index to Nursing and Allied Health Literature (CINAHL), Cochrane Library, Education Resources Information Centre (ERIC), Google Scholar, PsycINFO, PubMed, ScienceDirect, and Turning Research into Practice (TRIP) Database.

Results: The resulting practice parameter requires a personalised, family-centred process: (1) routine surveillance of speech-language, psychosocial, auditory, and academic or pre-academic development; (2) medical assessments for determination of aetiology of hearing loss; (3) assessment of hearing technologies; and (4) considerations for family-centred counselling.

Conclusions: This practice parameter provides guidance to clinical audiologists on individualising the management of children with UHL. In addition, the paper concludes with recommendations for research priorities.

Introduction

Unilateral hearing loss (UHL), once considered to be a nuisance and not taken seriously by hearing professionals, has been shown in recent decades to put children at risk for academic, speech and language, and social and/or behavioural deficits (e.g. Bess and Tharpe 1986; Lieu 2004; Lieu 2013). Despite increased understanding of these problems, there exists little evidence of effective interventions that ameliorate these deficits.

This practice parameter reviews the available empirical evidence, considers expert opinion, and provides specific recommendations for the management of children with UHL. The approach described requires a personalised, family-centred process: (1) routine developmental surveillance of speech-language, psychosocial, auditory, and academic or pre-academic development; and (2) assessments of hearing technologies specific to a variety of listening environments. In addition, although a number of important developments in the area of UHL have emerged over the past several decades, there remains a need for continued research. Recommendations are made for future research endeavours to enhance our understanding of and improve our management of children with UHL.

Much of the guidance for children with UHL applies to all children with hearing loss – those with unilateral or bilateral loss. Therefore, a brief summary of available published guidelines for assessment and management of children with hearing loss appears at the beginning of this document followed by more specific considerations for children with UHL and their families.

Definitions

a. Unilateral hearing loss (UHL) – any degree of permanent hearing loss in one ear (pure-tone average [0.5, 1.0, 2.0 kHz] > 15 dB for children), regardless of aetiology, with normal hearing in the opposite ear.

b. Paediatric population – for purposes of this guideline, refers to children birth through 18 years of age.

c. Contralateral routing of signal (CROS) hearing aid – a type of hearing aid that is intended for use by those with normal or near-normal hearing in one ear and an opposite side...
Table 1. Auditory-related subjective outcome evaluation tools.

<table>
<thead>
<tr>
<th>Measure</th>
<th>Purpose of instrument</th>
<th>Target population/ degree of HL</th>
<th>Respondent</th>
<th>Age range</th>
<th>Authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aural/oral performand of children PEACH</td>
<td>Evaluate effectiveness of amplification</td>
<td>Infants and children with mild to profound hearing loss</td>
<td>Parent interview</td>
<td>Preschool–7 years</td>
<td>Ching and Hill (2005)</td>
</tr>
<tr>
<td>Screenin Instrument for Targeting Educational Risk PRE SCHOOL SIFTER</td>
<td>Identify risk for educational delay</td>
<td>All</td>
<td>Teacher</td>
<td>3–5 years</td>
<td>Anderson and Matkin (1996)</td>
</tr>
<tr>
<td>屏蔽儿童的听力环境 Early Listening Function ELF</td>
<td>Evaluate functional use of hearing</td>
<td>Infants and toddlers with hearing impairment</td>
<td>Parent and audiologist</td>
<td>5 months–3 years</td>
<td>Anderson (2002)</td>
</tr>
<tr>
<td>Hearing Environments and Reflection on Quality of Life HEAR-QL-26</td>
<td>Measure condition-specific quality of life</td>
<td>All</td>
<td>Child</td>
<td>7–12 years</td>
<td>Umansky, Jeffe, and Lieu (2011)</td>
</tr>
<tr>
<td>Listening Inventory for Education LIFE-R</td>
<td>Identify challenging classroom listening situations</td>
<td>All</td>
<td>Child and teacher versions</td>
<td>6 + years</td>
<td>Anderson, Smaldino, and Spangler (2011)</td>
</tr>
<tr>
<td>Listening Situations Questionnaire LSQ</td>
<td>Identify benefit and satisfaction of amplification</td>
<td>All</td>
<td>Parent and child versions</td>
<td>7+ years</td>
<td>Grimshaw (1996)</td>
</tr>
<tr>
<td>PRS Educational Risk Targeting SIFTER</td>
<td>Identify risk for educational delay</td>
<td>All</td>
<td>Teacher</td>
<td>Children in grades 1–5 or 6</td>
<td>Anderson (1989)</td>
</tr>
<tr>
<td>Speech Spatial and Qualities of Hearing Questionnaire (SSQ) - Child</td>
<td>Investigate impact of hearing loss on a range of domains</td>
<td>All</td>
<td>Parent and Child versions</td>
<td>Child: children &gt; = 11 years</td>
<td>Galvin and Noble (2013)</td>
</tr>
<tr>
<td>- Child</td>
<td></td>
<td></td>
<td>Parents of children</td>
<td>&gt; = 5 years</td>
<td></td>
</tr>
</tbody>
</table>

un-aidable ear. Sound is transmitted from the side of the un-aidable ear to the ear with better hearing.

d. Bone-conduction devices – the class of bone-conduction devices that transmits vibration via transcutaneous or percutaneous means (surgical or non-surgical devices; transcranial).

e. Remote microphone system (RMS) – a wireless microphone system that converts audio signals into radio signals and transmits them to a receiver at the ear. Sounds can be transmitted via frequency modulation (FM) or digital modulation (DM).

f. Profound unilateral hearing loss – hearing loss in one ear with a pure tone average (PTA) of >90 dB HL.

g. Unaidable hearing loss – hearing loss that because of profound degree, very poor speech recognition, or intolerance for amplified sounds cannot be fitted with conventional hearing aids.

**Description of the process**

An international panel of experts on the screening, assessment, management, and monitoring of children with unilateral hearing loss and a parent advocate convened on October 24, 2017 following a day and a half of presentations on the same. Evidence reviewed for this parameter was gathered through literature searches using web search engines specifically designed for academic and health care resources and two recent systematic reviews of the literature were considered (Appachi et al. 2017; Anne, Lieu, and Cohen 2017). Eight electronic databases were searched using a series of keywords and expanded search terms related to children, unilateral hearing loss, cochlear implant, contralateral routeing of signal, bone conduction, BAHA, and single-sided deafness. Electronic databases included Cumulative Index to Nursing and Allied Health Literature (CINAHL), Cochrane Library, Education Resources Information Centre (ERIC), Google Scholar, PsycINFO, PubMed, ScienceDirect, and Turning Research into Practice (TRIP) Database.

**Basic principles of identification, assessment, and management of children with all degrees of hearing loss**

**Audiologic monitoring**

Numerous published guidelines by various national organisations have outlined recommendations for newborn hearing screening and assessment of hearing loss in children (e.g. Joint Committee on Infant Hearing [JCIH; AAP 2007]; American Academy of Audiology [AAA 2012]; American Speech-Language-Hearing Association [ASHA] 2004; American Academy of Paediatrics [AAP 2003]; Ministry of Children, Community and Social Services, 2018). This committee endorses the use of said guidelines in the provision of family-centered early intervention. Specifically, this committee supports the recommendation that hearing be screened by 1-month of age, hearing loss diagnosed by 3-months of age, and intervention be offered by 6-months of
age. These recommendations hold true for all forms of permanent hearing loss, including those that are unilateral.

**Developmental monitoring**

In addition to routine audiologic measures, a variety of tools are available to paediatric audiologists and other professionals charged with monitoring auditory, speech, language, and developmental milestones for children with permanent hearing loss, before and after the provision of various technologies. These tools are especially important when children with minimal degrees of loss are receiving limited or no intervention services from specialised providers (see Table 1). This committee endorses the use of these and other screening tools to assist families or teachers in identifying listening and/or speech or language concerns for children. Audiologists are encouraged to provide such tools to the families of children with UHL as deemed appropriate. Involving families and other professionals in the use of screening tools is expected to enhance communication among stakeholders, thus, leading to early suspicion, and perhaps intervention, of hearing-related problems.

The AAP recommends developmental and behavioural screening with a standardised developmental screening tool when a child is 9, 18, and 24 or 30 months of age. Although some of this screening will be done in the child’s medical home (AAP 2007), audiologists and other professionals working with children with hearing loss should be aware of available resources for developmental screening, ensure such screenings are ongoing and, if not, consider the use of such measures themselves to determine if additional referrals are needed. One such tool, ‘Birth to Five: Watch Me Thrive! A Compendium of Developmental Screening Measures,’ is produced through a coordinated effort by several U.S. agencies.

**Early intervention**

The JCIH (AAP 2007) states that families of infants with any degree of bilateral or unilateral permanent hearing loss should be considered eligible for early intervention services and that services for infants with confirmed hearing loss should be provided by professionals who have expertise in hearing loss. Furthermore, the JCIH Supplement (AAP 2013) states that children identified with hearing loss of any degree, including those with unilateral or mild hearing loss, receive appropriate monitoring, and follow-up intervention services when appropriate.

**Technology management**

For children with hearing loss, guidelines provide recommendations for hearing aids and other hearing technologies, including remote microphone systems (RMS; AAA 2011a, 2011b, 2013; Ontario Infant Hearing Program Audiologic Assessment Protocol 2008).

**Consensus-based principles of identification and audiologic assessment and monitoring of children with UHL**

**Identification**

The identification of UHL among children might be improved by monitoring children who present with risk factors for progressive or late-onset hearing loss, as recommended by JCIH (AAP 2007). The culmination of several reports demonstrate that the majority (>50%) of cases of UHL are due to cochlear malformations such as enlarged vestibular aqueduct syndrome (EVA) and Mondini dysplasia (Dodson et al. 2012; Friedman et al. 2013; Haffey, Fowler, and Anne 2013; Masuda, Usui, and Matsunaga 2013; Paul 2016; Fitzpatrick et al. 2017; van Beeck Calkoen et al. 2017). Environmental causes, such as cytomegalovirus (Paul 2016) and genetic causes are also implicated as common aetiologies of UHL (Dodson et al. 2012; Paul 2016; Fitzpatrick et al. 2017) although no specific genes for UHL have been identified to date. Additional risk factors for UHL include a stay in a neonatal intensive care unit (NICU), in utero infections, craniofacial anomalies, postnatal infections, and syndromes (Friedman et al. 2013; Fitzpatrick et al. 2017). Given the association between UHL and specific medical conditions, this committee supports etiologic assessment following confirmation of UHL. Specifically, audiologists should ensure that recommendations are made for complete otologic evaluation, including imaging, whether testing is completed within one’s own institution or requires referrals to outside care providers.

**Audiologic assessment and monitoring**

Recent estimates (2013–2014) show the prevalence of UHL among newborns to be approximately .6 to .7 per 1000 births in the U.S. (Centers for Disease Control Early Hearing Detection and Intervention [CDC] Database). Further evidence shows that UHL progresses to bilateral hearing loss in 7.5 to 11% of cases (Paul et al. 2017; Haffey, Fowler, and Anne 2013) and the number of children with UHL increases considerably after the newborn period. In fact, investigators have reported that approximately 2.5–3% of school-age children are reported to have UHL (Bess, Dodd-Murphy, and Parker 1998; Shargorodsky et al. 2010). This increase in prevalence underscores the need for ongoing vigilance throughout childhood.

The immediate consequence of UHL is loss of binaural function, which has a negative impact on localisation (Humes, Allen, and Bess 1980; Johnstone, Nabelek, and Roberston 2010) and speech perception in noise (Bess, Tharpe, and Gibler 1986; Russetta, Arjmand, and Pratt 2005). There is also evidence that suggests UHL negatively impacts balance (Wolter et al. 2016), early auditory behaviour and preverbal vocalisation (Kishon-Rabin et al. 2015), speech and language development (Ead et al. 2013; Lieu 2013), academic attainment (Bess and Tharpe 1986; Lieu 2004; Lieu 2013), and even cognition (Ead et al. 2013; Lieu 2013). Therefore, children with UHL will benefit from referral for comprehensive speech and language diagnostics, and academic monitoring. Measures that focus on speech-in-noise ability are recommended to assess the need for hearing assistive technologies (HAT) that enhance the signal-to-noise ratio (SNR).

A formal assessment of localisation is also needed to determine degree of difficulty. The ability to localise sounds on the horizontal plane is ideally measured in an anechoic chamber using a large number of speakers arranged in an array (Besing and Koenhke 1995; Johnstone, Nabelek, and Robertson 2010). This set up is not feasible for clinical use, though some clinics might choose to set up a speaker array with a smaller number of speakers. A more clinically feasible approach is to measure localisation using functional surveys or questionnaires. Localisation is a target skill of the Auditory Behaviour in Everyday Life (ABEL; Purdy et al. 2002) and a major domain of the Speech Spatial and Qualities of Hearing Questionnaire (SSQ parent and child...
versions; Galvin and Noble 2013). See Table 1 for these and other functional assessment tools.

Consensus-based principles of medical management of children with UHL

Medical considerations, although not the direct responsibility of paediatric audiologists, can impact the audiologic management of a child with UHL. Recommendations for the medical evaluation of children with hearing loss vary widely. However, we can draw upon international and interdisciplinary statements of recommendations that are relevant to children with UHL (e.g. International Paediatric Otorhinolaryngology Group [IPOG; Liming et al. 2016]; AAp 2007). These reports include several recommendations that are relevant to children with UHL, including temporal bone imaging (specifically with magnetic resonance imaging [MRI]); genetic testing (in patients with known or suspected syndromic hearing loss or in children with unilateral or bilateral Auditory Neuropathy Spectrum Disorder [ANSN] when imaging for cochlear dysplasia is negative); ophthalmology (because of the increased risk of ocular abnormalities in nonsyndromic sensorineural hearing loss [SNHL]); and testing for cytomegalovirus (CMV), which causes 20–25% of all congenital sensory hearing loss and is a frequent cause of progressive hearing loss (Fowler 2013). A specific link between CMV and UHL has not been determined at the time of this writing. However, with the nascent introduction of CMV screenings that are being initiated in some newborn nurseries, these data might soon be available.

Consensus-based principles of technology management for children with UHL

Conventional hearing aids

For permanent hearing loss on the affected side, conventional air-conduction hearing aid fitting is recommended as a first-line treatment if the affected side is moderate to severe in degree, regardless of child’s age. If the affected side is minimal to mild in degree, it is recommended that audiologists refer to the Clinical Decision Support Guideline (Bagatto and Tharpe 2014) for guidance. If the affected side is profound in degree, the ear might be unaidable and an RMS is recommended as first-line treatment for certain listening situations. Currently, there are no adjustments for prescriptive targets for paediatric unilateral hearing aid fittings. When appropriate, speech perception abilities, speech-in-noise testing, and spatial hearing assessments can be conducted to determine benefit with a device. Also, note that for some children, detection of sound – not just speech perception – might be a desirable outcome of an aided ear for safety purposes.

Bone-conduction devices

For infants and young children with unilateral microtia or atresia and for whom conventional air-conduction hearing aids cannot be fitted, a bone-conduction device should be considered. Children under the age of 5 years are not eligible for surgical bone-conduction devices in most countries and, therefore, can be offered devices on a soft headband. Surgical repair of the affected side often does not occur until age 7 or 8 years of age, so a non-surgical bone conduction device can be considered to support access to sounds to the affected side.

Remote microphone systems (RMS)

RMS provides a means of overcoming the deleterious effects of noise, distance, and reverberation by placing a microphone/transmitter close to the mouth of a talker or connecting the transmitter to audio sources such as televisions or computers (e.g. Wolfe, Lewis, and Eiten 2016). Remote receiver options include

- small receivers coupled to the listener’s ear(s) in a variety of ways (personal ear level RMS);
- one or more loudspeakers placed at strategic locations in a room (classroom audio distribution systems, aka sound field RMS);
- a small loudspeaker placed on a tabletop close to the listener (personal desktop RMS).

Guidelines for eligibility, implementation, and validation of RMS are available (AAA 2011a, 2011b). According to these guidelines, children are considered potential candidates for RMS when they demonstrate documented evidence of hearing, listening, and/or learning problems. Children with UHL exhibit poorer speech perception in noise and reverberation than peers with normal hearing (Bess, Tharpe, and Gliber 1986; Bovo et al. 1988; Hartvig Jensen, Johansen, and Borre 1989; Ruscetta, Arjmand, and Pratt 2005), thus, are candidates for RMS. However, there is limited research examining the efficacy of ear level RMS specifically for children with UHL and no research that has examined the benefits of desktop or classroom audio distribution systems for these children. In two studies conducted in the 1990s (Kenworthy, Klee, and Tharpe 1990; Updike 1994), children with UHL demonstrated better speech recognition with personal RMS than with either CROS or conventional hearing aids. However, the current applicability of those studies is limited by the small number of subjects as well as more recent advances in all classes of amplification technologies. At present, determining the efficacy of RMS for children with UHL relies on evidence of benefit from cohorts that include children with bilateral hearing loss and children with normal hearing who have special listening needs (Moeller et al. 1996; Pittman et al. 1999; Tharpe, Ricketts, and Sladen 2004; Anderson and Goldstein 2004; Iglehart 2004; Johnston et al. 2009; Wolfe et al. 2013; Mulla and McCracken 2014).

Decisions regarding RMS use for a child with UHL should be made on an individual basis. Considerations include the child’s age, the degree and configuration of hearing loss in the poorer hearing ear, whether the child uses personal amplification, and where the RMS will be used (National Workshop on Mild and Unilateral Hearing Loss: Workshop Proceedings 2005; AAA 2013; McKay, Gravel, and Tharpe 2008). When deciding whether to fit a receiver to the poorer hearing ear, it is important to consider aided speech discrimination ability; if significantly poorer than in the normally hearing ear, it is possible that RMS input to the personal amplification device might not improve outcomes.

Contralateral routing of signal (CROS)

Currently, CROS systems are typically used when no benefit is expected from fitting amplification to the ear with hearing loss. There is limited information about either the efficacy of CROS systems for children or the relative merits of the various options available. Results in children with simulated UHL suggest CROS systems can improve sentence recognition and story comprehension performance in noisy classroom situations, relative to unaided and a remote microphone condition. Benefits of the CROS system over RMS were most apparent for signals originating from the side of a
child when the remote microphone stays with a talker located in front of the child (Picou, Lewis, Angley, and Tharpe, in press).

Contralateral routing of signal can be achieved using
- a satellite microphone on the ear with hearing loss, coupled to an air-conduction hearing aid worn on the normal hearing ear (‘conventional’ CROS) or
- bone conduction to transmit stimuli detected on the side with the UHL to the cochlea of the normally hearing ear. Transcranial CROS fittings use a high-powered air-conduction hearing aid fitted to the ear with hearing loss, such that the amplified signal is transmitted via bone conduction to stimulate the normally hearing cochlea.

Transcranial CROS aids can use either a behind-the-ear aid or custom aid (in the ear or in the canal), all of which require a tightly fitted earmold that sits in the bony portion of the ear canal for optimal sound transmission (Valente et al. 1995; Hayes and Chen 1998; Hol et al. 2010).

Bone-conduction devices can also be used to transmit sounds from the ear with the UHL to the ear with normal hearing. They can be worn on a headband or coupled to a surgically implanted component, but limited data are available on the use of implantable technology in children (Christensen, Richter and Dornhoffer 2010).

Considerations when considering CROS fittings include the following.

**Inability to localise sound**
CROS fittings are unlikely to improve localisation because they do not facilitate the use of binaural hearing.

**Inability to hear speech on the side with the UHL**
CROS fittings improve detection of speech on the side of the UHL under quiet listening situations, but also enable the presentation of interfering noise to the normal hearing ear when the noise would previously have been attenuated due to head shadow effects in the unaided condition. Therefore, the CROS aid might have an adverse effect in complex listening situations if children are unable to control their device or listening environment, and, for this reason, CROS aids have not previously been recommended for young children (McKay, Gravel, and Tharpe 2008).

**Unaidable hearing due to poor speech discrimination**
If a CROS system is being considered because of poor speech perception in the ear with UHL, a conventional CROS aid is recommended to avoid adverse effects of stimulating that ear via bone conduction. When fitting a conventional CROS system, occlusion of the normal hearing ear by an earmold should be avoided to prevent reduced benefit from natural hearing.

**Cochlear implantation**
Some children with severe-to-profound UHL and their families might seek hearing restoration of the affected side via cochlear implantation (CI) with the ultimate goal of achieving some level of binaural hearing. Though limited in number, preliminary studies of CI use in children with UHL are encouraging. Investigators have reported that older children with post-lingual onset of UHL wear their devices consistently, and demonstrate that a normal hearing contralateral ear does not prevent device use (Távora-Vieira and Rajan 2015; Polonenko, Papsin, and Gordon 2017). However, it is worth noting that there are also examples of children with congenital hearing loss and longer periods of auditory deprivation who choose to discontinue device use after implantation (Távora-Vieira and Rajan 2015; Sladen et al. 2017a; Thomas et al. 2017).

Studies of adults and children with post-lingual onset of UHL show that patients demonstrate improved word and sentence recognition on the affected side (Friedman et al. 2016; Sladen et al. 2017a, 2017b; Finke et al. 2017), improved localisation on the horizontal plane (Firszt et al. 2012; Arndt et al. 2015; Dorman et al. 2015; Rahne and Plontke 2016), and modest improvements in overall speech recognition in noise (Mertens et al. 2015; Friedman et al. 2016; Rahne and Plontke 2016; Sladen et al. 2017b; Finke et al. 2017). Studies have also used disease-specific measures to demonstrate improvements in HRQoL (Härkänen et al. 2015; Sladen et al. 2017b; Thomas et al. 2017).

See Table 2 for a listing of hearing technologies available for children with UHL.

The Committee endorses consideration of these technologies in the context of the child’s and family’s needs and desires. Figure 1 provides a recommended clinical care plan for aiding in management decisions by the audiologist.

**Consensus-based principles of information counselling for families**
Despite technological progress and enhanced professional understanding of UHL in children and its implications, there remain numerous challenges to effective communication between audiologists and families of these children. This is in large part the result of a lack of management standards for UHL, and in part reflective of the difficulty audiologists face in talking with families about the variety of impacts UHL might have on their children. Services need to be family focussed, allowing for modifications based on unique family-identified concerns, priorities, goals, and desires (Moeller et al. 2013). Reactions of families to a hearing loss diagnosis are not necessarily related to the degree of the loss. Parents of children who have UHL can be just as concerned for their child’s future as are parents of a child with a severe or profound bilateral hearing loss. Moreover, families of these children are faced with a variety of intervention options that range from watchful waiting to classroom systems, and surgical as well as non-surgical technology options.

The following are consensus-based considerations on the essential components of effective communication to be included in counselling families of children with UHL:
- What are the possible consequences of their child’s hearing loss? Are all children affected the same way?
- What are the potential benefits, limitations, and risks of the technology for helping their child’s hearing and communication? What do we know and what do we not know?
- Are there other health concerns impacting the child with UHL that will influence success with technology?
- What are the consequences of proceeding or not proceeding with a particular technological intervention?
- If others will also have to engage with the technology (e.g. an RMS), what support is available to help with this?
- If the UHL is known to be progressive, consider how the various technology options are applicable if the hearing changes.
Table 2. Hearing technologies for children with unilateral hearing loss.

<table>
<thead>
<tr>
<th>Device</th>
<th>Benefit</th>
<th>Disadvantage</th>
<th>Considerations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conventional</td>
<td>Improved detection of sound on the affected side for mild to severe UHL</td>
<td>Might not provide benefit for profound SNHL. Not clear whether prescriptive</td>
<td>Environmental sound awareness might be the goal. Speech perception scores alone</td>
</tr>
<tr>
<td></td>
<td></td>
<td>targets for binaural paediatric fittings require adjustments for UHL fittings</td>
<td>should not be used to determine whether a hearing aid recommendation should be</td>
</tr>
<tr>
<td>Bone-Conduction</td>
<td>Suitable for unilateral microia/ataresia.</td>
<td>No fitting protocols for these devices. Sound quality of bone-conducted</td>
<td>Surgical BCD is not available for children under the age of 5 years in many</td>
</tr>
<tr>
<td></td>
<td>For profound UHL, the device can be considered in lieu of conventional</td>
<td>signal for profound UHL is inferior to routed signal in a CROS system</td>
<td>jurisdictions</td>
</tr>
<tr>
<td></td>
<td>CROS device</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CROS</td>
<td>Improved detection of speech on side with UHL in quiet.</td>
<td>Reduced speech understanding when noise is the dominant signal on the side</td>
<td>Requires ability to manage device and listening environment – especially for</td>
</tr>
<tr>
<td></td>
<td>Improved speech understanding in noise when speech is the dominant</td>
<td>with hearing loss.</td>
<td>young children</td>
</tr>
<tr>
<td></td>
<td>signal on side with hearing loss</td>
<td>Unlikely to help localisation</td>
<td></td>
</tr>
<tr>
<td>CI</td>
<td>Improved speech recognition on affected side, improved overall speech</td>
<td>Might prevent candidacy from future advancements in hearing restoration</td>
<td>Length of auditory deprivation can have a negative impact on performance</td>
</tr>
<tr>
<td></td>
<td>recognition in noise, improved localisation (adults and children) and</td>
<td>Performance might depend on amount of aural rehabilitation</td>
<td>Must have compelling audiologic data showing that the ear to be implanted</td>
</tr>
<tr>
<td></td>
<td>improved health related quality of life (adults)</td>
<td></td>
<td>will not benefit from other non-surgical forms of technology</td>
</tr>
<tr>
<td>Personal RMS</td>
<td>Addresses effects of noise, distance and reverberation on speech</td>
<td>Multiple microphones are needed for multiple talkers.</td>
<td>Cost is prohibitive and reimbursement is an obstacle in non-socialized health</td>
</tr>
<tr>
<td></td>
<td>understanding</td>
<td>Requires the talker’s cooperation</td>
<td>care models</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Use of a personal receiver could affect compliance for some users</td>
<td>All children considered for CI must have appropriate imaging to establish</td>
</tr>
<tr>
<td>Classroom Audio</td>
<td>Improved access to primary auditory signal.</td>
<td>Multiple microphones are needed for multiple talkers.</td>
<td>normal temporal bone anatomy</td>
</tr>
<tr>
<td>Distribution System</td>
<td>Delivers primary talker’s voice evenly throughout the learning space,</td>
<td>Requires the talker’s cooperation</td>
<td>Deciding which ear to fit depends on multiple factors, including degree of</td>
</tr>
<tr>
<td></td>
<td>through one or more loudspeakers. Benefits all listeners in learning</td>
<td>Limited portability and flexibility for use across a range of environments</td>
<td>hearing loss in the poorer ear and the ability to ensure open fitting in</td>
</tr>
<tr>
<td></td>
<td>space</td>
<td></td>
<td>better ear</td>
</tr>
<tr>
<td>Desktop RMS</td>
<td>Improved access to the primary auditory signal.</td>
<td>Multiple microphones are needed for multiple talkers.</td>
<td>Systems are most effective when classroom acoustics have been optimised</td>
</tr>
<tr>
<td></td>
<td>Addresses effects of noise, distance, and reverberation on speech</td>
<td>Requires the talker’s cooperation.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>understanding</td>
<td>Flexibility for use across a range of environments is limited</td>
<td></td>
</tr>
</tbody>
</table>


- Is it possible for parents to change their minds in the future (e.g. proceed with a surgical versus a non-surgical intervention)?
- When and how might the family begin to see a benefit (or a disadvantage) from technology use? Because it can be difficult to observe the impact of a UHL in young children, it is important to help families to have realistic expectations of progress.
- Families who choose a ‘watchful waiting’ approach should be equipped with information about key developmental milestones including knowing when and how to act if they are concerned. The use of functional assessment tools such as those listed in Table 1 can assist in engaging families in the monitoring of their child’s progress.
- If device fitting is declined or terminated by a family, ensure that the family is comfortable that they have received enough information to help with their decision and whether other support or information might be needed. For example,
  - Is the device itself the concern, or the feelings associated with its use?
  - Are there other factors impacting successful device use (e.g. influential others, bullying)?
  - What is the cost a concern? Are they considering what to buy and when (e.g. waiting on newer technology)
and are there hidden costs such as repairs, spare parts, and compatibility with other technology?

- Is the parent able to use the complicated features associated with the technology?
- Which HAT features benefit children in which environments and at what age? Ensure that families consider their child’s future needs as well as current concerns, so their decisions will provide the best value for their investment (both time and money).
- When is it time to invest in new technology or wait until a future feature or version comes out?
- What are the most advantageous and useful features that a particular child in his or her unique setting might need (DesGeorges 2015)?

Families receive information from a variety of sources including professionals, other families who have children who are deaf or hard of hearing, adults who are deaf or hard-of-hearing, and written and online information and resources. Audiologists can provide families with information to help them identify reputable online resources and to be aware that because the management of UHL is not clearly defined, they might encounter a variety of opinions about the best options for this type of hearing loss.

**Consensus-based priorities for research**

Recent systematic reviews have identified that studies characterising the consequences of UHL and evaluating the effectiveness of different management options have included different UHL characteristics (i.e. varying degrees and types of UHL), used different definitions of UHL, and used different outcome measures (Anne, Lieu, and Cohen 2017; Appachi et al. 2017). This heterogeneity in study design limits our ability to compare and contrast study findings. The following are consensus-based recommendations for further research in children with UHL designed to address treatment uncertainties and promote consistency in research design. These recommendations identify key issues in terms of PICOS (i.e. Participants, Interventions, Comparators, Outcomes, and Study designs/settings).
**Participants**

**Compare yield and accuracy of diagnostic tests**

Universal newborn hearing screening (UNHS) has lowered the average age of diagnosis for UHL by several years (Fitzpatrick, Whittingham, and Durieux-Smith 2014; Ghogomu, Umansky, and Lieu 2014). However, uncertainty remains over how to optimise audiological monitoring or medical evaluations based on known risk factors to detect late-onset or progressive cases of UHL.

**Separate reporting of children with UHL**

Synthesising evidence across studies is impeded by inconsistent definitions of the participant group such as the combined reporting of outcomes for children with UHL and mild bilateral losses (e.g. Appachi et al. 2017). Characteristics and outcomes of children with UHL should be reported separately from children with bilateral losses. Where effects of UHL or management options can be assumed to vary by degree of severity (e.g. Lieu et al. 2010), baseline characteristics and outcomes should be stratified into clearly defined categories of severity.

**Describe samples/cohorts in detail**

Standardised reporting frameworks consistently emphasise the need to report detailed information on sample characteristics (Schulz, Altman, and Moher 2010; Von Elm et al. 2014). A non-exhaustive list of key information that should be reported includes gender, ear with UHL, age at diagnosis, age at assessment, aetiology, degree of hearing loss in the poorer ear, age at intervention (e.g. HAT fitting), type of intervention, and any changes in intervention. Consideration should be given to self-selection bias in the design of the research and, where possible, an assessment of self-selection bias should be undertaken as part of the research design (Lieu and Dewan 2010).

**Interventions/comparators**

**Compare effectiveness of existing and emerging HATs**

There remains a lack of high-level evidence for the relative effectiveness of both established and emerging technologies and a paucity of evidence for the effectiveness of contemporary non-surgical interventions. Additional research should determine the effectiveness of these management options, report how the device was fitted, and consider assessing the influence of early versus late fitting.

**Determine what degrees of UHL are aidable with conventional hearing aids**

At the extremes of UHL are cases where the hearing loss is sufficiently minimal to raise questions about the benefits of aiding (McKay, Gravel, and Tharpe 2008) or so severe as to prompt the question as to whether aiding is likely to be appropriate and effective (Arndt et al. 2015). Future research should evaluate the effectiveness of HATs at or close to these extremes at a variety of ages (i.e. infants, school-aged children) to establish the points at which using a conventional hearing aid is unnecessary or ineffective.

**Outcomes**

**Ensure consistent selection, measurement, and reporting of outcomes**

Outcome measurement and reporting in studies of UHL are highly variable (Anne, Lieu, and Cohen 2017; Appachi et al. 2017). Research methodologies have been developed to identify what is most important to measure (Williamson et al. 2012) and for determining how those important outcomes can be measured (Mokkink et al. 2010). Further research should use consensus-based methods that involve clinicians, children with UHL, and the parents of children with UHL to identify the most important domains of the outcome when studying children with UHL and seek international consensus on the choice of measurement tools and instruments.

**Determine what a ‘successful’ intervention is and what predicts success**

As the impact of UHL can be highly variable across individual children (Reeder, Cadieux, and Firszt 2015), so too will be their needs, the choice of optimal intervention, and the desired effects of any intervention. A necessary step in designing studies to evaluate the effectiveness of an intervention is to define clearly what a ‘successful’ outcome is. This definition also needs to account for the smallest change that would be considered clinically meaningful, referred to as the minimal clinically important difference (MCID; Copay et al. 2007).

**Exploit data logging features of contemporary hearing technologies**

Contemporary HATs now include the capability to gather continuous and real-time information on device usage and the characteristics of the acoustic environments to which a child is exposed. Future research should exploit data logging as a tool to characterise the needs of children with UHL based on the environments in which they listen and use HAT.

**Study the relationship between behavioural and neuro-imaging outcomes**

Recent work has suggested that children with UHL form different connections and activate different neural networks than children with normal hearing (Jung et al. 2017). Future research should seek to combine detailed behavioural assessments of the impact of UHL and HATs with measures of central changes that might underpin longer-term effects of cognitive and emotional functioning, or provide predictive markers for the longer-term effects of early intervention.

**Study design/setting**

**Adopt multi-site designs**

Single-site studies are more likely to report larger effects than multi-site studies, an effect that can arise because single-site studies tend to have smaller sample sizes, be at greater risk of bias, and recruit more homogeneous samples (Bafeta et al. 2012). Multi-site designs should be adopted whenever possible and research teams should consider publishing study protocols either in scientific journals or online databases (e.g. clinicaltrials.gov).
Consider conducting replication studies

Replicating the methodology of key studies should be considered where the research question being answered is important and a limited number of existing studies have been conducted. To assist with replication, all teams conducting interventional research on UHL should ensure that their studies are registered prospectively and comply with established reporting standards (Schulz, Altman, and Moher 2010).

Conduct randomised control trials to compare effects of interventions

Using randomisation to determine the choice of intervention for an individual child within a study reduces the influence of selection bias, and allows one to assume that differences in outcomes between groups are likely to be due to the differences in the intervention they received (Akobeng 2005). A systematic review of surgical and non-surgical interventions for UHL in children identified no studies that used randomisation (Appachi et al. 2017).

Exploit longitudinal study designs to characterise long-term effects of untreated UHL and effects of interventions

Few studies have followed cohorts of children with UHL over long periods of time (Anne, Lieu, and Cohen 2017). Future research should consider the use of longitudinal designs to characterise the specific deficits that UHL imposes on a child’s developmental profile.

Leverage a ‘big data’ approach to identify risk factors for UHL

Identifying key risk factors requires access to a wide range of information including the results of audiometric evaluations and medical imaging (Friedman et al. 2013), and genetic testing (Dodson et al. 2012). The uncertainty around potential risk factors and the numerous potential aetiologies means that a large sample of cases and controls will be necessary to identify key risk factors with a high degree of certainty. Future research should address the identification of these risk factors, and in doing so consider whether existing large-scale population databases (‘biobanks’) could be leveraged and/or extended to address questions related to UHL.

Notes

1. For purposes of this document, normal hearing refers to an ear that is audiologically normal to near normal, not requiring audiologic intervention.
2. Although commonly referred to as bone-anchored hearing aids (BAHA), Cochlear Corporation now holds a trademark on that term for their specific devices. For purposes of this document, the term for this class of device is bone-conduction devices.
3. Profound unilateral hearing loss has been referred to as single-sided deafness, which is a non-audiologic term that emerged in recent years (Cire 2017). For purposes of this guideline, the term profound unilateral hearing loss is used.

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