Chapter 11

Cochlear Implants: Determining Candidacy for Young Children Who Are Deaf or Hard of Hearing

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Introduction

Families who have chosen a listening and spoken language outcome for their children who are deaf or hard of hearing (D/HH)—or desire to have sound be a meaningful part of communication—have a variety of options to help their children access speech and environmental sounds. With significant advancements in hearing aid technologies, real-ear fitting techniques, and the use of other hearing assistive technologies (HAT), children can hear better than ever before. There are times, however, that even with an appropriately fitted hearing aid technology, children cannot access critical speech information that can help them with the development of spoken language. At this critical juncture, cochlear implant(s) may be recommended. For Part C coordinators, Early Hearing Detection and Intervention (EHDI) coordinators, and early interventionists, it is critical to understand the cochlear implantation process, especially as more parents choose this procedure for their children who are D/HH.

A primary goal for all children who are D/HH is to obtain communicative competence (Ganek et al., 2012) and minimize the effects of hearing loss on the child's development. For those children who receive limited or no benefit from amplification, cochlear implantation is
often a viable option with associated positive outcomes in listening, spoken language, literacy, and social/emotional well-being (Fryauf-Bertschy, Tyler, Kelsay, Gantz, & Woodworth, 1997; Geers, 2008; Geers & Moog, 1994; Geers, Tobey, & Moog, 2008). Because each child must be evaluated from a variety of perspectives, an interdisciplinary approach to determine candidacy is the existing standard of care. To arrive at a candidacy decision, the child undergoes medical, audiological, and speech-language evaluations. These evaluation results, along with the long-term communication and educational goals of the parents, lead to candidacy decisions that are family centered and in alignment with the parents' desired outcomes.

The Process: An Interdisciplinary Approach

The decision to pursue cochlear implantation for a child who is D/HH requires careful consideration and thorough counseling. The success of an interdisciplinary approach depends upon collaboration among an effective team that includes the family as equal partners in the decision-making process. There are a number of considerations that may be unique to the child and family. The process includes the collection and consideration of medical and audiological findings and is further supported by evaluation by a speech-language pathologist (SLP), input from other interventionists and educators and importantly from the family. Counseling and discussion with the family about the process, as well as short-and long-term goals, is essential. When undergoing the evaluation, the following questions should be addressed:

- Are there other noninvasive technologies available that can make sounds accessible to develop listening, spoken language, literacy, and social skills?
- Are there qualified intervention providers and family support services to help maximize the child's ability to learn to listen and communicate?
- Are there aspects of the child that will require consideration of other forms of communication, and if so, how will they be implemented to supplement benefits from the cochlear implant?
- Are there any safety issues that should be considered to minimize any potential risk for this surgery, programming, and/or intervention?

As the child and the family progress through the cochlear implant candidacy process, in addition to the above questions, the interdisciplinary team members are trying to determine:

- Does the child meet the criteria for a cochlear implant based on the Food and Drug Administration (FDA) labeling (see Table 1)?
- If not, based on research and clinical observations, could the child receive more benefits if he or she received the cochlear implant? What are those benefits?
- Does the family have the information needed to plan for the best possible outcome?

To answer these questions, the cochlear implant team will not only consider the audilogic and medical results but will be evaluating the “whole” child and family unit (see Table 2; Winter & Phillips, 2009). The first step is to obtain a comprehensive history, including:

- Information on the incidence of hearing loss in the family.
- Birth history.
- Review of complications or concerns.
- Results of newborn hearing screening.

Results of previous assessments will dictate the need for further evaluation by the physician, audiologist, and SLP.

With a family's consent, the early intervention program, child care program, and/or school of choice should be consulted to review the child's response to current services and determine the
<table>
<thead>
<tr>
<th>Company</th>
<th>Device Name</th>
<th>Pediatric Approval Guidelines</th>
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</thead>
</table>
| Advanced Bions | Implant: HiRes90K                    | **12 months to 17 years**  
• Profound, bilateral sensorineural hearing loss (>90dB).  
• Used appropriately fit hearing aids in children under 2 for at least 6 months and for children 2-7 years of age for at least 3 months with little to no benefit.  
**<4 years**  
• Failure to reach appropriate developmental milestones as measured by IT-MAIS or MAIS and/or <20% word recognition testing.  
**>4 years**  
• <12% on word recognition testing.  
• <30% on sentence recognition testing. |
| Cochlear™      | Implant: Cochlear Nucleus® Profile Implant | **2 to 17 years**  
• Severe-to-profound bilateral sensorineural hearing loss.  
• Limited benefits from binaural hearing aid trial with word recognition scores ≤30%.  
**12 to 24 months**  
• Profound sensorineural hearing loss.  
• Limited benefits from binaural hearing aid trial.  
**Older Children**  
• ≥30% on MLNT or LNT.  
**Young Children**  
• Lack of progress for 3-9 months with amplification and intensive aural rehabilitation. |
| MED-EL         | Implant: MED-EL Cochlear Implant System—Synchrony | **12 months to 17 years, 11 months**  
• Bilateral, profound sensorineural hearing loss with at least a 90dB loss at 1000Hz.  
• 3-6 month hearing aid trial.  
• Little or no benefits from appropriately fit binaural hearing aids.  
• Lack of progress in developing auditory skills with amplification and intensive aural rehabilitation.  
• Scoring <20% on speech recognition tests MLNT or LNT. |
### Table 2
**Description of Cochlear Implant Candidacy Evaluations**

<table>
<thead>
<tr>
<th>Candidacy Evaluation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Audiological Evaluation</td>
<td>A comprehensive hearing assessment completed with and without the child’s hearing aids. This may require more than one visit. It is essential that the child brings his hearing aids and ear molds to the evaluation.</td>
</tr>
<tr>
<td>Auditory Brainstem Response and Otoacoustic Emissions Tests</td>
<td>Per the JCIH statement, the child should have at least one objective measure of hearing sensitivity. Some children require sedation to obtain these test results. If an ABR has not been completed, then one may be recommended by the cochlear implant team.</td>
</tr>
<tr>
<td>CT Scan/MRI</td>
<td>A specialized X-ray to evaluate the anatomy of the inner ear. Some children are sedated for this procedure. It is important to determine the status of the cochlea and the internal auditory meatus.</td>
</tr>
<tr>
<td>Medical Examination</td>
<td>The otologist/otolaryngologist will take a medical history, review the CT scan, and determine if there are any medical contraindications to surgery and make referrals to other medical specialties, as needed.</td>
</tr>
<tr>
<td>Speech-Language Evaluation</td>
<td>Formal and informal assessment of the child’s communication abilities with his/her hearing aids. Communication goals are usually discussed at this appointment.</td>
</tr>
<tr>
<td>Developmental/Cognitive/Psychological Evaluation</td>
<td>Formal and informal assessment of the child’s developmental milestones and capacity to learn.</td>
</tr>
<tr>
<td>Social Work Evaluation</td>
<td>To evaluate parent stressors and family support, the social worker will work with the family to navigate services needed to maximize the child’s outcomes. Family expectations also will be discussed.</td>
</tr>
<tr>
<td>Educational Assessment</td>
<td>The child’s school will be contacted regarding educational placement, support, and the need, if any, for inservice on cochlear implants.</td>
</tr>
<tr>
<td>Other Assessments</td>
<td>A genetic evaluation and ophthalmology examination may also be recommended. Since 40% of children with hearing loss may have additional special needs, genetic testing may assist the family in making a decision about how to proceed.</td>
</tr>
</tbody>
</table>
educational needs of the child. There may be additional assessments required to determine candidacy for the cochlear implant—depending on the age of the child and the needs of the child or the family. Table 2 provides a brief description of the evaluations the child and family may undergo to determine candidacy. The family—and the child (if old enough)—may be asked to complete an expectation questionnaire to assist the professionals in realistic counseling. Once the evaluations are completed, the cochlear implant team members review the findings and make a recommendation to the family.

There are many tools that can guide the cochlear implant team to help families understand the potential benefit of cochlear implantation and participate in the planning of ongoing support and intervention. While cochlear implants have been approved by the FDA based on published research guidelines, families and cochlear implant team members can decide to pursue cochlear implantation, even if the child performs outside of the FDA guidelines. Obtaining a complete medical and audiological history and gaining an understanding of the family’s resources and challenges is essential. Counseling that addresses many factors that affect outcomes can be reviewed and discussed. These may include aspects of the child, including:

- Anatomical, physiological, cognitive, developmental, and behavioral characteristics.
- The child’s hearing history, including age at onset of hearing loss, degree of loss, and age at diagnosis.
- The use of technology, including age at hearing aid fitting and consistency of use.
- The educational and therapeutic services that have been in place.
- Considerations of the family and environment.

The interdependence of these predictors is summarized in Figure 1 (Teagle & Eskridge, 2010). Using a candidacy checklist can help identify factors that may influence the outcomes. The Graded Profile Analysis (GPA; Daya et al., 1999), Children’s Implant Profile (CHIP; Hellman et al., 1991), the Cochlear Implant Candidacy–Children (CICC; Bradham, Lambert, Turick, & Swink, 2003), or the Modified ChIP (Barnes, Lundy, Schuh, Foley, & Maddern, 2000) are tools that guide the team in their discussions to identify strengths and needs as the family considers cochlear implantation. It is important to note that these tools are not meant to “grade” the family but to identify potential issues that could negatively impact meeting the family’s goals and expectations. Furthermore, in the era of having to justify payment for services, these measures can serve as an “objective” tool in making the case for reimbursement for services rendered.

The decision to pursue cochlear implantation is made once the evaluation is complete and families have been counseled on the potential benefits and risks of proceeding. The decision to pursue cochlear implantation is made once the evaluation is complete and families have been counseled on the potential benefits and risks of proceeding. With a thorough understanding of the process, the need for ongoing intervention, and the potential benefits to the child, families should be well equipped to make a decision that will have a profound impact on their child’s future. The recommendations fall into three categories:

- Proceed with the cochlear implant.
- Do not proceed (and why).
- Wait (and why).

It is not uncommon for families to want a second opinion. Every effort should be made to assist the family when seeking additional advice from other health care providers.

Medical/Physical Component

The role of the neuro-otologist/pediatric otolaryngologist/otologist precedes and extends far beyond performing the surgery. It includes, in collaboration with the audiologist, the diagnosis of hearing loss, the degree and type of loss, and etiology.
In addition, based on the physical evaluation and medical history, the surgeon considers the need for other laboratory tests as imaging (Computerized Tomography [CT] scan and/or Magnetic Response Imaging [MRI]) to evaluate the anatomical structures of the ear and brain. Other medical interventions and referrals may be indicated. The surgeon then discusses with the parents treatment options and ways to prevent further hearing loss or other related complications. The search for etiology and the identification of other medical conditions can impact the sequence and timing of treatment. For children with complex medical histories and co-morbid conditions or syndromes, referrals to neurology, genetics, ophthalmology, and other specialists are common (Buchman et al., 2008).

Radiographic imaging is an important topic both before and after surgery. With a combination of CT scanning and MRI, it is possible for the surgeon to visualize both the bony and soft tissues of the ear and neural anatomy. A cochlear implant cannot overcome the limitations of a severe cochlear malformation or an absent or diminished auditory nerve. Therefore, it is critical for the surgeon to obtain and share this information with the team and parents, as it can significantly affect cochlear implant outcome (Adunka et al., 2006; Adunka et al., 2007). Because the presence of an implanted device can impact future imaging needs, discussion of the contraindications to future imaging studies must take place with the parents.
The risks of surgery are typically discussed with the family by the surgeon (see Table 3). While it is rare to have complications in the hands of an experienced surgeon, parents must consider the possibilities during the decision-making process, and informed consent requirements dictate this discussion. Of particular importance is the increased risk of meningitis. Bacterial meningitis is a serious infection of the brain and the fluid surrounding it. Children who are D/HH and have cochlear implants have a higher risk for meningitis, so additional vaccines are recommended. The Centers for Disease Control and Prevention (CDC) provides detailed information on this topic (http://www.cdc.gov/vaccines/pubs/vis/downloads/vis-pcv.pdf).

As the evaluation of candidacy proceeds, discussion among the team includes the ear of implant, type of electrode array, and determination of whether the child should be a unilateral, bimodal (i.e., a hearing aid in one ear and a cochlear implant in the other ear), or a bilateral recipient (i.e., receive cochlear implants in both ears). If it is decided to proceed with two cochlear implants, the family and surgeon will need to discuss sequential versus simultaneous cochlear implantation. Factors that will determine how to proceed include:

- Age of the child.
- Degree of residual hearing.
- Family choice.
- Financial coverage/reimbursement rates.

### Table 3

**Risks of Surgery**

<table>
<thead>
<tr>
<th>Cochlear Implant</th>
<th>Any Ear</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loss of remaining hearing in the implanted ear.</td>
<td>Numbness/tenderness around implant site.</td>
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<tr>
<td>Facial nerve stimulation/involuntary facial movement.</td>
<td>Loss of feeling in face.</td>
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<tr>
<td>Inflammation/extrusion/swelling.</td>
<td>Change in taste.</td>
</tr>
<tr>
<td>Soreness, redness, or breakdown of skin in area around the implant, which may need more medical treatment, surgery, and/or removal of device.</td>
<td>Fluid leak.</td>
</tr>
<tr>
<td>Failure of surgery, possibly requiring removal of the implant.</td>
<td>Dizziness (vertigo).</td>
</tr>
<tr>
<td>Failure of implanted pieces, which may need replacing.</td>
<td>Tinnitus or “ringing in the ears.”</td>
</tr>
<tr>
<td>The CI may not work correctly, or it may cause your child to feel or hear odd or uncomfortably loud sounds.</td>
<td>Blood, fluid, or infection at the site or close to the site of surgery.</td>
</tr>
<tr>
<td></td>
<td>Skin reactions (rashes).</td>
</tr>
<tr>
<td></td>
<td>Pain, scarring, bleeding, and infection.</td>
</tr>
<tr>
<td></td>
<td>Anesthetic risks (medicines used to put the child to sleep) associated with the heart, lungs, kidneys, liver, and brain.</td>
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</tbody>
</table>
Recent studies suggest that outcomes for bilateral cochlear implantation are impacted by the child's age and time between surgeries (Galvin et al., 2014; Spareboom et al., 2014). In the presence of severe-to-profound hearing loss, earlier implantation—whether the first or second ear—yields better results.

For most pediatric cochlear implant recipients, once postoperative recovery is complete, the surgeon has less frequent interactions with the child and family relative to the SLP and audiologist. It is important, however, to maintain this relationship should concerns about ear and hearing health or the need for future surgeries arise. Of course, everyone likes to share and celebrate individual progress and the opportunity to have a meaningful role in the child and family's life. Team dynamics and practices are shaped by retrospective knowledge of each child's outcome.

**Audiology Component**

While all members of the interdisciplinary team interact with a family, an audiologist often serves as the initial or primary point of contact once a child has been identified as deaf or hard of hearing. For children who are identified with significant hearing loss at birth through a newborn hearing screening, there may be several audiologists involved in diagnosis, hearing aid fitting, and objective and ongoing behavioral assessment of hearing. The general goal of audiological management is to determine and monitor hearing thresholds and to provide the best possible access to sound. If degree of hearing loss is severe to profound, and the development of early communication milestones is delayed, a cochlear implant evaluation should be recommended, so parents can begin to consider this option as the child approaches the first birthday. While the FDA guidelines recommend cochlear implantation after 1 year of age, there are times when a cochlear implant will be recommended prior to the first birthday (e.g., child develops hearing loss as a result of bacterial meningitis). Children who are older with progressive or acquired hearing loss are often referred when communication challenges become difficult to address through the use of conventional amplification. An audiologist who serves on a cochlear implant team is typically responsible for:

- Collecting information about the child and family.
- Assessing hearing loss and benefit from amplification.
- Providing counseling about the implantation process, the technology, and the considerations for device use and follow-up care.

As mentioned previously, the FDA-approved criteria for pediatric cochlear implantation, which has been unchanged since 1990 (see Table 1), includes children who are 1 year of age or older, have severe-to-profound hearing loss (often interpreted as a pure tone average [PTA] of 90 dB HL or poorer), and/or demonstrate a lack of development in audition skills. Less conservative criteria have been advocated for and supported by several studies. Not only are children with lesser degrees of hearing loss and better speech perception performance being considered (Carlson et al., 2015; Dettman et al., 2004; Gantz et al., 2000), but children under 12 months of age are being implanted (Tajudeen, 2010). As more children receive cochlear implants, and the benefits are documented, the candidacy criteria have expanded in practice. Consideration of the individual child and his or her unique circumstances and implementation of best clinical practices should drive decision making.

The audiological assessment should include both physiologic and behavioral assessments to determine ear-specific degree and type of hearing loss. A diagnostic auditory brainstem response (ABR) assessment can provide a good estimate of hearing levels for children with sensorineural hearing loss. Typically, reliable behavioral testing of babies is possible using Visual Reinforcement Audiology (VRA) techniques starting at about 6 months of age. Hearing aids can be fit on the basis of ABR results and refined once behavioral information is obtained. Cochlear implantation is usually
deferred until a hearing aid trial has been completed. However, there is evidence that children who have no response ABR results are very likely to become cochlear implant recipients (Hang et al., 2015). Ideally, the family has the opportunity to explore the child’s use of noninvasive technologies in an environment that includes auditory intervention by a qualified therapist. For children with very limited residual hearing, the length of the hearing aid trial should not be extended beyond the time it takes to resolve other considerations addressed in the cochlear implant evaluation, including acquiring medical information, treatment, and counseling.

Depending on the child’s age and abilities, a battery of speech perception tests are used to document benefit from amplification. While no standard universal pediatric test battery has been recognized among cochlear implant teams, a number of tests have been developed or are routinely used in cochlear implant assessment. The commonly used tests are listed and briefly described in Table 4. Importantly, speech perception assessments should be selected that are appropriate for the child and can serve as a baseline to measure future progress. Because many children are too young and lack the communication skills to participate in speech perception assessments during candidacy evaluation, the audiologic assessment should include baseline auditory functional assessments. These functional assessments can include questionnaires, such as the Infant Toddler-Meaningful Auditory Integration Scales (IT-MAIS; Zimmerman-Phillips, Robbins, & Osberger, 2000) and/or LittEARS (Coninx et al., 2009), as well as aided testing in the sound booth and real-ear measures. The collaborative efforts of the audiologist, SLP, and early interventionist can combine to determine the benefit from amplification.

If the child is considered to be a cochlear implant candidate, a determination must be made regarding which device to use. Currently there are three manufacturers with established histories who produce the technology (see Table 1). Some centers only offer the option of a cochlear implant system available from one manufacturer, while others offer systems from multiple manufacturers. In some cases, the surgeon may recommend a particular device based on the medical and radiologic examination.

It is incumbent on the cochlear implant team to ensure that the family has access to unbiased information about each of the cochlear implant systems available and approved by the FDA. There is an abundance of information available to families via the Internet, including manufacturer websites and social networking sites. Support groups and other cochlear implant recipients can also share personal experience and perspective. It is important to note that not all sources of information will provide accurate and unbiased information. In the interest of preparing effectively for surgery, device programming, and for achieving outcomes that meet the family’s expectations, counseling from members of the cochlear implant team and shared decision making with the family is essential.

Beyond assessment, the cochlear implant team audiologist provides extensive counseling and information. In the process, he or she establishes a relationship with the child and parents and gains some insight about the family’s acceptance of the diagnosis and the stage at which they are entering the decision-making process.

• Is the family responding from grief or anger?
• Have they idealized the process and created expectations of normal hearing?
• Are they cognizant of other developmental or medical issues the child might have, and does the team appreciate what these might be?
• Is the family’s preference for communication mode realistic, and are services in place to support this plan?
### Table 4

**Description of Speech Perception Tests**

<table>
<thead>
<tr>
<th>Text</th>
<th>Age Recommendation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Consonant Nucleus Consonant (CNC) Test (Peterson &amp; Lehiste, 1962)</td>
<td>Recommended for older children and teens. This test is used to determine adult CI candidacy and includes less common vocabulary, which makes it more challenging than PB-k or LNT monosyllable word tests.</td>
<td>This test includes 10 lists of 50 monosyllabic words with equal phonemic distribution across lists, with each list having approximately the same phonemic distribution as the English language.</td>
</tr>
<tr>
<td>Early Speech Perception Test (ESP) (Moog &amp; Geers, 1990)</td>
<td>Recommended for children with limited vocabulary who cannot participate in open-set word testing. Minimum of 2 years for low verbal version and minimum of 6 years for standard version per test developers but can be attempted for younger ages.</td>
<td>Two versions, including low verbal and standard—both closed-set. Can be presented via live voice or a recording. Lo-verbal test materials consist of objects (toys) instead of pictures. The standard version includes 36 words presented as 3 subtests of 12.</td>
</tr>
<tr>
<td>Hearing in Noise Test (HINT-C)</td>
<td>Sentence material that requires child to have vocabulary and auditory memory to repeat. Recommended once these skills exhibited. HINT is used for adult CI candidacy determination.</td>
<td>HINT-C includes multiple lists of 10 sentences that are five to seven words in length. Can be presented in competing noise for more challenging assessment.</td>
</tr>
<tr>
<td>Ling Six Sound Test (Ling &amp; Ling, 1978)</td>
<td>Appropriate for any age once the child has learned to repeat on demand. These sounds (Learning to Listen Sounds) are used very often in therapy and therefore familiar to children.</td>
<td>The sounds used in this test are the vowels /a/ as in all, /u/ as in who, and /i/ as in be, and the consonants /m/ as in me, /S/ as in she, and /s/ as in so. These sounds include low-, mid-, and high-frequency components of speech. The ability to detect and discriminate these phonemes is the basis of scoring.</td>
</tr>
<tr>
<td>Lexical Neighborhood Test (LNT) (Kirk, Pisoni, &amp; Osberger, 1993)</td>
<td>Appropriate for children age 4-5 and older who can repeat words on demand.</td>
<td>This is a recorded open-set test of monosyllabic word recognition. The word list consists of 25 lexically “easy” words (high-frequency occurring words) and hard words (low-frequency occurring and more confusable). It is scored by both number of words and phonemes correct.</td>
</tr>
<tr>
<td>Multisyllabic Lexical Neighborhood Test (MLNT) (Kirk, Pisoni, &amp; Osberger, 1996)</td>
<td>For children age 3 and older who can repeat on demand. It is often used before the LNT, as vocabulary is easier because of redundant cues of multisyllable words.</td>
<td>This is recorded open-set test of multisyllabic word recognition. The word list consists of 12 lexically “easy” words and 12 lexically “hard” words scored by both number of words correct and number of phonemes correct.</td>
</tr>
<tr>
<td>Phonetically Balanced Kindergarten Test (PBK-50) (Haskins, 1949)</td>
<td>Recommended age is 4+ years, but children who will repeat what they hear regardless of comprehension can be tested to determine speech sounds perceived.</td>
<td>This is an open-set test of monosyllabic word recognition. Can be presented live voice or recorded. A full list consists of 50 phonetically balanced, one syllable, kindergarten words that the examiner phonetically transcribes to obtain a word and phoneme score.</td>
</tr>
</tbody>
</table>
Counseling and support needs can be shared with the team to help resolve these and other important issues.

Finally, plans for appointments and services for the next year and beyond should be discussed. During the first year following surgery, frequent device programming visits are necessary to optimize the program and ensure audibility is maximized. The typical child adapts to the electrical signal over time; tolerance increases; and as experience in hearing grows, children can play a larger role in providing feedback about hearing. At minimum, the following schedule is recommended for children:

- Initial stimulation (IS) occurs approximately 2 to 4 weeks after surgery
- 2 weeks post IS
- 1 month post IS
- 3 months post IS
- 6 months post IS
- 9 months post IS
- 1 year post IS
- Semiannual visits thereafter

During these appointments, hearing tests and speech perception assessments are completed to guide programming, validate settings, and ensure appropriate progress is being made. Families gain experience and confidence in managing the technology over time, but the audiologist continues to be a source for new information and problem solving on issues related to device use, such as troubleshooting and device retention.

**Speech-Language Pathology Component**

For children who are D/HH and are being evaluated for possible cochlear implantation, it is vital that the SLP have the knowledge and skills to accurately assess the child’s present level of functioning and determine whether the child’s communication development can be enhanced with cochlear implants.
If the child is delayed or is at risk for delayed language development, cochlear implantation may be the best option available.

When determining candidacy, most experienced SLPs serving children who are D/HH will use speech and language assessments that are standardized on normal hearing children—with only a few exceptions. If the child who is D/HH is acquiring spoken language, the SLP should use assessments that compare the child's performance to what is considered to be typical development for the child's age and cognitive abilities.

In addition to obtaining current and accurate audiological assessments on the child who is D/HH, SLPs also must obtain measures of functional listening skills, especially if the expectation is to use audition to develop spoken language. SLPs must document how the child is using his or her aided hearing in conjunction with amplification, hearing assistive technology (e.g., digital hearing aids and/or personal FM system), as well as how the child is responding to both environmental sounds and speech. For infants and toddlers, these auditory skills can be measured through play activities and in conjunction with parental or caregiver interviews, questionnaires, and informal assessments. For children ages 3 and above, more formal speech perception measures can be used, such as the Early Speech Perception Test for Profoundly Hearing-Impaired Children (ESP) developed by Moog and Geers (1990). In addition, clinicians may wish to use the Auditory Perceptual Test for the Hearing Impaired-Revised (Allen, 2008).

In conjunction with standardized measures, informal assessments are useful in determining how the child who is D/HH is functioning in their everyday environment, such as school, home, and in the community. Informal assessments can be given to the family; teacher; child, depending on age; caregiver; or completed by the SLP. Informal assessments include, but are not limited to:

- LittlEars Auditory Questionare

- PEACH
  Ching & Hill, 2007

- Early Listening Function (ELF)
  Anderson, 2002; Oticon, 2007

- Children’s Home Inventory for Listening Difficulties (CHILD)
  Anderson & Smaldino, n.d.

- Child Auditory Performance Scale (CHAPS)
  Smoski, Brunt, & Tannahill, 1998

- Listening Inventory for Education—Revised (LIFE-R)
  Anderson, Smaldino, & Spangler, 2012
  • LIFE-R Student Appraisal
  • Teacher LIFE-R

- Functional Listening Evaluation (FLE)
  Johnson & Von Almen, 1993

- Preschool Screening Instrument for Targeting Educational Risk in Preschool Children (Preschool SIFTER)
  Anderson & Matkin, 2004

- Informal Assessment of Fatigue and Learning
  Anderson, 2014; Fukuda et al., 2010

- Minnesota Social Skills Checklist for Students Who Are Deaf/Hard of Hearing
  Minnesota Department of Education, n.d.

- Social Communication Skills Pragmatics Checklist
  Goberis, 1999; Simon, 1984

- Placement and Readiness Checklists (PARC)
  Johnson, 2011

Speech intelligibility is another way to assess speech development and can be measured in terms of overall intelligibility, including segmental and suprasegmental errors (Tye-Murray, 1994). Speech intelligibility is a critical area of assessment that may be overlooked by SLPs. Formal measures of speech intelligibility are limited. The most common assessment is the CID Picture SPINE: Speech Intelligibility Evaluation (Monsen, Moog, & Geers, 1988). The Pediatric Speech Intelligibility (PSI) test can also be given as an assessment of speech intelligibility for children who are D/HH (Jerger & Jerger, 1984; Jerger et al., 1980, 1981). However, due to the limited formal
While this list of language assessments is not exhaustive, most practitioners who assess language acquisition of children who are D/HH to determine candidacy for cochlear implantation will use at least some of these evaluations in their preferred diagnostic protocol.

Assessments of speech intelligibility, many SLPs develop their own. These assessments obtain a percent of words, phrases, and sentences that are correctly spoken by the child who is D/HH and understood by familiar and unfamiliar listeners.

The acquisition of suprasegmental and segmental skills can be assessed using instruments that were designed to evaluate the spoken language of children who are D/HH. The Ling Phonetic-Phonological Speech Evaluation (Ling, 2002) is commonly used for this purpose.

Another assessment developed specifically for children who are D/HH and acquiring spoken language is Identifying Early Phonological Needs in Children with Hearing Loss (Paden & Brown, 1992). If the child has acquired some spoken language, most clinicians will use standard assessments, such as:

- Goldman-Fristoe Test of Articulation, 2nd Edition (Goldman & Fristoe, 2000)
- Arizona Articulation Proficiency Scale, 3rd Edition (Fudala, 2000)
- MacArthur-Bates Communicative Development Inventories (Fenson et al., 1993)
- Rossetti Infant-Toddler Language Scale (Rossetti, 1990)

As Tye-Murray (1994) notes, the assessment of a child’s language by an SLP usually involves the evaluation of syntax, morphology, semantics, vocabulary, and pragmatics. For infants and toddlers who are D/HH, practitioners may use assessments that measure performance across several developmental domains, such as:

- Peabody Picture Vocabulary Test, 4th Edition (PPVT-4) (Dunn & Dunn, 2006)
- Test of Auditory Comprehension of Language, 3rd Edition (Carrow-Woolfolk, 1999)
- Expressive Vocabulary Test, 2nd Edition (EVT-2) (Williams, 2006)
- Oral-Written Language Scales (Carrow-Woolfolk, 1995)

While this list of language assessments is not exhaustive, most practitioners who assess language acquisition of children who are D/HH to determine candidacy for cochlear implantation will use at least some of these evaluations in their preferred diagnostic protocol. Preferences based on clinical and professional experiences, as well as other factors related to a child’s unique case history and learning needs, also influence the selection of communication measures and assessments.
Conclusion

Determining if a child is a candidate for cochlear implantation requires an interdisciplinary team approach that places the family at the center of the decision-making process. Once the child is identified as being D/HH, parents should be informed about all of the technological options available, including cochlear implants. The cochlear implant team—comprised of at least a surgeon (i.e., otolaryngologist, otologist), audiologist, and SLP—will complete comprehensive medical, audiological, and speech-language assessments to ascertain if cochlear implantation would be beneficial to the child. The team discusses these findings with family and provides information about candidacy and potential outcomes, including the support that will be essential to achieve the desired communication and academic outcomes that were expressed by the family.

Once the child recovers from cochlear implant surgery, and the device(s) are activated, the real journey begins. Consistent audiological support with cochlear implant programming is required to ensure the speech processor program has been optimized, and the implant is working properly. The child must receive appropriate early intervention services that will focus on teaching the child to associate meaning with the auditory information provided by the cochlear implant(s). As hearing with a cochlear implant(s) is quite different than listening with hearing aids, the newly implanted child and family should receive speech-language therapy that has a strong auditory component. These services should be provided by an early interventionist or clinician who is well trained and experienced. The goal is to assist the family to integrate listening and communication into the daily routines that occur in the home. Through consistent, timely, and well-coordinated early intervention, young children with cochlear implants often can achieve language outcomes that rival their hearing peers.
References


