Outcomes for Children Who Are Deaf-Blind After Cochlear Implantation

Research Brief: Basic Information on Children with Deaf-blindness After Cochlear Implants

This research brief is the result of a study, Outcomes for Children Who Are Deaf-Blind after Cochlear Implants. The study collected information from 2006 – 2009 on 115 children who are deaf-blind and had recently received cochlear implants. The children span 24 states and have a variety of additional health challenges.

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A cochlear implant is a hearing device that is surgically implanted by a medical team to help people learn to hear. The cochlear implant has “an electronic system that stimulates the hearing nerves directly sending a sensation of sound to the brain” (Wheeler, 2008, p.10). Several research studies have been published concerning the “post-implant patterns of progress for children who are deaf; however few exist about children who are deaf-blind” (Bashinski, Durando, & Stremel Thomas, in press).

The intent of this research brief is to provide information to service providers about the progress children with deaf-blindness demonstrate after cochlear implantation, as well as families’ reactions and perceptions of the impact a cochlear implant had on the children in the study. The desired outcome of this brief is to provide meaningful information to educators and other services providers supporting children with cochlear implants and their families.

Children with deaf-blindness are often not totally deaf and totally blind; however over 85% of the population experience physical or cognitive disabilities, complex medical needs, or behavior challenges (Killoran, Davies, & McNulty, 2006). The December 2008 the National Consortium on Deaf-Blindness child count reported 763 children and young adults (birth – 21 years of age) have at least one cochlear implant (NCDB, 2009). This number is likely to continue to climb resulting in a need for educators, medical professionals, and service providers to have quality information about cochlear implants for children with both vision and hearing loss.

An initial study, Outcomes for Children Who Are Deaf-Blind after Cochlear Implants (CIDB), collected information from 2006 to 2009 about children between the ages of 10 months
and 12 years who are deaf-blind who had recently received cochlear implants. The research study was interested in seeking information to determine: (1) What progress did the children demonstrate, post-implant, in a variety of development domains? and (2) How do participating families feel about their child’s implant, and how do they perceive its impact—on both the child and the family?

The CIDB study included 115 children of 12 years of age and younger resulting in a file containing 495 variables of data. This number was both wonderful and conflicting because of the challenge it presented in sifting through the endless amount of “variability in nearly every aspect of the individuals who participated in the research” (Durando, 2010, p.1).

The outcomes of the children in the CIDB study from the scores on the Reynell-Zinkin Scales: Developmental Scales for Young Visually Handicapped Children (1981) assessment and interviews conducted of the families supported previous research. Both “the amount of time a child had been ‘in sound’ and the age at which the child received the cochlear implant was related to progress” (Bashinski, et al, in press). The younger the children were when the implant was activated, the higher the child tended to score on the Reynell-Zinkin Scales assessment, particularly on the subscales: exploration of the environment, social adaptation skills, expressive language vocabulary and content, and response to sound and verbal comprehension (Bashinski, Stremel Thomas, & Durando, 2009).

Results from this initial study suggest there are a number of factors that contribute to whether a child will benefit from a cochlear implant. The continued increase in the number of families thinking about cochlear implants for their child supports the need for increased information in order to help families make a more informed decision. Examining how families feel about the process of obtaining an implant and the outcomes their child experienced is critical to families who contemplate a cochlear implant as an option.

The CIDB research team decided that a family survey would be a useful way to gather information about families’ perspectives. After a review of the Children with Cochlear Implants: Parental Perspectives (Archibold & Lutman, n.d.) the team decided to develop a survey that was shorter in length and more targeted on their specific areas of interest. A new family survey was developed and piloted with families of children who are deaf-blind. A total of 60 families of the 71 contacted completed the new survey. Each of the participating families has or had a child with deaf-blindness with at least one cochlear implant. Ethnic diversity of the families varied, 19
states were represented, and the most common causes of the child’s deaf-blindness were
CHARGE syndrome or complications of prematurity.

The findings presented in this brief are the preliminary results of what the researchers
learned about outcomes for children with deaf-blindness after cochlear implantation and the
family survey conclusions. It is important to remember when reviewing the findings of this study
that they derive from a first investigative effort with a group of children who are known to be
exceptionally unique. Therefore, it cannot be assumed that the results would apply equally to
every child with deaf-blindness.

Key Findings from the Family Survey include:

➢ Families told us –
  ▪ 85% of parents agreed that, “knowing what they now know, they would again make
    the decision to seek a cochlear implant for their child”.
  ▪ 63% of parents said “my child enjoys wearing the implant”.
➢ When families were asked about the spoken language of their children –
  ▪ 53% felt the spoken language had not yet improved in the home
  ▪ 67% stated their children “demonstrated improvement in responding to feelings
    expressed through vocal inflections”
  ▪ 62% shared their children were “entertaining themselves by listening to music,
    watching television, or playing games”.

Findings from the Family Survey emphasized the need for:

➢ Effective strategies for ensuring the implants stay in place for children with motor
  challenges / cerebral palsy.
➢ Support from schools and support centers – families told us
  ▪ “The center does not have knowledge of children with multiple disabilities.”
  ▪ “The school does not encourage wearing of the implant; the teacher seems to think
    it an unnecessary hassle.”
➢ When families were asked about what is needed, they said.
  ▪ “Teachers are interested in supporting the child and family, but they don’t feel they
    have the skills.”
  ▪ “Effective staff training leads to more successful outcomes.”
  ▪ One family’s response – “Since getting a new teacher of the deaf, who is trained in
    auditory training, our son’s progress with his cochlear implant has skyrocketed.”

Information about outcomes for children with deaf-blindness who have received a
cochlear implant is still preliminary. Results of the family survey represent only the opinions and
perceptions of families who did choose to pursue a cochlear implant for their child with deaf-
blindness. Families of children who inquired about an implant for their child with deaf-blindness
but did not proceed were not surveyed. Also, families of children with deaf-blindness who discontinued wearing their cochlear implants were not included in the survey research.

The families of the children who have been implanted and the researchers conducting the cochlear implant studies are discovering many small details that over time may lead to significant conclusions. Findings to date, with children who are deaf-blind and the majority of whom have additional disabilities, reflect even greater variability than research findings for children with only deafness who have received cochlear implants. The presence of cognitive disabilities, physical disabilities, and/or complex health issues, in addition to varying levels of vision impairments, are factors that impact outcomes to varying degrees across the age span of the children included in this initial CIDB study.

Resources


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