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₁ ELSSP

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ELSSP

Introduction

In the United States, 1-2 children are born with hearing loss, per 1,000 births (CDC, 9 2018). This translates to 114,000 Deaf or Hard of Hearing (DHH) children born in the U.S. 10 per year (Martin, Hamilton, Osterman, & Driscoll, 2019). Of these 114,000, ~90\% will be 11 born to hearing parents (Mitchell & Karchmer, n.d.), in a home where spoken language is 12 likely the dominant communication method. Depending on the type and degree of hearing 13 loss and whether the child uses amplification, spoken linguistic input will be partially or totally inaccessible. Some of these children will develop spoken language within the range of 15 their hearing peers (Geers, Mitchell, Warner-Czyz, Wang, & Eisenberg, 2017; Verhaert, 16 Willems, Van Kerschaver, & Desloovere, 2008), but many will face persistent spoken 17 language deficits (???; Luckner & Cooke, 2010; Moeller, Tomblin, Yoshinaga-Itano, Connor, & Jerger, 2007; Sarchet et al., 2014), which may later affect reading ability (???) and academic achievement (Karchmer & Mitchell, 2003; Qi & Mitchell, 2012).

Despite many excellent studies examining language development in DHH children,
there is still a gap in the literature describing and analyzing spoken language development
across the full range of children receiving state services for hearing loss, with many studies
focusing in on specific subgroups (e.g. children under age X with Y level of hearing loss and
Z amplification approach, e.g. (Vohr et al., 2008; Yoshinaga-Itano, Sedey, Wiggin, & Mason,
2018)). In what follows, we first summarize the previous literature on predictors of spoken
language outcomes in DHH children. We then provide a brief overview of a common
vocabulary measure used in the current study, the MacArthur-Bates Communicative
Development Inventory (CDI). Finally, we turn to an empirical analysis of early vocabulary
in a wide range of young children receiving state services in North Carolina. We have two
broad goals in what follows. First, we aim to provide a comprehensive description of a
heterogeneous group of young children who receive state services for hearing loss. Second, we

aim to connect the intervention approaches and child characteristics of this sample with children's vocabulary, with the broader goal of considering the success of early diagnosis and

35 intervention initiatives.

Goldin-Meadow, 2010).

36 Predictors of Language Outcomes

Though the literature points towards spoken language delays and deficits for DHH 37 children, this is a highly variable population with highly variable outcomes (Pisoni, Kronenberger, Harris, & Moberly, 2018). Previous research indicates that gender (Ching et al., 2013; C Kiese-Himmel & Ohlwein, 2002), additional disability (Ching et al., 2013; Verhaert et al., 2008; Yoshinaga-Itano, Sedey, Wiggin, & Chung, 2017), degree and configuration of hearing loss (Ching et al., 2013; de Diego-Lázaro, Restrepo, Sedey, & Yoshinaga-Itano, 2018; Vohr et al., 2011; Yoshinaga-Itano et al., 2017), amplification (Walker et al., 2015), communication (Geers et al., 2017), and early diagnosis/intervention (Yoshinaga-Itano et al., 2017, 2018) predict language outcomes in DHH children. 45 **Gender.** For hearing children, the literature points to a female gender advantage in 46 early language acquisition. Girls speak their first word earlier (Macoby, 1966), have a larger 47 (Bornstein, Hahn, & Haynes, 2004; Fenson et al., 1994; Frank, Braginsky, Yurovsky, & 48 Marchman, 2017) and faster-growing vocabulary (???), and stronger grammatical and phonological skills (Lange, Euler, & Zaretsky, 2016; Özçalışkan & Goldin-Meadow, 2010). This finding appears to be consistent across studies (Wallentin, 2009), various spoken languages (Frank, Braginsky, Marchman, & Yurovsky, 2019), and gesture (Özçalışkan &

The DHH literature presents a more mixed (though rather understudied) picture. On one hand, DHH girls, like hearing girls, have been found to have a larger spoken vocabulary than DHH boys (Ching et al., 2013; C Kiese-Himmel & Ohlwein, 2002). However, in contrast to their hearing peers, DHH children do not seem to show a gender-based difference for some aspects of syntactic development (???).

Comorbidities. Additional co-occurring disabilities occur frequently in the DHH population, perhaps as much as three times more than in the hearing population (???).

Incidence estimates for co-occurring disabilities in DHH children range from 25-51% (???; ???; ???; Guardino, 2008; Luckner & Carter, 2001; Picard, 2004; Schildroth & Hotto, 1996; Soukup & Feinstein, 2007), with approximately 8% children living with 2 or more co-occurring disabilities (Schildroth & Hotto, 1996).

Some of these conditions, particularly those which carry risk of developmental delay

(e.g., Down syndrome), result in language delays independent of hearing loss (???; Chapman,

1997; Weismer, Lord, & Esler, 2010), with cognitive ability more predictive of language

outcomes than presence or absence of a specific disability (Meinzen-Derr, Wiley, Grether, &

Choo, 2011; Sarant, Holt, Dowell, Richards, & Blamey, 2008). Disability and hearing loss

likely each contribute to a given child's language development (Ching et al., 2013; Rajput,

Brown, & Bamiou, 2003, @vannierop2016), with differential effects of each (Vesseur et al.,

2016). In some cases, additional disabilities appear to interact with hearing loss to intensify

developmental delays (Birman, Elliott, & Gibson, 2012; Pierson et al., 2007).

Furthermore, incidence of hearing loss is higher among children born premature

(defined as 37 weeks gestational age). Compared to an incidence 0.2% in full-term infants,

incidence of hearing loss in extremely premature infants (defined as 33 weeks gestational age)

ranges 2–11%, with increased prematurity associated with increased rates of hearing loss

(???).

Independently of hearing status, prematurity is linked to increased risk of language
delay and disorder (Barre, Morgan, Doyle, & Anderson, 2011; Carter & Msall, 2017; Cusson,
2003; Rechia, Oliveira, Crestani, Biaggio, & de Souza, 2016; Van Noort-van Der Spek,
Franken, & Weisglas-Kuperus, 2012; Vohr, 2014). Unfortunately, research on language
development in premature DHH children is scant (Vohr, 2016), so it remains unclear how
hearing loss and prematurity may interact within spoken language skills. One study of

premature infants finds that auditory brainstem response during newborn hearing screening predicts language performance on the PLS-4 at age 3 (Amin, Vogler-Elias, Orlando, & Wang, 2014), suggesting a link between prematurity and hearing loss in early childhood, though 87 further research is needed in this domain. 88

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In extremely premature DHH children, incidence of additional disabilities may be as

high as 73% (Robertson, Howarth, Bork, & Dinu, 2009). Indeed, pre-term infants with comorbidities have been found to be more likely to also have hearing loss than those without comorbidities (Schmidt et al., 2003), further complicating language development for this population. Audiological Characteristics. Hearing loss varies in severity, ranging from slight 94 to profound (Clark, 1981). More severe hearing loss (less access to spoken language) 95 typically results in more difficulty with spoken language in infancy (Vohr et al., 2008), early 96 childhood (???; Ching et al., 2010, 2013; Sarant et al., 2008; Tomblin et al., 2015) and 97 school-age (Wake, Hughes, Poulakis, Collins, & Rickards, 2004). Although profound hearing 98 loss is associated with more pronounced spoken language difficulty, even mild to moderate hearing loss is associated with elevated risk of language disorders (Blair, Peterson, & 100 Viehweg, n.d.; Delage & Tuller, 2007). Hearing loss also varies in whether it affects one ear 101 or both. Bilateral hearing assists speech perception, sound localization, and loudness 102 perception in quiet and noisy environments (Ching, Van Wanrooy, & Dillon, 2007). The 103 literature on hearing aids and cochlear implants points to benefits for bilateral auditory 104 input (Lovett, Kitterick, Hewitt, & Summerfield, 2010; Sarant, Harris, Bennet, & Bant, 2014; 105 Smulders et al., 2016). At school-age, 3-6% of children have unilateral hearing loss (Ross, Visser, Holstrum, Qin, & Kenneson, 2010). Although children with unilateral hearing loss 107 have one "good ear," even mild unilateral hearing loss has been tied to higher risk of language delays and educational challenges relative to hearing children (Lieu, 2004, 2013; 109 Lieu, Tye-Murray, & Fu, 2012, @kiese-himmel2002; Vila & Lieu, 2015). That is, just as in 110 the bilateral case, more severe hearing loss leads to greater deficits in language and

educational outcomes for children with unilateral hearing loss (Anne, Lieu, & Cohen, 2017; 112 Lieu, 2013). Many DHH children receive hearing aids (HAs) or cochlear implants (CIs) to 113 boost access to the aural world. These devices have been associated with better speech 114 perception and spoken language outcomes (Niparko et al., 2010; Walker et al., 2015; 115 Waltzman et al., 1997). In turn, aided audibility predicts lexical abilities with children in 116 HAs (Stiles, Bentler, & McGregor, 2012). For both hearing aids and cochlear implants, 117 earlier fit leads to better spoken language skills, if the amplification is effective. For hearing 118 aids, some studies find that children with milder hearing loss who receive hearing aids earlier 119 have better early language achievement than children who are fit later (Tomblin et al., 2015), 120 but this finding does not hold for children with severe to profound hearing loss (Christiane 121 Kiese-Himmel, 2002; Watkin et al., 2007) (for whom hearing aids are generally ineffective). 122 Analogously, children who are eligible and receive cochlear implants earlier have better speech perception and spoken language outcomes than those implanted later (???; Dettman, Pinder, Briggs, Dowell, & Leigh, 2007; Miyamoto, Hay-McCutcheon, Kirk, Houston, & Bergeson-Dana, 2008; Svirsky, Teoh, & Neuburger, 2004, @yoshinaga-itano2018), with best 126 outcomes for children receiving implants before their first birthday (Dettman et al., 2007). 127 Communication. Total Communication (TC) refers to communication that 128 combines speech, gesture, and elements of sign (but not a full sign language, such as 129 American Sign Language), sometimes simultaneously. Clinicians currently employ TC as an 130 alternative or augmentative communication method for children with a wide range of 131 disabilities (Branson & Demchak, 2009; Gibbs & Carswell, 1991; Mirenda, 2003). 132

Compared to total communication, DHH children using an exclusively oral approach have better speech intelligibility (Dillon, Burkholder, Cleary, & Pisoni, 2004; Geers et al., 2017; Geers, Spehar, & Sedey, 2002; Hodges, Dolan Ash, Balkany, Schloffman, & Butts, 1999) and auditory perception (???; Geers et al., 2017). That said, there is some debate as to whether an oral approach facilitates higher spoken language performance, or whether children who demonstrate aptitude for spoken language are steered towards the oral

approach rather than TC (Hall, Hall, & Caselli, 2017).

1-3-6 Guidelines. Early identification (Apuzzo & Yoshinaga-Itano, 1995; Kennedy
et al., 2006; Robinshaw, 1995; White & White, 1987; Yoshinaga-Itano, Sedey, Coulter, &
Mehl, 1998; Yoshinaga-Itano et al., 2018) and timely enrollment in early intervention
programs (Ching et al., 2013; Holzinger, Fellinger, & Beitel, 2011; Vohr et al., 2008,

@watkin2007, 2011) are associated with better language proficiency. Indeed, DHH children
who receive prompt diagnosis and early access to services have been found to meet
age-appropriate developmental outcomes, including language (Stika et al., 2015).

In line with these findings, the American Academy of Pediatricians (AAP) has set an initiative for Early Hearing Detection and Intervention (EHDI). Their EHDI guidelines recommend that DHH children are screened by 1 month old, diagnosed by 3 months old, and enter early intervention services by 6 months old. We refer to this guideline as 1-3-6.

Meeting this standard appears to improve spoken language outcomes for children with HL (Yoshinaga-Itano et al., 2017, 2018) and the benefits appear consistent across a range of demographic characteristics.

At a federal level in the U.S., the Early Hearing Detection and Intervention Act of 154 2010 (Capps, 2009) was passed to develop state-wide systems for screening, evaluation, diagnosis, and "appropriate education, audiological, medical interventions for children 156 identified with hearing loss," but policies for early diagnosis and intervention vary by state. As of 2011, 36 states (including North Carolina, ("15A NCAC 21F .1201 - .1204," 2000)] 158 mandate universal newborn hearing screening [national conference of state legislatures, 2011] 159 FIX CITE. All states have some form of early intervention programs that children with 160 hearing loss can access (NAD, n.d.), but these also vary state-by-state. For instance, half of 161 the states in the US do not consider mild hearing loss an eligibility criterion for early 162 intervention (Holstrum, Gaffney, Gravel, Oyler, & Ross, 2008). 163

In evaluating the success of this initiative, the AAP (EHDI, n.d.) finds that about 70%

of US children who fail their newborn hearing screening test are diagnosed with hearing loss before 3 months old, and that 67% of those diagnosed (46% of those that fail newborn hearing screening) begin early intervention services by 6 months old. These findings suggest that there may be breaks in the chain from screening to diagnosis and from diagnosis to intervention, and the effect may be further delays in language development for children not meeting these guidelines.

171 Quantifying vocabulary growth in DHH children

The MacArthur Bates Communicative Development Inventory (CDI, Fenson et al. 172 (1994)] is a parent-report instrument that gathers information about children's vocabulary 173 development. The Words and Gestures version of the form (CDI-WG) is normed for 174 8-18-month-olds, and includes 398 vocabulary items that parents indicate whether their child 175 understands or produces, along with questions about young children's early communicative 176 milestones. The Words and Sentences version of the form (CDI-WS) is normed for 177 16-30-month-olds, and includes 680 vocabulary items that parents indicate whether their 178 child produces, along with some questions about grammatical development. The CDI has 179 been normed on a large set of participants across many languages (Anderson & Reilly, 2002; 180 Frank et al., 2017; Jackson-Maldonado et al., 2003). 181

The CDI has also been validated for DHH children with cochlear implants (Thal, 182 Desjardin, & Eisenberg, 2007). More specifically, in this validation, researchers asked parents 183 to complete the CDI, administered the Reynell Developmental Language Scales, and 184 collected a spontaneous speech sample. All comparisons between the CDI and the other 185 measures yielded significant correlations ranging from 0.58 to 0.93. Critically, the children in this study were above the normed age range for the CDI, and thus this validation helps to 187 confirm that the CDI is a valid measurement tool for older DHH children. In further work, 188 Castellanos, Pisoni, Kronenberger, and Beer (2016) finds that in children with CIs, number 189 of words produced on the CDI predicts language, executive function, and academic skills up 190

to 16 years later. Building on this work, several studies have used the CDI to measure vocabulary development in DHH children [Ching et al. (2013); Yoshinaga-Itano et al. (2017); Yoshinaga-Itano et al. (2018); de Diego-Lázaro et al. (2018); Vohr et al. (2008); Vohr et al. (2011); summarized in table XXX].

Table 1
Summary of findings of CDI studies in DHH children

Study	Population	Gender	1-3-6	Laterality	Degree	Amplification	Communication	Comorbidities
Ching et al., 2013	3 year old children receiving services in Australia	Female +	Did not study	Did not study	More severe -	No effect	No effect	Comorbidities -
Yoshinaga-Itano et al., 2017	8-39 month children with bilateral hearing loss	No effect	1-3-6 +	Did not study	More severe -	Did not study	Did not study	Comorbidities -
Yoshinaga-Itano et al., 2018	Children with cochlear implants	Did not study	1-3-6 +	Did not study	Did not study	Earlier CI activation +	Did not study	Did not study
De Diego-Lazaro et al., 2018	Spanish speaking children with bilateral hearing loss	No effect	Earlier intervention +	Did not study	Milder +	More functional hearing +	Did not study	Did not study
Vohr et al., 2011	18-24 month olds with hearing loss	Did not study	Earlier intervention +	Did not study	Milder +	Did not study	Did not study	NICU stay -; Comorbidities -

 $^{^{\}rm a}$ + equals bigger vocab, - equals smaller vocab

95 Goals and Predictions

This study aims to 1) characterize the demographic, audiological, and intervention
variability in the population of DHH children receiving state services for hearing loss; 2)
identify predictors of vocabulary delays; and 3) evaluate the success of early identification
and intervention efforts at a state level. We include two subgroups of DHH children
traditionally excluded from studies of language development: children with additional
disabilities and children with unilateral hearing loss (e.g., Yoshinaga-Itano et al., 2018).

For the first and third goal above, we did not have specific hypotheses and sought to provide descriptive information about a diverse sample of DHH children receiving state services. For the second, we hypothesized that male gender, more severe degree of hearing loss, bilateral hearing loss, no amplification use, prematurity, and presence of additional disabilities would predict larger spoken vocabulary delay. We did not have strong predictions regarding communication method, language background, or presence of other health issues (e.g., congenital heart malformation).

209 Methods

Clinical evaluations were obtained through an ongoing collaboration with the North
Carolina Early Language Sensory Support Program (ELSSP), an early intervention program
serving children with sensory impairments from birth to 36 months. ELSSP passed along
deidentified evaluations to our team after obtaining consent to do so from each family. No
eligibility criteria beyond hearing loss and receiving an ELSSP evaluation were imposed,
given our goal of characterizing the full range of DHH children with hearing loss in North
Carolina.

The clinical evaluations included demographic and audiological information, CDI vocabulary scores, and the results of any clinical assessments administered (e.g., PPVT), all detailed further below. For some children (n=43), multiple evaluations were available from different timepoints. In these cases, only the first evaluation was considered for this study due to concerns regarding within-subjects variance for statistical analysis.

While this collaboration is ongoing, we opted to pause for this analysis upon receiving 222 data from 100 children. Thus, the reported sample below consists of 98 children (55 male/43 223 female) ages 4.20–36.17(M=21.60, SD=9.01). Race and SES information was not available. 224 Families were administered either the WG or WS version of the CDI based on clinician 225 judgement. Children who were too old for WG, but who were not producing many words at 226 the time of assessment, were often given WG (n=37). Families for whom Spanish was the 227 primary language (n = 14) completed the Spanish version of the CDI (Jackson-Maldonado et 228 al., 2003). 229

Table 2

CDI details

CDI version	Average Age (SD)	Average Comprehension (SD)	Average Production (SD)	% Developmental Delays
WG (n=73)	20.07 (8.87) months	107 (99.9) words	32 (53.7) words	17.81%
WS (n=25)	26.05 (8) months	NA	132 (172.7) words	4%

Children in this sample were coded as yes/no for cognitive development concerns (e.g.,
Down syndrome, global developmental delays; Cornelia de Lange syndrome), yes/no for
prematurity (i.e., more than 3 weeks premature), yes/no for health issues (e.g., heart defects,
kidney malformations, VACTERL association), and yes/no for vision loss (not corrected to
normal by surgery or glasses)

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Condition	Specific Condition	
Premature		17
	Extremely Premature	10
	NICU stay	16
Health Issues		34
	Heart	10
	Lung	6
	Illness	15
	Feeding Issues	14
	Pregnancy/Birth Complications	11
	Musculoskeletal	10
	Cleft Lip/Palate	4
	Other	15
Developmental Concerns		16
	Down Syndrome	5
	Chromosomal Issues	3
	Neural Tube Defects	3
	Other	15
Vision Loss		5
	Retinopathy of Prematurity	2
	Nearsightedness	2

 $\ensuremath{\mathsf{end}} \{ \mathsf{table} \}$

Degree of hearing loss was most often reported with a written description (e.g., "mild sloping 240 to moderate" or "profound high frequency loss"). We created 3 variables: hearing loss in the 241 better ear, hearing loss in the worse ear, and average hearing loss (average of better and 242 worse ear). Using the ASHA hearing loss guidelines, each of these was coded with a dB HL 243 value corresponding with the median dB HL for the level of hearing loss (e.g., moderate 244 hearing loss was coded as 48dB HL), and sloping hearing loss was coded as the average of 245 the levels (e.g. mild to moderate was coded as 40.5 dB HL). Participants were also coded for 246 unilateral or bilateral hearing loss; presence or absence of Auditory Neuropathy Spectrum 247 Disorder; sensorineural, conditive, or mixed hearing loss. Amplification was recorded as the 248 device the child used at the time of assessment-either hearing aid, cochlear implant, or none. 249

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\tab:amp_info)Audiological Characteristics of the Sample for Unilateral /
Bilateral Hearing Loss}

	n	Average HL (better ear)	Average HL (worse ear)	Average Age at Amplification
Hearing Aid (n=53)	11 / 42	3.89 / 48.49 dB	58.54 / 56.09 dB	9.73 / 8.79 months
Cochlear Implant (n=17)	0 / 17	NA / 85.6 dB	NA / 89.79 dB	NA / 14.12 months
No Amplification (n=26)	14 / 12	2.5 / 49.67 dB	73.9 / 53.65 dB	NA
Total (n=98)	25 / 71	3.11 / 57.84 dB	67.57 / 63.97 dB	NA

 $^{^{\}rm a}$ N.B. Age Amplification for children with CIs represents age at implantation $\label{eq:condition} \\ \label{eq:condition} \\ \label{eq:condition}$

Communication method was recorded as spoken language, total communication, or cued speech. One participant had a parent fluent in sign language, but the reported communication method in the home was total communication. No child in our sample used sign language. Participants were also coded as monolingual or multilingual based on whether families reported using more than one language at home. Total communication was not

counted as multilingualism.

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\caption{(#tab:comm_info)Language and communication characteristics of the sample}

Communication Method	English	Spanish	Hindi
Spoken Language (n=79)	68	10	1
Total Communication (n=16)	13	3	0
Cued Speech (n=1)	1	0	0

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Age at screening was measured as the child's age in months at their first hearing screening. Age at screening was only available for (XXX) participants. If participants received their newborn hearing screening, age at screening was recorded as 0 (months). Age at diagnosis was taken as the age in months when children received their first hearing loss diagnosis. All children were enrolled in birth-to-three early intervention services through NC ELSSP, and the date of enrollment was listed on the clinician evaluation. From the clinician report, we calculated the number of hours of early intervention services received per month (including service coordination, speech therapy, and occupational therapy, among others). Because of the sparse data on screening age, if participants had an age at diagnosis <= 3 mo. and an age of intervention <= 6 mo., they were recorded as meeting 1-3-6. It is possible that a participant did not receive screening by 1 month, but did receive diagnosis by 3 months and services by 6 months. This special case would be coded as meeting 1-3-6 by our criteria.

 \begin{table}

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Diagnosis by 3 months	$\boxed{72.04\%}$
Average Age Diagnosis (SD)	4.11 (7.34) months
Intervention by 6 months	37.89%
Average Age Intervention (SD)	11.32 (8.75) months
Meets 1-3-6	36.56%

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Variable	Scale	Range		
Age Continuous		4.2-36.16666667 months		
Age at Amplification	Continuous	2-31 months		
Age at Diagnosis	Continuous	0-30 months		
Age at Implantation	Continuous	7-32 months		
Age at Intervention	Continuous	1-33 months		
Amplification	Categorical	Hearing Aid / Cochlear Implant / None		
Communication	Categorical	Spoken / Total Communication / Cued Speech		
Degree Hearing Loss (worse ear)	Continuous	17.75-100 dB HL		
Developmental Delay	Categorical	Yes / No		
Gender	Categorical	Female / Male		
Health Issues	Categorical	Yes / No		
Language in Home	Categorical	English / Other		
Laterality	Categorical	Unilateral / Bilateral		
Meets 1-3-6	Categorical	Yes / No		
Prematurity	Categorical	Full-term / Premature		
Services Received Per Month	Continuous	0-43 services / month		
Type of Hearing Loss	Categorical	Sensorineural / Conductive / Mixed		
CDI - Words Produced Continuo		0-635 words		
\end{table}				

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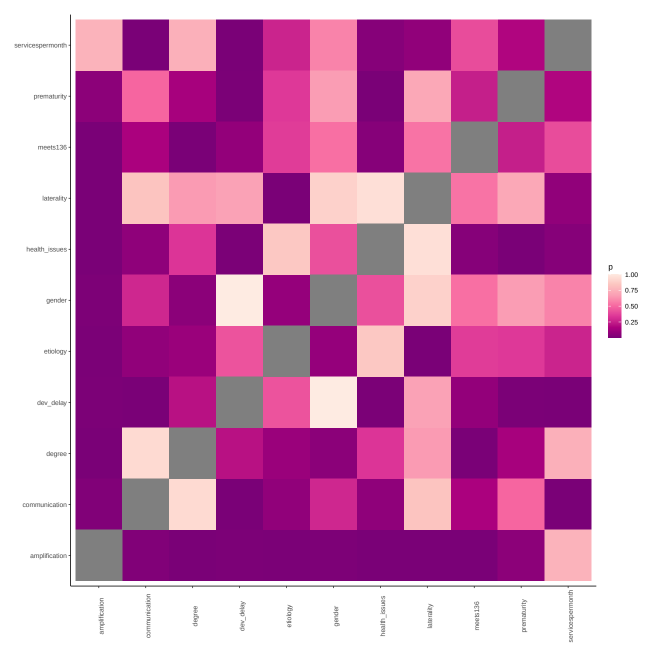
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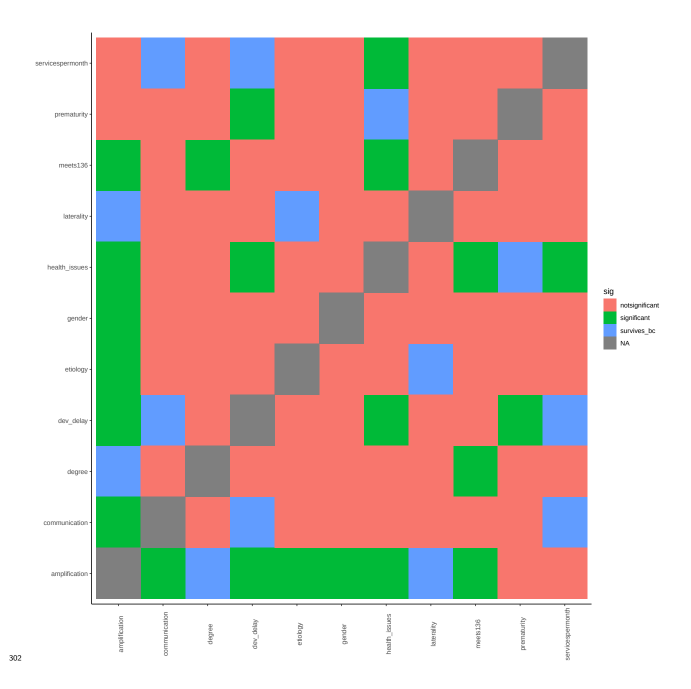
Results

All analyses were conducted in R. All code is available on Github. In the first section, we explore relationships among child demographic, audiological, and clinical variables. In the second section, we examine the influence of these factors on vocabulary development. In the third section, we describe the implementation of the EHDI 1-3-6 guidelines and predictors of early diagnosis and intervention.

Part I: Interactions Among Variables

Shapiro–Wilk tests revealed that all of our continuous measures (i.e. degree of hearing loss, services received per month, vocabulary delay) significantly differed from a normal distribution (ps <.05), so we used nonparametric tests to explore relationships among our variables. For categorical-categorical relationships, we used chi square tests; for continuous-categorical tests, we used mann-whitney U tests (2 levels for categorical variable) or kruskal-wallis tests (>2 levels for categorical variable; for continuous-continuous relationships, we used Of the fifty-five combinations of variables, p < .05 for sixteen, and seven survived bonferroni correction (p < 0.00). The full set of comparisons is shown in figure XXX.





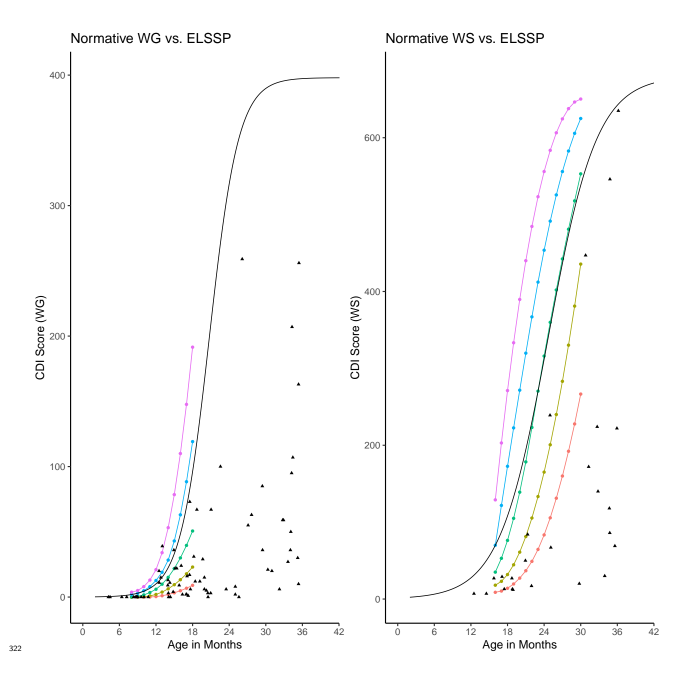
From this analysis, we found that children born premature were more likely to also have health issues (X2 (1, N = 96) = 25.69, p = 0.00.) Children with conductive hearing loss were more likely to have unilateral hearing loss (X2 (2, N = 86) = 14.84, p = 0.00). Children with unilateral hearing loss were unlikely to receive a cochlear implant and more likely to use no amplification (X2 (2, N = 96) = 17.19, p = 0.00). Children with more severe hearing loss were more likely to use a cochlear implant than children with milder hearing loss

 309 (H(2)=23.80, p=0.00). Children with developmental delays received more services per month than typically developing DHH children (H()=134.50, p=0.00)and were more likely to use total communication (X2 (2, N = 96)) = 19.38, p = 19.38). Children who used total communication received more services per month than children using spoken language (H(1)=15.60, p=0.00).

Part II: Influence on vocabulary

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We first constructed a binary logistic growth curve for vocabulary from the 50th percentile
data for typically developing children from Wordbank. With this function, each participant's
CDI score yielded a predicted age from the normative data. For each child, we subtracted
this predicted age (given the score) from the child's actual age to give us a measure of delay
in months. Descriptively, we found widespread vocabulary delays on both Words and
Gestures and Words and Sentences, with the majority of DHH children testing around or
below the 25th percentile for hearing children.



We next explored the effect of the different audiological, demographic, and intervention characteristics on vocabulary delay. Vocabulary delay did not meet the assumption of normality, so we used non-parametric tests for the following set of analyses.

Mann-Whitney-Wilcoxen tests were conducted to examine the effects of gender, laterality, developmental delay, health issues, prematurity, meeting 1-3-6 guidelines, and communication on vocabulary delay. We used kruskal-wallis tests for amplification and etiology, and Kendall's rank correlations for degree of hearing loss (worse ear) and services

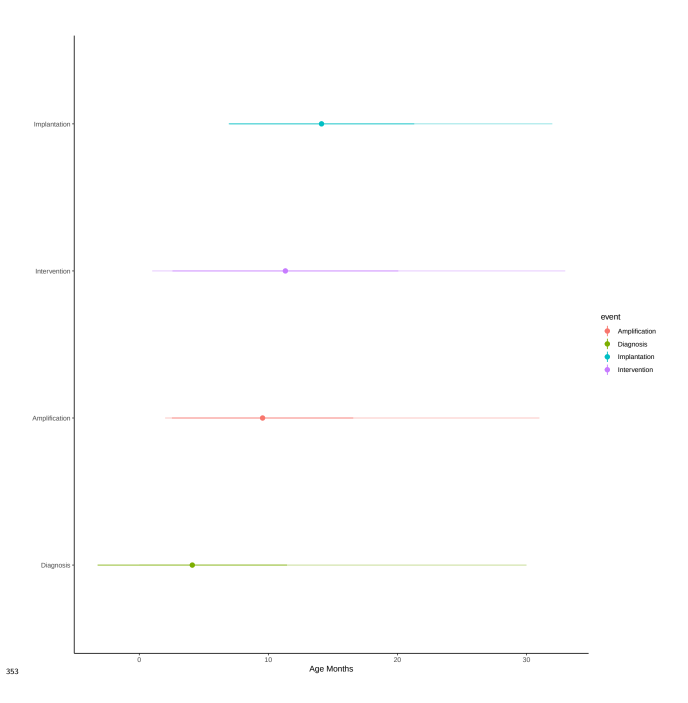
received per month. These results are exploratory and descriptive, and their interpretation should be tempered accordingly.

Boys were significantly more delayed than girls on Words and Sentences but not Words and 332 Gestures. Children with developmental delays had larger vocabulary delays than children 333 without developmental delays on Words and Gestures. Because only one child with a 334 developmental delay took the Words and Sentences form, we did not perform the analysis for 335 Words and Sentences. Premature children and children with health issues had smaller 336 vocabularies than typically developing children on Words and Gestures but not Words and Sentences. Children who met 1-3-6 guidelines had larger vocabulary than children who did 338 not on Words and Gestures but not Words and Sentences. On Words and Gestures but not Words and Sentences, receiving more early intervention services was correlated with lower vocabulary. We did not observe an effect of laterality, communication, degree, or etiology on 341 vocabulary delay on either form of the CDI. For communication, we omitted cued speech 342 from the analysis because only one child in our sample used this method of communication 343 (shown on graph anyway for the curious). A kruskal-wallis test showed a significant effect of 344 amplification on vocabulary delay on Words and Gestures, such that children with no 345 amplification were more delayed than children without amplification. 346

Part III: Meets136 success

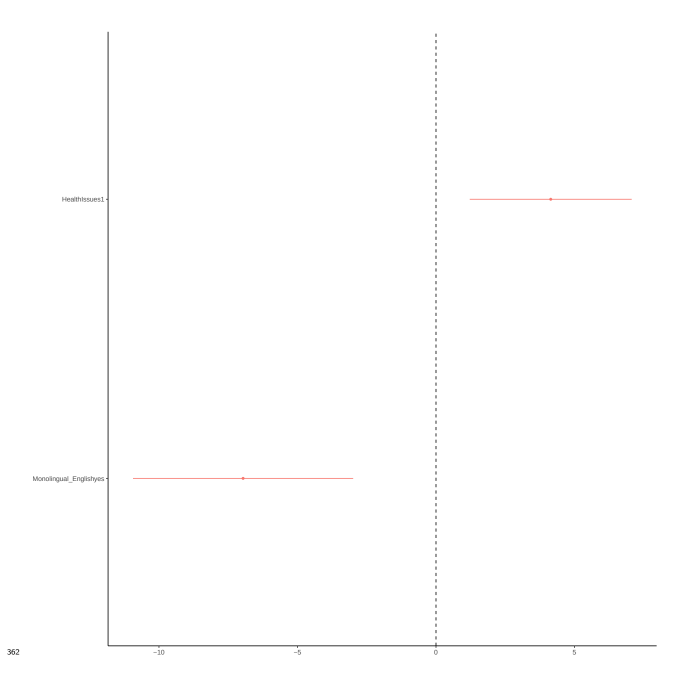
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Lastly, we looked at the ages at which children received diagnosis and intervention, and how
this mapped onto the 1-3-6 guidelines. 36.56% of our sample met 1-3-6 guidelines for early
diagnosis and intervention. Of children with comorbidities (developmental concerns,
prematurity, health issues), only 15% met 1-3-6 guidelines, compared to 40% of typically
developing children. See table XXX for complete breakdown.



We created linear regression models for age at diagnosis and age at intervention. Models were paired down using stepwise regression by AIC using the stepAIC function (cite MASS package). For age at diagnosis, we included the set of child-specific factors that would be relevant before diagnosis of hearing loss. We began with: Age diagnosis \sim gender + laterality + degree (worse ear) + developmental delay + health issues + prematurity + laterality + language background + etiology The best fit model (R2=0.19 , p=0.00)included health

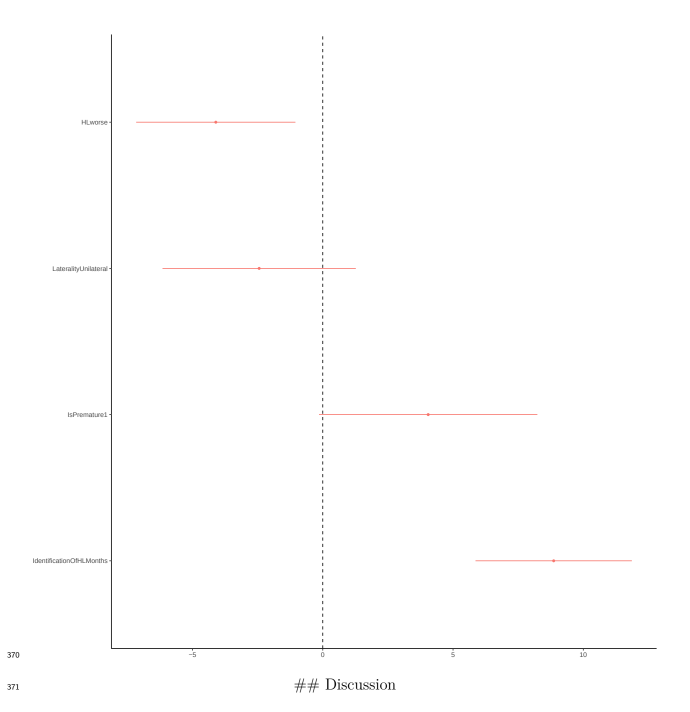
issues (beta weight = 4.14, p = 0.01) and language background (beta weight = -6.96, p = 0.00). Age diagnosis \sim health issues + language background



For age at intervention, we first included the variables potentially relevant prior to intervention: Age intervention \sim gender + degree (worse ear) + developmental delay + health issues + prematurity + laterality + language background + etiology + age diagnosis

The best fit model (R2=0.41, p=0.00) included prematurity (beta weight = 4.05, p = 0.06),

laterality (beta weight = -2.44, p = 0.19), degree of hearing loss (beta weight = 4.14, p = 0.01), and age at diagnosis (beta weight = 0.62, p = 0.00). Age intervention \sim laterality + degree (worse ear) + prematurity + laterality + age diagnosis



372 Conclusion

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Footnotes: Despite exciting, increasing, and converging evidence for benefits of early sign language exposure (e.g., Schick, De Villiers, De Villiers, & Hoffmeister, 2007; ???; Clark et al., 2016; Hrastinski & Wilbur, 2016; Magnuson, 2000; Spencer, 1993), the majority of DHH children will not be raised in a sign language environment. This is particularly true for North Carolina, which does not have a large community of sign language users, relative to states like Maryland or areas like Washington D.C. or Rochester, NY. For this reason, we focus on spoken language development.

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