

1 Exploring variation in infants' preference for infant-directed speech: Evidence from a
2 multi-site study in Africa

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37

Abstract

38 Infants show a preference for infant-directed speech (IDS) over adult-directed speech
39 (ADS). This preference has been linked to infants' language processing and word learning
40 in experimental settings, and also correlates with later language outcomes. Recently, the
41 cross-cultural consistency of infants' IDS preference has been confirmed by large-scale,
42 multisite replication studies, but conclusions from these studies were primarily based on
43 participants from North America and Europe. The current study addressed this sampling
44 bias via a large-scale, multisite study of infants (3-15 months) across communities in
45 Africa. We investigated whether participants showed a preference for IDS over ADS, and if
46 so, whether the magnitude of their preference differs from effects documented in other
47 populations of infants. Across six sites (total N = 200), we observed a preference for IDS
48 over ADS ($\hat{\beta}_{IDS \text{ vs. } ADS} = 0.06$), suggesting that infants look on average 6% longer on the
49 IDS trials than the ADS trials. There was no significant difference between African infants
50 in this study and a method-matched subsample of infants from prior studies of IDS
51 preference. This study provides new evidence on the generalizability of IDS preference and
52 looking-time methods more broadly, while also highlighting some of the challenges of global
53 big team science.

54 *Keywords:* infant-directed speech; reproducibility; Africa; infants; generalizability

55 Word count: 9412

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57 multi-site study in Africa

58 Adults often speak to infants differently than to other adults, using a speech register
59 known as infant-directed speech (IDS). Infant-directed speech tends to have exaggerated
60 prosodic characteristics, including higher pitch, greater pitch variation, longer pauses,
61 simplified grammatical structure, and shorter and slower utterances as compared to
62 adult-directed speech (ADS; e.g. Fernald et al., 1989; Trainor & Desjardins, 2002). Even
63 very young infants from a variety of language backgrounds have a preference for listening
64 to IDS over ADS (Cooper, Abraham, Berman, & Staska, 1997; Cooper & Aslin, 1994;
65 Fernald, 1985; Hayashi, Tamekawa, & Kiritani, 2001; Kitamura & Lam, 2009; Newman &
66 Hussain, 2006; Pegg, Werker, & McLeod, 1992; Santesso, Schmidt, & Trainor, 2007; Singh,
67 Morgan, & Best, 2002; Werker & McLeod, 1989). Infants' preference for IDS over ADS has
68 also been demonstrated in a meta-analysis; across 34 studies, IDS preference had a fairly
69 large average effect size with a value of Cohen's d 0.72 (Dunst, Gorman, & Hamby, 2012).

70 Why do infants prefer IDS? Perhaps IDS is intrinsically salient to infants because of
71 its perceptual characteristics (e.g., higher pitch, greater pitch variability). Or perhaps, as
72 infants are exposed to IDS, familiarity leads to preference. These explanations have
73 different developmental predictions: while the intrinsic view would suggest an early
74 preference (e.g., Cooper & Aslin, 1990), the exposure account would predict developmental
75 increases in preference. Further, these explanations are not mutually exclusive: infants'
76 early preference for IDS may motivate their parents to use more IDS, which in turn could
77 lead infants to show a stronger IDS preference. Regardless of its origins, infants' preference
78 for IDS may benefit their early language development. For example, in experimental
79 studies, infants can segment words better in fluent speech produced in IDS than ADS
80 (Thiessen, Hill, & Saffran, 2005), show better recognition of words introduced in IDS after
81 a 24-hour delay (Singh, Nestor, Parikh, & Yull, 2009), and more successfully learn words

82 from IDS than ADS (Graf Estes & Hurley, 2013).

83 Further evidence comes from correlational studies, which have found that the amount
84 of IDS in the language environment is positively related to children's language outcomes,
85 such as vocabulary size (e.g., Ramírez-Esparza, García-Sierra, & Kuhl, 2014; L. A.
86 Shneidman & Goldin-Meadow, 2012; L. Shneidman, Arroyo, Levine, & Goldin-Meadow,
87 2013; but cf., Casillas, Brown, & Levinson, 2020; Casillas, Brown, & Levinson, 2021, who
88 found similar timing of language development milestones even in a population that hears
89 very limited IDS). Together, this work suggests that infants' preference for IDS over ADS
90 may support their language development, which explains why infants' IDS preference
91 continues to be an important topic in the literature on early childhood.

92 However, it is important to note that almost all prior studies, including the
93 meta-analysis by Dunst et al. (2012) have included mainly infants learning English in
94 Western, educated, industrialized, rich, and democratic (WEIRD) societies (Henrich,
95 Heine, & Norenzayan, 2010), with only a few studies extended to non-Western infant
96 populations learning languages other than English (Hayashi et al., 2001; Werker, Pegg, &
97 McLeod, 1994). As such, there is a large sampling bias in the existing data about infants'
98 preference for IDS, as in many other research topics in developmental psychology (see
99 Nielsen, Haun, Kärtner, & Legare, 2017). This sampling bias is a problem for generalizing
100 findings about infants' IDS preference to infants growing up in different cultures and
101 learning different languages. In light of this generalizability issue – as well as the recent
102 replication crisis in psychology (e.g., Open Science Collaboration, 2015) – infant
103 researchers have begun to collaborate on large-scale, multi-site studies to replicate key
104 developmental findings (Frank et al., 2017).

105 One of these multi-site projects investigated infants' preference for IDS over ADS: the
106 ManyBabies1 study (MB1; ManyBabies Consortium, 2020). MB1 collected monolingual
107 data from 67 laboratories, with a total sample of 2329 monolingual infants 3 – 15 months

108 old. The protocol for this experiment was simple: infants listened to alternating audio clips
109 of IDS and ADS while viewing an uninformative visual stimulus (a colored checkerboard).
110 Their looking time was measured over the course of up to 16 trials, 18s each in length (8
111 IDS and 8 ADS). Notably, all participants in the study listened to stimuli that were
112 constructed from naturalistic speech by North American mothers (speaking either to
113 another adult or to their own infant). The mismatch between the stimuli and the native
114 language of many infants in the study allowed inferences about native language effects and
115 also minimized variability due to differences in the stimuli (a follow-up project now in
116 progress seeks to measure native-language preferences in a subset of MB1 labs). Overall,
117 older infants showed a stronger preference for IDS than younger infants. There was also an
118 effect of infants' language backgrounds: North American infants exhibited a stronger IDS
119 preference than infants who were not exposed to North American English (NAE).
120 Although infants' ages and language backgrounds affected the magnitude of IDS
121 preference, essentially all groups of infants preferred NAE IDS over ADS.

122 Despite the breadth of its sample relative to previous work, the MB1 study still
123 constitutes a biased sample of infant populations in the world. Most of the data in MB1
124 were contributed by laboratories in economically-advantaged areas, accessing relatively
125 high socio-economic status participant populations. Further, although this large-scale
126 study had a diverse sample from 17 countries, 60 out of the 67 participating laboratories
127 were from Europe and North America, only a handful of laboratories were from Australia
128 and Asia, and none were from Africa or South America. Thus, the sample studied in MB1
129 came almost exclusively from Western, educated, affluent populations who heard
130 Indo-European languages, limiting the generalizability of the findings to infants growing up
131 in other cultural and linguistic contexts. This lack of evidence on generalizability of a key
132 finding about infants' preference restricts our ability to build robust developmental theories
133 of language learning across cultural contexts. Our current study takes a step towards
134 addressing this gap.

We investigate whether infants growing up in a variety of African cultures show an IDS preference, using the paradigm developed by the MB1 study. Our study has both a theoretical goal and a practical goal. Theoretically, we are interested in whether IDS preference is a culturally and linguistically invariant developmental pattern (Nielsen et al., 2017). The inclusion of infants across many African cultures (who are acquiring many different languages, see Table 1) provides an important test of generalizability of the IDS preference. Practically, increasing sample diversity also promotes diversity among researchers engaged in developmental science and hopefully increasing exchanges between researchers across cultures. Thus, one goal of our study is building research networks to facilitate further studies with the communities represented in the current study.

Our study builds on a foundation of prior descriptive work investigating the generality of IDS across cultures. Although this work has investigated a variety of different cultures and languages, it can be (and often is) crudely summarized via the distinction between WEIRD and non-WEIRD cultures discussed above. We follow this convention here without endorsing this distinction as necessarily being meaningful in the context of our study, as IDS in WEIRD and non-WEIRD cultures shares similar prosodic properties. For example, Broesch and Bryant (2015) reported that IDS produced by North-American mothers, as well as by Kenyan and Fijian mothers, is produced with higher pitch, greater pitch variation, and is spoken at a slower rate than ADS. This finding is consistent with past work reporting that IDS shares some common exaggerated prosodic features (e.g., higher pitch, larger pitch variation) across diverse languages, which include French, Italian, German, Japanese, British English, American English (Fernald et al., 1989), Mandarin Chinese (Grieser & Kuhl, 1988), Thai, Australian English (Kitamura, Thanavishuth, Burnham, & Luksameeyanawin, 2001), Arabic (Farran, Lee, Yoo, & Oller, 2016).

IDS can also be recognized as being infant-directed by listeners from non-WEIRD cultures. Bryant, Liénard, and Clark Barrett (2012) reported that Turkana adults in Kenya can discriminate between NAE IDS and ADS (see similar results in Bryant &

162 Barrett, 2007 for Shuar hunter horticulturists from Amazonian Ecuador). These studies are
163 consistent with findings from the MB1 studies showing that children who are not learning
164 NAE, including children from Singapore and Korea, nonetheless show a preference for NAE
165 IDS over ADS. Taken together, the common acoustic properties of IDS across different
166 languages, and how NAE IDS can be recognized by non-native participants, raise the
167 possibility of infants' IDS preference over ADS being quite consistent across different
168 cultures and languages. However, it is possible that the strength of this preference would
169 nonetheless be influenced by similarity between the test language (English) and the
170 language(s) that each infant is learning, which could bolster the measured preferences to
171 the extent that test and native language are similar (as in the case of infants learning other
172 Indo-European languages with similar phonetic and acoustic properties). If this is the case,
173 we expect that phylogenetic similarity between Indo-European languages and our stimuli
174 would lead to comparable or stronger observed IDS preferences in samples of infants
175 learning Indo-European languages than those learning languages in other families (e.g.,
176 Bantu, the language family we expect to be most prevalent in our sample).

177 Despite evidence for general recognition of and preference for IDS across cultures, the
178 strength of IDS preferences is likely modulated by exposure. Exposure to IDS in the home
179 environment varies widely both within and between cultures (Casillas et al., 2020, 2021;
180 Cristia, Dupoux, Gurven, & Stieglitz, 2019; LeVine, 1994; L. A. Shneidman &
181 Goldin-Meadow, 2012; Vogt, Mastin, & Schots, 2015). Differences in IDS quantity have
182 also been hypothesized to reflect differences in child-rearing practices across cultures. For
183 example, direct verbal interaction between parents and infants can be rare in some societies
184 (Heath, 1983; LeVine, 1994; LeVine & LeVine, 2016; Schneidman & Goldin-Meadow, 2012;
185 Weber, Fernald, & Diop, 2017). Children in these societies – which are typically
186 non-WEIRD, though certainly not all non-WEIRD societies can be characterized this way
187 – are often expected to learn through observation and participation according to their skill
188 levels (see Legare, 2019, for a review). Thus, infants and young children in such societies

189 may hear less IDS directly from their caregivers than those in WEIRD societies in which
190 the norm involves a greater degree of direct address to parents. Of course, variation is also
191 present within as well as across cultures. Within-culture variation has primarily been
192 studied in North American contexts, where children from higher socioeconomic status
193 (SES) families tend to hear more IDS than children from lower SES families (e.g., Hart &
194 Risley, 1995; Hoff, 2006a; Huttenlocher, Waterfall, Vasilyeva, Vevea, & Hedges, 2010;
195 Rowe, 2012; Schneidman & Goldin-Meadow, 2012; Weisleder & Fernald, 2013).

196 By virtue of our broad sample of African cultures, we expect that our study can
197 capture substantial cultural variation in the average amount of IDS in children's
198 environments. The African sites we sample vary widely in their degree of urbanization,
199 their culture, their parenting values, and the average resources available in children's home
200 environments – all of which have been argued to be meaningful dimensions governing
201 children's early linguistic environment. For example, Keller (2012) suggested three
202 prototypical cultural environments for children based on the degree of urbanization of the
203 families in Western and non-Western societies. In this framework, in Western middle-class
204 urban societies, highly educated parents generally aim to help children develop individual
205 psychological autonomy. In contrast, in non-Western rural subsistence-based societies,
206 parents generally aim to help children develop communal action autonomy, so that children
207 have a strong sense of social responsibility and can contribute to the economic functioning
208 of the family (e.g., farming). Importantly, non-Western middle-class urban societies are a
209 hybrid of non-Western, rural and Western, urban societies, where parents generally want
210 their children to develop more individual autonomy but also emphasize the importance of
211 social responsibility in a large family. Broadly speaking, African families are from the
212 non-Western, urban and non-Western, rural groups in this taxonomy (see Table 1).

213 The confirmatory analyses of our study are designed to test whether there are
214 differences in the magnitude of IDS preferences measured in this sample and in the prior
215 samples of MB1. Although the average IDS production in the African sites we examine is

216 unknown, consistent differences along this dimension might plausibly lead to variation in
217 the magnitude of IDS preferences between our current study and MB1. In addition, our
218 exploratory analyses attempt to understand whether variation in IDS preference among
219 infants in our sample of African cultures is explained by demographic proxies related to
220 this taxonomy (e.g., urbanization and/or socioeconomic status).

221 Since multilingualism is common in Africa (e.g., Posel & Zeller, 2016; Rosenhouse &
222 Goral, 2004), many African children begin learning two or more different languages during
223 infancy. Does early multilingualism alter infants' preferences for IDS? The
224 ManyBabies1-Bilingual (MB1B) study provides some evidence that bilingual infants
225 showed a similar preference for NAE IDS when compared to monolingual infants
226 (Byers-Heinlein et al., 2021). MB1B examined bilingual infants' preference for NAE IDS at
227 6 to 9 months and 12 to 15 months and found that bilingual and monolingual infants did
228 not differ in terms of the magnitude of their IDS preferences. MB1B also found similar
229 results to MB1, that older bilingual infants and those bilinguals with higher exposure to
230 NAE show stronger IDS preference. However, as in the MB1 study, data collected in MB1B
231 mainly came from laboratories in WEIRD areas, such as North America and Europe, with
232 no laboratories from Africa, so the same caveats of generalizability apply to MB1B as to
233 MB1. Thus, in the current study, we included both monolingual and multilingual infants,
234 allowing us to assess the generalizability of MB1B's conclusions to our samples in Africa.

235 In sum, there are three primary (confirmatory) goals for the current study. First, we
236 aim to measure infants' preference for North-American English IDS across a range of
237 cultural and linguistic contexts in Africa. Second, we seek to measure developmental
238 changes in this preference. As we found that older infants show stronger IDS preferences
239 than younger infants in both MB1 and MB1B, we evaluate whether participants in our
240 study show the same developmental increases in IDS preference. Finally, we investigate
241 whether there are differences in IDS preferences between infants in Africa in our study and
242 those in Europe and Asia in MB1 and MB1B. As an exploratory aim, we also examine

Table 1

Test locations, participant ages, and languages by country.

| Country | Test location | Target age | Test language(s) |
|--------------|--|-------------|-----------------------------|
| Ghana | University of Ghana, Accra | 3–9 months | Akan, Ga, Ewe |
| Kenya | Nanyuki | 9–15 months | Kikuyu, Kiswahili, English |
| Malawi | University of Malawi – Chancellor College, Zomba | 6–15 months | Chichewa, English, or both |
| Nigeria | University of Jos, Plateau | 3–15 months | Hausa, English, Birom, Ngas |
| Rwanda | Kigali | 6–15 months | Kinyarwanda |
| Senegal | Dakar | 3–15 months | French, Wolof |
| South Africa | Pretoria | 3–15 months | Setswana, Xitsonga |
| Uganda | Makerere University, Kampala | 3–15 months | Luganda, English |

²⁴³ relationships between parents' demographics, their responses to survey items regarding
²⁴⁴ subjective use of IDS, and their child's IDS preference.

²⁴⁵

Methods

²⁴⁶ All deviations from the preregistration in the Stage 1 Registered Report are given in
²⁴⁷ Appendix A.

²⁴⁸ Participation Details

²⁴⁹ **Time-frame.** On July 23, 2018, we issued an open call for participation by African
²⁵⁰ researchers via listservs and social/professional networks. Prior to submission, 11 labs
²⁵¹ committed to data collection, but some labs were unable to collect data, in part due to
²⁵² changing professional obligations and delays due to Covid-19 closures. We received data

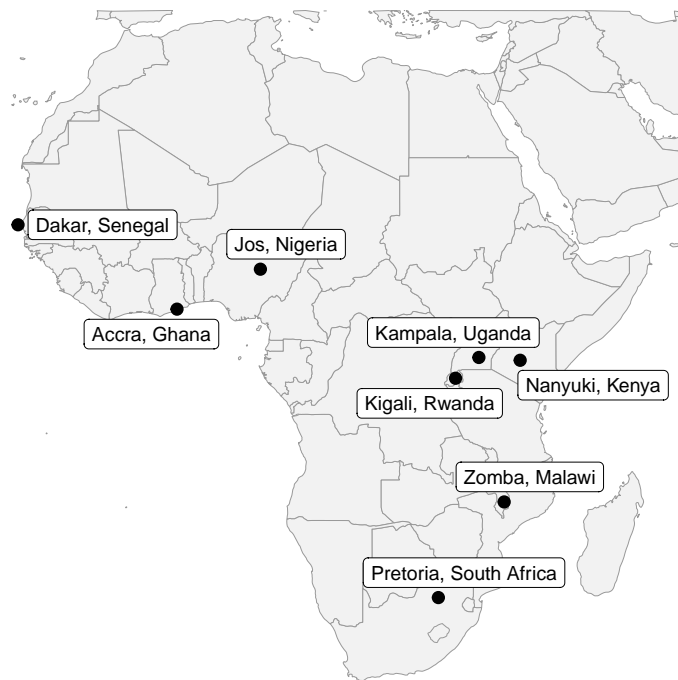


Figure 1. Map shows the location of the eight participating labs.

from 8 labs (See Table 1 for target sample characteristics of each site; Figure 1 for the location of the lab). Our participating laboratories would recruit infants living in eastern (e.g., Kenya), western (e.g., Senegal) and southern (e.g., South Africa) regions of Africa. Because many of our participating laboratories are located in East Africa, thus East African participants are disproportionately represented in our sample. Data collection began September 21, 2021. We initially anticipated finishing data collection a year later, but labs encountered a wide variety of unforeseen circumstances due to the COVID-19 pandemic, challenges with receiving IRB approval, and equipment and staffing issues. Thus, data collection was extended through 2023. Unfortunately, due to experimental setup issues, data from two labs was not analyzable, leading to a final sample of 6 datasets. In both excluded datasets, looking times were not recorded: this was due to a software bug in one lab and experimenter error in the other.

Age distribution. Each participating laboratory was asked to recruit participants in two age bins: 3;0 – 9;0 and 9;1 – 15;0 months. Similar to MB1, each laboratory was

267 asked to collect data spanning the age bin window, but aiming for the mean of the age bin.

268 **Sample size determination.** We estimated the effect size of infants' IDS
269 preference on the basis of the data from MB1. We used data from laboratories in MB1 that
270 used the single-screen central visual-fixation preference procedure (which we also use here:
271 see below) and that tested infants with no exposure to North American English (similar to
272 our population of interest). In a mixed-effects model, we examined the effect of test trial
273 type (IDS vs. ADS) on infants' looking time (log-transformed seconds), while controlling
274 for normally-distributed random intercepts by infant and laboratory. The intercept,
275 representing infants' average log-looking time across ADS trials, was 1.91; the variances of
276 the random intercepts were 0.074 and 0.022 at the infant and laboratory levels respectively.
277 The fixed-effect coefficient representing infants' preference for IDS over ADS was 0.080 and
278 the residual variance was 0.33.

279 In the first power analysis, we simulated datasets based on the above coefficient
280 estimates and variances. Using the simr package in R (Green & MacLeod, 2016), we ran a
281 power analysis for a mixed-effect analysis with the above-mentioned simulated datasets
282 (number of simulations = 1000). We were uncertain exactly how many labs to assume but
283 settled on 10, given the likelihood of some later signups as well as some lab attrition.
284 Assuming that we had 240 infants across 10 laboratories in each simulated dataset and an
285 alpha level of 0.05, we found that the average power was 99.40% [95% confidence interval:
286 98.70% – 99.78%] to detect the fixed ADS vs. IDS coefficient of 0.08. This first power
287 analysis was based on very small random-effect variances estimated from MB1 and MB1B
288 datasets. Given that most of the laboratories that participated in MB1 and MB1B had
289 more resources and more extensive experience in running infancy studies in comparison to
290 the participating laboratories in Africa, we planned for potentially higher variances in the
291 data collected in the current project. Thus, we ran a conservative second power analysis by
292 doubling the values of the random intercept and residual variances reported in the datasets
293 from MB1 and MB1B, while holding constant the intercept and the fixed-effect coefficient

representing infants' preference for IDS over ADS. With larger variances, the average power estimate dropped to 87.20% [95% confidence interval: 84.97% – 89.21%] for a total sample of 240 infants. The power analysis can be found at <https://osf.io/jgr79>.

Given that MB1 reported around 15% data excluded in the final analysis, we expect the exclusion rate for our project is around 15% to 20%. Thus, each laboratory agreed to contribute a minimum of 32 infants (16 infants in each age bin), including infants tested but excluded for reasons not related to the demographic and age inclusion criteria (e.g., fussiness). Further, we encouraged each laboratory to contribute additional data beyond that minimum. We propose that our projected sample size of 352 would have sufficient power, as 80% of this sample size exceeds our targeted final sample size ($n = 240$) based on the power analysis described above.

Ethics. All laboratories collected data under their own independent IRB protocol. Videos of individual infant participants during the experiment were recorded and stored at each laboratory. However, these videos were not shared with the central data analysis team. Laboratories were instead asked to only submit de-identified data for central data analyses.

Exclusion Criteria

All data collected for the study (i.e., every infant for whom a data file was generated, regardless of how many trials were completed) were uploaded to a central database for data analysis. Every laboratory followed the protocol to report any infants who were tested in this study, including those who were excluded from the analysis. Furthermore, each laboratory followed the protocol to make note of the reasons that infants were excluded from the study. A total number of 274 infants were tested in this study, and 74 infants were excluded from the final analysis.

Typically, participants were only included in the analysis if they met all of the criteria below. However, we allowed parents to choose not to answer some of the questions

319 (e.g., about full-term gestation and developmental disorders) because disclosures might
320 violate cultural norms in some areas of Africa. Thus, participating laboratories may have
321 included infants who did not fully meet the inclusion criteria defined here

322 **Full-term.** We defined full term as gestation times greater than or equal to 37
323 weeks. 11 (4.01%) of infants tested did not meet this criterion, and were excluded from
324 further analysis. To maximize parents' comfort in participating in the experiment, they
325 were given the option of not responding to questions about gestation.

326 **No developmental disorders or hearing loss.** We excluded infants with
327 parent-reported developmental disorders (e.g., chromosomal abnormalities, etc.) or
328 diagnosed hearing impairments. Developmental disorders and delays are stigmatized in
329 some cultures in Africa (e.g., negative attitudes towards children with disorders or delays),
330 therefore some parents may decline to answer the question about children's developmental
331 disorders. In this case, we still tested the infants and included the infants' data in the
332 analysis. This inclusion criterion was chosen to allow us to retain as much data as possible
333 while ensuring our questionnaire accommodates cultural norms. Further, we noted that
334 only 2 participants (i.e. less than 0.1%) in MB1 were excluded based on parents' report of
335 developmental disorders. Accordingly, we do not expect that including children whose
336 parents decline to answer this question would lead to an inclusion of large numbers of
337 children with developmental disorders that could potentially skew the results in the study.
338 4 (1.46%) of the infants tested did not meet this criterion. (We did not plan exclusions
339 based on self-reported ear infections unless parents reported medically-confirmed hearing
340 loss.)

341 **Trial-level and session-level errors.** Following MB1 and MB1B, we adopted a
342 relatively liberal inclusion criterion for this study. To be included in the study, a child must
343 have contributed non-zero looking time on at least one pair of test trials (i.e., one trial each
344 of IDS and ADS from a particular stimulus pair). We asked laboratories to identify two
345 different types of errors when uploading their data: trial-level errors and session-level

346 errors. Trial-level exclusions were based on whether we could use infants' data from a
347 particular test trial. For example, if an infant only completed the first six test trials of the
348 experiment, we entered this infant's data from the first six trials and discarded data from
349 all other trials. In this case, laboratories would identify this infant's data from the first to
350 sixth trials as "no trial errors" and any trials from the seventh trial onwards would be
351 identified as "trial errors". In contrast, session-level errors were errors that occurred when
352 running a particular participant. This type of error is different from the trial-level error
353 exclusions because it indicates that errors occurred which affected an entire session (e.g.,
354 failure to save data in the experiment). If a laboratory indicated a session-level error for a
355 particular infant, all data from this infant was excluded from the analysis. In sum, infants
356 who can contribute at least one pair of test trials (i.e., one IDS trial and one ADS trial)
357 would have some data excluded at the trial level whereas infants who cannot contribute
358 one pair of test trials would be excluded at the session level. In general, errors included the
359 following: equipment error (e.g., no sound or visuals on the first pair of trials),
360 experimenter error (e.g., an experimenter was unblinded in setups where infant looking was
361 measured by live button press), or evidence of parent interference or other types of
362 interference (e.g., talking or pointing by parents, construction noise, sibling pounding on
363 door), and infants being uncooperative or fussy (e.g., crying, not willing to do the
364 experiment).

365 Overall, at the trial level, 18 trials (0.39% of all trials) were excluded; relatively few
366 trials were marked by labs as having errors. Due to experimental setup errors, data from
367 two sites were unusable, leading to the exclusion of 60 infants (21.58% of all tested
368 participants). No additional test sessions were excluded.

369 Participants

370 **Final sample.** Our final sample included 200 infants (see Table 2 for more specific
371 sample demographic information) from 6 laboratories (mean sample size per laboratory:

Table 2

Final sample's demographics and language background by country. Mean age in months.

| Country | N | Mean Age (SD) | Sex | Language background |
|--------------|----|---------------|--------------------------|--|
| Ghana | 32 | 8.66 (3.72) | F: 17; M: 15 | Monolingual: 3; Bilingual: 8; Other: 21 |
| Kenya | 27 | 11.14 (2.88) | F: 15; M: 12 | Monolingual: 17; Bilingual: 5; Other: 5 |
| Malawi | 38 | 8.82 (4.10) | Unknown | Monolingual: 36; Other: 2 |
| Rwanda | 31 | 9.83 (4.23) | F: 16; M: 15 | Monolingual: 31 |
| South Africa | 31 | 9.00 (3.71) | F: 16; M: 14; Unknown: 1 | Monolingual: 23; Other: 3; Unknown: 5 |
| Uganda | 41 | 12.27 (2.17) | F: 21; M: 20 | Unknown |

³⁷² 33.33, SD: 5.16, range: 27 – 41). The mean age of infants included in the study was 296.49
³⁷³ days (range: 71 – 606). Similar to MB1, each laboratory was asked to collect data
³⁷⁴ spanning the two target age ranges (3;0 – 9;0 and 9;1 – 15;0 months); however, in practice,
³⁷⁵ many laboratories recruited participants outside the intended windows (younger than
³⁷⁶ 3-month-old: $N = 4$; older than 15-month-old: $N = 16$). We did not exclude these infants
³⁷⁷ from our analyses. See Figure 2 for a distribution of the age of the included participants in
³⁷⁸ each site. An additional 60 infants were tested but excluded (see the full details on
³⁷⁹ exclusions above).

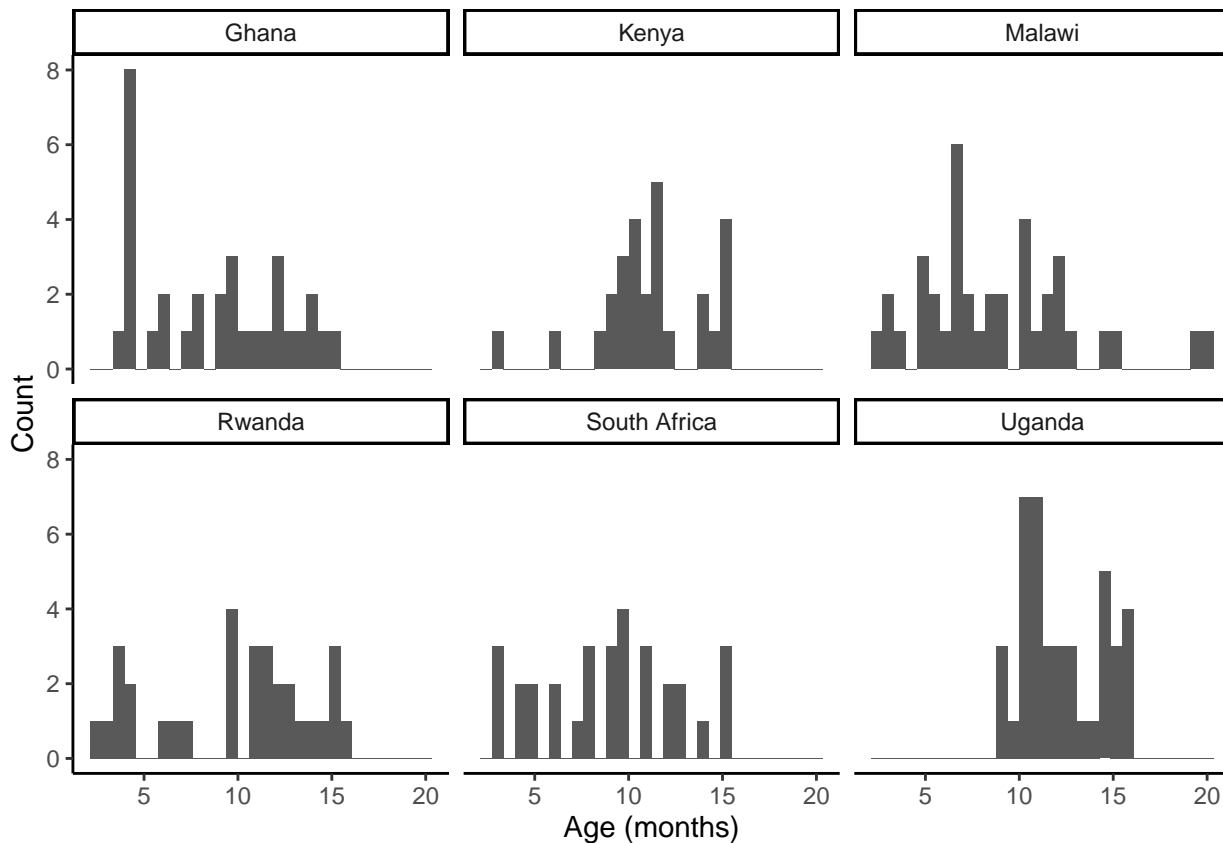


Figure 2. Histograms of the age distribution in each lab. X-axis represents the age in months.

As mentioned in the Introduction, multilingualism is common in Africa. Thus, many

infants in the final sample are likely to have been exposed to more than one language. To

assess infants' language backgrounds, each laboratory completed a family questionnaire

with the participating parents (see materials in linked repository:

https://osf.io/jgr79/?view_only=5ee43f58762742daaa2caa21b85e3780). Our family

language background questionnaire was created based on the family language background

questionnaire in the MB1 and MB1B studies, and included questions asking parents to

estimate the number of hours that their infants heard different languages. We calculated

the percentage of time that infants were exposed to a given language as the number of

hours they hear that language (per day) divided by the total number of hours the infant

hears any language each day. This method is simpler than the traditional interview method

391 used in assessing bilingual infants' language exposure (Byers-Heinlein et al., 2019), but in
392 order to minimize the burden on participating laboratories and families, we decided to use
393 a short questionnaire method to assess infants' language backgrounds.

394 In this paper, we define bilingualism following the criteria established in MB1B
395 (Byers-Heinlein et al., 2021). Monolingual infants are defined as those who have a minimum
396 of 90% exposure to one language. Simultaneous bilingual infants are defined using the
397 following criteria: (i) infants are regularly exposed to two or more languages beginning
398 within the first month of life; (ii) they have a minimum of 25% exposure to each of their
399 languages. In other words, bilingual infants are exposed to two languages between 25% to
400 75% of their time. Based on these criteria, it is possible that bilingual infants in our paper
401 were exposed to multiple languages. For example, an infant with 45% English, 45% French,
402 and 10% Spanish exposure would be regarded as a bilingual infant. Infants who did not
403 meet the bilingual or monolingual criteria were designated as "other language background."
404 All infants were included in the main, confirmatory analyses regardless of language
405 background. Language background groupings were treated as a covariate in the analyses.

406 Based on the above-mentioned criteria, 87 infants were classified as monolingual
407 infants, 13 infants were classified as bilingual infants, and 28 infants were classified as
408 other. The remaining 72 infants' language background was unknown.

409 Materials

410 Visual stimuli. All visual stimuli were the same as those used in the MB1 study. We
411 used a brightly colored static checkerboard as the fixation stimulus, and an animation with
412 shrinking concentric multi-colored circles to ensure infants were attending to the screen at
413 the start of each trial. All of the stimuli can be found at <https://osf.io/wh7md/>.

414 Auditory stimuli. All auditory stimuli were identical to those used in the MB1 study.
415 The stimuli were recordings of North-American English mothers either speaking with

416 experimenters (ADS) or with their infants whose ages ranged from 122 to 250 days in a
417 laboratory setting. Mothers were provided with a set of objects and were asked to talk
418 about the objects with the experimenters and their infants in separate recording sessions.
419 In total, two sets of auditory stimuli were created: one set consisted of 8 IDS stimuli and
420 the other set consisted of 8 ADS stimuli. Each stimulus lasted for 18 seconds. The details
421 of stimulus creation can be found in the report of MB1 (ManyBabies Consortium, 2020).

422 Volume. Each laboratory measured stimulus volume level using a smartphone app
423 (e.g., the Android app “Sound Meter”). Labs kept the stimulus volume close to 63 – 65 dB
424 SPL. According to the protocol, labs would measure and report the background noise level
425 and the stimulus level. However, this information was not collected.

426 Procedure

427 Apparatus. Each laboratory used a laptop computer that had the experiment
428 programmed in Habit 2.26 (Oakes, Sperka, DeBolt & Cantrell, 2019). Moreover, each
429 laboratory used a computer monitor to present the visual stimuli, a speaker for audio
430 stimuli, a webcam for the experimenter to observe and record infants’ performance,
431 curtains/room dividers that separated the experimenter from the infant and parent during
432 the experiment, and two sets of headphones: one for the experimenter and one for the
433 parent.

434 Experimental procedure. The procedure was identical to the single-screen central
435 visual fixation preference procedure reported in the MB1 study (ManyBabies Consortium,
436 2020). Using the single-screen central fixation method, researchers measured in real time
437 the duration of infants’ looking time to the computer monitor while they listened to the
438 audio recordings. Infants’ looking time to the computer monitor indicated their preference
439 for the audio recordings (i.e., IDS/ADS). Each laboratory followed procedural instructions
440 closely (based on pre-recorded videos illustrating the procedures, which were shared with

441 all participating laboratories) to maintain the consistency of the experimental procedure
442 across laboratories.

443 The experimenter explained the study to the parent and obtained consent from the
444 parent before running the experiment. After completing the consent form, the
445 experimenter led the participant to the testing room. To minimize distraction, the
446 experimenter was separated from the infant and parent by curtains or a room divider.
447 During the experiment, the infant sat on the parent's lap. To minimize any bias introduced
448 by the experimenter or parent hearing the stimuli, each of them wore headphones and
449 heard masking music during the experiment.

450 Parents were instructed not to speak to the infant during the experiment and not to
451 point to the screen. Infants' performance was recorded by a webcam that was placed in
452 front of and below the computer monitor. Infants' looking time to each trial was measured
453 online by the experimenter, who observed the infant's behavior via the webcam. At the
454 beginning of each trial, a short video of a colorful circle was presented to orient the infant's
455 attention to the screen. Once the infant fixated on the screen, the experimenter started the
456 trial. The first two trials of the session were warm-up trials that accustomed infants with
457 the procedure of the experiment, so the infant's looking time during warm-up trials was
458 not analyzed. The auditory stimuli for the warm-up trials was piano music that lasted 18
459 seconds on each trial and the visual stimulus was the same as in the test trials (i.e., a
460 colorful checkerboard). After the first two warm-up trials, the infant was tested with 16
461 trials presenting the IDS and ADS stimuli. Each infant was randomly assigned to one of
462 four pseudo-random orders to counterbalance the order of presentation of IDS and ADS
463 stimuli. Within each order, there were four blocks and each block presented 2 IDS and 2
464 ADS trials in alternating order. The presentation of the trials within each block were
465 counterbalanced such that two blocks started with an IDS trial, and the other two blocks
466 started with an ADS trial. On each trial, the auditory stimulus would continue to play
467 until the infant looked away for 2 consecutive seconds or reached the maximum length of

468 the auditory stimulus (18 seconds). Experimenters used the Habit program to record all
469 looking time for every trial. There was no minimum looking time per trial that was
470 required for continuation of the experiment. However, as in the MB1 study, any looking
471 time that was less than 2 seconds was not analyzed. We excluded 335 (11.46%) trials that
472 had less than 2 seconds looking time in total.

473 After the main looking-time task, the parents answered questions from the
474 experimenter about participant and family demographic information, such as infant sex,
475 date of birth, language exposure, and preterm/full term status. The questionnaire was
476 translated into the appropriate language(s) for participants from each data collection site.
477 See supplementary materials for the English template and adaptations.

478 **General Lab Practices**

479 **Training of the experimenters.** Three of the authors conducted a 2-day training
480 workshop in Nairobi, Kenya on January 28 – 29, 2020, which was attended by lead
481 researchers from 8 of the participating laboratories. The training session provided an
482 overview of the experimental procedure, advice on setting up the apparatus at the
483 researcher's institution, and training, instructions and guidelines for running the
484 experiment. Further, the first author sent instructions for experiment set-up and the
485 workshop materials to all participating laboratories, and kept close contact with all lead
486 researchers in the participating laboratories to provide technical support for the
487 experiment.

488 **Training of research assistants.** Each laboratory was responsible for maintaining
489 good experimenter training practices. We extended an invitation for the training workshop
490 to one research assistant in each laboratory, so that the researcher primarily responsible for
491 data collection could receive training directly as well. Following the MB1 study, each
492 laboratory reported on which research assistant ran each infant using pseudonyms or
493 numerical codes. After data collection, each laboratory completed a questionnaire

494 regarding their training practices, the experience and academic status of each
495 experimenter, and their basic participant greeting practices.

496 **Results**

497 **Confirmatory Analyses**

498 Data processing and analytic framework. Our primary dependent variable of interest
499 was infants' looking time (LT). Infants' looking time was defined as time spent fixating on
500 the computer screen during test trials. We did not count LT when infants looked away
501 from the screen, though the trial was discontinued if an infant looked away and did not
502 look back to the screen within 2 seconds. Following MB1 and MB1B, we log-transformed
503 looking times prior to statistical analysis (Csibra, Hernik, Mascaro, Tatone, & Lengyel,
504 2016). We made this decision because we wanted to compare the data of the current study
505 with those in MB1 and MB1B.

506 We tested our research questions via general linear mixed effects models. We fit all
507 models using a maximal random effects structure (Barr, Levy, Scheepers, & Tily, 2013).
508 Under this approach, we first specified all random effects that are appropriate for the
509 experimental design (e.g., IDS/ADS trial type varied within subjects in our experimental
510 design, thus it can be specified as a random effect by subject; see below for the full list of
511 effects considered). If any of these mixed-effects models failed to converge, we used an
512 iterative pruning strategy: first removing random slopes nested within subjects, next
513 removing random slopes nested within labs, and finally removing random intercepts from
514 groupings in the same order, retaining effects of trial type as these were of greatest
515 theoretical interest. Following MB1 and MB1B, we fit all models using the lme4 package
516 with the bobyqa optimizer, version 1.1-35.3 (Bates, Mächler, Bolker, & Walker, 2015) and
517 computed confidence intervals and p values using the lmerTest package (Kuznetsova,
518 Brockhoff, & Christensen, 2017).

In addition to the mixed-effect models, we assessed the reliability of measurement in our study by reporting the reliability of the infants' looking time difference to the IDS vs ADS stimuli across different trials. We reported the intraclass correlation coefficient (ICC) as our reliability measure. The ICC was computed using the psych package in R (Revelle, 2017). We reported an ICC3k measure, on the basis of a two-way random effects model, a mean-rating of 8 (i.e., we had 8 pairs of IDS and ADS trials) and consistency agreement (Koo & Li, 2016; Parsons, Kruijt, & Fox, 2019). The estimated ICC was 0.18, 95% CI [0.00, 0.34]. This relatively low value is consistent with previous work showing limited test-retest reliability in IDS studies (Schreiner et al., 2024).

Below is a description of variables in our mixed-effect models:

- Log_lt: Dependent variable. Log-transformed looking time in seconds.
- Trial_type: a dummy coded variable with two levels: ADS (reference) and IDS. A positive coefficient means that infants look longer to IDS trials compared to ADS trials.
- Age_months: a continuous variable measuring the infant's age in months (centered).
- Trial_num: An index for the current trial (1-16 for infants who completed the experiment). Excluded trials were reflected as missing trial numbers.
- Language_background: this consisted of two dummy coded variables that represented infants from three different language backgrounds: monolinguals ($\geq 90\%$ exposure to one's native language); bilinguals ($\geq 25\%$ to each of their languages); other (any infants who were not categorized as monolinguals or bilinguals). Using monolinguals as the reference level, the two dummy-coded variables are: (i) bilingual – infants who were categorized as bilinguals would be coded as 1 and all other infants would be coded as 0; (ii) Other (any infants who are not monolinguals or bilinguals) – infants who were categorized as other would be coded as 1 and all other infants would be coded as 0. In this case, monolingual

infants would be coded as 0 in the above-mentioned dummy-coded variables.

- Infant Type: a dummy coded variable with two levels, representing infants living in Africa in our current study (coded as 1) and infants living in Europe, Australia and Asia who were not hearing North American English, with data from MB1(B) (coded as 0).

As a reminder, we examined the following research questions in our paper: (1) IDS

preference: whether infants in our multi-site African sample showed a preference for IDS

and what is the corresponding effect size of this preference; (2) Age effect: whether there

were changes in the infants' IDS preference across different ages; (3) Population

comparison: examine whether the magnitude of infants' IDS preference in our study

differed from infants in MB1 and in MB1B (comparing only infants in these three samples

who were not exposed to North American English).

Research questions 1 and 2: Infants' IDS preference and age effect. We addressed our

first two research questions using only the data collected in the current paper from

laboratories in Africa. We specified the following model: `log_lt ~ trial_type +`

`trial_num + age_months + trial_type * trial_num + age_months * trial_num +`

`age_months * trial_type + (trial_type * trial_num | subid) + (trial_type |`

`lab).`

The fixed-effects structure of this model included main effects of trial type (IDS vs

ADS), age, and trial number. This structure controls for the effects of each independent

variable on infants' looking time (e.g., longer looking times for IDS, shorter looking times

on later trials). In addition, we included several two-way interaction terms: trial type

interacting with trial number to model the possibility of infants' faster habituation to ADS,

age interacting with the trial type to model the developmental trajectory of infants' IDS

preference, and age interacting with trial number to model faster habituation for older

children. The random effects structure of the model controlled for subject-level and

571 lab-level grouping. For subject-level grouping, we added random intercepts and random
 572 effects of trial type, trial number, and their interaction to model the possibility that each
 573 infant may have different rates of habituation for IDS and ADS trials. For lab-level
 574 grouping, we added a random effect trial type to model differences in IDS preferences
 575 across labs.

576 After pruning for non-convergence, our final model specification was: `log_lt ~`
 577 `trial_type + trial_num + age_months + trial_type * trial_num + age_months *`
 578 `trial_num + age_months * trial_type + (1 | subid) + (1 | lab)`.

579 As in MB1 and MB1B, the fixed effect estimate for trial type corresponds to the
 580 predicted infant-directed speech preference effect in units of log looking time (research
 581 question 1). The fixed effect estimate for the interaction of trial type and age indicates the
 582 estimated age-related change in infant-directed speech preference in log seconds per month
 583 (research question 2).

Table 3

Model estimates for Research Questions 1 and 2. The baseline for trial type is adult-directed speech. Trial number is mean-centered. Age is measured in months and standardized.

| Term | Estimate [95% CI] | std.error | t | p |
|---------------------------|----------------------|-----------|-------|-------|
| (Intercept) | 1.89 [1.66, 2.12] | 0.09 | 21.34 | < .01 |
| Trial Type | 0.06 [0.02, 0.10] | 0.02 | 2.76 | 0.01 |
| Trial Number | -0.03 [-0.04, -0.02] | 0.00 | -9.72 | < .01 |
| Age | -0.09 [-0.14, -0.04] | 0.03 | -3.40 | < .01 |
| Trial Type * Trial Number | 0.00 [-0.01, 0.01] | 0.00 | -0.22 | 0.83 |
| Trial Number * Age | 0.00 [0.00, 0.01] | 0.00 | 1.60 | 0.11 |
| Trial Type * Age | 0.02 [-0.02, 0.06] | 0.02 | 0.94 | 0.35 |

584 The model revealed a significant main effect of trial type, such that infants looked

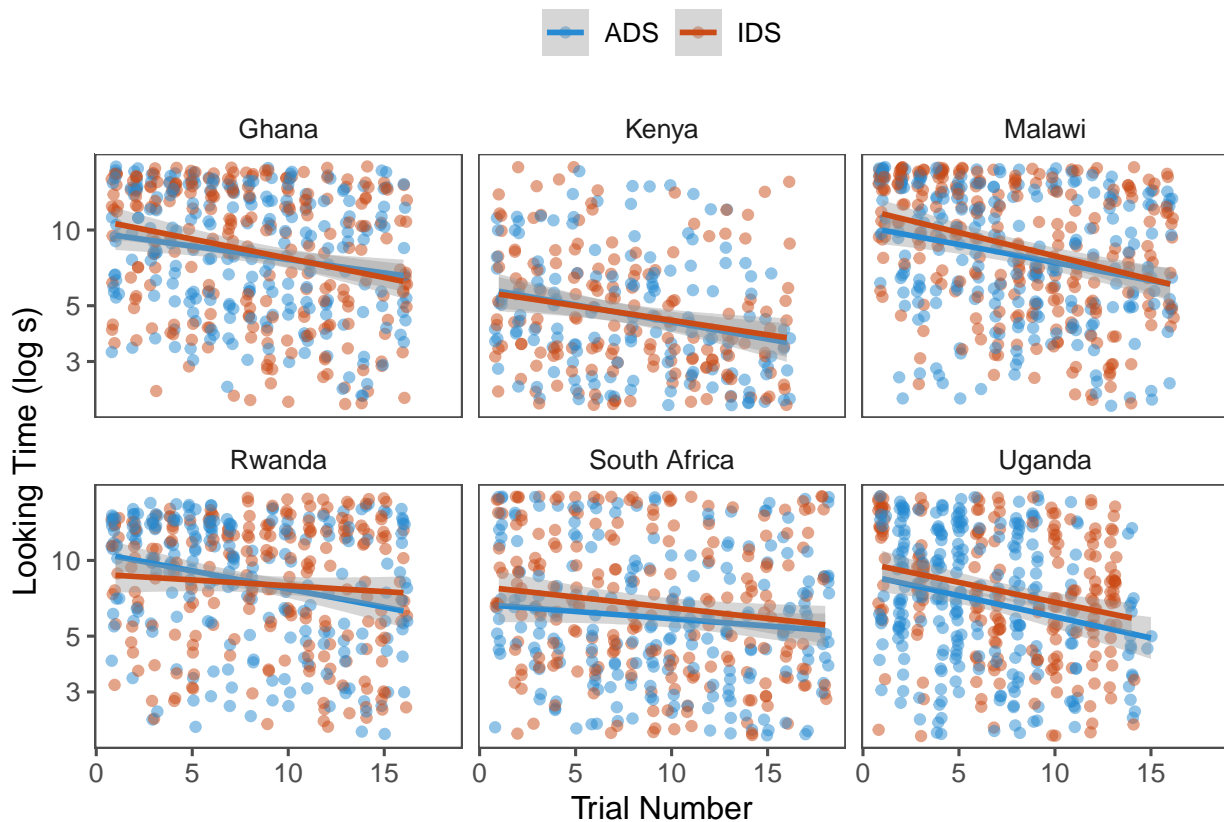


Figure 3. By lab results. Each dot represents one trial of looking time. X-axis represents the trial number. Y-axis represents the log looking time in seconds. Red represents IDS trials. Blue represents ADS trials.

longer at IDS trials than ADS trials (Figure 3, Figure 4; Table 3; $\beta = 0.06 [0.02, 0.10]$; $SE = 0.02$; $t = 2.76$; $p = 0.01$). There was also a significant negative effect of trial number, indicating that looking times decreased over the course of the session ($\beta = -0.03 [-0.04, -0.02]$; $SE = 0$; $t = -9.72$; $p < 0.01$). Age in months was also a significant predictor, with older infants showing shorter looking times overall ($\beta = -0.09 [-0.14, -0.04]$; $SE = 0.03$; $t = -3.40$; $p < 0.01$). None of the interaction terms reached statistical significance, including the interaction between trial type and age, suggesting that the magnitude of IDS preference did not change reliably with age.

Research question 3: Population comparison. In this analysis, we compare the data

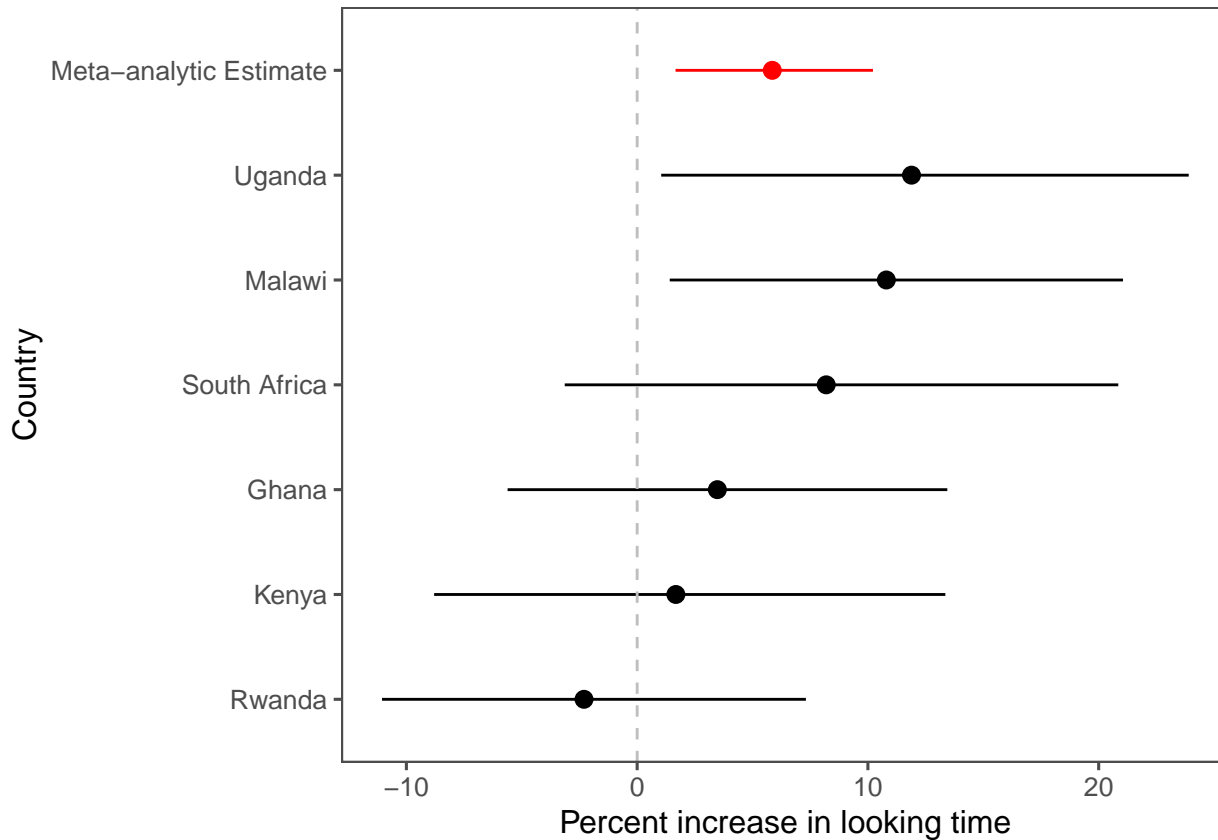


Figure 4. Percent increase in infants' looking time to infant-directed speech (IDS) relative to adult-directed speech (ADS) across six African sites and the meta-analytic estimate. While the meta-analytic estimate (red point) reflects the fixed-effect coefficient from the preregistered model, individual site estimates (black points) were obtained by fitting site-specific mixed-effects models predicting looking time using the same specification without the by-site random effect. Percent changes were computed from the fixed effect estimate, and horizontal lines represent 95% confidence intervals. Positive values indicate longer looking to IDS.

⁵⁹⁴ collected from the laboratories in Africa to data collected in MB1 and MB1B in Germany,

⁵⁹⁵ Italy, New Zealand, Turkey, United Kingdom. We selected the subset of data from MB1

⁵⁹⁶ and MB1B that was collected using central fixation procedures (to match methods across

⁵⁹⁷ studies) and from infants who were not exposed to North American English (non NAE) (to

598 match stimulus un-familiarity due to language background). While we could have
599 controlled the methodological and demographic variables statistically (and hence included
600 all data from MB1 and MB1B in the full model), we believed that the increase in model
601 complexity – and comparable decrease in interpretability – outweighed the benefits of this
602 strategy.

603 We examine whether our sample of infants' IDS preference is different from those in
604 MB1 and MB1B with the following model: `log_lt ~ trial_type + trial_num +`
`age_months + infant_type + language_background + trial_type * trial_num +`
`age_months * trial_num + age_months * trial_type + trial_type * infant_type`
`+ trial_num * infant_type + trial_type * language_background + (trial_type *`
`trial_num | subid) + (trial_type | lab)`

609 In this mixed-effects model, the fixed-effects included main effects of trial type,
610 language background, age, trial number, infants in our study/non NAE infants in MB1(B)
611 and language background. In addition, we included several two-way interaction terms in
612 the fixed effects structure: (i) trial type interacted with trial number, modeling the
613 possibility of infants' faster habituation to ADS, (ii) age interacted with trial number,
614 modeling faster habituation for older children, (iii) age interacted with trial type, modeling
615 the developmental trajectory of infants' IDS preference, (iv) trial type interacted with
616 infants in our sample, modeling the possible difference in IDS preference between infants in
617 Africa and infants tested in MB1 and MB1B, (v) trial num interacted with infants in our
618 sample, modeling the possible difference in habituation between our sample of infants and
619 infants tested in MB1 and MB1B, and (vi) trial type interacted with language background,
620 modeling the possible difference in IDS preference from infants with different language
621 backgrounds. We adopted the same baseline random effects as in the previous model.

622 After pruning for non-convergence, our final model specification was: `log_lt ~`
623 `trial_type + trial_num + age_months + infant_type + language_background +`

624 trial_type * trial_num + age_months * trial_num + age_months * trial_type +
625 trial_type * infant_type + trial_num * infant_type + trial_type *
626 language_background + (1 | subid) + (1 | lab). The fixed effect estimate
627 corresponding to our research question is the trial_type * infant_type, which captures
628 differences in measured IDS preference between the current data and data from
629 MB1/MB1B in units of log seconds of looking time.

630 The model revealed no significant difference in IDS preference between infants tested
631 in our sample and those tested in MB1/MB1B, as indicated by the non-significant
632 interaction between trial type and infant type (Table 4; $\beta = 0 [-0.06, 0.06]$; $SE = 0.03$; $t =$
633 0.03; $p = 0.97$). However, there was a significant main effect of trial type ($\beta = 0.02 [-0.04,$
634 0.07]; $SE = 0.03$; $t = 0.62$; $p = 0.53$), suggesting that infants showed a reliable preference
635 to IDS over ADS.

636 Consistent with expectations, looking times decreased over the course of the session
637 ($\beta = -0.05 [-0.05, -0.05]$; $SE = 0$; $t = -31.08$; $p < 0.01$), and older infants looked for less
638 time overall ($\beta = -0.12 [-0.15, -0.09]$; $SE = 0.01$; $t = -8.26$; $p < .001$). We also found that
639 older infants habituated more quickly, as indicated by a significant negative interaction
640 between age and trial number ($\beta = 0 [-0.01, 0]$; $SE = 0$; $t = -3.79$; $p < .001$). The
641 interaction between trial number and infant type was also significant ($\beta = 0.02 [0.01, 0.03]$;
642 $SE = 0$; $t = 6.64$; $p < .001$), indicating that looking times declined more slowly across
643 trials for infants in our sample compared to those in MB1/MB1B. No other interactions
644 reached significance.

645 Exploratory Analyses

646 SES. Previous research in North America (e.g., Hart & Risley, 1995; Hoff, 2006b;
647 Weisleder & Fernald, 2013) has shown that the quantity and quality of child-directed
648 speech vary across families with different SES backgrounds. These differences in language

Table 4

Model estimates for Research Questions 3. The baseline for trial type is adult-directed speech (ADS), trial number is mean-centered, age is measured in months and standardized, the baseline for infant type is infants from MB1/MB1B (non-NAE sample), and the baseline for language background is bilingual infants.

| Term | Estimate [95% CI] | SE | t | p |
|--|----------------------|------|--------|-------|
| (Intercept) | 1.76 [1.62, 1.91] | 0.07 | 24.16 | < .01 |
| Trial Type | 0.02 [-0.04, 0.07] | 0.03 | 0.66 | 0.51 |
| Trial Number | -0.05 [-0.05, -0.05] | 0.00 | -30.82 | < .01 |
| Age | -0.12 [-0.15, -0.09] | 0.02 | -8.38 | < .01 |
| Infant Type | 0.13 [-0.09, 0.36] | 0.12 | 1.32 | 0.21 |
| Language Background (Monolingual) | -0.02 [-0.12, 0.07] | 0.05 | -0.48 | 0.63 |
| Language Background (Other) | 0.00 [-0.14, 0.14] | 0.07 | 0.24 | 0.81 |
| Trial Type * Trial Number | 0.00 [-0.01, 0] | 0.00 | -0.81 | 0.42 |
| Trial Number * Age | 0.00 [-0.01, 0] | 0.00 | -4.39 | < .01 |
| Trial Type * Age | 0.02 [0, 0.04] | 0.01 | 1.86 | 0.06 |
| Trial Type * Infant Type | 0.00 [-0.06, 0.06] | 0.03 | -0.26 | 0.80 |
| Trial Number * Infant Type | 0.02 [0.01, 0.03] | 0.00 | 4.96 | <.01 |
| Trial Type * Language Background (Monolingual) | 0.04 [-0.02, 0.11] | 0.03 | 1.42 | 0.16 |
| Trial Type * Language Background (Other) | 0.01 [-0.08, 0.10] | 0.05 | 0.07 | 0.95 |

649 input may drive differences in infants' preference for IDS. Thus, we explored how SES
 650 affects infants' preference for IDS. SES was measured by primary caregiver's formal
 651 education (number of years). We entered primary caregiver's formal education (in years) as
 652 a predictor in the regression model specified for RQ1 and RQ2, along with its interaction
 653 with trial type.

The interaction between trial type and primary caregiver education was not significant ($\beta = -0.01 [-0.05, 0.03]$; $SE = 0.02$; $t = -0.46$; $p = 0.65$), not allowing rejection of the null hypothesis that SES does not moderate infants' preference for IDS. In other words, the magnitude of IDS preference was similar regardless of caregivers' years of formal education. At the same time, we did observe a significant main effect of trial type, with infants looking longer at IDS than ADS trials overall ($\beta = 0.06 [0.01, 0.10]$; $SE = 0.02$; $t = 2.55$; $p = 0.01$). Looking times decreased significantly across trials ($\beta = -0.04 [-0.04, -0.03]$; $SE = 0$; $t = -9.92$; $p < 0.01$), and older infants looked for less time overall ($\beta = -0.10 [-0.15, -0.05]$; $SE = 0.03$; $t = -3.62$; $p < 0.01$). No other interactions were significant. As a robustness check, we re-ran the model on the subset of infants with female primary caregivers (80.52% infants). The pattern of results was qualitatively unchanged.

Meta-analysis. For comparison with ManyBabies 1 and 1B, we computed standardized effect sizes for each lab using looking time, following the method used in ManyBabies Consortium (2020). We also confirmed that the effect size was similar when computing using log-transformed looking time. The resulting meta-analytic plot is shown in Appendix B: Figure B1. The meta-analytic effect size is 0.17[-0.03, 0.37] (log transformed: 0.14[-0.06, 0.34]), which is numerically smaller than the .35 [0.29, 0.42] reported in MB1 (Manybabies Consortium, 2020) and the .26 [0.09, 0.43] for bilingual infants reported in MB1B (Byers-Heinlein et al., 2021). But these estimates are not directly comparable due to the different method and age distribution in the current project (cf., Zettersten et al., 2024).

General Discussion

Infants' preference for IDS is both an important phenomenon for understanding language learning and a case study for infant methods more broadly. The MB1 study investigated variation in IDS preference across laboratories and across countries and found a small but reliable effect such that infants preferred IDS over ADS. Although there was

680 substantial variation across labs, our analyses suggest that much of it reflected random
681 sampling error, as the moderators we tested accounted for little of the between-lab
682 variance. Nevertheless, the effect was moderated by infant age, language background, and
683 experimental method. Though MB1 and its sister project MB1B investigated children from
684 a wide variety of language backgrounds, no sites from Africa were included in this initial
685 group of participating laboratories. The current study was designed to fill this gap.

686 We summarize our findings with respect to three research questions in the paper.
687 First, consistent with MB1 and MB1B, we found evidence for a significant IDS preference
688 in a sample of 200 African infants. Second, and unlike MB1, we did not find significant
689 age-related variation in IDS preference, but given the relatively small magnitude of the
690 overall effect, we may not have had sufficient power to detect an interaction with age in our
691 primary analytic model. In addition, the variability in age across sites was limited, with
692 four of the six labs having mean ages clustered around 8 to 9 months, leaving us little age
693 variation to detect such an effect.

694 Finally, we were interested in comparing the magnitude of the IDS preference in the
695 current study to the estimates obtained in MB1 and MB1B (with multilingual data
696 providing an important comparison because of the diverse language backgrounds in our
697 current sample). We did not find significant study effects in a model comparing data from
698 the current study to a method- and language-background matched sample of infants from
699 MB1 and MB1B. The magnitude of the IDS preference in African infants in the current
700 study ($d = 0.17$) was numerically smaller than the overall estimate reported by MB1
701 ($d = 0.35$) and the estimates for bilingual infants reported by MB1B ($d = 0.26$). But the
702 two estimates are not directly comparable; among other things, the current study was
703 conducted with infants who were not growing up learning North American English and it
704 used the central fixation method, both of which were associated with overall lower effect
705 sizes in MB1 compared with other groups (Zettersten et al., 2024).

The current study provides important evidence on the generalizability of the IDS preference and of looking-time methods in infancy more broadly. Despite the diversity of MB1, as noted above, the vast majority of labs were from Western countries where Indo-European languages are spoken. Thus, the findings of the current study provide evidence that the IDS preference observed in MB1 is present in infants growing up in a diverse array of non-WEIRD environments. More broadly, there is limited work using looking-time methods for infancy research in the African context (cf. Pyykkö et al., 2019). Recent studies in Ghana have begun to build this foundation, demonstrating that Ghanaian infants show vowel-harmony preferences and can use harmony cues for speech segmentation using the same central-fixation paradigm (Omane, Benders, & Boll-Avetisyan, 2024; Omane, Boll-Avetisyan, & Benders, 2025). The current work complements these studies by extending the approach across multiple African sites and testing a different domain, thereby providing a broader demonstration of methodological feasibility across cultures.

Importantly, our findings also suggest that the IDS preference extends to infants who grow up in highly multilingual environments. Many infants in our study were likely exposed to more than one language from birth, a common characteristic in African contexts. For instance, infants in Ghana may receive input from two to six different languages within a single household (Omane, Benders, & Boll-Avetisyan, 2025). The persistence of the IDS preference at the group level despite such linguistic diversity underscores the robustness of this effect.

A second goal of the current study was to build a team of laboratories in Africa collecting data with infants. We were successful in accomplishing this goal and we believe that the current study represents the largest experimental study of African infants to date. Despite this success, we encountered a number of substantial challenges. Some of these were unique to the African context (e.g. variable internet connectivity) and to the specific time-period of our study (e.g., spanning the Covid-19 pandemic) while others were more general to the project of conducting “big team science” investigations across institutions

733 with a wide range of resources.

734 Although we received commitments for data collection from 11 teams, three of these
735 teams were unable to collect data due to a variety of challenges relating to personnel and
736 resources. One source of these challenges was the initiation of Covid-19 lockdowns soon
737 after our initial project training in the winter of 2020. In some cases, personnel left the
738 project or priorities changed, leading teams to lose the ability to participate. These
739 lockdowns were also difficult even for teams who stayed involved in the project. Due to
740 turnover and the long delay between initial training and setup for data collection after
741 restrictions were eased, a number of procedural deviations were introduced. In two cases,
742 these deviations were so severe that we could not analyze data from the site. In an ideal
743 world, our group would have been able to conduct additional site visits and training after
744 sites began collecting data; unfortunately this was not possible due to budget and
745 personnel limitations.

746 In addition to logistical and training challenges, it is important to acknowledge the
747 broader structural differences between the participating laboratories in this project and the
748 Western laboratories that contributed to MB1 and MB1B. Most Western labs had access
749 to established infrastructure for infant research, such as participant databases, mailing
750 lists, as well as stable funding and institutional traditions of developmental
751 experimentation. In contrast, many of the African sites in our study were building these
752 capacities for the first time. Recruiting families without existing participant pools required
753 extensive community engagement and word-of-mouth networks. The success of this effort
754 therefore reflects not only careful coordination and training, but also the commitment and
755 creativity of local researchers who developed sustainable pathways for infant recruitment
756 and testing in settings with limited prior infrastructure.

757 These efforts resonate with broader calls to diversify and globalize infant research.
758 Recent commentaries have highlighted that developmental science continues to draw

759 disproportionately from Western contexts, and have urged the field to address the
760 structural barriers that prevent broader participation (Kidd & Garcia, 2022; Singh, Cristia,
761 Karasik, Rajendra, & Oakes, 2023). Equitable global collaboration requires attention not
762 only to sampling diversity but also to the power dynamics and infrastructural inequities
763 that shape how research is conducted and credited. Encouragingly, recent initiatives such
764 as the /L+/ Global School on Language Acquisition (Aravena-Bravo et al., 2024), co-led
765 by several members of the MB1-Africa team, demonstrate how capacity building can foster
766 sustainable participation from underrepresented regions.

767 The current study has a number of further scientific limitations, including some
768 shared with prior studies including MB1 and MB1B. First, although we attempted to
769 estimate IDS preference, we did so using a specific set of speech stimuli and a specific
770 paradigm. It is likely that the stimuli used here are less extreme than many used in prior
771 studies, and further, they are produced in North American English, making them
772 linguistically unfamiliar to one degree or another to all of the infants in our study.
773 Followups using native language stimuli are needed to measure the importance of this
774 choice to the IDS preference (i.e., MB1N, an in-progress follow up study to MB1 examining
775 native language IDS preferences). Second, although we invited broad participation, our
776 samples are convenience samples in at least two ways: both of the sites who participated
777 and the infants who participated at each site. Thus our effect size estimates cannot be
778 treated as population effects but rather “proof of concept” that an IDS preference can be
779 observed in African infants across a diverse set of sites. Finally, although we did not
780 observe major demographic variation, we caution against over-interpretation of any
781 demographic differences in IDS preference given that IDS preference has not been shown to
782 be individually predictive of any later outcomes (Soderstrom et al., 2024).

783 In sum, our study offers a case study of “big team science” (Coles, Hamlin, Sullivan,
784 Parker, & Altschul, 2022) carried out via a collaboration between African researchers and
785 the ManyBabies Consortium. Although it faced a variety of logistical challenges (many

786 shared with other grass roots efforts; Baumgartner et al., 2023), it nevertheless yields
787 important evidence on generalizability, of a key phenomenon in early language learning, of
788 looking time methods, and finally of a broad-based collaborative model for studying infant
789 development.

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Appendix A

Discrepancies between pre-registration and final analyses

1. ICC comparison

Pre-registration: We planned to compare the estimated ICC in our study with the ICC reported in MB1(B).

Final: This comparison was not conducted because MB1(B) did not report ICC values.

2. Metrics of effect heterogeneity

Pre-registration: We planned to report distributional metrics describing heterogeneity across labs (Mathur & VanderWeele, 2020) using the *MetaUtility* package.

Specifically, we proposed to estimate: (1) the percentage of effects greater than 0, (2) the percentage greater than Cohen's $d = 0.2$, and (3) the percentage less than $d = -0.2$. These metrics were to be reported only if random slopes of trial type by lab were included in the final model, their variance was estimated as greater than 0, and at least 10 labs contributed data.

Final: Random slopes of trial type by lab were not included in the final model, and only six labs contributed data. As a result, we did not estimate these metrics.

3. Subset of MB1 and MB1(B) data

Pre-registration: We planned to subset MB1 and MB1(B) data to include only infants tested with central fixation procedures and infants not exposed to North American English (NAE), to maximize comparability with the present study.

Final: In MB1 and MB1(B) data, information about English exposure often did not distinguish between American and British English. Therefore, we excluded infants based on country of residence (United States/Canada) rather than reported language exposure.

4. Urban–rural exploratory analysis

1029 *Pre-registration:* We planned an exploratory analysis testing whether infants' IDS
1030 preference differed between urban and rural areas, motivated by prior findings of
1031 differences in parental speech input across these contexts (e.g., Keller, 2012; Vogt et
1032 al., 2015). *Final:* We did not collect information on urban versus rural residence, so
1033 this analysis was not conducted.

1034 **5. Socioeconomic status (SES) analyses**

1035 *Pre-registration:* We planned to measure SES using both mothers' years of formal
1036 education and the MacArthur Scale of Subjective Social Status (MacSSS), including
1037 both variables in regression models.

1038 *Final:* We did not have MacSSS data for most participants, and information on
1039 mothers' education was incomplete. We therefore used primary caregiver education
1040 as the main SES variable and conducted a robustness check using the subset of infant
1041 whose primary caregiver's sex is female.

Appendix B

By-lab Meta-analysis

1042 We computed a single effect size per lab and fit an intercept-only mixed-effect
1043 meta-regression to estimate the overall IDS preference across sites. This approach provides
1044 a comparable summary of results across sites. Even with a standardized protocol, sites
1045 differ in their cultures, recruitment pools, equipment, and experimenter behavior. A
1046 random-effects meta-analysis treats those differences as legitimate heterogeneity rather
1047 than noise, yielding a conservative estimate of the cross-lab mean and its uncertainty.

1048 To do so, we calculated each infant's mean IDS–ADS difference score ¹, standardized
1049 these within lab to obtain effect sizes, and estimated their sampling variances. These
1050 lab-level estimates were then entered into a REML random-effects model to produce the
1051 pooled effect size and 95% confidence interval (Figure 1). The meta-analytic effect size is
1052 0.17[-0.03, 0.37], which is numerically smaller than the .35 [0.29, 0.42] reported in MB1
1053 (Manybabies Consortium, 2020) but these estimates are not directly comparable due to the
1054 different method and age distribution in the current project (cf. Mathur et al., 2024).

¹ Due to an experimental procedure error, infants at the South Africa site were not always presented with complete IDS–ADS stimulus pairs; in some cases, the same stimulus was played multiple times. While this issue did not affect the random-effects model in the main analysis, it does impact the present meta-analysis. We therefore trimmed the data by (1) retaining only the first presentation of each trial and (2) including only trials in which both the IDS and ADS versions were presented. The trimmed data includes 176 trials from 28 infants.

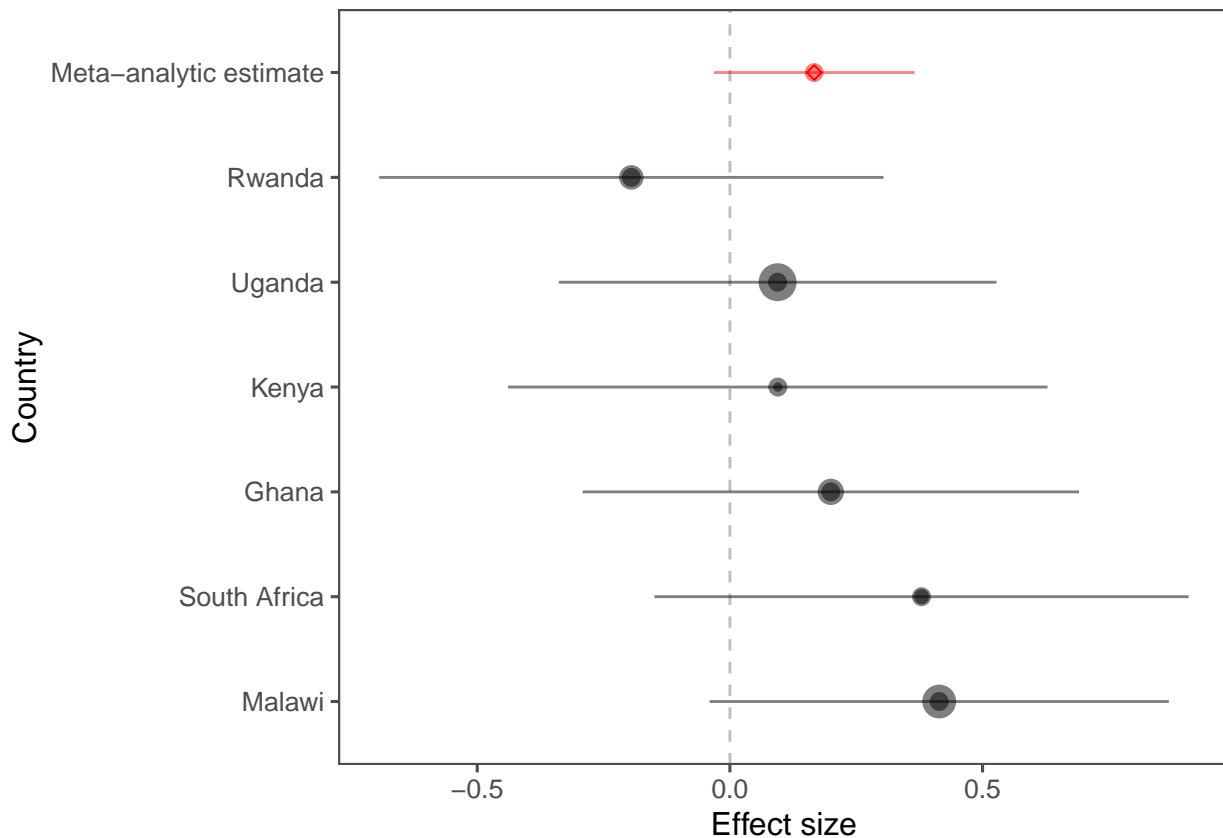


Figure B1. Forest plot of lab-level standardized effect sizes (for the IDS–ADS preference). Points represent individual country estimates, with size proportional to the inverse of their sampling variance; horizontal bars show 95% confidence intervals. The meta-analytic aggregate (top, red) is from an intercept-only random-effects model.