Limited evidence of test-retest reliability in infant-directed speech preference in a large 1 pre-registered infant sample 2 Melanie S. Schreiner^{1,2}, Martin Zettersten^{3,4}, Christina Bergmann⁵, Michael C. Frank⁶, Tom Fritzsche⁷, Nayeli Gonzalez-Gomez⁸, Kiley Hamlin⁹, Natalia Kartushina¹⁰, Danielle J. Kellier¹¹, Nivedita Mani^{1,2}, Julien Mayor¹⁰, Jenny Saffran³, Mohinish Shukla¹², Priya Silverstein^{13, 14}, Melanie Soderstrom¹⁵, & Matthias Lippold^{1,2} ¹ University of Goettingen ² Leibniz Science Campus PrimateCognition ³ University of Wisconsin-Madison ⁴ Princeton University 10 ⁵ Max Planck Insitute for Psycholinguistics 11 ⁶ Stanford University 12 ⁷ University of Potsdam 13 ⁸ Oxford Brookes University ⁹ University of British Columbia ¹⁰ University of Oslo ¹¹ University of Pennsylvania 17 ¹² Università di Padova 18 ¹³ Institute for Globally Distributed Open Research 19

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39 Abstract

Test-retest reliability — establishing that measurements remain consistent across multiple testing sessions — is critical to measuring, understanding, and predicting individual 41 differences in infant language development. However, previous attempts to establish 42 measurement reliability in infant speech perception tasks are limited, and reliability of frequently-used infant measures is largely unknown. The current study investigated the test-retest reliability of infants' preference for infant-directed speech (hereafter, IDS) over adult-directed speech (hereafter, ADS) in a large sample (N=158) in the context of the ManyBabies1 collaborative research project (hereafter, MB1; Frank et al., 2017; ManyBabies Consortium, 2020). Labs of the original MB1 study were asked to bring in participating infants for a second appointment retesting infants on their IDS preference. This approach allows us to estimate test-retest reliability across three different methods used to investigate preferential listening in infancy: the head-turn preference procedure, 51 central fixation, and eye-tracking. Overall, we find no consistent evidence of test-retest 52 reliability in measures of infants' speech preference (overall r = .09, 95% CI [-.06,.25]). 53 While increasing the number of trials that infants needed to contribute for inclusion in the analysis revealed a numeric growth in test-retest reliability, it also considerably reduced the 55 study's effective sample size. Therefore, future research on infant development should take into account that not all experimental measures may be appropriate for assessing 57 individual differences between infants.

Keywords: language acquisition; speech perception; infant-directed speech;
 adult-directed speech; test-retest reliability

Word count: 3998

Limited evidence of test-retest reliability in infant-directed speech preference in a large

pre-registered infant sample

Obtaining a quantitative measure of infants' cognitive abilities is an extraordinarily difficult endeavor. The most frequent way to assess what infants know or prefer is to track overt behavior. However, measuring overt behavior at early ages presents many challenges: participants' attention span is short, they do not follow instructions, their mood can change instantly, and their behavior is often inconsistent. Therefore, most measurements are noisy and the typical sample size of an infant study is small (around 20 infants per group), resulting in low power (Oakes, 2017). In addition, there is individual and environmental variation that may add even more noise to the data (e.g., Johnson & Zamuner, 2010). Despite these demanding conditions, reliable and robust methods for assessing infants' behavior are critical to understanding development.

In order to address these challenges, the ManyBabies collaborative research 74 consortium was formed to conduct large-scale, conceptual, consensus-based replications of 75 seminal findings to identify sources of variability and establish best practices for 76 experimental studies in infancy (Frank et al., 2017). The first ManyBabies collaborative 77 research project (hereafter, MB1, ManyBabies Consortium, 2020) explored the reproducibility of the well-studied phenomenon that infants prefer infant-directed speech 79 (hereafter, IDS) over adult-directed speech (hereafter, ADS, Cooper & Aslin, 1990). Across 80 many different cultures, infants are commonly addressed in IDS, which typically is characterized by higher pitch, greater pitch range, and shorter utterances, compared to the language used between interacting adults (Fernald et al., 1989). A large body of behavioral studies finds that infants show increased looking times when hearing IDS compared to ADS stimuli across ages and methods (Cooper & Aslin, 1990; see Dunst, Gorman, & Hamby, 2012 for a meta-analysis). This attentional enhancement is also documented in neurophysiological studies showing increased neural activation during IDS compared to

ADS exposure (Naoi et al., 2012; Zangl & Mills, 2007). IDS has also been identified as facilitating early word learning. In particular, infants' word segmentation abilities (Floccia et al., 2016; Schreiner & Mani, 2017; Singh, Nestor, Parikh, & Yull, 2009; Thiessen, Hill, & Saffran, 2005) and their learning of word-object associations (Graf Estes & Hurley, 2013; Ma, Golinkoff, Houston, & Hirsh-Pasek, 2011) are enhanced in the context of IDS. In sum, several lines of evidence suggest that IDS is beneficial for early language development.

Within MB1, 67 labs contributed data from 2,329 infants showing that babies 94 generally prefer to listen to IDS over ADS. Nevertheless, the overall effect size of d=0.3595 was smaller than a previously reported meta-analytic effect size of d = 0.67 (Dunst et al., 2012). The results revealed several additional factors that influenced the effect size. First, 97 older infants showed a larger preference of IDS over ADS. Second, the stimulus language was linked to IDS preference, with North American English learning infants showing a larger IDS preference than infants learning other languages. Third, comparing the different 100 methods employed, the head-turn preference procedure yielded the highest effect size, while 101 the central fixation paradigm and eye-tracking methods revealed smaller effects. Finally, 102 exploratory analyses assessed the effect of different inclusion criteria. Across methods, 103 using stricter inclusion criteria led to an increase in effect sizes despite the larger proportion of excluded participants (see also Byers-Heinlein, Bergmann, & Savalei, 2021). 105

However, there is a difference between a result being reliable in a large sample of 106 infants and the measurement of an individual infant being reliable. In studies tracking 107 individual differences, the measured behavior during an experimental setting is often used 108 to predict a cognitive function or specific skill later in life. Individual differences research of this kind often has substantial implications for theoretical and applied work. For example, 110 research showing that infants' behavior in speech perception tasks can be linked to later 111 language development (see Cristia, Seidl, Junge, Soderstrom, & Hagoort, 2014 for a 112 meta-analysis) has the potential to identify infants at risk for later language delays or 113 disorders. However, a necessary precondition for this link to be observable is that

individual differences between infants can be measured with high reliability at these earlier stages, in order to ensure that measured inter-individual variation mainly reflects differences in children's abilities rather than measurement error. How reliable are the measures used in infancy research?

Previous attempts to address the reliability of measurements have typically been 119 limited to adult populations (Hedge, Powell, & Sumner, 2018), or have been conducted 120 with small sample sizes (e.g., Houston, Horn, Qi, Ting, & Gao, 2007). For example, 121 Colombo, Mitchell, and Horowitz (1988) used a paired-comparison task, in which infants 122 were familiarized with a stimulus and presented with the familiarized and a novel stimulus 123 side-by-side at test. Results indicated that infants' novelty preference was extremely 124 variable from task to task. Assessing infants' performance from one week to another 125 revealed that infants' attention measures were moderately reliable. However, reliability 126 seemed to increase with the number of tasks infants completed in the younger age group, 127 suggesting that reliability is influenced by the number of assessments. In addition, infants' 128 performance from 4 to 7 months was longitudinally stable but somewhat smaller than 129 week-to-week reliability. Cristia, Seidl, Singh, and Houston (2016) also retested infant 130 populations by independently conducting 12 different experiments on infant speech perception at three different labs with different implementations of the individual studies. Hence, it was only after completed data collection that the data was pooled together by the different labs revealing potential confounds. Nevertheless, the results showed that 134 reliability was extremely variable across the different experiments and labs and low overall 135 (meta-analytic r = .07).

Against this background, the current study investigates test-retest reliability of infants' performance in a speech preference task. Within MB1, a multi-lab collaboration, we examine whether infants' preferential listening behavior to IDS and ADS is reliable across two different test sessions. We also investigate the influence of various moderators on the reliability of IDS preference (e.g., time between test and retest; infants' language

background).

Our study was faced with a critical design choice: what stimuli to use to assess 143 test-retest reliability. One constraint on our study was that, since it was a follow-on to 144 MB1, any stimulus we used would always be presented after the MB1 stimuli. One option 145 would be simply to bring back infants and have them hear exactly the same stimulus 146 materials. A weakness of this design would be the potential for stimulus familiarity effects, 147 however, since infants would have heard the materials before. Further complicating 148 matters, infants might show a preference for or against a familiar stimulus depending on 149 their age (Hunter & Ames, 1988). The ideal solution then would be to create a brand new 150 stimulus set with the same characteristics. Unfortunately, because of the process of how 151 MB1 stimuli were created, we did not have enough normed raw recordings available to 152 make brand new stimulus items that conformed to the same standards as the MB1 stimuli. 153 We therefore chose an intermediate path: we reversed the ordering of MB1 stimuli. 154 Average looking times in MB1 were always lower than 9s per trial, even for the youngest 155 children on the earliest trials (the group who looked the longest on average), so most 156 children in MB1 did not hear the second half of most trials. Thus, by reversing the order, 157 we had a perfectly matched stimulus set that was relatively unfamiliar to most infants. The disadvantage of this design was that infants who looked longer might be more likely to hear a familiar clip heard in the previous study. If infants then showed a familiarity 160 preference — an assumption which might not be true — the end result could be to inflate 161 our estimates of test-retest reliability slightly, since longer lookers would on average look 162 longer at retest due to their familiarity preference. We view this risk as relatively low, but 163 do note that it is a limitation of our design. 164

The current study also explores whether there are any differences in test-retest reliability between three widely used methods: central fixation (CF), eye-tracking (ET), and the head-turn preference procedure (HPP). Exploring differences in CF, ET, and HPP, Junge et al. (2020) provide experimental and meta-analytic evidence in favor of using the

HPP in speech segmentation tasks. Similarly, the MB1 project reported an increase in the effect size for HPP compared to CF and ET (ManyBabies Consortium, 2020). HPP 170 requires gross motor movements relative to other methods, such as CF and ET paradigms, 171 for which subtle eye movements towards a monitor located in front of the child are 172 sufficient. One possible explanation for the stronger effects with HPP may be a higher 173 sensitivity to the contingency of the presentation of auditory stimuli and infants' head 174 turns away from the typical forward-facing position. While these findings suggest that 175 HPP may be a more sensitive index of infant preference, they do not necessarily imply 176 higher reliability for individual infants' performance using HPP. For example, Marimon 177 and Höhle (2022) found no evidence for test-retest reliability when testing infants' prosodic 178 preferences using the HPP method. It remains an open question whether the same 179 measures that produce larger effect sizes at the group-level also have higher test-retest reliability for individual infants (Byers-Heinlein, Bergmann, et al., 2021). Therefore, 181 assessing the test-retest reliability of the different preference measures is crucial, so that 182 researchers can make informed decisions about the appropriate methods for their particular 183 research question. Critically, only measures with high test-retest reliability should be used 184 for studies of individual differences.

186 Method

187 Preregistration

Prior to the start of data collection, we preregistered the current study on the Open Science Framework (https://osf.io/v5f8t; see S1 in the Supplementary Materials for details).

91 Data Collection

A call was issued to all labs participating in the original MB1 study on January 24th,
2018 (ManyBabies Consortium, 2020). The collection of retest session data was initially set
to end on May 31st, 2018, one month after the end date of the original MB1 project. Due
to the fact that the original MB1 project extended the time frame for data collection and
the late start of data collection for the MB1 test-retest study, we also allowed participating
labs to continue data collection past the scheduled end date.

198 Participants

Contributing labs were asked to re-recruit their monolingual participants between the ages of 6 to 12 months who had already participated in the MB1 project. If participating labs had not committed to testing either of these age groups, they were also allowed to re-recruit participants from the youngest age group of 3- to 6-month-olds and/or the oldest age group of 12- to 15-month-olds. Labs were asked to contribute half (n=16) or full samples (n=32); however, a lab's data was included in the study regardless of the number of included infants. The study was approved by each lab's respective ethics committee and parental consent was obtained for each infant prior to participation in the study.

Our final sample consisted of 158 monolingual infants from 7 different labs (Table 1).

In order to be included in the study, infants needed a minimum of 90% first language

exposure, to be born full term with no known developmental disorders, and normal hearing

and vision. We excluded 11 participants due to session errors and 11 participants who did

not have at least one valid trial per condition (IDS and ADS) at their first or second

session. The mean age of infants included in the study was 245 days (range: 108 – 373

days).

Materials

Visual stimuli. The visual stimuli and instructions were identical to MB1. For the
CF paradigm and ET, labs used a multicolored static checkerboard as the fixation stimulus
as well as a multicolored moving circle with a ringing sound as an attention-getter between
trials. For the HPP method, labs used their standard procedure, as in MB1.

Speech stimuli. We used the identical training stimuli of piano music from MB1.

A second set of naturalistic IDS and ADS recordings of mothers either talking to their

infant or to an experimenter was created for the retest session by reversing the order of

clips within each sequence of the original study. This resulted in eight new sequences of

natural IDS and eight new sequences of natural ADS with a length of 18 seconds each.

Procedure. Infants were retested using the identical procedure as during the first testing day: CF, HPP, or ET. Participating labs were asked to schedule test and retest sessions 7 days apart with a minimum number of 1 day and a maximum number of 31 days. However, infants whose time between test and retest exceeded 31 days were still included in the analyses (n = 3). The mean number of days between test and retest was 10 (range: 1 - 49).

A total of 18 trials, including two training, eight IDS, and eight ADS trials, were
presented in one of four pseudo-randomized orders. Trial length was either infant-controlled
or fixed depending on the lab's standard procedure: a trial stopped either if the infant
looked away for 2 seconds or after the total trial duration of 18 seconds. The online coding
experimenter and the parent listened to music masked with the stimuli of the study via
noise-cancelling headphones. If the experimenter was in an adjacent room separate from
the testing location, listening to masking music was optional for the experimenter.

Data exclusion. A child was excluded if they had a session error, i.e., an
experimenter error (e.g., inaccurate coding, or presentation of retest stimuli on the first
test session) or equipment failure (visual stimuli continued to play after the end of a trial).

Table 1
Statistics of the included labs. n refers to the number of infants included in the final analysis.

Lab	Method	Language	Mean age (days)	N
babylab-potsdam	HPP	German	227	22
babyling-oslo	eye-tracking	Norwegian	249	10
brookes-babylab	central fixation	English	267	18
InfantCog-UBC	central fixation	English	147	7
infantll-madison	HPP	English	230	30
lancslab	eye-tracking	English	236	16
wsi-goettingen	central fixation	German	280	39
wsi-goettingen	HPP	German	242	16

Trials were excluded if they were marked as trial errors, i.e., if the infant was reported as
fussy, an experimental or equipment error occurred, or there was parental interference
during the task (e.g., if the parent spoke with the infant during the trial). Trials were also
excluded if the minimum looking time of 2 s was not met. If a participant was unable to
contribute at least one IDS and one ADS trial for either test or retest, all data of that
participant was excluded from the test-retest analyses.

Results

IDS preference

First, we examined infants' preference for IDS in both sessions. Two-samples t-tests comparing the difference in average looking time between IDS and ADS to zero revealed that infants showed a preference of IDS over ADS in Session 1, t(157) = 6.47, p < .001, and

Trial type	Session 1 Mean	Session 1 SD	Session 2 Mean	Session 2 SD
ADS	7.72	2.77	6.96	2.92
IDS	8.76	2.85	7.75	2.75

Table 2 Average looking times (in seconds) for each session and condition

Session 2, t(157) = 4.19, p < .001, replicating the main finding from MB1 (Table 2). 68.35% of infants in Session 1 and 63.29% of infants in Session 2 showed a preference for 252 IDS. In order to test whether there was a difference in the strength of the preference effect across sessions, we fit a linear mixed-effects model predicting infants' average difference in looking time between IDS and ADS from test session (1 vs. 2), including by-lab and 255 by-participant random intercepts. There was no significant difference in the magnitude of 256 infants' preference between the two sessions, β =-0.30, SE=0.24, p=.208.

Reliability

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We assessed test-retest reliability in two ways. First, we fit a linear mixed-effects 250 model predicting IDS preference in Session 2 from IDS preference in Session 1, including a 260 by-lab random intercept. The results revealed no significant relationship between IDS 261 preference in Session 1 and 2 (Table 3). Second, we calculated the Pearson correlation 262 coefficient. While a simple correlation coefficient might overestimate the test-retest 263 reliability in our sample because it does not control for the differences between different labs and methods (HPP, CF, and ET), we felt it was important to also conduct a Pearson correlation as it is commonly used to assess reliability. The size of the correlation 266 coefficient was not statistically different from zero and the estimate was small, r = .09, 95%267 CI [-.06, .25], t(156) = 1.19, p = .237. Moreover, no significant correlations emerged in 268 each sample considered separately (Figure 1; see Supplementary Materials S3 for a 269

Table 3

Coefficient estimates from a linear mixed effects model predicting IDS preference in Session 2.

	Estimate	SE	t	р
Intercept	0.87	0.46	1.92	0.10
IDS Preference Session 1	0.04	0.09	0.41	0.68

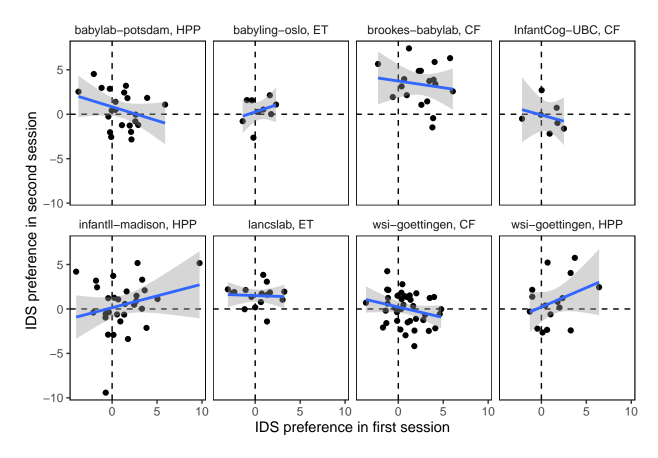


Figure 1. Correlation between IDS Preference in Session 1 and Session 2 in each lab and method. Dots indicate individual participants. Error bands represent 95 percent confidence intervals. The dashed line indicates no preference (i.e., a value of zero) for the first and second session, respectively.

Table 4

Coefficient estimates from a linear mixed effects model predicting IDS preference in Session 2 and Pearson correlation coefficient for each method separately.

Method	beta	SE	p	Pearson r
HPP	0.15	0.14	0.28	0.13
ET	0.03	0.16	0.84	0.02
CF	-0.20	0.12	0.12	0.08

meta-analytic approach). 41.77% of the infants reversed their direction of preference for IDS versus ADS from the test to the retest session.

To investigate the test-retest reliability of each specific method, we computed Pearson 272 correlation coefficients and the same mixed-effects model described above for HPP, CF, 273 and ET separately (Table 4). None of the three methods showed evidence of test-retest 274 reliability. Neither the Pearson correlation coefficients nor the coefficients of the multilevel 275 analysis were significant, all p-values > 0.12. In planned secondary analyses, we found that 276 time between test sessions, participant age, and language background did not moderate the 277 relationship between IDS preference in session 1 and session 2 (see Supplementary 278 Materials S2). Taken together, we find no significant evidence of test-retest reliability 270 across our preregistered analyses. 280

Results with different inclusion criteria

To this point, all analyses were performed using the inclusion criteria from MB1,
which required only that infants contribute at least one trial per condition for inclusion
(i.e., one IDS and one ADS trial). However, more stringent inclusion criteria yielded larger

effect sizes in MB1. We therefore conducted exploratory analyses assessing test-retest 285 reliability after applying progressively stricter inclusion criteria, requiring two, four, six, 286 and eight valid trials per condition. Applying stricter criteria — and thereby increasing the 287 number of test trials — increased reliability numerically from r = 0.07 to r = 0.34 (Figure 288 2). In part due to a decrease in sample size, only one of these correlations was statistically 280 significant (when requiring six trial pairs): two valid trial pairs, t(152) = 0.90, p = .367; 290 four valid trial pairs, t(143) = 1.03, p = .306; six valid trial pairs, t(98) = 2.23, p = .028; 291 eight valid trial pairs — all trials in both sessions — t(22) = 1.68, p = .108. The analyses 292 provide tentative evidence that stricter inclusion criteria may lead to higher test-retest 293 reliability, but at the cost of substantial decreases in sample size (see Supplementary 294 Materials S5 for additional analyses). 295

General Discussion

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The current study investigated the test-retest reliability of infants' preference for IDS 297 over ADS. We retested the IDS preference of infants participating in the original MB1 298 project to assess the extent to which their pattern of preference would remain consistent 299 across multiple testing sessions. While we replicated the original effect of infants' speech 300 preference for IDS over ADS for both the test and retest session on the group-level, we 301 found that infants' speech preference measures showed no evidence of test-retest reliability. 302 In other words, we were unable to detect stable individual differences in infants' preference 303 for IDS. This finding is consistent with past research suggesting low test-retest reliability in 304 other infant paradigms (Cristia et al., 2016). Given that most experimental procedures conducted in infant research are interested in the comparison of groups, individual differences between participants within a specific condition are usually minimized by the 307 experimental procedure while differences between conditions are maximized. Therefore, 308 infant preference measures may be a good approach for capturing group-level phenomena, 309 but may be less appropriate for examining individual differences in development. 310

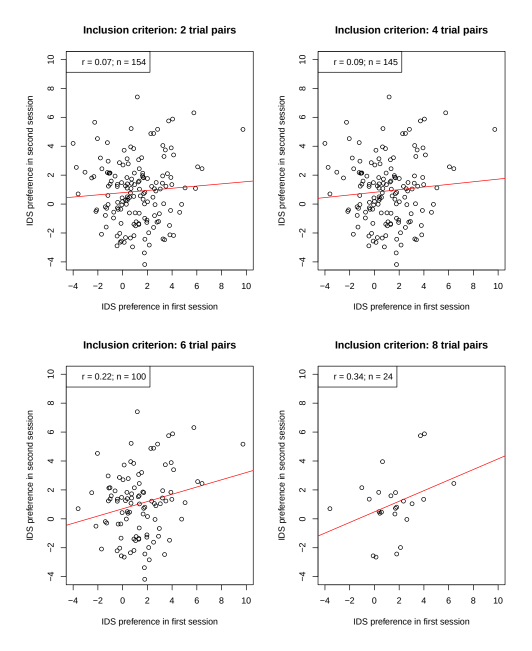


Figure 2. IDS preferences of both sessions plotted against each other for each inclusion criterion. n indicates the number of included infants, r is the Pearson correlation coefficient as the indicator for reliability.

Consistent with general psychometric theory (e.g., DeBolt, Rhemtulla, & Oakes, 311 2020), stricter inclusion criteria — and consequently a larger number of included test trials 312 per participant — tended to increase the magnitude of the correlation between test 313 sessions. However, this association was based on exploratory analyses and was in part only 314 observed descriptively, and hence should be interpreted with caution. A similar effect on 315 the group-level was found in the MB1 project, where a stricter inclusion criterion led to 316 bigger effect sizes (ManyBabies Consortium, 2020). As in MB1, higher reliability through 317 strict exclusions came at a high cost. In particular, with the strictest criterion, only a small 318 portion of the original sample size (24 out of 158 infants) could be included in the final 319 sample. In other words, applying stricter criteria leads to a higher drop-out rate and can 320 dramatically reduce the sample size. In the case of studies in the field of developmental 321 science, where there are many practical restrictions in collecting large samples of infants (e.g., birth rate in the area, restricted lab capacities, budget restrictions), a strict drop-out 323 criterion may often be difficult to implement. Note that studies in developmental science already have above-average drop-out rates (Miller, 2017). In addition, drop out may not be 325 random, and so having high drop-out rates can further limit the generalizability of a study. 326 In fact, the number of trials individual infants contributed was highly correlated between 327 test sessions in the current study (see Supplementary Materials S6). Particularly in the 328 context of turning individual differences measures into diagnostic tools, high drop-out rates 329 have an additional limitation of not being broadly usable. 330

An alternative approach to increasing the number of valid trials is to increase the
number of experimental trials. This approach seeks to increase the likelihood that
participants will contribute sufficient trials (after trial-level exclusions) to allow for precise
individual-level estimates (DeBolt et al., 2020; see also Silverstein, Feng, Westermann,
Parise, & Twomey, 2021). While this approach is promising, it may not always be feasible,
because the attention span of a typical infant participant is limited. Therefore, prolonging
the experimental procedure to maximize the absolute number of trials is often challenging

in practice. Other avenues for obtaining higher numbers of valid trials may include changes in the procedure (e.g., Egger, Rowland, & Bergmann, 2020) or implementing multi-day test sessions (Fernald & Marchman, 2012).

As our results are only based on the phenomenon of IDS preference (albeit, with 341 three widely used methods: HPP, CF, ET) it is essential to further assess the underlying 342 reliability of preferential looking measures within other areas of speech perception (Marimon & Höhle, 2022). While most infants prefer IDS over ADS (Dunst et al., 2012), patterns of preferential looking in other tasks (e.g., speech segmentation) are often inconsistent and difficult to predict (Bergmann & Cristia, 2016). These inconsistencies in looking behavior are especially important to consider in the context of relating a direction of preference to later language development, and can sometimes lead to seemingly contradictory findings. That is, both familiarity and novelty responses have been suggested 349 to be predictive of infants' later linguistic abilities (DePaolis, Vihman, & Keren-Portnoy, 350 2014; Newman, Ratner, Jusczyk, Jusczyk, & Dow, 2006; Newman, Rowe, & Ratner, 2016). 351 In light of our findings, researchers conducting longitudinal studies with experimental data 352 from young infants predicting future outcomes should be cautious, as there may be large 353 intra-individual variability affecting preference measurement. 354

355 Limitations

While we had an above-average sample size for a study in infant research, we were
unable to approach the number of participants collected within the original MB1 study. In
addition to a delayed call, the extra effort of having to schedule a second lab visit for each
participant and the fact that there were already other collaborative studies taking place
simultaneously (MB1B, Byers-Heinlein, Tsui, Bergmann, et al., 2021; MB1G,
Byers-Heinlein, Tsui, Van Renswoude, et al., 2021), might have contributed to a low
participation rate. A higher sample size and a larger number of participating labs from
different countries would have enabled us to conduct a more highly-powered test of

differences in test-retest reliability across different methods, language backgrounds, and participant age.

A further limitation concerns the stimuli. While the order of the audio recording clips 366 presented to infants within a given trial differed between the first and second session, the 367 exact same stimulus material as in MB1 was used in both sessions. In particular, all 368 children heard the exact same voices in Session 1 and in Session 2. From a practical point of view, this was the most straightforward solution for coordinating the experiment within 370 the larger MB1 project. However, familiarity effects might have influenced infants' looking behavior. Infants with longer looking times in their first session might have had more 372 opportunity to recognize familiar audio clips in their second session. For infants with short looking times, familiar audio clips would only occur towards the end of second-session trials, thus offering infants less opportunity to recognize voices from their first session. 375 Therefore, inconsistent familiarity with the stimulus material in the second session across 376 infants might have artificially lowered test-retest reliability. 377

378 Conclusion

Following the MB1 protocol, the current study could not detect test-retest reliability
in measures of infants' preference for IDS over ADS. Subsequent analyses provided
tentative evidence that stricter criteria for the inclusion of participants may enhance
test-retest reliability at the cost of high drop-out rates. Developmental studies relying on
stable individual differences between their participants need to consider the underlying
reliability of their measures, and we recommend a broader assessment of test-retest
reliability in infant research.

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Manybabies1 Test-Retest Supplementary Materials

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S1. Notes on and deviations from the preregistration

Below, we have compiled a list of notes on and deviations from the preregistered methods and analyses available at https://osf.io/v5f8t.

- All infants with usable data for both test and retest session were included in the
 analyses, regardless of the number of total infants a lab was able to contribute after
 exclusion. This decision is consistent with past decisions in ManyBabies projects to
 be as inclusive about data inclusion as possible (ManyBabies Consortium, 2020).
 - A small number of infants whose time between sessions exceeded 31 days were still included in the analyses (n = 3).
- Consistent with analytic decisions in ManyBabies 1 (ManyBabies Consortium, 2020), total looking times were truncated at 18 seconds (the maximum trial time) in the small number of cases where recorded looking times were slightly greater than 18s (presumably due to small measurement error in recording infant looking times).
- In assessing differences in IDS preference between test and retest sessions, we preregistered an additional linear mixed-effects model including a by-lab random slope for session. This model yielded qualitatively equivalent results (see R markdown of the main manuscript). However, the model resulted in a singular fit, suggesting that the model specification may be overly complex and that its estimates should be interpreted with caution. We therefore focused only on the first preregistered model (including only by-lab and by-participant random intercepts) in reporting the analyses in the main manuscript.
 - In assessing the reliability of IDS using a linear mixed-effects model predicting IDS preference in session 2 from IDS preference in session 1, we also assessed the robustness of the results by fitting a second preregistered model with more complex random effects structure, including a by-lab random slope for IDS preference in session 1. This model is included in the main R markdown script and yields

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- qualitatively equivalent results to the model reported in the manuscript that includes
 a by-lab random intercept only.
- We report a series of secondary planned analyses in the Supplementary Materials
 exploring potential moderating variables of time between test sessions (S2.1), the
 language background of the participants (S2.2.), and participant age (S2.3.).
 - We did not fit all models (in particular, the models investigating interactions between moderators) described in the secondary analyses of the preregistration, because our final sample size was smaller than we anticipated, which made it less feasible to investigate more complex relationships between moderators.

S2. Secondary analyses investigating possible moderating variables

S2.1. Time between test sessions

The number of days between the first and second testing session varied widely across participants (mean: 10 days; range: 1 - 49 days). We therefore tested for the possibility that the time between sessions might have an impact on test-retest reliability. We fit a linear mixed-effects model predicting IDS preference in Session 2 from IDS preference in Session 1 (mean-centered), number of days between testing sessions (mean-centered), and their interaction, including a by-lab random intercept and random slope for IDS preference in Session 1. A more complex random effects structure including additional random slopes for number of days between test sessions and its interaction with IDS preference in Session 1 did not converge. We found no evidence that the number of days between test sessions moderated the relationship between IDS preference in Session 1 and 2. Neither the main effect of time between sessions, β =-0.01, SE=0.03, t(148.70)=-0.41, p=.684, nor the interaction term, β =-0.01, SE=0.02, t(149.10)=-0.73, t=-465, showed significant effects.

69 S2.2. Language background

NAE-learning infants showed greater IDS preferences than their non-NAE counterparts in MB1. We therefore also assessed whether test-retest reliability interacted with children's language background. A linear mixed-effects model predicting IDS preference in Session 2 based on IDS preference in Session 1 (mean-centered), NAE (centered), and their interaction, including Lab as a random intercept, revealed no interaction, β =0.29, SE=0.18, t(151.30)=1.59, p=.115 (Figure 1).

⁷⁶ S2.3. Participant age

To investigate the possibility that age moderated test-retest reliability, we fit a linear mixed-effects model predicting IDS preference in Session 2 from IDS preference in Session 1

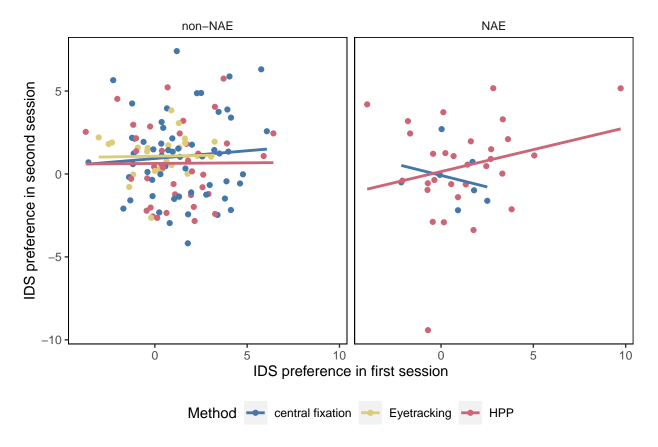
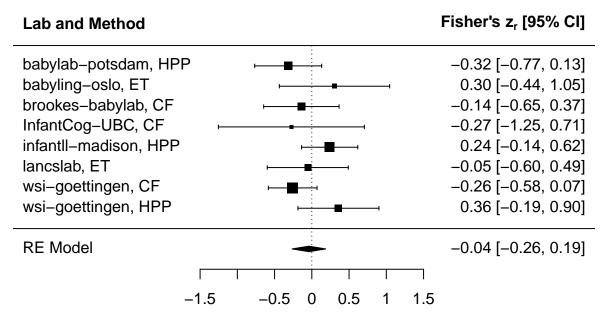


Figure 1. Infants' preference in Session 1 and Session 2 with individual data points and regression lines color-coded by method (CF, ET, or HPP). Results are plotted separately for North American English-learning infants (right panel) and infants learning other languages and dialects (right panel).

- (mean-centered), participant age (mean-centered) and their interaction. The model
- 80 included a by-lab random intercept and a by-lab random slope for IDS preference in
- Session 1. We found no evidence that age influenced test-retest reliability as indicated by
- the interaction between IDS preference in Session 1 and age, $\beta=0.00$, SE=0.00,
- t(76.60) = -0.85, p = .398.

S3. Meta-analysis of test-retest reliability



Fisher's z Transformed Correlation Coefficient

Figure 2. Forest plot of test-retest reliability effect sizes. Each row represents Fisher's z transformed correlation coefficient and 95% CI for a given lab and method (HPP = head-turn preference procedure; ET = eye-tracking; CF = central fixation). The black diamond represents the overall estimated effect size from the mixed-effects meta-analytic model.

In addition to the methods for assessing test-retest reliability reported in the main manuscript, we also investigated test-retest reliability across labs using a meta-analytic approach. We used the metafor package (Viechtbauer, 2010) to fit a mixed-effects meta-analytic model on z-transformed correlations for each combination of lab and method using sample size weighting. The model included random intercepts for lab and method. The overall effect size estimate was not significantly different from zero, b = -0.04, 95% CI = [-0.26, 0.19], p = 0.73. A forest plot of the effect sizes for each lab and method is shown in Figure 2.

Table 1

Coefficient estimates from a linear mixed-effects model predicting

Log LT IDS preference in Session 2.

	Estimate	SE	t	р
Intercept	0.14	0.07	2.05	0.09
Log LT IDS Preference Session 1	-0.06	0.09	-0.68	0.50

S4. Alternative dependent variables

To check the robustness of our results, we also investigated whether we obtained similar results with other possible dependent measures: average log-transformed looking times and a proportion-based preference measure. For each alternative dependent variable, we conducted the main analyses of test-retest reliability reported in the manuscript: the overall Pearson correlation, the test-retest linear mixed-effects model, and an inspection of applying stricter inclusion criteria for number of trials contributed.

∞ S4.1. Log-transformed looking times

In these analyses, we calculated IDS preference by first log-transforming looking 101 times for each trial, computing the average log-transformed looking time for IDS and ADS 102 for each participant, and calculating the difference between average IDS and ADS 103 log-transformed looking times. We fit a linear mixed-effects model predicting IDS preference in Session 2 from IDS preference in Session 1, including a by-lab random 105 intercept. As in the analyses using average raw looking times, the results revealed no 106 significant relationship between IDS preference in Session 1 and 2 (Table 1). The Pearson 107 correlation coefficient was also not statistically significant, r = .03, 95% CI [-.12, .19], 108 t(156) = 0.43, p = .670. Applying successively stricter inclusion criteria — by requiring a 109

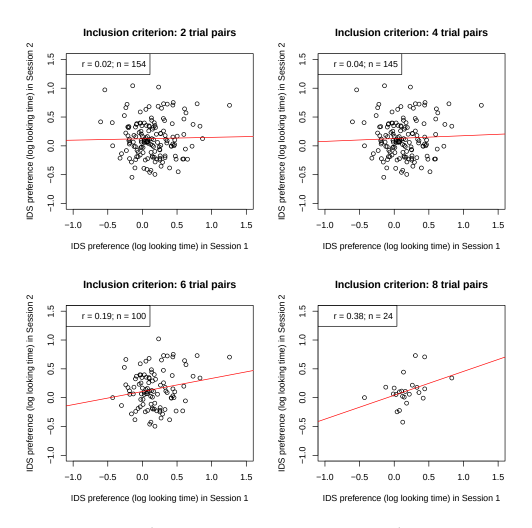


Figure 3. IDS preferences (based on average log-looking times) of both sessions plotted against each other for each inclusion criterion. n indicates the number of included infants, r is the Pearson correlation coefficient as the indicator for reliability.

higher number of valid trials per condition in each session — showed a similar pattern to the main manuscript, such that correlations increased somewhat with stricter inclusion 111 criteria, but substantially reduced the sample size at the same time (Figure 3). 112

S4.2. Proportion looking to IDS

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Next, we calculated a proportion-based IDS preference measure by computing the average proportion (raw) looking time to IDS relative to total (raw) looking time to IDS 115 and ADS for each subject (i.e., IDS looking time / (ADS looking time + IDS looking 116

time)). We fit a linear mixed-effects model predicting proportion-based IDS preference in Session 2 from proportion-based IDS preference in Session 1, including a by-lab random intercept. As in the analyses using other measures of IDS preference, the results revealed no significant relationship between IDS preference in Session 1 and 2 (Table 2). The Pearson correlation coefficient based on proportional IDS looking was also not statistically significant, r = .01, 95% CI [-.15, .16], t(156) = 0.09, p = .927. Stricter inclusion criteria increased the correlation somewhat, as in previous analyses (Figure 4).

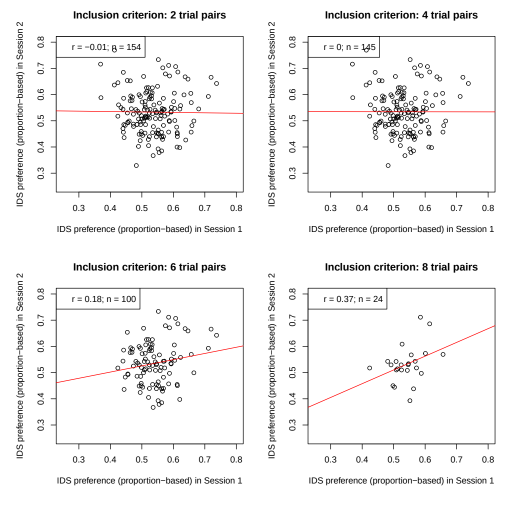


Figure 4. IDS preferences (based on proportion IDS looking) of both sessions plotted against each other for each inclusion criterion. n indicates the number of included infants, r is the Pearson correlation coefficient as the indicator for reliability.

Table 2

Coefficient estimates from a linear mixed-effects model predicting IDS preference

(based on proportion IDS looking) in Session 2.

	Estimate	SE	t	р
Intercept	0.59	0.05	10.70	0.00
IDS Preference (proportion measure) Session 1	-0.10	0.10	-1.01	0.31

S5. Sensitivity of test-retest reliability to trial number inclusion criteria

To conduct a more fine-grained analysis of how stricter trial inclusion criteria affect
test-retest reliability, we computed correlations while gradually increasing the number of
total valid trials required for inclusion. For this analysis, we required a minimum of one
IDS and one ADS trial and gradually increased the number of total valid trials required in
both sessions (irrespective of IDS and ADS condition) from 2 to 16 (the maximum number
of total trials). Figure 5 depicts the Pearson correlation coefficients for increasingly stricter
requirements for the overall trial numbers of a given participant in both sessions.

Correlations only increase and reach conventional levels of significance once the number of
total required trials for both sessions is greater than 12.

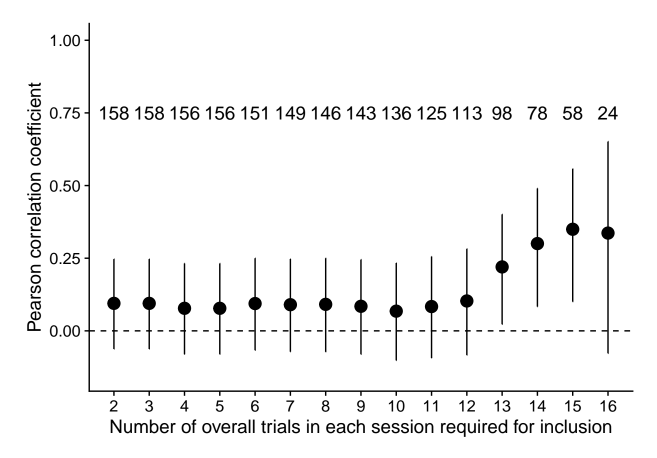


Figure 5. Pearson correlation coefficient with increasingly strict trial-level inclusion criteria. The x-axis depicts the required number of overall valid trials in both session 1 and session 2. Dots represent corresponding correlation coefficients, with 95 percent CIs. The sample size is shown above each dot.

S6. Patterns of preference across sessions

We also conducted analyses to explore whether there were any patterns of preference reversal across test sessions. While there was no strong correlation in the magnitude of IDS preference between test session 1 and test session 2, here we asked whether infants consistently expressed the same preference across test sessions. Overall, 58.20% of the infants had a consistent preference from test to retest session. Of the 158 total infants, 44.90% of infants showed a consistent IDS preference and 13.30% showed a consistent ADS preference. 23.40% of infants switched from an IDS preference at test session 1 to an ADS

 $_{142}$ preference at test session 2 and 18.40% switched from an ADS preference to an IDS $_{143}$ preference.

Next, we explored whether we could detect any systematic clustering of infants with 144 distinct patterns of preference across the test and retest session. We took a bottom-up 145 approach and conducted a k-means clustering of the test-retest difference data (here using 146 log-transformed looking time data). We found little evidence of distinct clusters emerging 147 from these groupings: the clusterings ranging from k=2 (2 clusters) to k=4 (4 clusters) 148 appear to mainly track whether participants are approximately above or below the mean 149 looking time difference for test session 1 and test session 2 (Figure 6A). The diagnostic elbow plot shows little evidence of a qualitative improvement as the number of clusters is 151 increased, which suggests little evidence for a distinctive set of clusters of participants who 152 showed similar patterns of looking across the test and retest sessions (Figure 6B). 153

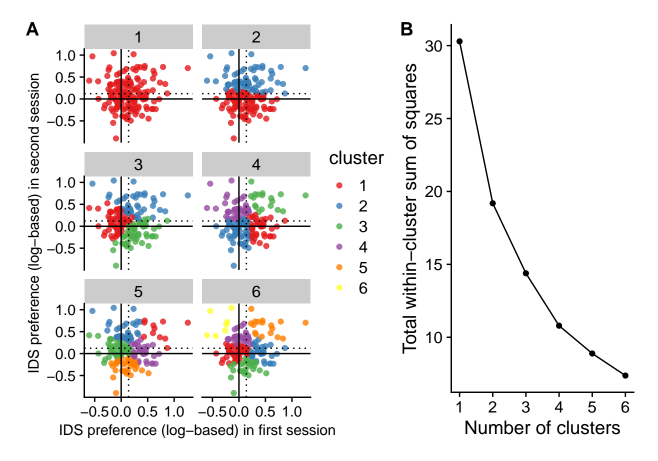


Figure 6. (A) Results from the k-means clustering analysis of IDS preference (based on average log looking times) in session 1 and 2 for different numbers of k and (B) the corresponding elbow plot of the total within-cluster sum of squares. In (A), points represent indvidual participants' magnitude of looking time difference at test sessions 1 (x-axis) and 2 (y-axis). The solid line indicates no preference for IDS vs. ADS, the dotted lines indicate mean IDS preference at test session 1 and 2, respectively. Colors indicate clusters from the k-means clustering for different values of k.

S7. Relation between number of contributed trials in each session

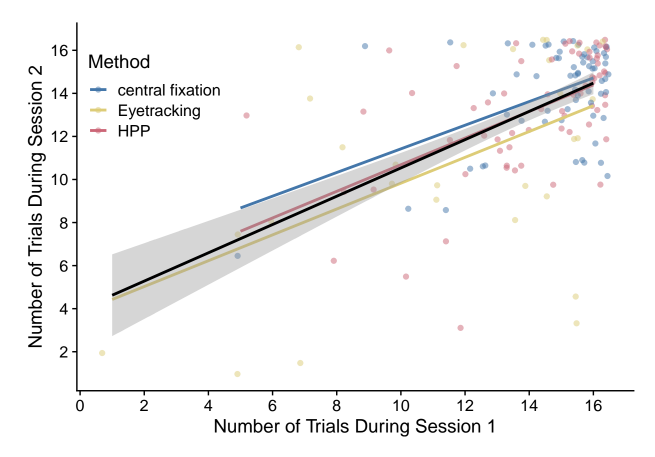


Figure 7. Correlation between the number of trials contributed in Session 1 and Session 2. Each data point represents one infant. Colored lines represent linear fits for each method.

Are there stable individual differences in how likely an infant is to contribute a high 155 number of trials? To answer this question, we conducted an exploratory analysis 156 investigating whether there is a relationship between the number of trials an infant 157 contributed in Session 1 and Session 2. Do infants who contribute a higher number of trials 158 during their first testing session also tend to contribute more trials during their second testing session? A positive correlation between trial numbers during the first and second session would indicate that there is some stability in a given infants' likelihood of 161 remaining attentive throughout the experiment. On the other hand, the absence of a 162 correlation would indicate that the number of trials a given infant contributes is not 163 predictive of how many trials they might contribute during their next session. 164

We found a strong positive correlation between number of trials contributed during 165 the first and the second session r = .58, 95% CI [.47, .68], t(159) = 9.05, p < .001 (Figure 166 7). This result suggests that if infants contribute a higher number of trials in one session, 167 compared to other infants, they are likely to contribute a higher number of trials in their 168 next session. This finding is consistent with the hypothesis that how attentive infants are 169 throughout an experiment (and hence how many trials they contribute) is a stable 170 individual difference, at least for some infant looking time tasks. Researchers should 171 therefore be mindful of the fact that decisions about including or excluding infants based on 172 trials contributed may selectively sample a specific sub-set of the infant population they are 173 studying (Byers-Heinlein, Bergmann, & Savalei, 2021; DeBolt, Rhemtulla, & Oakes, 2020). 174

S8. Correlations in average looking times between sessions

To what extent are participants looking times between the two sessions related? To 176 test this question, we first investigated whether participants' overall looking times — 177 irrespective of condition — were correlated between the first and second session. There was 178 a robust correlation between average looking time in Session 1 and Session 2: infants with 179 longer looking times during their first session also tended to look longer during their second 180 session, r = .45, 95% CI [.31, .57], t(156) = 6.28, p < .001. This relationship held even after 181 controlling for number of trials in the first and second session, suggesting that the relation 182 between average looking in Session 1 and 2 could not be entirely explained by the 183 correlation in the number of trials contributed between the two sessions (S7), b = 0.42, 95%CI [0.27, 0.58], t(154) = 5.52, p < .001 (Figure 8A). The result is also similar when 185 controlling for participants' average age across the two test sessions, b = 0.44, 95% CI 186 [0.30, 0.59], t(155) = 6.16, p < .001.187

Next, we explored the extent to which average looking times for IDS and ADS stimuli 188 were related. First, we found similar correlations in average looking time to IDS stimuli in 189 Session 1 and 2, r = .38, 95% CI [.24, .51], t(156) = 5.19, p < .001, and ADS stimuli in 190 Session 1 and 2, r = .40, 95% CI [.26, .53], t(156) = 5.49, p < .001 (Figure 8B). To test 191 whether these correlations were specific to looking times for IDS or ADS stimuli alone, we 192 fit linear regression models predicting average looking to IDS (or ADS) stimuli in Session 2 193 from average looking to IDS and ADS stimuli in Session 1. We found that average looking 194 to IDS stimuli in Session 2 could be predicted from average looking to IDS stimuli in Session 1, even after controlling for average looking to ADS stimuli in Session 1, b = 0.21, 95% CI [0.01, 0.41], t(155) = 2.11, p = .037. Conversely, average looking to ADS stimuli in 197 Session 2 could be predicted from average looking to ADS stimuli in Session 1, even after 198 controlling for average looking to IDS stimuli in Session 1, b = 0.36, 95% CI [0.14, 0.58], 199 t(155) = 3.20, p = .002. These results suggest that the condition-specific correlations in 200

²⁰¹ average looking time cannot be fully explained by the fact that infants' overall looking times between sessions are correlated.

Finally, we inspected item-level correlations between the two test sessions.

Specifically, we investigated the relation between items composed of the same recording clips in Session 1 and Session 2 (but with a reversed order of clips between the two sessions). We fit a linear mixed-effects model predicting item-level looking time in Session 2 from item-level looking time in Session 1, including random intercepts for participant, item, and lab, as well as a random slope for item-level looking time in Session 1 for participant and lab. Item-level looking in Session 2 was related to item-level looking in Session 1, $\hat{\beta} = 0.17$, 95% CI [0.07, 0.27], t(5.52) = 3.38, p = .017 (Figure 8C). Similar results hold if looking times are log-transformed

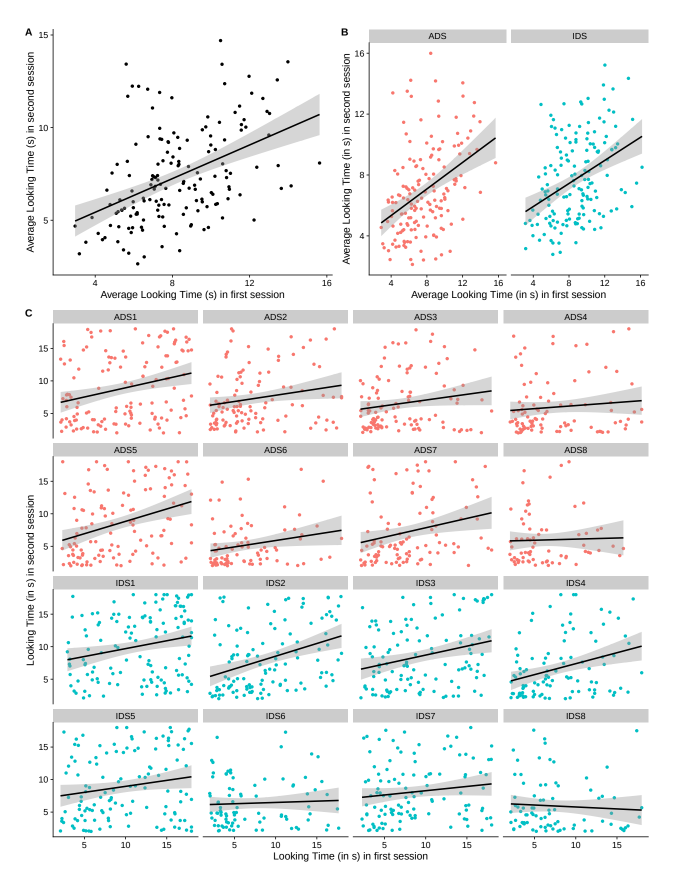


Figure 8. Correlations in average looking time (in s) between Session 1 and 2 (A) overall, (B) by condition, and (C) by item.

Table 3

Linear mixed-effects model results predicting IDS

preference in Session 2 from IDS preference in

Session 1 at the stimulus level.

Term	\hat{eta}	95% CI	t	df	p
Intercept	1.02	[0.14, 1.90]	2.27	6.55	.060
Diff 1	0.07	[-0.01, 0.14]	1.79	718.46	.074

S9. By-item-pair preference scores across sessions

Finally, we inspected on a more fine-grained item level whether IDS preference in 213 Session 1 was related to IDS preference in Session 2. To do so, we exploited the fact the specific IDS and ADS stimuli were paired together in test orders in both sessions, such that 215 one IDS stimulus (e.g., IDS1) always occurred adjacently to a specific ADS stimulus (e.g., 216 ADS1). We therefore computed stimulus-specific IDS preference scores by calculating the 217 difference in raw looking time for each of the eight IDS-ADS stimulus pairs for each 218 participant (whenever both trials in a given pair were available). We then fit a linear 219 mixed-effects model predicting stimulus-specific IDS preference in Session 2 from 220 stimulus-specific IDS preference in Session 1, including by-participant and by-lab random 221 intercepts (models with more complex random effects structure, including by-item random 222 effects, failed to converge). There was a marginal, but non-significant relation in 223 stimulus-specific IDS preference between the two test sessions (Table 3). 224

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