- A baseline for inferences about human cognitive evolution: structure, stability and
- predictability of great ape cognition
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22 Abstract

Uniquely human cognition is often defined by invoking comparisons to non-human great 23 apes. This approach requires a comprehensive way of describing great ape cognition in order to specify how it differs from that of humans: a great ape baseline. DEfining such a baseline 25 entails that a) group-level results are stable, b) individual differences are measured reliably, 26 and c) cognitive performance is predictable. These prerequisites, however, are rarely tested 27 empirically. This study addresses this shortcoming. We repeatedly tested a comparatively large sample of great apes in five tasks covering a range of cognitive domains. We found group-level performance to be relatively stable so that different testing occasions licensed the same conclusions. Most of the tasks showed good re-test reliability and were thus suited to 31 study individual differences. Using structural equation models, we found that individual differences in performance were largely explained by stable differences in cognitive abilities between individuals. Furthermore, we found systematic relations between cognitive abilities. When predicting cognitive performance by variables capturing individual characteristics (e.g., 35 group, sex or age) or aspects of everyday experience (e.g., life events, testing arrangements or 36 sociality), we found stable individual characteristics to be the most important. Taken 37 together, this study provides an empirical basis for assumptions implicit in much 38 comparative research. Based on the results, great ape cognition can be described as being structured by stable differences in cognitive abilities between individuals that develop in 40 response to different – put partially overlapping – external conditions. 41

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Introduction

In their quest for understanding the evolution of the human mind, psychologists and 47 cognitive scientists face one major obstacle: cognition does not fossilize. Instead of directly 48 studying the cognitive abilities of our extinct ancestors, we have to rely on backward inferences. We can study fossilized skulls and crania to approximate brain size and structure and use this information to infer cognitive abilities (Coqueugniot, Hublin, Veillon, Houët, & Jacob, 2004; Gunz et al., 2020). We can study the material culture left behind by our ancestors and try to infer its cognitive complexity experimentally (Coolidge & Wynn, 2016; Currie & Killin, 2019; Haslam et al., 2017). Yet, the archaeological record is sparse and only goes back so far in time. Thus, one of the most fruitful approaches to cognitive evolution is the comparative method. By studying extant species of primates, we can make backward inferences about the last common ancestor. If species A and B both show cognitive ability X, the last common ancestor of A and B most likely also had ability X (Burkart, Schubiger, & Schaik, 2017; MacLean et al., 2012; Martins & Martins, 1996). To make inferences about the most recent events in human cognitive evolution, we have to study and compare humans and the non-human great apes. Such an approach has been highly productive and provides the empirical basis for numerous theories about human cognitive uniqueness (Dean, Kendal, Schapiro, Thierry, & Laland, 2012; Dunbar & Shultz, 2017; Heyes, 2018; Laland & Seed, 2021; Penn, Holyoak, & Povinelli, 2008; Tomasello, 2019).

However, using the comparative method in this way requires a strong great ape
baseline. That is, it takes a solid and robust way of describing the great ape mind in order
to map out how it differs from that of humans. What kind of empirical evidence is required
to define such a baseline? First, group-level results should be stable. Our inferences about
the cognitive abilities that great apes – as a group or species – do or do not have based on

the data we collect today should not change if we repeat the study tomorrow. Second, individual differences in cognitive abilities should be reliable. That is, methods and procedures should also reliably measure cognitive abilities on an individual level. This is a prerequisite for investigating the relations between different tasks in order to map out the internal structure of great ape cognition (Matzel & Sauce, 2017; Shaw & Schmelz, 2017; Thornton & Lukas, 2012; Volter, Tinklenberg, Call, & Seed, 2018). Finally, individual differences should be predictable. Understanding great ape cognition means that we can point to variables that describe individual characteristics or aspects of everyday experience that induce variation in cognitive performance and development.

Recently, a number of concerns have been voiced, questioning whether the prototypical way of conducting comparative research is suited to provide the empirical basis for inferring the great ape baseline (Farrar & Ostojic, 2019; ManyPrimates, Altschul, Beran, Bohn, Caspar, et al., 2019; Schubiger, Fichtel, & Burkart, 2020; Stevens, 2017). A key point in this criticism is that most research simply assumes that the three requirements outlined above are met without testing them empirically. The work reported here directly addresses this fundamental problem.

There are, however, several notable exceptions that undertook great effort to provide a more comprehensive picture of one or more aspects of the nature and structure of great ape cognition (Beran & Hopkins, 2018; Hopkins, Russell, & Schaeffer, 2014; MacLean et al., 2014; Wobber, Herrmann, Hare, Wrangham, & Tomasello, 2014). Herrmann and colleagues (Herrmann, Call, Hernández-Lloreda, Hare, & Tomasello, 2007) tested a large sample of great apes (chimpanzees and orangutans) and human children in a range of tasks from different cognitive domains. The results indicated pronounced group-level differences between great apes and humans in the social, but not the physical domain. Furthermore, relations between the tasks pointed to a different internal structure of cognition, with a distinct social cognition factor for humans but not great apes (Herrmann, Hernández-Lloreda, Call, Hare, & Tomasello, 2010). Völter and colleagues (Völter et al.,

2022) focused on the structure of executive functions. Based on a multi-trait multi-method approach they developed a new test battery to assess memory updating, inhibition, and attention shifting in chimpanzees and human children. Overall, they found low correlations between tasks and thus no clear support for any of the structures put forward by theoretical models built around adult human data.

Despite their seminal contributions to the field, these studies suffer from one or more of
the three shortcomings outlined above. It is unclear if the results are stable. That is, if the
same individuals were tested again, would we see the same results and arrive at the same
conclusions about absolute differences between species. Furthermore, the psychometric
properties of the tasks are unknown and it is thus unclear if, for example, low correlations
between tasks reflect a genuine lack of shared cognitive processes or simply measurement
imprecision. Finally, it remains unclear what causes individual differences – which individual
characteristics and experiences predict cognitive performance and development.

The studies reported below seek to solidify the empirical grounds of the great ape 110 baseline. For one-and-a-half years, every two weeks we administered a set of five cognitive 111 tasks (see Figure 1)) to the same population of great apes (N=43). The tasks spanned 112 across cognitive domains and were based on published procedures widely used in the field of 113 comparative psychology. As a test of social cognition, we included a gaze following task 114 (Brauer, Call, & Tomasello, 2005). To assess reasoning abilities, we included a causal 115 reasoning and an inference by exclusion task (Call, 2004). Numerical cognition was tested 116 using a quantity discrimination task (Hanus & Call, 2007). Finally, as a test of executive 117 functions, we included a delay of gratification task (Rosati, Stevens, Hare, & Hauser, 2007). 118

In addition to the cognitive data, we continuously collected 14 variables that capture stable and variable aspects of our participants' life and used this to predict inter- and intra-individual variation in cognitive performance. Data collection was split into two phases.

After Phase 1 (14 data collection time points), we analysed the data and registered the

results (https://osf.io/7qyd8). Phase 2 lasted for another 14 time points and served to replicate and extend Phase 1. This approach allowed us to test a) how stable group level results are, b) how reliable individual differences are, c) how individual differences are structured and d) what predicts cognitive performance.

127 Results

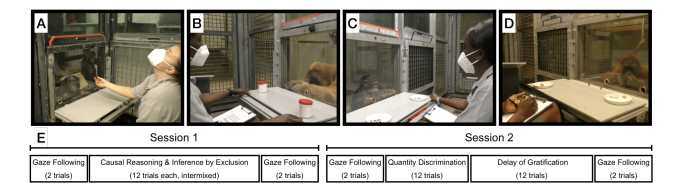


Figure 1. Setup used for the six tasks. A) Gaze following: the experimenter looked to the ceiling. We coded if the ape followed gaze. B) Causal reasoning: food was hidden in one of two cup, the baited cup was shaken (food produced a sound) and apes had to choose the shaken cup to get food. Inference by exclusion: food was hidden in one of two cups. The empty cup was shaken (no sound) so apes had to choose the non-shaken cup to get food. C) Quantity discrimination: Small pieces of food were presented on two plates (5 vs. 7 items); we coded if subjects chose the larger amount. D) Delay of gratification (only Phase 2): to receive a larger reward, the subject had to wait and forgo a smaller, immediately accesible, reward. E) Order of task presentation and trial numbers

Stability of group-level performance

Group-level performance was largely stable or followed clear temporal patterns (see Figure 2). The causal inference and quantity discrimination tasks were the most robust: in

both cases performance was clearly different from chance across both phases with no apparent change over time. The rate of gaze following declined in the beginning of Phase 1 132 but then settled on a low but stable level until the end of Phase 2. This pattern was 133 expected given that following the experimenters gaze was never rewarded – neither explicitly 134 with food or by bringing something interesting to the participant's attention. The inference 135 by exclusion task showed an inverse pattern with group-level performance being at 136 chance-level for most of Phase 1, followed by a small, but steady, increase throughout Phase 137 2. These temporal patterns most likely reflect training (or habituation) effects that are a 138 consequence of the repeated testing. Performance in the delay of gratification task (Phase 2 139 only) was more variable, but within the same general range for the whole testing period. In 140 sum, performance was very robust in that time points generally licensed the same group-level 141 conclusions. The tasks appeared well suited to study group-level performance. In the supplementary material, we report additional analysis – latent state structural equation models – that corroborate this interpretation.

Reliability of individual differences

Stable group-level performance does not imply stable individual differences. In fact, a 146 well-known paradox in human cognitive psychology states that some of the most robust – on 147 a group level – cognitive tasks do not produce reliable individual differences (Hedge, Powell, 148 & Sumner, 2018). In a second step, we therefore assessed the re-test reliability of our five 149 tasks. For that, we correlated the performance at the different time points in each task. 150 Figure 3 visualizes these raw re-test correlations. Correlations were generally high – 151 exceptionally high for animal cognition standards (Cauchoix et al., 2018). As expected, 152 values were also higher for time points closer in time (Uher, 2011). The quantity 153 discrimination task was less reliable compared to the other tasks.

What stands out in this is that *stability does not imply reliability* - and vice versa. The quantity discrimination task showed robust group-level performance above chance but

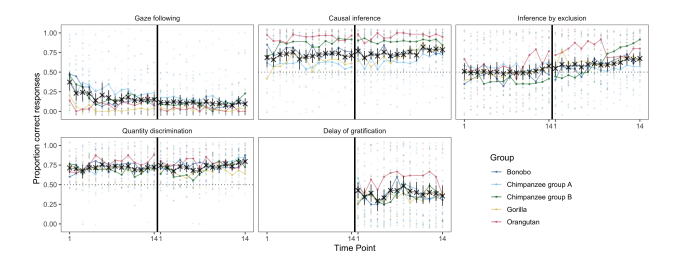


Figure 2. Results from the five cognitive tasks across time points. Black crosses show mean performance at each time point across species (with 95% CI). Colored dots show mean performance by species, light dots show individual means per time point. Dashed line shows chance level whenever applicable. The vertical back line marks the transition between phase 1 and 2.

relatively poor re-test reliability. Group-level performance in the inference by exclusion and gaze following tasks changed over time but were highly reliable on an individual level. Taken together, the majority of tasks were well suited to study individual differences.

Structure of individual differences

Next, we investigated the structure of these individual differences. First, we asked to
what extent individual differences reflect stable differences in cognitive abilities. We used
structural equation modelling – in particular latent state-trait models (LSTM) – to partition
the variance in performance into latent traits (*Consistency*) and states (*Occasion specificity*)
(Geiser, 2020; Steyer, Ferring, & Schmitt, 1992; Steyer, Mayer, Geiser, & Cole, 2015). In the
present context, one can think of a latent trait as a stable cognitive ability (e.g. ability to
make causal inferences) and states as time-specific, variable psychological conditions
(e.g. variations in performance due to being more or less attentive or motivated). These

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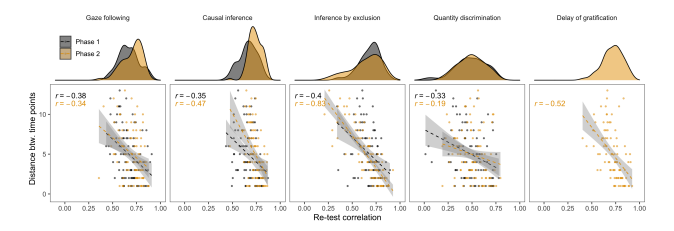


Figure 3. Top: Distribution of re-test correlation coefficients between time points for each task. Bottom: Correlations between re-test reliability coefficients and temporal distance between the testing time points.

latent variables are measurement-error free because they are estimated taking into account
the reliability of the task. In the LSTM context, reliability is the correlation between task
and occasion specific test-halves. We report additional models that account for the temporal
structure of the data in the supplementary material.

Individual differences were largely explained by stable differences in cognitive abilities. Across tasks, more than 75% of variance was accounted for by latent trait differences and less than 25% by state differences (Figure 4A). The high reliability estimates show that these latent variables accounted for most of the variance in raw test scores – with the quantity discrimination task being, once again, an exception. The estimates for consistency and occasion specificity were also remarkably similar for the two phases.

For inference by exclusion, we could not fit an LST model to the data from Phase 2.

Instead, we had to divide Phase 2 in two parts (time point 1-8 and 9-14) and estimate a

separate trait for each part. The estimates for the proportion of variance explained by states

and traits were similar for both parts (Figure 4A) and the two traits were highly correlated (r = .82). Together with additional latent sate models which we report in the supplementary

material, this suggests that the increase in group-level performance in Phase 2 was driven by

a relatively sudden improvement of a few individuals, mostly from the chimpanzee B group (see Figure 2). These individuals "rose through the ranks" half-way through Phase 2 and then retained this position for the rest of the study. Some of the orangutans changed in the opposite direction – though to a lesser extend.

As the second step, we investigated the relations between latent traits. That is, we 189 asked whether individuals with high abilities in one domain also have higher abilities in 190 another. We fit pairwise LST models that modeled the correlation between latent traits for 191 two tasks (two models for inference by exclusion in Phase 2). In Phase 1, the only correlation that was reliably different from zero was the one between quantity discrimination 193 and inference by exclusion. In Phase 2, this finding was replicated and, in addition, four more correlations turned out to be substantial (see Figure 4B). One reason for this increase 195 was the inclusion of the delay of gratification task. Across phases, correlations involving the 196 gaze following task were the closest to zero, with quantity discrimination in Phase 2 being an 197 exception. Taken together, the overall pattern of results suggests substantial shared variance 198 between tasks – except for gaze following. 199

200 Predictability of individual differences

The results thus far suggest that individual differences originate from stable differences 201 in cognitive abilities. In the last set of analysis, we sought to explain the origins of these 202 differences. That is, we analysed whether inter- or intra-individual variation in cognitive 203 performance in the tasks could be predicted by non-cognitive variables that capture a) stable 204 differences between individuals (group, age, sex, rearing history, time spent in research), b) differences that vary within and between individuals (rank, sickness, sociality), c) differences that vary with group membership (time spent outdoors, disturbances, life events), and d) differences in testing arrangements (presence of observers, study participation on the same 208 day and since the last time point). We collected these predictor variables using a 200 combination of directed observations and caretaker questionnaires. 210

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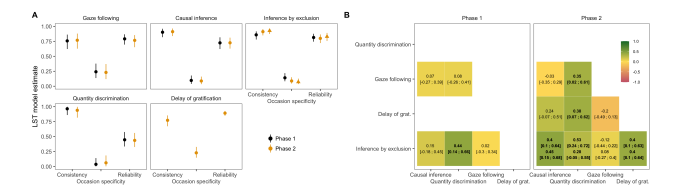


Figure 4. A. Estimates from latent state-trait model for Phase 1 and 2. Consistency: proportion of (measurement-error free) variance in performance explained by stable trait differences. Occasion specificity: variance explained by variable states. Reliability: proportion of variance in raw scores explained by the trait and the state. For inference by exclusion: different shapes show estimates for different parts of Phase 2 (see main text for details). B. Correlations between latent traits based on pairwise LST models between tasks with 95% Credible Interval. Bold correlations are reliably different from zero. Inference by exclusion has one value per part in Phase 2. The models for quantity discrimination and causal inference showed a poor fit and are not reported here (see supplementary material for details).

This large set of potentially relevant predictors poses a variable selection problem. In 211 our analysis we sought to find the minimal set of predictors that allowed us to accurately 212 predict performance in the cognitive tasks. We chose the projection predictive inference 213 approach because it provides and excellent trade-off between model complexity and accuracy 214 (Pavone, Piironen, Bürkner, & Vehtari, 2020; Piironen, Paasiniemi, & Vehtari, 2020; Piironen 215 & Vehtari, 2017). The outcome of this analysis is a ranking of the different predictors in 216 terms of how important they are to predict performance in a given task. Furthermore, for 217 each predictor, we get a qualitative assessment of whether it makes a substantial 218 contribution to predicting performance in the task or not. 219

Predictors capturing stable individual characteristics were ranked highest and selected 220 as relevant most often (Figure 5). Of the five highest-ranked predictors, four belonged to this category. This result aligned well with the SEM results reported above, in which we saw that
most of the variance in performance could be traced back to stable trait differences between
individuals. The tasks with the highest occasion specific variance (gaze following and delay of
gratification, see Figure 4) were also those for which the most time point specific predictors
were selected. The quantity discrimination task did not fit this pattern in Phase 2; even
though the LSTM suggested that only a very small portion of the variance in performance
was occasion specific, three time point specific variables were selected to be relevant.

The most important predictor was group. Interestingly, differences between groups 229 were not systematic in that one group would consistently outperform the others across tasks. Furthermore, group differences could not be collapsed into species differences as the two chimpanzee groups varied largely independent of one another. The way in which the 232 predictors that were selected more than once influenced performance was largely coherent. 233 The presence of observers always had a negative effect on performance. The more time an 234 individual had been involved in research, the better performance was. Higher-ranking 235 individuals outperformed lower-ranking ones. Exceptions were age and sex: while the rate of 236 gaze following increased with age, performance in the inference by exclusion task decreased. 237 Females were more likely to follow gaze than males, but males were more likely to wait for 238 the larger reward in the delay of gratification task (Figure 5B). 230

In sum, of the predictors we recorded, those capturing stable individual characteristics
were most predictive of cognitive performance. However, the the largest portion of the
variance in cognitive performance was accounted for by a random intercept term in the
model (not shown in Figure 5) that simply captured the identity of the individual (see
supplementary material for details). This suggests that idiosyncratic developmental
processes or genetic pre-dispositions, which operate on a much longer time-scale than what
we captured in the present study, were responsible for most of the variation in cognitive
performance.

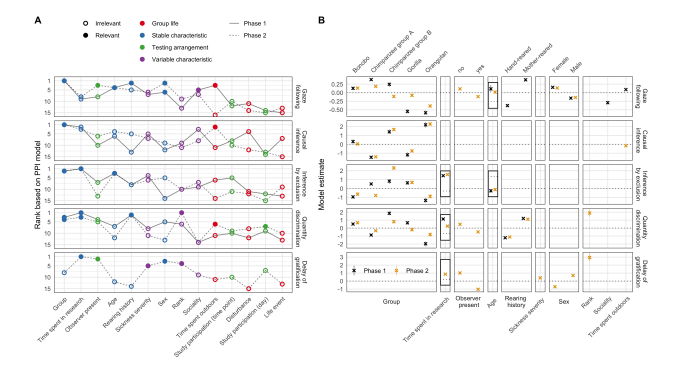


Figure 5. A. Ranking of predictors based on the projection predictive inference model for the five tasks in the two phases. Order (left to right) is based on average rank across phases. Solid points indicate predictors selected as relevant. Color of the points shows the category of the predictor. Line type denotes the phase. B. Posterior model estimates for the selected predictors for each task. Crosses show the mean of the posterior distribution and error bars the 95% Credible Interval. Color denotes phase. Black rectangels zoom in on the perdictors that are numerically small to show whether they have a positive or negative influence on performance.

248 Discussion

The goal of this study was to test the assumptions underlying much of comparative 249 research and theorizing about human cognitive evolution. We repeatedly tested a large 250 sample of great apes in five tasks covering a range of different cognitive domains. We found 251 group-level performance to be relatively stable so that conclusions drawn based on one 252 testing occasion mirrored those on other occasions. Most of the tasks showed good re-test 253 reliability and were thus suited to study individual differences. Using structural equation 254 models, we found that individual differences in performance was largely explained by stable differences in cognitive abilities between individuals. Furthermore, we found systematic relations between cognitive abilities. When predicting cognitive performance by variables capturing individual characteristics or aspects of everyday experience, we found stable individual characteristics (e.g., group) to be the most important. However, most of the 259 variance in performance remained unexplained. 260

At first glance, the results send a reassuring message to the field: most of the tasks that 261 we used produced stable group-level results and captured individual differences in a reliable 262 way. However, this did not apply to all tasks. In the supplementary material, we report on a 263 rule-switching task (Haun, Call, Janzen, & Levinson, 2006) that produced neither stable nor 264 reliable results. The quantity discrimination task was stable on a group level but less reliable. 265 We draw two conclusions based on this pattern. First, replicating studies – even if it is with 266 the same animals – should be an integral part of primate cognition research (Farrar, Boeckle, 267 & Clayton, 2020; ManyPrimates, Altschul, Beran, Bohn, Caspar, et al., 2019; Stevens, 2017). Second, for individual differences research, it is crucial to assess the reliability of the measures involved (Fried & Flake, 2018). If this step is skipped, it is difficult to interpret studies, especially when they produce null-results. It is important to note that the sample 271 size in the current study was large compared to other comparative studies (ManyPrimates, 272 Altschul, Beran, Bohn, Caspar, et al., 2019). With smaller sample sizes, group-level and

reliability estimates are more likely to be more variable and thus more likely to produce
false-positive or false-negative conclusions (Forstmeier, Wagenmakers, & Parker, 2017; Oakes,
2017). Small samples in comparative research usually reflect resource limitations of
individual labs. Pooling resources in large-scale collaborative projects like *ManyPrimates*(ManyPrimates et al., 2021; ManyPrimates, Altschul, Beran, Bohn, Call, et al., 2019) will
thus be vital to corroborate findings. Some research questions – for example the distinction
between group- vs. species-level explanations of primate cognitive performance (Van
Leeuwen, Cronin, & Haun, 2018) we hinted at – cannot even be addressed by a single lab.

Given their good psychometric properties, our tasks offer insights into the structure of 282 great ape cognition. We used structural equation modelling to partition variance in 283 performance into stable (trait) and variable (state) differences between individuals. We 284 found traits to explain more than 75% of the variance across tasks. This suggests that stable 285 differences in cognitive abilities and not variable differences in e.g. attention and motivation 286 are responsible for the patterns we observed. This finding does not mean that there is no 287 developmental change over time. In fact, for the inference by exclusion task we saw a 288 relatively abrupt change in performance for some individuals which stabilized on an elevated 289 level, suggesting sustained change in cognitive abilities. With respect to structure, we found systematic relations between traits estimated via LSTMs for the different tasks. Correlations tended to be higher for the non-social tasks compared to when gaze following was involved which could be taken to hint at shared cognitive processes. However, we feel such a 293 conclusion would be premature and would require additional evidence from more tasks and 294 larger sample sizes (Herrmann et al., 2010). Furthermore, cognitive modeling could be used to explicate the processes involved in each task. Shared processes could be probed by 296 comparing models that make different assumptions about overlapping processes (Bohn, 297 Liebal, & Tessler, n.d.; Devaine et al., 2017). 298

The finding that stable differences in cognitive abilities explained most of the variation
between individuals was also corroborated by the analyses focused on the predictability of

performance. We found that predictors that captured stable individual characteristics 301 (e.g. group, time spent in research, age, rearing history) were more likely to improve model 302 fit. Aspects of everyday experience or testing arrangements that would influence 303 performance on particular time points and thus increase the proportion of occasion specific 304 variation (e.g. life events, disturbances, participating in other tests) were ranked as less 305 important. Despite this general pattern, there was, however, variation across tasks in which 306 individual characteristics were selected to be relevant. For example, rearing history turned 307 out to be an important predictor for quantity discrimination and gaze following but less so for the other three tasks (Figure 5A). Group – the overall most important predictor – 309 exerted its influence differently across tasks. Orangutans, for example, outperformed the 310 other groups in causal inference but were the least likely to follow gaze. Together with the 311 finding that the random intercept term improved model fit the most across tasks, this pattern suggests that the cognitive abilities underlying performance in the different tasks 313 respond to different – though sometimes overlapping – developmental conditions.

Our results also speak to very general issue. Comparative psychologists often worry – 315 or are told they should worry – that their results can be explained by rapid associative 316 learning processes (Hanus, 2016). Oftentimes such explanations are theoretically plausible 317 and hard to disprove. The present study speaks to this issue in so far as we created optimal 318 conditions for such associative learning processes to happen. Great apes were tested by the 319 same experimenter, in the same tasks, using differential reinforcement and the same 320 counterbalancing for hundreds of trials. The fast and steady increase in performance – uniform for all individuals – that an associative learning account would predict did not show. 322 Instead, when we saw change over time, performance either decreased (gaze following) or increased at a late point in time for only a few individuals (inference by exclusion). This does not take away they the theoretical possibility of associative learning accounts, it just 325 makes them less useful given that their predictions do not bear out.

327 Conclusion

Uncovering what makes human cognition unique requires a solid understanding of 328 great ape cognition. The present study puts the assumptions underlying much of 329 comparative research to an empirical test. While we found reassuring results in terms of 330 group-level stability and reliability of results, we also pointed out the importance of 331 explicitly questioning and testing these assumptions, ideally in large-scale collaborative 332 projects. Our results paint a picture of great ape cognition in which variation between individuals is predicted and explained by stable individual characteristics that respond to different – though sometimes overlapping – developmental conditions. We hope these results contribute to a more solid and comprehensive understanding of the nature and origins of great ape and human cognition. 337

338 Methods

A detailed description of the methods and results can be found in the supplementary material available online. All data and analysis scripts can be found in the associated online repository (https://github.com/ccp-eva/laac).

342 Participants

A total of 43 great apes participated at least once in one of the tasks. This included 8
Bonobos (3 females, age 7.30 to 39), 24 Chimpanzees (18 females, age 2.60 to 55.90), 6
Gorillas (4 females, age 2.70 to 22.60), and 6 Orangutans (4 females, age 17 to 41.20). The
sample size at the different time points ranged from 3 to 18 for the different species.

Apes were housed at the Wolfgang Köhler Primate Research Center located in Zoo
Leipzig, Germany. They lived in groups, with one group per species and two chimpanzee
groups (group A and B). Research was noninvasive and strictly adhered to the legal
requirements in Germany. Animal husbandry and research complied with the European

Association of Zoos and Aquaria Minimum Standards for the Accommodation and Care of
Animals in Zoos and Aquaria as well as the World Association of Zoos and Aquariums
Ethical Guidelines for the Conduct of Research on Animals by Zoos and Aquariums.

Participation was voluntary, all food was given in addition to the daily diet, and water was
available ad libitum throughout the study. The study was approved by an internal ethics
committee at the Max Planck Institute for Evolutionary Anthropology.

357 Material

Apes were tested in familiar sleeping or observation rooms by a single experimenter.
Whenever possible, they were tested individually. The basic setup comprised a sliding table
positioned in front of a clear Plexiglas panel with three holes in it. The experimenter sat on
a small stool and used an occluder to cover the sliding table (see Figure 1).

362 Procedure

The tasks we selected are based on published procedures and are commonly used in the field of comparative psychology. Example videos for each task can be found in the associated online repository.

Gaze Following. The gaze following task was modeled after Brauer et al. (2005). 366 The experimenter sat opposite the ape and handed over food at a constant pace. That is, 367 the experimenter picked up a piece of food, briefly held it out in front of her face and then 368 handed it over to the participant. After a predetermined (but varying) number of food items had been handed over, the experimenter again picked up a food item, held it in front of her 370 face and then looked up (i.e., moving her head up - see Figure 1A). The experimenter looked to the ceiling, no object of particular interest was placed there. After 10s, the experimenter 372 looked down again, handed over the food and the trial ended. We coded whether the 373 participant looked up during the 10s interval. Apes received eight gaze following trials. We assume that participants look up because they assume that the experimenter's attention is focused on a potentially noteworthy object.

Causal inference. The causal inference task was modeled after Call (2004). Two 377 identical cups with a lid were placed left and right on the table (Figure 1B). The 378 experimenter covered the table with the occluder, retrieved a piece of food, showed it to the 379 ape, and hid it in one the cups outside the participant's view. Next, the experimenter 380 removed the occluder, picked up the baited cup and shook it three times, which produced a 381 rattling sound. Next, the cup was put back in place, the sliding table pushed forwards, and 382 the participant made a choice by pointing to one of the cups. If they picked the baited cup, 383 their choice was coded as correct, and they received the reward. If they chose the empty cup, 384 they did not. Participants received 12 trials. The location of the food was counterbalanced; 385 six times in the right cup and six times in the left. Causal inference trials were intermixed 386 with inference by exclusion trials (see below). We assume that apes locate the food by 387 reasoning that the food – a solid object – causes the rattling sound and therefore must be in 388 the shaken cup. 389

Inference by exclusion. Inference by exclusion trials were also modeled after Call 390 (2004) and followed a very similar procedure compared to causal inference trials. After 391 covering the two cups with the occluder, the experimenter placed the food in one of the cups 392 and covered both with the lid. Next, they removed the occluder, picked up the empty cup 393 and shook it three times. In contrast to the causal inference trials, this did not produce any 394 sound. The experimenter then pushed the sliding table forward and the participant made a 395 choice by pointing to one of the cups. Correct choice was coded when the baited (non-shaken) cup was chosen. If correct, the food was given to the ape. There were 12 inference by exclusion trials, intermixed with causal inference trials. The order was counterbalanced: six times the left cup was baited, six times the right. We assume that apes reason that the absence of a sound suggests that the shaken cup is empty. Because they saw 400 a piece of food being hidden, they exclude the empty cup and infer that the food is more

likely to be in the non-shaken cup.

Quantity discrimination. For this task, we followed the general procedure of
Hanus and Call (2007). Two small plates were presented left and right on the table (see
Figure 1C). The experimenter covered the plates with the occluder and placed five small
food pieces on one plate and seven on the other. Then they pushed the sliding table
forwards, and the participant made a choice. We coded as correct when the subject chose
the plate with the larger quantity. Participants always received the food from the plate they
chose. There were 12 trials, six with the larger quantity on the right and six on the left
(order counterbalanced). We assume that apes identify the larger of the two food amounts
based on discrete quantity estimation.

Delay of gratification. This task replaced the switching task in Phase 2. The 412 procedure was adapted from Rosati et al. (2007). Two small plates including one and two 413 pieces of pellet were presented left and right on the table. E moved the plate with the 414 smaller reward forward allowing the subject to choose immediately, while the plate with the 415 larger reward was moved forward after a delay of 20 seconds. We coded whether the subject 416 selected the larger delayed reward (correct choice) or the smaller immediate reward 417 (incorrect choice) as well as the waiting time in cases where the immediate reward was 418 chosen. Subjects received 12 trials, with the side on which the immediate reward was 419 presented counterbalanced. We assume that, in order to choose the larger reward, apes 420 inhibit choosing the immediate smaller reward. 421

We collected data in two phases. Phase 1 started on August 1st, 2020, lasted until
March 5th, 2021 and included 14 time points. Phase 2 started on May 26th, 2021 and lasted
until December 4th, 2021 and also had 14 time points. Phase 1 also included a strategy
switching task. However, because it did not produce meaningful results, we replaced it with
the delay of gratification task. Details and results can be found in the supplementary
material available online.

One time point meant running all tasks with all participants. Within each time point, 428 the tasks were organized in two sessions (see Figure 1E). Session 1 started with two gaze 429 following trials. Next was a pseudo randomized mix of causal inference and inference by 430 exclusion trials with 12 trials per task, but no more than two trials of the same task in a row. 431 At the end of Session 1, there were again two gaze following trials. Session 2 also started 432 with two gaze following trials, followed by quantity discrimination and strategy switching 433 (Phase 1) or delay of gratification (Phase 2). Finally, there were again two gaze following 434 trials. The order of tasks was the same for all subjects. So was the positioning of food items 435 within each task. The two sessions were usually spread out across two adjacent days. The 436 interval between two time points was planned to be two weeks. However, it was not always 437 possible to follow this schedule so that some intervals were longer or shorter. Figure S1 in 438 the supplementary material shows the timing and spacing of the time points.

440 Analysis

We had two overarching questions. First, we were interested in the cognitive measures 441 and the relations between them. That is, we asked how stable performance in a given task 442 was on a group-level, how stable individual differences were, how reliable the measures were. 443 We also investigated relations between the different tasks. To assess re-test reliability, we 444 computed computed pearson correlations between the data from two time points, aggregated 445 for each individual and task. We used Structural Equation Modeling (SEM) (Bollen, 1989; 446 Hoyle, 2012) to address the stability and structure of each task as well as relations between 447 tasks. 448

SEMs allowed us to partition the variance in performance into latent traits (stable over time), latent states (time varying) and measurement error. Because the latent variables are estimated on multiple indicators (here: test halves), they are assumed to be measurement-error free (Geiser, 2020; Steyer et al., 1992, 2015). In the present context, one can think of a trait as a stable psychological ability (e.g. ability to make causal inferences)

and states as time-specific, variable psychological conditions (e.g. variations in performance due to being attentive or inattentive).

We used Bayesian estimation techniques to fit the models. In the supplementary
material available online we report the prior settings for our models as well as the structural
restrictions we imposed on the model and the parameter estimation process. We justify these
settings and restrictions via simulations studies also included in the supplementary material.

Our focal Latent Trait-State (LST) model was defined as follows. The latent state 460 variables S_{it} were decomposed into a latent trait variable T_{it} and a latent state residual 461 variable ζ_{it} . The latent trait variables T_{it} are time-specific dispositions, that is, trait scores 462 capture the expected value of the latent state (i.e., true score) variable for an individual at 463 time t across all possible situations the individual might experience at time t (Eid, 464 Holtmann, Santangelo, & Ebner-Priemer, 2017; Steyer et al., 2015). The state residual 465 variables ζ_{it} captured the deviation of a momentary state from the time-specific disposition 466 T_{it} . We assumed that latent traits were stable across time. In addition, we assumed common 467 latent trait and state residual variables across the two test halves, which leads to the 468 following measurement equation for parcel i at time point t:

$$Y_{it}^* = T + \zeta_t + \epsilon_{it} \tag{1}$$

Here, T is a stable (time-invariant) latent trait variable, capturing stable
interindividual differences. The state residual variable ζ_t captures time-specific deviations of
the respective true score from the trait variable at time t, and thereby captures deviations
from the trait due to situation or person-situation interaction effects. ϵ_{it} denotes a
measurement error variable, with $\epsilon_{it} \sim N(0,1) \ \forall i,t$. The following variance components can
be computed for the presented LST model.

Consistency: Proportion of true variance (i.e., measurement-error free variance) that is
due to true inter-individual stable trait differences.

$$Con(Y_{it}^*) = \frac{Var(T)}{Var(T) + Var(\zeta_t)}$$
(2)

Occasion specificity: Proportion of true variance (i.e., measurement-error free variance)
that is due to true inter-individual differences in the state residual variables (i.e.,
occasion-specific variation not explained by the trait).

$$OS(Y_{it}^*) = 1 - Con(Y_{it}^*) = \frac{Var(\zeta_t)}{Var(T) + Var(\zeta_t)}$$
(3)

As state residual variances $Var(\zeta_t)$ were set equal across time, $OS(Y_{it}^*)$ is constant across time (as well as across item parcels i).

To investigate associations between cognitive performance in different tasks, the LSTMs were extended to multitrait models. Due to the small sample size, we could not combine all tasks in a single, structured model. Instead, we assessed relations between tasks in pairs.

Our second question was, which predictors explain variability in cognitive performance.

Here we wanted to see which of the predictors we recorded were most important to predict

performance over time. This is a variable selection problem (selecting a subset of variables

from a larger pool) and we used *Projection Prediction Inference* for this (Piironen et al.,

2020).

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