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Apes are intuitive statisticians

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ABSTRACT

Inductive learning and reasoning, as we use it both in everyday life and in science, is characterized by flexible inferences based on statistical information: inferences from populations to samples and vice versa. Many forms of such statistical reasoning have been found to develop late in human ontogeny, depending on formal education and language, and to be fragile even in adults. New revolutionary research, however, suggests that even preverbal human infants make use of intuitive statistics. Here, we conducted the first investigation of such intuitive statistical reasoning with non-human primates. In a series of 7 experiments, Bonobos, chimpanzees, gorillas and Orangutans drew flexible statistical inferences from populations to samples. These inferences, furthermore, were truly based on statistical information regarding the relative frequency distributions in a population, and not on absolute frequencies. Intuitive statistics in its most basic form is thus an evolutionarily more ancient rather than a uniquely human capacity.

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1. Introduction

Much research has suggested that reasoning about probabilities develops late in ontogeny, depends on language and formal education (Piaget & Inhelder, 1975), remains fragile even in adulthood (Tversky & Kahneman, 1974, 1981), and only works under special circumstances (Cosmides & Tooby, 1996; Gigerenzer & Hoffrage, 1995). Exciting new research, however, suggests that such reasoning capacities might well be in place in the absence of language. Even preverbal infants engage in some intuitive statistics: they expect randomly drawn samples to reflect the distribution in the population drawn from and vice versa (Denison & Xu, 2010b; Téglás, Girotto, Gonzalez, & Bonatti, 2007; Xu & Garcia, 2008).

This early intuitive statistics seems to be a cognitive capacity that is functionally integrated in humans with other cognitive domains from very early on: For example, infants already integrate information about physical and psychological background conditions into their statistical inferences when judging whether sampling processes are random or non-random. Regarding physical information, for example, infants understand that mechanical constraints (e.g. some kinds of objects in a population cannot be drawn physically in the same way as others) can turn a sampling process into a non-random one such that the sample need not reflect the distribution in the population (Denison & Xu, 2010a; Téglás et al., 2007). Moreover, statistical information is combined with geometrical and temporal information in rather systematic ways to form predictions about future events (Téglás et al., 2011). Regarding psychological information, infants appreciate that when a person draws from a population but has both a preference regarding the different kinds of objects in the population and visual access, her sampling will probably be non-random and her sample will thus not match the distribution of the population (Xu & Garcia, 2008).

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Conversely, infants draw inferences in the other direction, from statistical to psychological states of affairs: when confronted with a person who draws samples that are absolutely non-representative of the populations, infants assume the person must have informational access and corresponding preferences (Kushnir, Xu, & Wellman, 2010; Ma & Xu, 2011). Convergent findings have shown such intuitively statistical expectations in infants and toddlers with a number of different measures tapping different types of behaviors: looking time in response to violations of expectations (e.g. Xu & Garcia, 2008), active choice measures (of samples drawn from different populations; (Denison & Xu, 2010b) and actions directed towards others (such as giving them the kind of item they prefer; (Kushnir et al., 2010; Ma & Xu, 2011).

Compared to the information available for human infants, nothing is currently known about the phylogenetic origins and distributions of such intuitive statistics. We do not know how old evolutionarily capacities for intuitive statistics are, and we do not know whether they are shared by any non-human animals. Studies on optimal choice and foraging (Balci, Freestone, & Gallistel, 2009; Kamil, Krebs, & Pulliam, 1987; Stephens, 2008; Stüttgen, Yildiz, & Güntürkün, 2011) and numerosity discrimination (Brannon & Terrace, 1998; Hanus & Call, 2007) have demonstrated that non-human animals share with humans basic cognitive capacities to maximize the amount of food rewards on the basis of perceptual information. Those studies, however, lack some of the crucial features present in intuitive statistics research. Unlike subjects in optimal choice studies, subjects in intuitive statistics studies form expectations and select optimally based on statistical information without any prior training to associate the stimuli and their reinforcement contingencies or any other reliance on past sampling (Téglás et al., 2007, 2011).

It is true that such good first trial performance in the absence of training can also be found in primates' discrimination of absolute set sizes. First, from numerous comparative studies we know that many non-human animals, notably primates, share with humans an analog magnitude system that allows for the approximate discrimination between arbitrarily large sets (Cantlon & Brannon, 2006, 2007; Flombaum, Junge, & Hauser, 2005). The signature limit of this capacity, following Weber's Law, is constituted by the ratios of the sizes of two sets to be discriminated: if a subject can discriminate 4 from 8 objects, it can discriminate 10 from 20, 150 from 300, etc. Second, humans and other primates share an object individuation system that allows for the exact parallel individuation ("subitizing") of small sets (Hauser, MacNeilage, & Ware, 1996). The signature limit here is defined by the absolute set sizes: only sets smaller than 3 (infants) or 4 (monkeys and apes) can be discriminated, such as 1:2, 2:3, and 1:3 (see (Carey, 2009), for review). In contrast to such tasks, however, intuitive statistical problems crucially require representing truly statistical matters, namely relative rather than absolute frequencies – that is, frequencies of items of a given kind in a population (say, winner tickets in a lottery) relative to the frequencies of all kinds of items in the population (all tickets). It is thus an open question whether intuitive statistical reasoning, understood as the capacity

to flexibly draw inferences from populations to samples and vice versa, is evolutionarily recent and uniquely human or evolutionarily ancient and shared with other animals. Here we report a series of studies that speaks to that question. These studies with our closest relatives, the great apes, investigated one of the most basic forms of such intuitive statistical capacities: the ability to draw inferences from information about a population to a randomly drawn sample. We used tasks modeled after those developed in recent infant studies (Denison & Xu, 2010b). In these tasks, subjects are confronted with two visible populations with different distributions of items of two kinds (one preferable over the other) and the experimenter randomly draws from each population a 1-object-sample that the subject cannot see. Subjects are then given a choice between the two samples. These tasks thus require the subjects, first, to distinguish between the two populations according to the ratios of the two kinds of objects in their distributions and, second, to form expectations about the probability of sampling events accordingly, that is, expectations as to which sample is more likely to contain an object of the more desirable kind.

Control experiments ruled out alternative explanations such as simpler choice heuristics (Exp. 2 and 3), Clever Hans effects (Exp. 5 and 6) and use of olfactory information (Exp. 7). Most importantly, two experiments (Exp. 4 and 6) tested whether such inferences were truly based on probability information and not just on information about absolute frequencies.

2. Experiment 1: inferences from populations to samples

2.1. Subjects

Participants in all experiments were recruited from a group of four species of Great Apes ($N = 33$; Female $N = 24$): Chimpanzees (*Pan troglodytes verus*, $N = 17$), Gorillas (*Gorilla gorilla*, $N = 5$), Orangutans (*Pongo pygmaeus*, $N = 6$) and Bonobos (*Pan paniscus*, $N = 5$) housed at the Wolfgang Köhler Primate Research Center (WKPRC) in the Leipzig Zoo. Mean age of animals was 16;10 (years; months) with a range of 6;2–30;6. About one third were hand-reared and the remaining two thirds were mother-reared. All subjects had experience in cognitive studies and were used to receiving food-items as reinforcement (see SI Table 1 for a detailed description of the animals' demographics and background). 28 apes (15 Chimpanzees, 2 Gorillas, 6 Orangutans and 5 Bonobos) were included in the final sample of this experiment. Four further apes (2 Chimpanzees and 2 Gorillas) were tested but excluded from data analysis due to inconsistent item preference during the Preference Test ($N = 1$) or because they did not complete all trials due to lack of motivation ($N = 3$).

2.2. Design and procedure

Populations of banana pellets and carrot pieces were presented in two transparent buckets. Both buckets contained the same absolute amount of food items (80),

193 with distribution of banana pellets to carrots of 4:1 in
194 bucket A and 1:4 in bucket B. Each ape participated in 12
195 trials in which an experimenter drew one item from each
196 bucket (always of the majority type). Apes were tested
197 individually by two experimenters in special testing cages
198 or their sleeping quarters. Stimuli were presented on a table
199 (35 × 78 cm) mounted to one side of the testing cages.
200 A Plexiglas panel mounted on the cage mesh and perpendicular
201 to the table separated the ape from the experimenters.
202 Two small holes (∅2 cm; distance between
203 holes 59 cm) drilled into the Plexiglas panel allowed the
204 apes to indicate their choices by inserting a finger into
205 one of them.

206 2.3. Preference test

207 For each experiment, a preference test was administered
208 place before the first test trial session. One banana
209 pellet and one carrot piece were placed in front of the holes
210 on the experimenters' side of the panel. Apes indicated
211 their choice with their finger and immediately received the
212 selected food item as reinforcement. This preference test
213 was carried out twice in succession to establish whether
214 apes' preference was consistent. Virtually all apes
215 (with the exception of one subject in Exp. 1, and one subject
216 in Exp. 4 who were excluded from the experiments)
217 consistently chose the banana pellet over the carrot piece
218 on both trials.

219 2.4. Test trials

220 Apes participated in a total of 12 test trials split evenly
221 between two testing sessions. Depending on the availability
222 of animals, the delay between sessions was 1–11 days.
223 Apes were confronted with the two transparent buckets
224 containing the different populations consisting of banana
225 pellets and carrots in each bucket. Items of the two types
226 of food were of roughly equal size, but differed clearly in
227 color and shape and could thus be easily distinguished.
228 To aid the apes in gaining an overview of the two populations,
229 the first experimenter (E1) shook the buckets several
230 times and slightly tilted them forward. She then placed the
231 buckets on the table and drew one item from each bucket
232 (in such a way that the animal could not see which item it
233 was because the drawing hand and the drawn object were
234 occluded by other objects in the bucket), kept it invisibly in
235 one hand, and then moved forward both hands simultaneously
236 so that the ape could choose one of them. Apes
237 chose an item by inserting their finger through one of the
238 holes and touching the desired hand with the concealed
239 food item inside. If the animal pointed to both hands
240 simultaneously, the experimenter responded by saying:
241 "Just one, [Name]" until the animal clearly chose a single
242 hand. Apes then immediately received the food item as
243 reinforcement. After the ape had made her decision, E1
244 handed her the chosen food item. The trial was over and
245 E1 then removed the buckets from the table and out of
246 the ape's sight so they could be refilled by E2 and placed
247 them back on the table to start the next trial.

248 The side on which the more favorable population was
249 positioned in a given session was counterbalanced across

250 sessions and subjects. To rule out that low-level side preferences
251 might suffice to solve the task (for the side with the more
252 favorable population), E1 crossed her hands in half of the
253 trials before offering the ape a choice. Trials with and
254 without such crossing were administered in alternating
255 order (it was counterbalanced across subjects which kind
256 of trial came first) (see SI for details) (see Fig. 1). Q5

257 2.5. Observational and coding procedure

258 For this and all following experiments, a second blind
259 observer coded 25% of trials from video. Inter-rater reliability
260 was excellent for all experiments ($\kappa > .86$).

261 2.6. Results

262 The mean proportion of trials in which apes chose the
263 samples drawn from the two populations is depicted in
264 Fig. 2. Apes as a group chose the hand from the bucket with
265 the 4:1 distribution in 71% of the trials, significantly more
266 often than expected by chance, $t(27) = 6.43$, $p = .001$ (Cohen's
267 $d = .55$). (We detected no differences between the species,
268 $F(3,24) = 1.58$, $p = .22$). This pattern cannot be due to
269 learning over trials, as it was also reflected in trial 1
270 performance where 20 (71%) of the apes chose the hand
271 from the 4:1 bucket, significantly more than expected by
272 chance (Binomial test, $p = .02$; Cohen's $g = .21$).

273 3. Experiments 2 and 3: ruling out simple choice
274 heuristics

275 3.1. Subjects

276 31 Apes (16 Chimpanzees, 4 Gorillas, 6 Orangutans and
277 5 Bonobos) were included in the final samples of Experiments
278 2 and 3. One further Chimpanzee was tested but
279 failed to complete all trials due to lack of motivation.

280 3.2. Design and procedure

281 The general procedure was identical to Exp. 1 with the
282 following exceptions: The aim of Experiments 2 and 3
283 was to rule out explanations in terms of superficial choice
284 heuristics to the effect that apes' choices were simply
285 based on a preference for buckets where the preferred kind
286 of objects are in the majority. To do so, E1 sampled from
287 two populations which had in common that banana pellets
288 were in the minority (Exp. 2) or in the majority (Exp. 3),
289 but which still differed in their frequency distributions
290 (Exp. 2: 16:64 vs. 0:80/Exp. 3: 64:16 vs. 80:0).

291 3.3. Results

292 In Exp. 2, confronted with two buckets each containing
293 80 food items, with banana pellets: carrots distributions of
294 0:5 and 1:4, apes chose the latter in 84% of the trials,
295 significantly more often than expected by chance, $t(30) = 11.42$,
296 $p = .001$ (Cohen's $d = 1.06$), with a similar pattern on trial
297 1 where 87% of the apes chose from the 1:4 bucket,
298 significantly more than expected by chance (Binomial test,

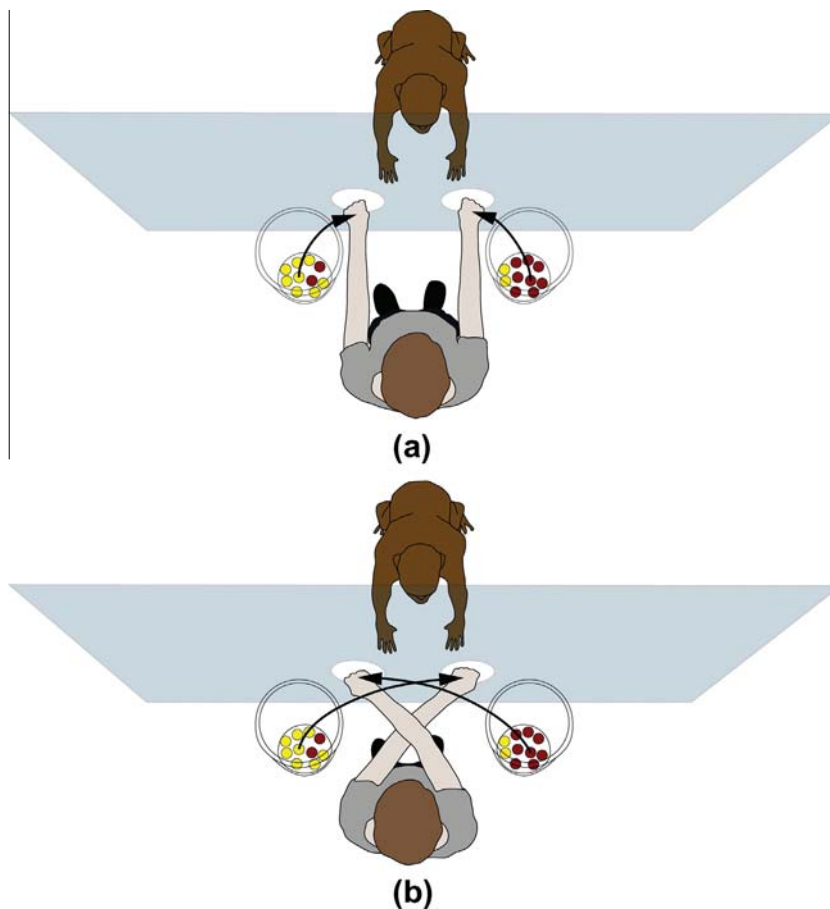


Fig. 1. Basic setup of the studies. An experimenter drew one item from each of the transparent buckets, kept the object invisibly in her hand, and then simultaneously moved both hands towards the ape, each behind a small hole in the plexiglas panel separating the ape and the experimenter. The ape then chose by inserting a finger through one of the holes. To rule out simple spatial heuristics (such as simply choosing the side where more attractive objects are), it was counterbalanced whether the hand with the object from one bucket was presented on the same side as the bucket (a) or on the opposite side (b).

299 $p = .001$, Cohen's $g = .37$). Our data revealed no differences
300 between the species, $F(3,27) = 0.82$, $p = .49$.

301 In Exp. 3, confronted with two buckets with banana pel-
302 lets: carrots distributions of 5:0 and 4:1, apes chose sam-
303 ples from the former in 62% of the trials, significantly
304 more often than expected by chance, $t(30) = 3.84$, $p = .001$
305 (Cohen's $d = .36$), with a similar pattern on trial 1 where
306 61% of the apes chose from the 5:0 bucket (Binomial test,
307 $p = .14$, Cohen's $g = .11$). One again, we detected no differ-
308 ences between the species, $F(3,27) = 1.40$, $p = .27$.

309 4. Experiment 5: ruling out Clever Hans effects

310 4.1. Subjects

311 26 Great apes (13 Chimpanzees, 3 Gorillas, 5 Orangu-
312 tans and 5 Bonobos) were tested.

313 4.2. Design and procedure

314 Experiment 5 was designed to test whether the findings
315 of Exp. 1 can be replicated when ruling out Clever Hans

316 effects. To this end, any information (visual, tactile, audi-
317 tory) about the items sampled on the part of the experi-
318 menter drawing and offering the objects was removed in
319 the following way:.

320 4.2.1. Blocking of visual access

321 Black cardboard was used to line the backside of buck-
322 ets and to create visual occluders that were attached to
323 either side of the top of the buckets to ensure that E1
324 had no visual access to the population distribution.

325 4.2.2. Blocking of tactile access

326 In order to prevent E1 from feeling from which popula-
327 tion she was drawing, E1 actually moved her hand into a
328 hidden compartment at the back of each bucket that was
329 invisible to the apes. In order to ensure that the experi-
330 menter had no tactile information about the identity of
331 each item sampled, she had two small plastic tubes at-
332 tached to her palm which were pre-baited before each trial
333 by E2. E1 then pretended to draw from the bucket, moving
334 her hands into the hidden compartments in such a way
335 that the tubes remained invisible to the subjects.

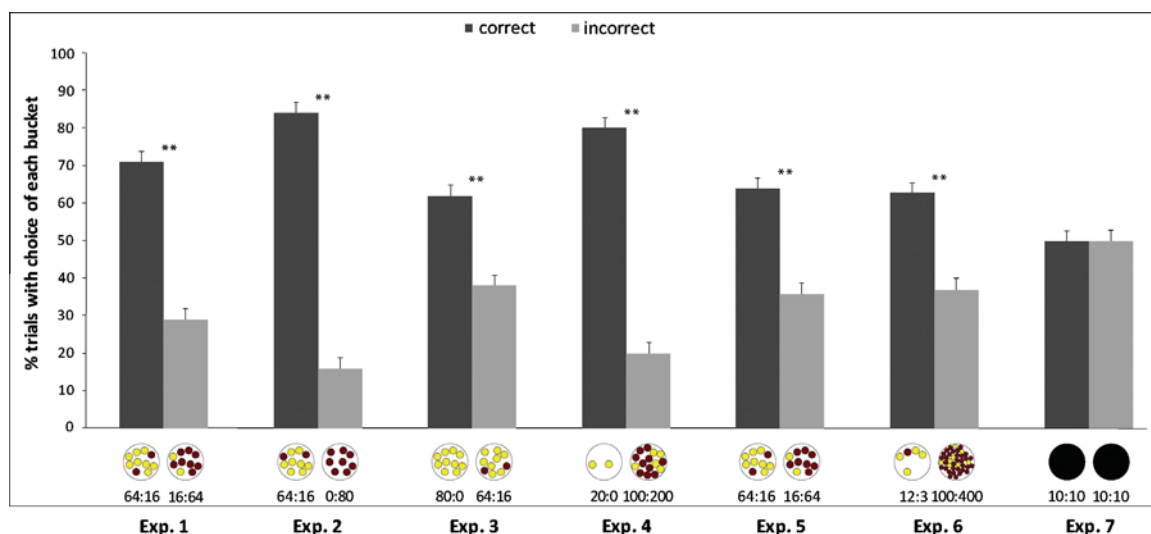


Fig. 2. Mean percentage of trials (with standard errors) in which the subject chose the correct/incorrect buckets. Below the graphs there are schematic representations of the distributions in the populations in both buckets in each study (yellow balls represent banana pellets (the preferred food items), orange balls represent carrots (less preferred). The ratios refer to the banana pellets: carrots ratio. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

336 4.3. Results

337 Controlling for Clever Hans effects, and with the same
338 distributions of objects in the two buckets as in Experiment
339 1, the findings were replicated: apes ($N = 26$) chose
340 the hand from the bucket with the 4:1 distribution in
341 64% of the trials, $t(25) = 4.84$, $p = .001$ (Cohen's $d = .52$),
342 with a similar pattern on trial 1 where 69% of the apes
343 chose from the 4:1 bucket (Binomial test, $p = .05$, Cohen's
344 $g = .19$).

345 5. Experiments 4 and 6: do apes really represent relative
346 (rather than absolute) frequencies?

347 5.1. Subjects

348 31 Apes (17 Chimpanzees, 3 Gorillas, 6 Orangutans and
349 5 Bonobos) were included in the final sample of Exp. 4. One
350 further Gorilla was excluded from analysis due to inconsis-
351 tency of item preference during the preference test. 26
352 great apes (13 Chimpanzees, 3 Gorillas, 5 Orangutans and
353 5 Bonobos) were tested in Exp. 6.

354 5.2. Design and procedure

355 Experiments 4 and 6 tested whether the inferences
356 from populations to samples found in Exp. 1–3 were truly
357 statistical inferences, based on relative frequencies, or
358 whether they could be explained more parsimoniously in
359 terms of representing absolute frequencies – absolute
360 and relative frequencies were confounded in all but one
361 previous infant studies (Denison & Xu, 2012) and in Exp.
362 1–3. In Exp. 4, therefore, apes had to choose from a bucket
363 with 20 food items with a 20 banana pellet: 0 carrot distri-
364 bution and a bucket with 300 food items and a 100 banana
365 pellet: 200 carrots distribution. While the latter bucket

366 contained a higher absolute number of banana pellets
367 (100 vs. 20), what matters for statistical inferences from
368 such a population to a sample is that the former was prefer-
369 able in terms of the relative frequency of the desired
370 items (20/20 = 1 vs. 100/300 = 0.33).

371 Exp. 6 administered a similar design but included a Cle-
372 ver Hans control like Exp. 5 such that E1 pretended to sam-
373 ple from a secret hidden compartment with a special tube
374 attached to her hand in such a way that she had no visual,
375 tactile or other cues as to the identities of the populations
376 or the samples. Two populations (buckets) containing
377 items of each kind were used: apes had to choose from a
378 bucket with 15 food items with a 12 banana pellets: 3 car-
379 roots distribution and a bucket with 500 food items and a
380 100 banana pellets: 400 carrots distribution, the latter
381 bucket containing a higher absolute but lower relative
382 number of banana pellets.

383 5.3. Results

384 In Exp. 4, apes chose the 20:0 bucket in 80% of the trials,
385 significantly more often than expected by chance,
386 $t(30) = 11.17$, $p = .001$ (Cohen's $d = 1.15$) (with a difference
387 between species $F(3,27) = 3.212$, $p = .04$ due to the fact that
388 all species but the gorillas performed above chance as a
389 group). This pattern was also reflected in trial 1 perfor-
390 mance where 22 (71%) of the apes chose the hand from
391 the 20:0 bucket, significantly more than expected by
392 chance (Binomial test, $p = .02$, Cohen's $g = .21$).

393 Similarly, in Exp. 6, apes chose the 12:3 bucket in 63% of
394 the trials, significantly more often than expected by
395 chance, $t(25) = 3.85$, $p = .001$ (Cohen's $d = .38$). This pattern
396 was also reflected in trial 1 performance where 20 (77%) of
397 the apes chose the hand from the 12:3 bucket, significantly
398 more than expected by chance (Binomial test, $p = .01$,
399 Cohen's $g = .27$).

400	6. Experiment 7: ruling out the use of olfactory cues	80% avoidance responses) and 400 (Exp. 6, 64% avoidance responses) less desired food items.	451 452
401	6.1. Subjects		
402 403	20 Great apes were tested (7 Chimpanzees, 3 Gorillas, 5 Orangutans and 5 Bonobos).		
404	6.2. Design and procedure		
405 406 407 408 409 410	In order to test whether apes' choices in the previous experiments could have been based on olfactory information regarding which objects were in E1's hands, apes were confronted with E1 sampling banana pellets and carrots from two mixed populations (both with a banana pellet: carrot ration of 10:10) in two opaque buckets.		454 455 456 457 458 459 460 461
411	6.3. Results		
412 413 414 415 416	Without any visual or other information about the populations drawn from, and with olfaction as the only source of information, apes chose the hand with the banana pellet in 50% of the trials, no different from chance, $M = 6$, $SD = 1.68$, $t(19) = 0$, $p = 1$.		
417	7. Performance across experiments		
418 419 420 421 422 423 424 425 426 427 428 429 430 431 432 433 434 435 436 437 438 439 440 441 442 443 444 445 446 447 448 449 450	Overall, apes performed above chance in all six studies which could be solved on the basis of intuitively statistical inferences (Exp. 1–6), but were at chance in Exp. 7 which could not be solved in this way. Across experiments, individual performance patterns were highly consistent. 24 apes participated in all six experiments in which the animals could choose correctly based on statistical information regarding the two populations randomly drawn from (Exp. 1–6). A Fisher's Omnibus Test exploring whether the distribution of Binomial test scores in the individual experiments was compatible with chance performance, showed that the vast majority of apes (20 out of 24) participating in all 6 studies performed above chance according to this test (see SI for details). It is theoretically conceivable that apes solved each task by avoiding the sample drawn from the bucket with the higher absolute frequency of less desirable food items, not by taking into account relative frequencies. However, while this cannot be ruled out for each experiment taken by itself, we think that the results of all experiments taken together render this possibility unlikely. If apes' choices had been based on such avoidance they should have shown a greater avoidance for samples from those buckets with the higher absolute frequency of less desired items across experiments. This was not the case. For instance, there were 64 and 400 items of less desired food in the incorrect buckets (depicted on the right side in Fig. 2) of Experiments 5 and 6, respectively. However, subjects avoided both buckets at comparable levels (about 64% of the trials). Moreover, the incorrect bucket that received the greatest percentage of avoidance responses (about 84% of the trials) contained 80 items (Experiment 2), which is well below two other incorrect buckets that contained 200 (Exp. 4,		462 463 464 465 466 467 468 469 470 471 472 473 474 475 476 477 478 479 480 481 482 483 484 485 486 487 488 489 490 491 492 493 494 495 496 497 498 499 500 501 502 503 504 505 506
		8. Discussion	453
		The findings of the present experiments show that a basic form of drawing inferences from populations to samples is not uniquely human, but evolutionarily more ancient: It is shared by our closest living primate relatives, the great apes, and perhaps by other species in the primate lineage and beyond and it thus clearly antedates language and formal mathematical thinking both phylogenetically and ontogenetically.	454 455 456 457 458 459 460 461
		8.1. How should the findings be best interpreted?	462
		Yet, exactly what cognitive capacity do the present findings show? Do they reveal intuitive statistical reasoning properly so-called, that is, reasoning from relative frequencies (of favorable items in a given population relative to all items in the population) to predictions about random samples drawn from these populations? This is how adults would typically reason about such problems and this is how the cognitive capacities of infants recently amply documented in similar studies have been generally interpreted.	463 464 465 466 467 468 469 470 471
		Or might these findings be explained more parsimoniously by simpler cognitive strategies and heuristics? The most obvious alternative would be that apes (and infants) might not reason about relative frequencies, but solve the tasks simply by discriminating absolute frequencies. And the most obvious and plausible version of this alternative would be that they discriminate the absolute frequencies of preferred items in each population. This alternative, however, can be ruled out empirically by the findings of the present Experiments 4 and 6 (explicitly designed for that purpose) for apes (and by similar recent control studies for infants; (Denison & Xu, in press). A second possibility might be that apes engage in avoidance strategies involving the comparison of the absolute frequencies of dis-preferred items. Such a strategy, however, is not compatible with the present findings either, for the following reasons: If apes merely engaged in comparisons between the absolute frequencies of dis-preferred items, one would expect to find the discrimination functions and signature limits that is virtually always found in primates' (and humans') numerical discrimination of absolute set sizes exceeding the subitizing range (Cantlon, 2012). Discriminability of two populations should thus vary, following Weber's Law, as a function of the ratio of the absolute set sizes of dis-preferred items in each population. But this was clearly not the case in the present studies (for similar results regarding infants, see Denison and Xu (in press). In Experiments 1 and 5, for example, the ratio of carrot pieces in population 1 and population 2 was 1/4 whereas in Exp. 6 it was 3/400 – yet, despite these massive differences absolutely comparable rates of discrimination were found.	472 473 474 475 476 477 478 479 480 481 482 483 484 485 486 487 488 489 490 491 492 493 494 495 496 497 498 499 500 501 502
		A third possibility might be that apes did not only focus on the absolute frequency of preferred items, nor on the absolute frequency of dis-preferred items but went beyond this and did take into account the relation of the two	503 504 505 506

507 absolute frequencies within a given population to each
508 other. However, instead of representing the crucial propor-
509 tional relation (the relative frequency of preferred items
510 relative to the whole population consisting of preferred
511 and dis-preferred items), they might have used a much less
512 complex heuristics based on difference scores (between
513 the absolute frequencies of preferred and dis-preferred
514 items). The simplest form of such an alternative might be
515 a heuristic along the following lines: “If in a given popula-
516 tion the frequency of preferred items is bigger than the fre-
517 quency of dis-preferred items, choose samples from this
518 population”. This heuristics might explain some of the in-
519 fant findings, and some of the results of the present study,
520 but crucially it cannot explain the findings from Exp. 3
521 [64:16 vs. 80:0]. In this experiment, the frequency of pre-
522 ferred items is bigger than the frequency of dis-preferred
523 items in both populations, and thus the heuristics would
524 give the ape both the output “choose population 1” and
525 the output “choose population 2” and should thus lead to
526 chance behavior – yet apes still chose the population with
527 the more favorable relative frequency of bananas.

528 Alternatively, the heuristic might be more subtle such
529 that it is not only determined for each given population
530 whether there are more preferred than dis-preferred items
531 in that population, but the difference scores (between the
532 absolute frequencies of preferred and dis-preferred items)
533 might then be compared to each other. In other words, this
534 would amount to a strategy of engaging in numerical dis-
535 crimination of difference (rather than proportional) scores.
536 Again, if this were the case, one would expect that this
537 discrimination task would reveal the characteristics and
538 signature limits found in virtually all numerical discrimina-
539 tion tasks with set-sizes exceeding the subitizing range. Dis-
540 criminability of populations should thus vary, following
541 Weber’s Law, as a function of the ratios of the difference
542 scores to each other. But this was clearly not the case in
543 the present study. Take, for example, Experiments 1 and
544 5, on the one hand, and Exp. 6 on the other hand. In all of
545 the experiments, the ratio of the relative frequencies of pre-
546 ferred items to dis-preferred ones was kept constant (4/5 in
547 population 1 vs. 1/5 in population 2), yet the differences
548 scores between preferred and dis-preferred items varied.
549 In Exp. 1 and 5 the difference scores are $64 - 16 = 48$ and
550 $16 - 64 = -48$, whereas in Exp. 6 they are $12 - 3 = 9$ and
551 $100 - 400 = -300$. Despite these massive differences in
552 the relation of the difference scores to each other, however,
553 these different experiments show absolutely comparable
554 rates of discrimination across the experiments – and thus
555 clearly suggest that apes tracked relative frequency and
556 not the alternative differences scores. All in all, thus, the
557 findings from the present experiments taken together are
558 not compatible with any obvious simple heuristics but
559 seem rather best explained by the assumption that apes
560 are well capable of simple forms of intuitive statistical rea-
561 soning based on the representation of relative frequencies.

562 8.2. Implications for the comparative psychology of numerical 563 cognition

564 Up to now, the numerical cognitive capacities known to
565 be shared by humans and non-human primates comprised

566 two systems for dealing with absolute set sizes – for the
567 exact individuation of small sets (<4) and for approximate
568 set size discrimination for arbitrarily large sets. The pres-
569 ent findings are the first to show that beyond these two
570 systems for representing absolute frequencies, we share
571 with other apes the capacity to represent relative frequen-
572 cies – a core foundation of statistical reasoning.

573 One question for future research concerns the relation
574 of these capacities to each other: what roles do the systems
575 for representing absolute set sizes play in the representa-
576 tion of relative frequencies both for small and for large
577 sets? Relatedly, what are the properties and signature lim-
578 its of the ability to distinguish relative frequencies? The
579 system for approximate set size discrimination follows
580 Weber’s Law in humans and other primates (discriminabil-
581 ity of two sets depends on the ratio of the absolute set
582 sizes: if the system can discriminate 5 bananas from 10 ba-
583 nanas, it can discriminate 10 from 20, 30 from 60, etc.
584 (Cantlon & Brannon, 2007; Xu & Spelke, 2000). And recent
585 research with human children (McCrink & Wynn, 2007;
586 Sophian, 2000) and non-human animals (Emmerton,
587 2001; Wilson, Britton, & Franks, 2002; Woodruff & Pre-
588 mack, 1981) as well as work in cognitive neuroscience
589 (Jacob, Vallentin, & Nieder, 2012; Vallentin & Nieder,
590 2008; Yang & Shadlen, 2007) suggests that discrimination
591 of ratios might be subject to the same signature limit in
592 accordance with Weber’s Law (discrimination breaks down
593 as a function of the ratio of the ratios to be discriminated)
594 as the discrimination of sets (which breaks down as a func-
595 tion of the ratio of the absolute set sizes). Future studies
596 will thus need to test whether intuitive statistical reason-
597 ing behaves similarly one level up, by discriminating any
598 two relative frequencies of as a function of the ratio of
599 the relative frequencies to each other. If such a pattern
600 were found, this would be first evidence to suggest that
601 intuitive statistics might be based on similar or the same
602 processes as approximate absolute set size discrimination.

603 Another question concerns the relation of the capacity
604 for intuitive statistics as documented here –inferentially
605 relating populations and randomly drawn samples– to
606 the capacity to represent statistical information found in
607 many species in domains such as auditory pattern extrac-
608 tion (Abe & Watanabe, 2011; Fitch & Hauser, 2004;
609 Gentner, Fenn, Margoliash, & Nusbaum, 2006; Toro & Tro-
610 balón, 2005), risk assessment and decision making (Balci
611 et al., 2009) or optimal foraging (Stephens, 2008). Are all
612 of these phenomena manifestations of a common underly-
613 ing domain-general capacity to deal with information
614 regarding distributions and relative frequencies? Or are
615 they separate and fragmented, perhaps modular capacities
616 with little inter-connections? Auditory statistical pattern
617 extraction, for example, arguably is such a special and
618 potentially separate domain (possibly having to do with
619 learning the “grammar” of songs or other communicative
620 systems) that it is highly unclear what its relation is to a
621 more general capacity for flexibly drawing inferences from
622 populations to samples and vice versa.

623 A broader question, finally, concerns the relation of
624 such intuitive statistics to other kinds of reasoning: In hu-
625 mans, statistical information is systematically integrated
626 with other types of information from very early on: Even

627 infants, for example, understand that mechanical constraints (e.g. only some kinds of objects in a population a
628 physically drawable) or psychological factors (the person drawing likes one kind of item more) can turn a sampling
629 process into a non-random one such that the sample need not reflect the distribution in the population (Denison &
630 Xu, 2010a; Téglás et al., 2007, 2011). Whether such systematic integration of different cognitive domains can
631 be found in non-human animals is an exciting open question with potentially far-reaching theoretical ramifications:
632 some influential theories of comparative cognition view this very cross-domain integration as one of the hall-
633 marks of uniquely human cognition (Carruthers, 2002; Gopnik & Schulz, 2004; Penn, Holyoak, & Povinelli, 2008;
634 Spelke, 2003; Woodward, 2007).

642 Author contributions

643 H.R., A.C. and J.C. designed the experiments, L.S., N.S., A.G. and J.M. ran the experiments. All authors analyzed
644 the results and discussed the findings. H.R., A.C. and J.C. wrote the paper.

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659 Appendix A. Supplementary material

660 Supplementary data associated with this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.cognition.2013.12.011>.

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