# The influence of live-capture on the risk perceptions of habituated samango monkeys Katarzyna Nowak<sup>a,b</sup>\* ; knowak02@gmail.com Shane A. Richards<sup>c,d</sup>; shane.richards@csiro.au Aliza le Roux<sup>b</sup> ; leroux.aliza@gmail.com Russell A. Hill<sup>a,e</sup>; r.a.hill@durham.ac.uk <sup>a</sup> Evolutionary Anthropology Research Group, Department of Anthropology, Durham University, South Road, Durham DH1 3LE, United Kingdom <sup>b</sup> Department of Zoology and Entomology, University of the Free State, Owaqwa, Phuthaditihaba 9866, South Africa <sup>c</sup> Department of Biological and Biomedical Sciences, Durham University, South Road, Durham DH1 3LE, UK <sup>d</sup> Current address: CSIRO Oceans & Atmosphere, P.O. Box 1538, Hobart, Tasmania 7001, Australia <sup>e</sup> Primate & Predator Project, Lajuma Research Centre, P.O. Box 522, Makhado 0920, South Africa \* Corresponding Author (Katarzyna Nowak; <u>knowak02@gmail.com;</u> +44 07400482343)

- **Running Heading:** Measuring effects of live-capture on monkeys

#### 27 The influence of live-capture on the risk perceptions of habituated samango monkeys

28 Live-capture of animals is a widely used technique in ecological research, and previously 29 trapped individuals often respond to traps with either attraction or avoidance. The effects of trapping on animals' risk perception are not often studied, even though non-lethal effects of risk 30 31 can significantly influence animals' behavior and distribution. We used a combination of 32 experimental (giving-up densities: GUDs) and behavioral (vigilance rates) measures to gauge monkeys' perceived risk before and after a short live-trapping period aimed at ear-tagging 33 34 monkeys for individual recognition as part of on-going research. Two groups of arboreal 35 samango monkeys, Cercopithecus albogularis schwarzi, showed aversion to capture in the form 36 of generalized, group-level trap shyness after two individuals per group were cage-trapped. We 37 predicted that trapping would increase monkeys' anti-predatory behavior in trap vicinity, and 38 raise their GUDs and vigilance rates. However, live-capture led to no perceptible changes in 39 monkeys' use of space, vigilance or exploitation of experimental food patches. Height above 40 ground and experience with the experiment were the strongest predictors of monkeys' GUDs. By 41 the end of the experiment, monkeys were depleting patches to low levels at ground and tree 42 heights despite the trapping perturbation, while vigilance rates remained constant. The presence 43 of cage traps, re-introduced in the final 10 days of the experiment, likewise had no detectable influence on monkeys' perceived risk. Our findings, consistent for both groups, are relevant for 44 45 research that uses periodic live-capture to mark individuals subject to long-term study, and more 46 generally to investigations of animals' responses to human interventions.

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48 Keywords: Live-trapping; ear-tagging; giving-up density; vigilance; African guenon

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#### **INTRODUCTION**

50 Live-capturing is a necessary technique in many studies of wild mammals where external 51 markings are added to enable observers to distinguish individuals (Glander et al. 1991; Rocha et 52 al. 2007; Stone et al. 2015), where tracking collars are affixed to study animals' movement 53 patterns (Moehrenschlager et al. 2003), or biological samples are obtained (Fietz 2003). Seldom, 54 however, are the effects of live-capture and associated handling on study animals' behavior 55 explicitly investigated and even where data are available, they suggest inconsistent patterns of 56 responses ranging from avoidance to attraction. For example, adult and juvenile coyotes (Canis 57 latrans) become trap-shy after initial captures using box traps (Way et al. 2002). Coyotes that 58 were trapped and released without sedation strictly avoided traps in the future and, once an 59 individual from their social group had been trapped, other group members stayed away from 60 traps (Way et al. 2002). Trap aversion has also been reported during a related study on San 61 Nicolas Island, where the island fox, Urocyon littoralis dickeyi, avoided areas in which they had been trapped, altering their ranging behavior in favor of areas where trapping had not occurred 62 63 (Jolley et al. 2012). In contrast, some species become trap happy and excessive recaptures need 64 to be reduced. For example, recaptures of the endangered fox, U.l. clementae, were reduced by 65 using bait treated with odorless salt (Phillips and Winchell 2011).

Other responses to trapping include signs of short-term stress. Live-capture induces a 66 67 stress response in meadow voles (Microtus pennsylvanicus) (Fletcher and Boonstra 2006) and 68 ground squirrels (Delehanty and Boonstra 2009) but has no long-term effect on the stress 69 physiology of mouse lemurs (*Microcebus murinus*), which readily habituate to trapping and are 70 therefore easily re-trapped (Hämäläinen et al. 2014). Rhesus monkey (Macaca mulatta) mothers 71 that have experienced an extended period of trapping on Cayo Santiago were more likely to 72 maintain proximity with their infants, and less likely to encourage independence or reject infants 73 (Berman 1989). Recent research on red colobus monkeys (*Procolobus rufomitratus*) shows that 74 they responded similarly to darting and collaring as to a predatory attack by chimpanzees (Pan 75 *troglodytes*) (Wasserman et al. 2013) – with an acute but short-term stress response. This finding is consistent with the "risk-disturbance hypothesis," which stipulates that human disturbance can 76 77 be similar to, or even exceed natural predation risk (Frid and Dill 2002). In contrast, a study of the effects of trapping on baboons (*Papio hamadryas*) and vervet monkeys (*Cercopithecus aethiops*) found no obvious effects on individual or group behavior, nor did animals become more wary of traps following previous capture (Brett et al. 1982). The length and frequency of capture, as well as the type of species under study, all appear to influence response type and magnitude.

83 While animals may quickly learn the association between their captors, the captors' tools 84 (e.g., traps), and danger, it remains unclear if non-lethal human "predators" can influence the 85 perceived risk and therefore foraging costs of wild animals. At the most basic level, we expect 86 wild animals' threat-sensitive responses to be affected by persistent human activities (Frid and 87 Dill 2002). For example, where woolly monkeys (Lagothrix poeppigii) are hunted, they learn to distinguish between three types of humans: hunters, gatherers, and researchers, responding most 88 89 strongly to hunters (Papworth et al. 2013). Other mammals, such as ungulates, may not as 90 readily distinguish hunting from other human activities. Red deer (Cervus elephus) respond with 91 increased vigilance to both recreational park users and hunters, although overall vigilance levels 92 are higher in the hunting season (Jayakody et al. 2008). Roe deer (Benhaiem et al. 2008) and 93 mountain gazelle (Gazella gazelle) (Manor and Saltz 2003) become more vigilant when and 94 where they are hunted or exposed to "human nuisance behavior". Red deer (Cervus elaphus) 95 shift habitats, trading off feeding opportunities to avoid human hunters (Lone et al. 2015) and Nubian ibex (Capra nubiana) do the same in response to tourists (Tadesse and Kotler 2012). 96

97 The majority of studies investigating this risk-disturbance hypothesis have assessed 98 relatively crude changes in vigilance behavior and range use. To maintain optimal fitness, 99 animals could make smaller-scale behavioral adjustments and discriminate between objects or 100 contexts that vary in risk level. For example, monkeys are known to increase vigilance and 101 decrease foraging time in the lower forest strata (Makin et al. 2012). Our study aimed to 102 investigate short-term and local changes in microhabitat use (including vertical height) and rates 103 of vigilance in reaction to humans and traps. We assessed if a habituated primate will distinguish

104 between non-threatening human observers, who may actually be perceived as offering 105 protection from natural predators (Nowak et al. 2014) and potentially dangerous traps left by the 106 observers, adjusting their behavior accordingly. We measured the effects of live-trapping, aimed 107 at marking individuals as part of an on-going long-term study, and subsequent placement of cage 108 traps on monkeys' perceived risk and associated foraging cost for two groups of habituated 109 samango monkeys (Cercopithecus albogularis schwarzi). We employed two commonly adopted 110 approaches for quantifying risk perceptions: giving-up densities (GUDs) and rates of vigilance. 111 GUDs represent the amount of food a forager gives up in a food patch, with lower GUDs 112 predicted in areas where animals feel safe and have a higher food harvest rate (Brown 1999). In 113 contrast, higher vigilance interferes with feeding (Brown 1999; Benhaiem et al. 2008), and is 114 expected to raise GUDs. We predicted that both indirect measures of risk (GUDs and vigilance 115 rates) would increase relative to the baseline (pre-capture) rates, at least in the short-term (days), 116 following both live captures and the subsequent placement of traps within the experimental area 117 where monkeys forage on artificial feeding stations.

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# MATERIALS AND METHODS

120 Study site and subjects.---We conducted our study between May and September 2013 at the 121 Lajuma Research Centre (23°02'23''S, 29°26'05''E) in the western Soutpansberg Mountains, 122 Limpopo Province, South Africa (Nowak et al. 2014). The site is characterized by fragments of 123 tall moist forest (up to 20 m high) and short dry forest (up to 10 m high) (Coleman and Hill 124 2014a). Monkeys' natural predators include leopards (Chase Grey et al. 2013), crowned 125 (Stephanoaetus coronatus) and black (Aquila verreauxii) eagles, caracals (Caracal caracal), and 126 rock python (*Python sebae*). Sympatric diurnal primates are chacma baboons (*Papio ursinus*) 127 and vervet monkeys (Chlorocebus aethiops pygerythrus).

128 We studied two groups of samango monkeys, which belong to the polytypic 129 *Cercopithecus mitis* group widely distributed across Africa but rare in South Africa, where they 130 are limited to Afro-montane and coastal forest fragments (Dalton et al. 2015). They are 131 medium-sized (up to 70 cm; 4.4 kg for adult females, 7.6 kg for adult males; Harvey et al. 1987), 132 group-living arboreal monkeys with a mostly frugivorous diet (Coleman and Hill 2014b). Our 133 two study groups consisted of 40 and 60 individuals, respectively. Since the beginning of 2012, both groups – called Barn and House – have been followed by researchers (3-4 times per week) 134 135 as part of a long-term behavioral study and are thus well-habituated to human presence. Both 136 groups had previously experienced GUDs experiments and cage-trapping, but never in 137 combination or in temporal proximity as in this experiment.

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139 Giving-up densities.---Giving-up densities were measured for 20 days (4 consecutive days per 140 week for 5 weeks) both before and after the pre-baiting (8 days) and live-capture periods (5 141 days). Artificial foraging patches were established at 16 trees, randomly selected within groups' 142 known winter home ranges, in short forest adjoining tall evergreen forest. At each GUDs patch, 143 we suspended basins at four heights: 0.1, 2.5, 5, and 7.5 meters. Each basin (46 cm in diameter) 144 was filled with 4 liters sawdust and "baited" daily with 25 shelled raw peanut halves. We 145 counted remaining peanuts every day after 1600 h and topped up any spilled sawdust (Nowak et 146 al. 2014). The pre-baiting and live-capture took place within this experimental patch area. 147

*Live-trapping.*---The main purpose of the live capture was to ear-tag monkeys for identification
purposes in the context of a large on-going scientific research project at this site. We took
advantage of these captures to answer our questions about the specificity of monkeys' responses.
All trapping procedures were approved by the Limpopo Province Department of Economic
Development and Tourism, with ethical approval from Durham University's Life Sciences
Ethical Review Process Committee and the Anthropology Department's Ethical Sub-Committee.
Our research followed ASM guidelines.

155 The trapping period was preceded by eight days of pre-baiting, consisting of baiting 156 with orange quarters two custom-made cage traps (123 cm long x 60 cm wide x 80 cm tall) per 157 group. Monkeys (including previously ear-tagged individuals from an earlier trapping event in 158 2012) took oranges from traps on a regular basis during this period. Active trapping was then 159 initiated and four individuals (all untagged) were trapped and marked, two from each group, on 160 the first two days of the five-day trapping period (Fig. 1). Samango monkeys forage as a 161 cohesive group (Emerson and Brown, 2013) and other group members were moving through the 162 trapping area when individuals were trapped (mean neighbors within 5 m = 1.79 (SD = 1.89) 163 (Coleman 2013), and they dispersed in response to the capture events and/or other individuals 164 dispersing. Trapped individuals were hand-injected with Zoletil within minutes (<5 min) of 165 capture by an experienced veterinarian, and carried to a nearby area to be measured, ear-tagged, 166 and finally placed in a holding cage during recovery. Three of the four captured individuals 167 recovered quickly (one adult female reacted strongly to the anesthetic), and all four were 168 released within a mean of 3.7 (SD = 0.79) hours of capture back into their social group. All 169 animals subsequently were monitored and returned to typical activity patterns the following day. 170 The traps were supplemented with additional bait, consisting of bananas and passion fruits, in the ensuing three days of the trapping period to try and increase the probability of further captures. 171 172 These same cage traps were later re-placed, open and without bait, next to GUDs trees in the 173 final 10 days of the 20-day post-trapping experimental period.

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*Vigilance.---*Vigilance behavior of monkeys while foraging on artificial food patches was
recorded either by observers, standing with binoculars at no less than 20 m from GUDs trees
(during monkeys' first foraging bout of the day), or using camera traps (Cuddeback Attack IR
and Bushnell Trophy Cam) in the absence of observers (throughout the day). "Vigilance"
referred to a monkey looking or glancing up from an experimental basin to visually scan the area
in an upright posture. Vigilance data were recorded from the point at which a monkey began

181 foraging within an experimental food patch and ended when the monkey left the basin (n = 85182 records before trapping and 72 after trapping for Barn group; 220 before and 177 after for House 183 group). Vigilance was extracted from camera trap video footage based on the same criteria for 184 the start and end of a bout (n = 16 video clips before and 20 after capture for Barn group: 30 185 before and 108 after for House group) for a total of 728 records. The number of glance-ups per minute constituted "vigilance rate". For statistical purposes, data from direct observations and 186 187 camera traps were pooled following assessment that there were no statistical differences between 188 these two data sources.

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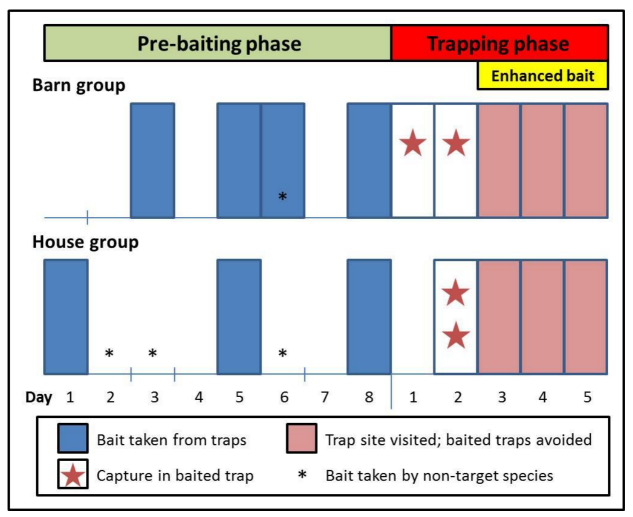
190 Analyses.---As prior research detected subtle effects of human followers on monkeys' perceived 191 risk of predation (Nowak et al. 2014), we analyzed only data from days on which monkeys were 192 not followed from dawn to dusk by researchers. To appropriately account for the structured 193 nature of data collection (i.e., repeated sampling at trees), and the fact that the data were highly 194 over-dispersed, we developed likelihood functions that incorporated these effects. This more 195 general approach also allowed us to better link our biological hypotheses with our study design 196 (Richards 2015). Specifically, we fitted discontinuous temporal models of GUDs and vigilance 197 rates to our data to investigate whether or not our two experimental disturbances (live-capture 198 and the presence of cages post-trapping) resulted in short-term changes in foraging behavior, 199 while still allowing for any underlying gradual changes in foraging behavior. For both the GUD 200 and the vigilance data sets we fitted models that incorporated up to three predictive factors: the 201 sampling day of the GUDs experiment (D: 1-40), the period of the study delineated by the two 202 imposed disturbances (P: pre-trap [days 1-20], post-trap without trap stimulus [days 21-30], and 203 post-trap with trap stimulus [days 31-40]), and the height of the food basin (H: ground and 204 aboveground, which included the three tree-level basins). Thus, D reflects long-term responses 205 (weeks), P reflects short-term responses (days) in the form of break-points, and H reflects local 206 responses (meters). Random variation in foraging behavior among basins, caused by unknown

207 differences among the trees sampled, and day-day site-wide differences (e.g. variation in 208 weather), were also explicitly incorporated into the models. In brief, our GUD model is a 209 generalized example of a logistic regression, and our vigilance model is a generalized example 210 of a non-linear regression, where P and H are treated as discrete factors, D is a covariate, and 211 day-day variation is a random effect. Also, for both models, we account for additional sources of 212 over-dispersion in the data. Full details of the statistical models can be found in Supporting 213 Information S1. Evidence that any of the three factors improved model parsimony and 214 explanatory power was evaluated by performing model selection using AIC (Richards 2015).

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### RESULTS

217 During the first phase of our study (sampling days 1-20), before animals were introduced to the 218 traps, we observed samango monkeys foraging within all of our basins. During the pre-baiting 219 phase, when traps were placed at two trees within both groups' foraging range but not set to 220 trigger (eight consecutive days), animals continued to forage at basins placed on trees associated with the trap, and also removed bait from the cage traps (Fig. 1). After traps were set to trigger 221 222 (trapping phase) only two animals per group were caught and catches occurred only on the first 223 two days of the trapping period (Fig. 1). No trapping location was successful on more than a 224 single day. Animals in both groups continued to feed near the areas where the captures took 225 place but individuals avoided approaching or entering the set traps despite the presence of 226 significantly enhanced bait in each of the traps.



228 Fig. 1. Samango monkey (C. albogularis schwarzi) responses to traps placed near feeding 229 stations during the pre-baiting and trapping phase of the study, carried out from May to September 2013 in the Western Soutpansberg Mountains, South Africa. Each day, from 6<sup>th</sup>-18<sup>th</sup> 230 231 July 2013, a trap was placed at two trees within the foraging range of House and Barn groups. 232 Traps always contained food but they were not set to trigger during the pre-baiting phase. Bars 233 depict days that samango monkeys were observed at one or more traps; blue bars indicate that 234 samango monkeys removed bait from traps, whereas red bars indicate that bait within traps was 235 avoided. On four days during pre-baiting baboons or vervet monkeys removed food from the 236 traps before samango monkeys arrived (asterix). Four samango monkeys were caught during the 237 trapping phase (red stars): two adult females (AF), one in each group, a juvenile female (JF) in 238 House group, and a sub-adult male (SM) in Barn group.

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240 GUDs (measured as number of peanuts remaining in basins) declined gradually over the course

- of the study and GUDs were lower for basins placed aboveground for both groups (Fig. 2a,b).
- However, there was no obvious short-term change in GUDs after live-capture for either group
- 243 (sampling days 21-40). Re-placing traps back in the foraging area (days 31-40) did not raise
- 244 monkeys' perceived risk as measured by GUDs and rates of vigilance. These conclusions are
- supported by our AIC analyses selecting the model including height and day (H+D) as the best

- 246 model for both groups (Table 1). While the model including sampling period and height
- 247 (P+H) also was considered parsimonious for Barn group (Table 1), temporal changes in GUDs
- could be better explained by assuming a gradual decline over time rather than a response to
- trapping. We found no evidence of a gradual change in the rate of vigilance for either group, nor
- 250 did we find any evidence of a short-term vigilance response to trapping (Fig. 2c,d). However,
- both groups significantly elevated vigilance behavior when foraging on the ground (only model
- H was selected for both groups: Table 1) and, in general, House group (the larger of the two
- 253 study groups) was less vigilant.
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255 Table 1. Summary of the AIC analyses of the GUD and vigilance data for two groups of 256 samango monkeys (C. albogularis schwarzi) studied in the Western Soutpansberg Mountains, South Africa from May to September 2013. Models incorporated up to three factors for both data 257 258 sets: day of sampling (D), period of sampling (P), and patch height (H). K is the number of 259 estimated model parameters, LL is the maximum log-likelihood, and  $\Delta AIC$  is the difference in 260 the AIC of the model compared to the lowest AIC model. Bold  $\triangle$ AIC values indicate that the 261 model is selected (i.e., is relatively more parsimonious, given the models considered). Models 262 were selected if they had  $\Delta AIC < 6$  and no simpler, nested model had a lower AIC score 263 (Richards 2015). 264

Model	GUD			Vigilance		
	K	$\mathbf{L}\mathbf{L}$	ΔΑΙC	K	LL	ΔΑΙΟ
Barn group						
Null	10	-1945.1	33.7	10	-152.4	16.2
D	11	-1940.0	25.6	11	-152.3	18.0
Р	12	-1938.9	25.2	11	-152.0	17.5
D+P	13	-1938.9	27.2	13	-151.7	20.9
Н	11	-1933.9	13.3	11	-143.3	0.0
H+D	13	-1925.2	0.0	12	-142.9	1.3
H+P	15	-1924.0	1.4	15	-140.7	2.9
H+D+P	17	-1923.6	4.7	16	-140.7	4.9
House group						
Null	10	-1528.9	35.2	10	-285.9	51.4
D	11	-1519.1	17.6	11	-285.8	53.2
Р	12	-1520.7	22.8	11	-285.1	51.8
D+P	13	-1518.6	20.6	13	-284.6	54.9
Н	11	-1522.1	23.6	11	-259.2	0.0
H+D	13	-1508.3	0.0	12	-259.1	1.8
H+P	15	-1511.0	9.3	15	-255.6	0.8
H+D+P	17	-1507.7	6.7	16	-255.2	1.9

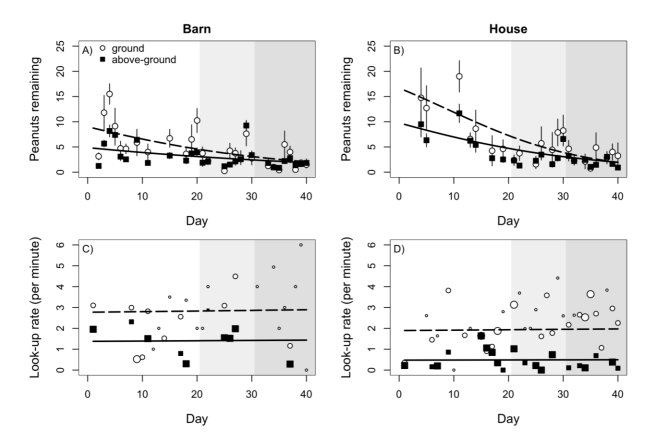


Fig. 2. Observed and predicted GUDs and vigilance rates for two groups of samango monkeys, 268 C. albogularis schwarzi, studied in 2013 in South Africa. Time is sectioned into three periods: 269 270 pre-trap (days 1-20, white), post-trap without trap stimulus (days 21-30, light grey), and posttrap with trap stimulus (days 31-40, grey). Panels A and B show observed GUDs averaged 271 272 across eight trees for two height categories, and error bars represent 1 se. Sloped lines indicate 273 the best AIC model predictions, which was model D (sampling day) + H (ground or tree) for 274 both groups. Panels C and D show the corresponding vigilance rates. Symbols size indicates the 275 period of the observations used to calculate the mean rate: < 5 minutes (small), 5-15 minutes 276 (medium), and > 15 minutes (large). Again, lines indicate the AIC-best model predictions, which 277 was model H for both groups.

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### DISCUSSION

- 280 We found no evidence for live trapping affecting the anti-predatory behavior (i.e., GUDs and
- vigilance rates) of these habituated samango monkeys, with the exception of observing the
- 282 monkeys' trap avoidance following initial successful captures. Their trap shy response may
- 283 represent long-term individual aversion to trapping and the trap stimulus in that no individual
- samangos have ever been re-captured at Lajuma (of 18 caught and tagged), while at Hogsback in
- the Eastern Cape, only 4 out of 64 samango monkeys were re-caught following successful

capture (Kirsten Wimberger, University of Cape Town, personal communication, May 2014).
This is in contrast to mouse lemurs (Hämäläinen et al. 2014) and galagos (Charles-Dominique
and Bearder 1979) which show no aversion to traps or being trapped and therefore re-enter traps
on successive occasions.

290 Despite samango monkeys' apparent trap aversion, we nevertheless found no further 291 evidence that live-capture or subsequent placement of traps in the GUDs experimental area 292 altered these samango monkeys' perceived risk, even in the short-term (neither in the days 293 immediately following trapping nor during the five weeks following trapping). Monkeys' typical 294 anti-predator behavior (e.g. vigilance, use of the ground stratum) remained unaltered after the 295 capture events, even while the trap stimulus was present in the experimental area. Monkeys did 296 not transfer their negative trap-response (trap shyness) to our experiment, i.e. the experimental 297 area or the artificial food sources (man-made containers/basins used in the experiment). This 298 suggests that monkeys likely distinguish between anthropogenic sources of risk, possibly 299 because they already had five weeks prior (positive and rewarding) experience with experimental 300 food patches before the live-capture. The food patches (raw unshelled peanuts) were also of high 301 quality and required no processing once found inside the sawdust.

Where samango monkeys face a variety of risks including conflict with people and depredation by domestic dogs, such as in Hogsback, Eastern Cape, South Africa, they will still capitalize on high-quality food in the form of fallen exotic oak acorns and seeds in people's gardens (Wimberger et al. in review). Supplementing the cage traps in this study with additional high-quality bait (bananas and passion fruits) in the final three days of the live capture, however, failed to attract the trap-averse monkeys.

308 Despite the absence of evidence suggesting behavioral changes in space-use and 309 vigilance in response to capture, animals did exhibit consistent, predictable variation in risk 310 responses in relation to foraging height and experience with the GUDs experiment. Monkeys had 311 higher GUDs at the start of the experiment, foraged less at ground level (Emerson et al. 2011;

312 Nowak et al. 2014) and had lower vigilance rates at higher canopy levels (MacIntosh and 313 Sicotte 2009; Campos and Fedigan 2014). The larger House group had lower vigilances rates 314 than the smaller Barn group, consistent with the group size effect (Hill and Cowlishaw 2002; 315 Makin et al. 2012; Campos and Fedigan 2014). Animals also showed a steady increase in 316 foraging proficiency over the course of the experiment, suggesting that practice and familiarity 317 may result in falling GUDs; GUDs decreased over time at ground and tree levels, indicating 318 monkeys' ability to quickly adapt to their current environment and efficiently exploit newly 319 available sources of food from which they were not easily deterred by a perturbation like live-320 capture.

321 We found no evidence of a trade-off between vigilance rates and GUDs; GUDs declined 322 but vigilance rates were fixed throughout the duration of the study. However, we have only 323 quantified vigilance rates and not duration of vigilance; it may be that look-up duration declined 324 over time, which freed up time for lowering GUDs. We had enough video data of House group 325 foraging to see if time spent at trees changed during the study, and, if it differed between basin 326 heights. We found no evidence of a day effect on the mean time spent at trees (ANCOVA;  $F_{1,31}$ = 2.67, P = 0.112); however, there was evidence of a height effect (ANCOVA;  $F_{1,31} = 9.40$ , P =327 328 0.004) with monkeys spending less time at ground than tree canopy level (Supporting 329 Information S2). On average, over the course of the day, animals spent shorter times at the 330 basins placed on the ground (3.26  $\pm$  0.75 minutes) compared with basins placed aboveground  $(8.56 \pm 1.67 \text{ minutes})$ . These additional findings suggest that animals improved their 331 332 proficiency at finding peanuts rather than spending more time at basins, given that the amount of 333 peanuts taken from basins increased over time but time spent at basins did not increase.

While we did not measure hormones or stress responses directly like Wasserman et al. (2013), we similarly did not find monkeys' behavior to be suggestive of a prolonged stress response as a result of the live-capture. The monkeys in our study appeared to be extremely apt at distinguishing among different forms of risk and clearly made trade-offs that optimized their exploitation of food-rich patches (Emerson and Brown 2013). While the monkeys, like coyotes and foxes (Way et al. 2002; Jolley et al. 2012), became trap shy, their trap aversion did not result in or extend to spatial avoidance of the area in which trapping took place as it did for the carnivores. This has important implications in management terms, as the use of trapping and release would not be a worthwhile approach to deterring primates from food sources. Our research indicates that primates are unlikely to show a generalized fear response following livecapture, particularly if carried out by humans to whom they are already habituated.

345 Where goals are to study primates long-term by habituating them, insights about the risk-346 disturbance hypothesis, specifically fear, risk avoidance and learned responses to humans and 347 their research tools, are important for conservation managers looking to monitor endangered 348 species. The methods we used here are generalizable to other longitudinal field studies that 349 employ live capture to mark and study animals. Further comparative data are essential to gauge 350 the relative differences among species and individuals in responses to capture and other 351 potentially stressful research practices, such as wearing of GPS collars. This study is important 352 for understanding how our research and management practices may distort animal behavior - or 353 even cause harm – and result in misinterpretation of wild animals' resilience to our presence and 354 activities.

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declare.

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### SUPPORTING INFORMATION

373 Supporting Information S1. Full details of statistical models and analysis of giving-up
 374 densities and vigilance.

375 Likelihood function describing GUDs.---Giving-up densities (GUDs) and vigilance behaviors 376 were recorded for two groups of samango monkeys: Barn and House. The two groups foraged at separate but nearby sites. At each site four basins were placed on 8 randomly selected trees. One 377 378 basin, termed the ground basin, was placed at the base of the tree (10 cm above the ground) and 379 the remaining three basins, termed aboveground basins, were placed at 2.5 m, 5 m and 7.5 m. At 380 the start of the day 25 peanuts were placed in each basin mixed in with sawdust and at the end of 381 the day the number of peanuts remaining (GUD) was recorded. GUDs were recorded during 382 three time periods: pre-trap (sample days 1-20), post-trap without a trap stimulus (sample days 21-30), and post-trap with a trap stimulus (sample days 31-40). Additional details of the 383 384 experimental protocol can be found in Materials and methods.

We assumed that the mean fraction of peanuts remaining in basins at the end of the day could be described by the following piecewise logistic function:

387 
$$\log \operatorname{it} \overline{y}(t) = \begin{cases} b_0 + a(t-1), & \text{if } 1 \notin t \notin 20; \\ b_1 + a(t-21), & \text{if } 21 \notin t \notin 30; \\ b_2 + a(t-31), & \text{if } 31 \notin t \notin 40; \end{cases}$$
(S1)

388 where

389 
$$b_1 = b_0 + 20a + d_1$$
 (S2)

390 and

$$391 b_2 = b_2 + 10a + d_2. (S2)$$

392  $\beta_0$  describes the degree to which peanuts are depleted on sample day t = 1,  $\alpha$  describes how 393 GUDs change gradually over sampling days, and  $\delta_1$  and  $\delta_2$  describe rapid changes in GUDs due 394 to the trapping event and the reintroduction of a trap stimulus, respectively.

395 To describe the patterns of GUDs in our data we needed to explicitly account for three 396 sources of random variation. Random day-day variation in the fraction of peanuts remaining in 397 basins on individual trees (within-tree variation) was accounted for by incorporating the beta-398 binomial distribution with variance parameter  $\phi$  (Richards 2008). Between-tree variation in 399 GUDs was accounted for by associating each tree with a parameter u, which were estimated 400 from the data; trees having a lower *u* tended to have more peanuts removed. Random day-day 401 variation in GUDs across trees caused by site-wide factors (e.g., weather conditions) were 402 incorporated by assuming that sampling days were associated with a random variate, Z, drawn 403 from a normal distribution with mean zero and standard deviation,  $\sigma_z$ ; days associated with a low 404 z-value resulted in all trees at the site experiencing fewer than the expected number of peanuts.

405 Let  $y_{ijt}$  be the number of peanuts remaining in basin *j* located on tree *i* on sample day *t*. The 406 assumptions described above define our model, which is described by the set of parameters,  $\theta$ . 407 The probability of observing all the data at a site, denoted **Y**, given our model, is:

408 
$$\Pr(\mathbf{Y}|\boldsymbol{\theta}) = \prod_{t=1}^{40} \int_{z=-\infty}^{\infty} f_{N}(z|0,\sigma_{z}) \prod_{i=1}^{8} \prod_{j=1}^{4} f_{BB}(y_{ijt}|25, T[\overline{y}_{j}(t), u_{i}+z], \phi) dz$$
(S4)

17

409 where

410 
$$T[p,w] = \frac{e^w p}{1 - p + e^w p},$$
 (S5)

411  $f_N$  is the probability density function of the normal distribution, and  $f_{BB}$  is the probability mass 412 function of the beta-binomial distribution with variance parameter  $\phi$  (see Richards 2008 for 413 details). Here we have added a subscript *j* to the population expectation  $\overline{y}(t)$  as we allow for the 414 associated parameters to vary depending on whether or not the basin is place on the ground. We 415 equated this probability of the data with the likelihood of the model and estimated the log-416 likelihood using

417 
$$LL(\boldsymbol{\theta} \mid \mathbf{Y}) = \sum_{t=1}^{40} \ln \sum_{k=-K}^{K} \exp\left[\ln f_k + \sum_{i=i}^{8} \sum_{j=1}^{4} \ln f_{BB}(y_{ijt} \mid 25, T[\overline{y}_j(t), u_i + \sigma_z z], \phi)\right]$$
(S6)

418 where 2K+1 is the number of intervals used to approximate the standard normal distribution,  $z_k = 419 = \frac{8k}{2K}$ , and

420 
$$f_k = \frac{e^{-z_k^2/2}}{\mathring{a}_{m^{--K}}^K e^{-z_m^2/2}}.$$
 (S7)

421 We found K = 20 gave an accurate estimate of LL.

422 Likelihood function describing vigilance.---The rate of looks performed per minute for both 423 groups was modelled in a very similar manner as the GUDs; however, the within-tree variation in the number of looks was assumed to have a negative-binomial distribution with variance 424 425 parameter  $\phi$  (see Richards 2008 for details), rather than a beta-binomial distribution. As the 426 number of looks is unbounded we modified the expected number of looks by replacing the logit 427 transformation on the left side of equation (S1) with the natural logarithm, ln. Also, because we 428 had less temporal resolution for the look data (Fig. 1) we forced  $\alpha$  to be equal for both the 429 ground and aboveground basins.

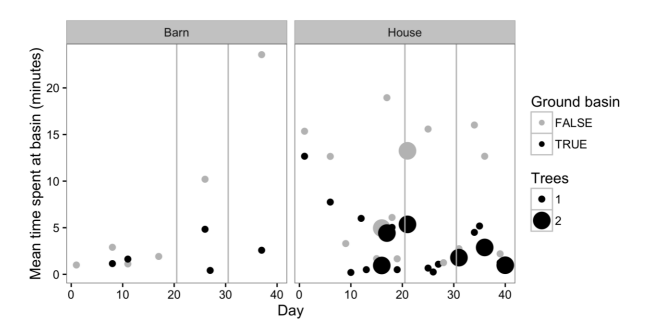
Model selection.---Eight models were considered for both groups when investigating both the 430 431 GUD and the vigilance data (Table 1). Models included zero or more of the following three 432 effects: (D) sampling day, (P) sampling period, and (H) basin height. Factors were removed from a model as follows: (D)  $\alpha = 0$ , (P)  $\delta_1 = \delta_2 = 0$ , (H)  $\beta_0$ ,  $\alpha$ ,  $\delta_1$  and  $\delta_2$  were set equal for both 433 434 heights. For all models we assumed random within-tree variation ( $\phi > 0$ ), between-tree variation  $(u_i \neq 0)$ , and day-day variation  $(\sigma_z > 0)$ . Models were selected using the recommendations of 435 436 Richards (2015); namely, all models with  $\triangle$ AIC within 6 of the minimum are initially selected, 437 but complex models with simpler, nested models having a lower AIC score, were subsequently 438 discarded.

439

440 **Supporting Information S2.** ANCOVA analysis of basin visit times.

441 ANCOVA analysis of basin visit times.---Vigilance behaviors were examined using both direct 442 observations and reviews of video. The video data provided a standardized approach for 443 measuring the time spent at basins during the day. Between one and two trees were videoed at a site on any given day. 90% of visits to basins occurred between 6:00am and 9:30am. 444 445 Unfortunately, we only had enough video data to statistically analyze House group. Foraging 446 times for House group did not appear to change over time, however animals appeared to spend 447 less time at the ground basins (figure S1). These patterns were investigated using an ANOVA, in 448 which we log-transformed the times to normalize residuals, treated sampling day as a covariate, 449 and treated basin height as a factor with two levels: ground, aboveground. The statistical analysis 450 was performed using the lm function in R v. 3.1.3 (www.r-project.org).

451



452

Figure S1. Mean time each day that experimental basins with peanuts established in the home ranges of two monkey groups were visited during our study in Western Soutpansberg Mountains, South Africa in 2013. Video data are presented for the two groups of monkeys we studied, and basins have been grouped according to whether or not they were placed on the ground, the most risky stratum for arboreal monkeys. The number of trees videoed each day is also indicated. Vertical blue bars delineate the two experimental manipulations that delineate the three periods: pre-trapping, post-trapping without trap stimulus, and post-trapping with trap stimulus.

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