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1	Sleep patterns, daytime predation and the evolution of diurnal sleep site selection
2	in lorisiforms
3	
4	Magdalena S Svensson ¹ *, KAI Nekaris ^{1,2} , Simon K Bearder ¹ , Caroline Bettridge ³ , Thomas M Butynski ^{1,4} ,
5	Susan M Cheyne ⁵ , Nabajit Das ^{1,6,7} , Yvonne A de Jong ^{1,4} , Averee M Luhrs ¹ , Lydia Luncz ⁸ , Simon T
6 7	Maddock ^{9, 10} , Andrew Perkin ^{1,11} , Elizabeth Pimley ^{1,12} , Stephanie A Poindexter ¹ , Kathleen D Reinhardt ¹ , Denise Spaan ^{1,13} , Danica J Stark ^{14,15} , Carly R Starr ¹⁶ , Vincent Nijman ^{1,2}
8	
9	¹ Nocturnal Primate Research Group, Oxford Brookes University, Oxford, UK
10	² Little Fireface Project, Rumah Hijau, Cipaganti, Garut, Indonesia
11	³ Manchester Metropolitan University, Manchester, UK
12	⁴ Eastern Africa Primate Diversity and Conservation Program, Nanyuki, Kenya
13	⁵ Borneo Nature Foundation, Palangka Raya, Indonesia
14	⁶ Primate Research Centre NE India, Guwahati, Assam, India
15	⁷ Department of Zoology, B.H. College, Gauhati University, Howly, Assam, India
16	⁸ Institute of Cognitive and Evolutionary Anthropology, University of Oxford, Oxford, UK
17	⁹ Faculty of Science and Engineering, University of Wolverhampton, Wolverhampton, UK
18	¹⁰ Department of Life Sciences, The Natural History Museum, London, UK
19	¹¹ Tanzania Forest Conservation Group, Dar es Salaam, Tanzania
20	¹² Department of Natural and Social Sciences, University of Gloucestershire, Cheltenham, UK
21	¹³ Instituto de Neuroetologia, Universidad Veracruzana, Xalapa, Mexico
22	¹⁴ Organisms and Environment Division, Cardiff School of Biosciences, Cardiff University, Cardiff, UK
23	¹⁵ Danau Girang Field Centre, c/o Sabah Wildlife Department, Sabah, Malaysia
24	¹⁶ Northern Gulf Resource Management Group, Georgetown, Queensland, Australia
25	
26	
27 28 29 30 31 32 33 34	Corresponding Author: Magdalena Svensson Oxford Brookes University Nocturnal Primate Research Group Oxford OX3 0BP, UK <u>m.svensson@brookes.ac.uk</u> Number of pages: 29
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41 ABSTRACT

Objectives: Synthesize information on sleep patterns, sleep site use, and daytime predation at sleep
sites in lorisiforms of Asia and Africa (ten genera, 36 species), and infer patterns of evolution of sleep site
selection.

45 **Materials and methods**: We conducted fieldwork in twelve African and six Asian countries, collecting 46 data on sleep sites, timing of sleep and predation during daytime. We obtained additional information 47 from literature and through correspondence. Using a phylogenetic approach, we established ancestral 48 states of sleep site selection in lorisiforms and traced their evolution.

49 Results: The ancestral lorisiform was a fur-clinger and used dense tangles and branches/forks as sleep 50 sites. Use of tree holes and nests as sleep sites emerged ~22 Mya (range 17-26 Mya) in Africa, and use 51 of bamboo emerged ~11 (7-14) Mya in Asia and later in Africa. Fur clinging and some sleep sites (e.g. 52 tree holes, nests, but not bamboo or dense tangles) show strong phylogenetic signal. Nests are used by 53 Galagoides, Paragalago, Galago and Otolemur, tree holes by Galago, Paragalago, Sciurocheirus and 54 Perodicticus; tangles by Nycticebus, Loris, Galagoides, Galago, Euoticus, Otolemur, Perodicticus and 55 Arctocebus; all but Sciurocheirus and Otolemur additionally sleep on branches/forks. Daytime predation 56 may affect sleep site selection and sleep patterns in some species of Nycticebus, Galago, Galagoides, 57 Otolemur and Perodicticus. Most lorisiforms enter their sleep sites around sunrise and leave around sunset; several are active during twilight or, briefly, during daytime. 58

59 Conclusion: Variations in sleep behavior, sleep patterns and vulnerability to daytime predation provide a 60 window into the variation that was present in sleep in early primates. Overall, lorisiforms use the daytime 61 for sleeping and no species can be classified as cathemeral or polycyclic.

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65 To understand broader evolutionary implications of sleep among vertebrates, including sleep architecture, 66 type of sleep, intraspecific variation in sleep, sleep duration, and the ecological pressures selecting for 67 sleep and sleep site selection, a comparative approach is required (Elgar, Pagel and Harvey, 1988; 68 Lesku, Roth II, Amlaner and Lima, 2006; Rattenborg, Martinez-Gonzalez and Lesku, 2009). Sleep can 69 comprise more than 50% of a primate's activity budget (Campbell and Tobler, 1984). Sleep can occur in 70 single continuous bouts or take the form of fragmented sleep with periods of non-sleep and activity amidst 71 otherwise continuous sleep bouts. Thus, knowledge of sleep site selection and sleep patterns can provide 72 valuable insights into a species' ecology, social behavior, and habitat requirements (Anderson, 2000; 73 Mueller and Thalmann, 2000; Gursky, 2003; Grow and Gursky-Doyen, 2010). Where primates choose to 74 sleep is not only related to their body size, degree of arboreality, competition, and pressure from 75 predation and/or parasites, but also to their activity pattern (Anderson, 2000; Eberle and Kappeler, 2004; 76 Lock and Anderson, 2013; Tagg, Willie, Petre and Haggis, 2013).

77 More than 50% of primate species are nocturnal, yet comparative information on the ecology of sleep is 78 lacking for many nocturnal taxa, vital for constructing scenarios about the evolution of primate sleep 79 (Capellini, Barton, McNamara, Preston and Nunn, 2008). The use of sleep sites in primates varies 80 substantially, ranging from the ground, rocky outcrops, tree branches/forks, dense clumps of herbs and 81 lianas, sleep platforms, tree cavities and nests that are self-constructed or constructed by other species. 82 Use of nests (either self-constructed or made in tree holes or hollows) and platforms as sleep sites is 83 common among strepsirhines and great apes, and, presumably, the earliest humans (Sabater, Veá and 84 Serrallonga, 1997; Bearder et al., 2003; Fultz, Brent, Breaux and Grand, 2013; Samson and Shumaker, 85 2015b), but are rarely used by other haplorhines. Samson and Nunn (2015) distinguished these 86 assembled nests, on the basis that for larger primates, tree hollows would not be a viable sleeping option, 87 and suggest that ancestral Paleocene and Eocene primates probably had galago-like fixed point nest 88 use. Since most monkeys do not use nests, nest use must have evolved multiple times. To be able to 89 infer potential sleep site patterns in early primates (i.e. the ones for which only morphological data are 90 available), we also must examine how body size, forelimb to hindlimb ratio, and hand dexterity combine to 91 assist living primates in their sleep site choices (Covert, 2002; Gebo and Dagosto, 2004).

92 To examine the question further, Kappeler (1998) reviewed several explanations for the use of nests and 93 tree cavities amongst primates, especially among lemurs. Nests may serve as concealment against 94 predators and/or provide thermoregulatory benefits to prevent heat loss, especially for small and solitary 95 primates (Charles-Dominique and Martin, 1972). Kappeler (1998) also posited that nests and tree cavities 96 particularly benefit species with neonates too altricial to cling to their mother's fur by allowing them to be 97 placed in a safe location. Through phylogenetic analyses of multiple primate taxa, he concluded that the 98 latter hypothesis received most support for nocturnal strepsirhines. Kappeler (1998) notably lacked any in 99 situ study of Asian lorises [instead citing Rasmussen (1986) and Ehrlich and MacBride, (1989)]. 100 Regarding the paucity of field data on many primate taxa, he urged further research of wild primates to 101 understand better the evolution of sleep site selection.

Five years after Kappeler's review, Bearder et al. (2003), focusing on the African lorisiforms, also noted the scarcity of data on sleep sites and sleep patterns even though such data are vital to understanding diversity within nocturnal primates. Most of Bearder et al.'s (2003) data were based on studies conducted in the latter part of the last century. The authors found similarities among species within the same genus, but clear differences among genera.

107 In the twenty-first century, substantial taxonomic changes have occurred for both African and Asian 108 lorisiforms. First, the dwarf galagos of the genus Galagoides were recognized as a polyphyletic clade 109 (Pozzi et al., 2015), and now are comprised of Galagoides (western and central Africa) and Paragalago 110 (eastern Africa). Paragalago is a sister taxon to the genus Galago, and Galagoides and is a sister taxon 111 to the clade containing Sciurocheirus, Otolemur, Paragalago and Galago (Masters et al., 2017). Second, 112 divergence among lorisiforms is estimated to be far more ancient than previously thought; for instance 113 Euoticus split from other galagos ~30 Mya and Arctocebus split from Perodicticus ~23 Mya (Pozzi et al., 114 2015). Third, and related to the previous two points, more species are recognized (i.e. two additional species of Perodicticus, four Nycticebus, one Galagoides, and one Sciurocheirus). Fourth, researchers 115 116 studying nocturnal primates have amassed substantial new field data from countries such as Angola, 117 Cameroon, Equatorial Guinea, The Gambia, Kenya, Malawi, South Africa, Tanzania, Cambodia, India, 118 Indonesia, Sri Lanka and Vietnam (Nekaris, 2003a,b; Nekaris and Jayewardene 2003; Butynski and De

119 Jong, 2004, 2007, 2017; Pimley, Bearder and Dixson, 2005a,b; Butynski, De Jong, Perkin, Bearder and 120 Honess, 2006; De Jong and Butynski, 2009; Svensson and Bearder, 2013; Nekaris, 2014; Kenyon et al., 121 2014; Bersacola, Svensson and Bearder, 2015; Engelbrecht, 2016; Génin et al., 2016; Ray, Wren and 122 Bowers, 2016; Kumara, Sasi, Chandran and Radhakrishna, 2016; Kappeler et al., 2017). Fifth, 123 primatologists working on diurnal primates have taken an interest in certain lorisiforms, as lorisiforms 124 share sleep sites with diurnal primates (Llorente, Sabater and Houle, 2003), or are hunted by them 125 (Nishida, Uehara and Nyundo, 1979; Boesch and Boesch, 1989; Pruetz and Bertolani, 2007; O'Malley, 126 2010; Hardus et al., 2012).

127 Combined, the recent advancements in our understanding of lorisiforms allow for an overview of sleep 128 sites, sleep patterns, sleep associations, and predation pressure faced by lorisiforms while sleeping. 129 Using new genetic data on the relationships within Lorisiformes, we predict when various sleeping 130 patterns emerged within this group. The deep evolutionary divergence times between various lorisiform 131 genera help us explicitly to address several questions. Do lorisiforms provide evidence that the early 132 primate ancestors were fixed point nest users? Did nest using evolve multiple times amongst the 133 lorisiforms? Does the ability of a neonate to cling to the mother's fur relate to the use of fixed point nests? 134 These data can be used as a basis to understanding ancestral sleep behavior of primates that can help to 135 inform sleep patterns that occurred later in primate evolution.

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MATERIAL AND METHODS

We follow the taxonomy of Nekaris (2013a,b), but recognize the genus *Paragalago* (Masters et al., 2017), *Nycticebus kayan, N. bancanus* and *N. borneanus* (Munds, Nekaris and Ford, 2013), *Sciurocheirus makandensis* (Ambrose, 2013), and *Galagoides kumbirensis* (Svensson et al., 2017). We treat the Mount Kenya potto (*Perodicticus ibeanus stockleyi*) as a subspecies of *P. ibeanus*, not *P. potto* (Butynski and De Jong 2017). As such, we include 10 genera with 36 species of lorisiform. In the subsequent text, we abbreviate *Galagoides* as *Gd.* to distinguish it from *Galago* (*G.*), and *Paragalago* as *Pg.* to distinguish it from *Perodicticus* (*P.*).

Data collection

146 Post-2003 (i.e. after the publication of Bearder et al.'s 2003 compendium) we conducted nocturnal field 147 work in Angola (SKB, MSS; 1 mo), Cameroon (AML, TMB, YdJ; 3 mo), Democratic Republic of the Congo 148 (TMB; 2 mo), Equatorial Guinea (Bioko: TMB; 12 mo), Ethiopia (TMB; 1 mo), The Gambia (SKB, MSS; 1 149 mo), Kenya (TMB, YdJ; 34 mo), Nigeria (AL; 2 mo), Malawi (SKB; 1 mo), Rwanda (SKB, MSS; 1 mo), 150 Tanzania (TMB, YdJ, CB, AP; 19 mo), Uganda (TMB, YdJ, MSS, AML; 19 mo), Cambodia (CRS, KAIN; 151 11 mo), India (KAIN, ND; 32 mo), Indonesia (Java: KAIN, VN, KDR, DS; 60 mo; Sumatra: KAIN; 1 mo), 152 Malaysia (Borneo: DJS; 60 mo), Sri Lanka (KAIN, EP; 22 mo) and Vietnam (SAP, KAIN; 9 mo). We 153 collected most data on populations where individuals could not be individually recognized, but in Borneo, 154 Cambodia, India, Sri Lanka, Vietnam and Java, we followed identified individuals with radio collars or 155 other markers. We obtained additional data from published studies and through correspondence with 156 researchers, including those working on great apes (bonobos Pan paniscus, common chimpanzees P. 157 troglodytes, Sumatran orangutan Pongo abelii, Bornean orangutan P. pygmaeus and Tapanuli orangutan P. tapanuliensis: Nater et al., 2017) to obtain data on predation events. 158

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Analyses

We used species as the unit of analysis. We pooled data from studies to provide a global picture. Based on previous research (Bearder et al., 2003), we placed sleep site types into five groups: nests, tree holes or hollows, dense tangles of vegetation, tree branches/ forks, and bamboo thickets. We ranked the use of sleep sites types from zero (no evidence of use), one (occasional use or mixed evidence) or two (regular use).

To typify social cohesion, we collected data on sleep group size. When transporting infants, these can be carried in the parent's mouth or they can cling on to their parent's fur. For each species we recorded whether they carried infants in the parent's mouth or if they can cling to their parent's fur, or whether they employed both methods. Regarding vocalizations, we included call types (audible to humans) used for social cohesion, advertisement and maintenance. We exclude the ultrasonic calls of *Perodicticus, Loris*

and *Nycticebus*. We ranked vocalizations as one (social cohesion vocalizations displayed at sleep site) or zero (social cohesion vocalizations not displayed at sleep site). Intermembral Index (IMI, a ratio of forelimb length to hindlimb length) for the different species was taken from Fleagle (2013) and for slow and slender lorises from measurements taken by KAIN and DJS on wild-caught live animals.

To gain insight into sleep patterns and the presence of fragmented sleep in the lorisiforms, we compiled data on when individuals entered and exited sleep sites. From selected sites, we added information on pre- or post-dusk waking and pre- or post-dawn sleeping. We added observations of sleep during the night or non-sleep behavior during the day.

179 We examined evidence of predation on lorisiforms and highlight those instances where the events 180 occurred while the animal was asleep, or where we could reasonably infer that predation had taken place 181 during the daytime. We excluded predation events by nocturnal predators such as owls, but included 182 events from cathemeral or crepuscular predators. While we acknowledge that most lorisiforms, at least 183 occasionally, sleep for brief periods during the night, and that they may be subject to predation by 184 nocturnal predators at these times, this form of rest is distinctly different from them selecting and using a 185 sleep site where they will sleep during day time. Additionally, we compiled information on anti-predator 186 strategies used by lorisiforms and which of these might be most effective at sleep sites.

187 We carried out reconstruction of ancestral states on a subset of species for which full sleep site and fur 188 clinging behaviour and published genetic sequences were available. We obtained cytochrome b 189 sequences (1,140 bp in length) of 23 species of lorisiform from GenBank (for accession numbers see Fig. 190 1) and we aligned them with MAFFT v.7 multiple sequence alignment (Katoh and Standley, 2013). This 191 formed the basis of our ancestral state reconstruction analyses. We constructed phylogenetic trees using 192 BEAST v.2.4.6 (Drummond and Rambaut, 2007; Suchard and Rambaut, 2009; Bouckaert et al., 2014). 193 We implemented a strict clock with the birth-death speciation tree prior for 100 million generations, 194 sampling every 10,000 iterations. We checked analyses for convergence using Tracer v.1.6. We then 195 used the posterior probability tree produced by BEAST to perform stochastic character mapping 196 (Huelsenbeck, Nielsen and Bollback, 2013) to infer ancestral states of sleep site and fur clinging using the 197 R package phytools v.0.6-20 (Revell, 2012). Phylogenetic signal was tested for discrete character

evolution of each character by comparing AICc scores with and without phylogenetic error structure using
the fitDiscrete function in the R package, geiger v.2.0.6 (Harmon, Weir, Brock, Glor and Challenger,
2008). This signal was estimated by testing a model with complete phylogenetic independence (lambda
fixed to 0) to one with phylogenetic non-independence (free lambda tree transformation).

We fitted stochastic character histories for each character set by executing continuous-time reversible Markov models, to sleep sites and fur clinging, over 99,900 simulations each. We used an equal rates of transition model to sample the state transition matrix "Q" from the posterior probability. We used ancestral character estimation ("ace") to demonstrate the probabilities of states at each node. To date the timing of the ancestral states of sleep site and fur clinging, we used the timed phylogeny of Pozzi et al. (2015) to calculate mean values and 95% highest probability estimates in millions of years ago (Mya).

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RESULTS

Physical characteristics of sleep sites and evolution of sleep site selection

211 The type of sleep site lorisiforms most commonly used was tangles (67% or 24 of 36 species), followed 212 by holes (44%, 16 species), branches/forks (44%, 16 species), nests (either self-built or built by other 213 species: 33%, 12 species) and bamboo (14%, 5 species) (Table 1). Of the 24 species that use tangles, 214 62% also use branches/forks, 46% also use holes, and 42% also use nests. Of the 16 species that use 215 branches/forks, 94% also use tangles. Of the 16 species that use holes, 69% also use nests, 69% also 216 use tangles, and 31% also use branches/forks. Twelve species use nests, 92% of which also use holes 217 and 83% also use tangles. Of the five species that use bamboo, the four Asian lorises also use 218 branches/forks and tangles, but none use nests or holes, whilst Gd. demidovii mainly uses nests in dense 219 undergrowth, and, to some extent, tree holes and tangles. Species in which infants cling to the adult's fur 220 do not tend to use nests or tree holes.

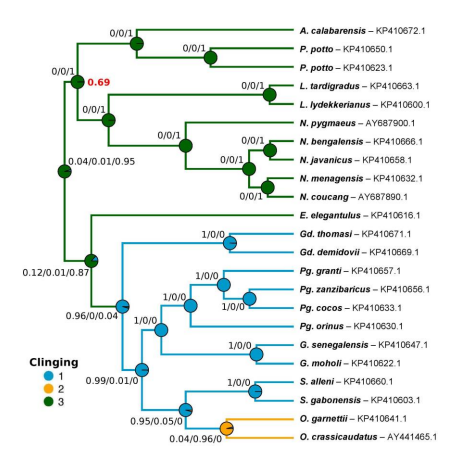
The IMI ranges from lows of around 50 in *Galago* spp, representing clear vertical clingers and leapers with legs twice as long as their arms, to an intermediate value of around 70 in *Otolemur* and *Galagoides*, and highs of over 90 in *Loris* and *Nycticebus*, with arms and legs being almost the same length. Species

with low IMIs tend to be those that carry their young in their mouth, and that use nests and tree holes, whereas species with intermediate IMIs tend to have infants that cling on the adult's fur and do not use nests or tree holes (Table 2).

At least four lorisiforms use human-made sleep sites. *Galago senegalensis* sleeps in traditional bee-hives (hollowed tree boles), birdhouses, and roofs of buildings while *G. moholi* uses ventilation pipes at some study sites. *Otolemur crassicaudatus* sleeps in traditional bee-hives and roofs of buildings, and *Pg. cocos* is also known to utilize human-made sleep sites.

231 Phylogenetic relationships showed strong support for all splits except for the sister group relationship 232 between Artocebus + Perodicticus and Nycticebus + Loris (bpp = 0.63) (Fig. 1 and 2). Fur clinging and 233 some sleep sites show strong phylogenetic signal under a lambda transformation model: fur clinging 234 (estimated lambda = 1, AICc = 25.175) is a better fit (Δ AICc = 22.08) than a model with no phylogenetic 235 signal (lambda fixed to 0, AICc = 47.255); tree hole (estimated lambda = 1, AICc = 33.64) is a better fit 236 $(\Delta AICc = 19.10)$ than a model with no phylogenetic signal (lambda fixed to 0, AICc = 52.74); 237 branches/forks (estimated lambda = 0.98, AICc = 28.96) is a better fit (\triangle AICc = 2.948) than a model with 238 no phylogenetic signal (lambda fixed to 0, AICc = 31.91). Nests showed some support for phylogenetic 239 signal (estimated lambda = 0.557, AICc = 47.01) and is a better fit (\triangle AICc = 0.744) than a model with no 240 phylogenetic signal (lambda fixed to 0, AICc = 47.75). Two sleep sites showed no support of phylogenetic 241 signal: bamboo (estimated lambda = 0.363, AICc = 35.559) is a worse fit (\triangle AICc = -0.144) than a model with no phylogenetic signal (lambda fixed to 0, AICc = 35.415); dense tangle (estimated lambda = 0.607, 242 243 AICc = 44.78) is a worse fit (Δ AICc = -1.462) than a model with no phylogenetic signal (lambda fixed to 0, 244 AICc = 43.318). Although stochastic character histories were estimated for all datasets, no information 245 about ancestral evolution should be drawn from sleep sites in bamboo and dense tangles and the use of 246 nests should be interpreted very loosely due to a lack of signal.

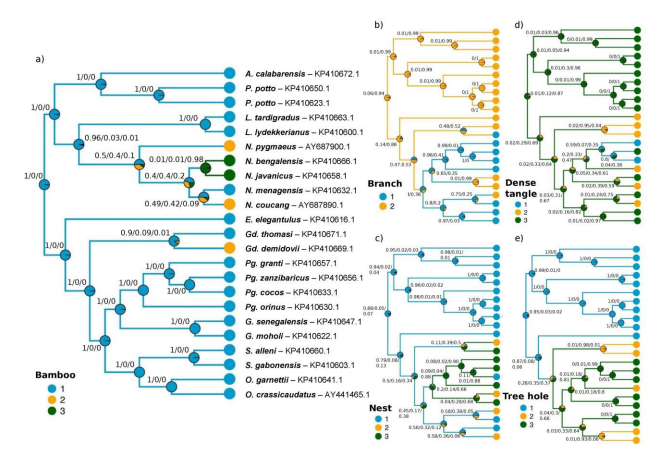
At ~40 Mya (range 36-44 Mya, nb. all dates used herein are taken from Pozzi et al., 2015), we suggest that the ancestral lorisiform infant was carried by clinging to the fur of its parent (Fig. 1). This ancestral state is retained in all Asian taxa as well as in some African taxa (e.g. *Perodicticus, Arctocebus,* and *Otolemur*). We propose that carrying infants in the mouth evolved ~22 (17-26) Mya in the ancestor of the African galagos. We assume that the ancestral lorisiform used dense tangles and branches/forks as sleep sites. Almost all extant species still use dense tangles as sleep sites, but we suggest that this trait was lost twice in the east African coastal *Paragalago* species. While the majority of species still use branches/forks as sleep sites, we propose that this trait changed at ~14 (12-18) Mya for *Paragalago* and ~12 (8-15) Mya, when *Otolemur* and *Sciurocheirus* split from the other galagos (Fig. 2).



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FIGURE 1 Ancestral state reconstructions of stochastic character mapping of lorisiform fur-clinging
 whereby infants cling onto the fur of their parents when being transported. Red numbers indicate the
 Bayesian posterior probabilities of the phylogenetic tree if <1. Branches above nodes (closer to tips) are
 coloured based on their ancestral state probability. Pie charts on nodes and black numbers (states 1/2/3)
 indicate the probability of the state in the common ancestor. The states are in the following order: 1 =
 absent, 2 = occasionally present, 3 = present.

The use of nests is restricted to the African lorisiforms and likely emerged ~22 (17-26) Mya, after *Euoticus* split from other galagos (Fig. 2). Use of bamboo as a sleep site appears to have emerged early on in their evolution at ~11 (7-14) Mya, after *Nycticebus* split from *Loris*. At present, all *Nycticebus* species, apart from *N. menagensis,* are known to use bamboo as sleep sites. Independently, *Gd. demidovii* uses bamboo as a sleep site but bamboos are absent over most of its geographic range; this behavior probably emerged in the last 5 million years.



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FIGURE 2 Ancestral state reconstructions of stochastic character mapping of lorisiform sleep site use: a) bamboo, b) branch, c) dense tangle, d) nest, e) tree hole. Branches above nodes (closer to tips) are coloured based on their ancestral state probability. Pie charts on nodes and black numbers (states 1/2/3) indicate the probability of the state in the common ancestor. The states are in the following order: 1 = absent, 2 = occasionally present, 3 = present; except for the branch sleep site where: 1 = absent, 2 = present.

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Sleep patterns

Most lorisiforms enter their sleep site between 0.5 hr before and 0.5 hr after sunrise, and leave their sleep site between 0.5-1.0 hr before and 0.5-1.0 hr after sunset. Several lorisiforms are active (moving, feeding and calling) during twilight: e.g. *N. javanicus* and *O. garnettii* up to 1.5 hrs before sunset, and *S. alleni*, *Pg. cocos*, *Pg. zanzibaricus* and *G. senegalensis* up to 1.0 hr before sunset.

286 In Africa, the number of daylight hours (time between sunrise and sunset), and thus the numbers of hours 287 available for sleep, varies between ~13 hrs (June) and ~11 hrs (December) in Senegal and Eritrea, ~10 288 hrs (June) and ~14 hrs (December) in southern Africa, and ~12 hrs (year-round) in East Africa (Kenya, 289 Tanzania and Uganda). We found no evidence that species in the more northern or southern regions 290 adjust their sleep pattern. In general, for most species, sleep is an equitable 12 hrs year-round. In Asia, 291 N. bengalensis in northeastern India, Myanmar and China, have ~10 hrs of daylight available for sleep in 292 December and ~14 hrs in June; again, we found no evidence to suggest that they adjust their sleep 293 pattern. The southern-most populations of lorisiforms in Asia are found in Sri Lanka (Loris tardigradus) 294 and Java (*N. javanicus*), both situated ~7° north and south of the equator, respectively. As such, annual 295 variation in daylight hours is small and sleep is equitable 12 hrs year-round.

296 Numerous lorisiforms, including N. javanicus, G. gallarum, G. senegalensis and O. garnettii are 297 sometimes active during the day, presumably only for short periods and possibly in response to being 298 disturbed by humans, adverse weather or because of (real or perceived) predator threats. Galago 299 senegalensis occasionally sleep in the middle of the night, but the lengths of these sleep bouts remain 300 unknown. Additionally, G. moholi, N. javanicus and N. pygmaeus occasionally sleep during the night. 301 These species have been known to use daily and multiday torpor, which may suggest they are indeed in 302 a state of torpidity, and not sleeping (Nowack, Mzilikazi and Dausmann, 2013a; Ruf, Streicher, Stalder, 303 Nadler and Walzer, 2015; Reinhardt, Wirdateti and Nekaris, 2016). Overall, however, the daytime is used 304 for sleeping and we could classify no species as cathemeral or polycyclic.

TABLE 1 Sleep site type: 0 - no evidence of use, 1 - irregular or occasional use or mixed evidence use from different studies, 2 - regular or habitual use of nests, ? - evidence is based on anecdotal information or when information is lacking. * - using man-made structures as sleep sites Social cohesion: 0 - no, 1 - yes.

	nformation	or when	informati				an-mad	e structu				sion: 0 – no		_	
Species				Slee	ep site ty	уре			Social	cohesion			Non-human diurnal and crepuscular predators (Confirmed	Predator avoidance strategies	Reference
	Intermembral Index	Mouth-carrying	Fur-clinging	Nest	Tree hole	Dense tangle	Branch/fork	Bamboo	Audible advertising calls	Audible contact maintenance calls	Allogrooming at sleep site	Sleep group size (range)	records in bold)		
Galagoides demidovii	68	2	0	2	1	1	0	1	1	1	?	2-10	Sooty mangabeys, bonobos		1; 2; 3; 4 ; 5; 6
Gd. thomasi	67	2	0	1	1	1	1	0	1	1	?	<5	Snakes, hawks, hornbill, viverrids, mongooses, blue monkey		7; 8; 9
Gd. kumbirensis		?	?	?	?	?	?	?	?	?	?	?	?		10
Paragalago orinus		1	0	2	1	1	?	0	1	?	1-5	1-9	Snakes, raptors, genets, Sykes's monkeys		11;12; 13
Pg. rondoensis		2	0	2	0	?	?	?	1	0	?	<3	Snakes e.g. forest cobras, boomslangs, green mambas)		13
Pg. granti		2	0	2	2	0	0	0	1	0	?	4-5	Snakes e.g. forest cobras, boomslangs, green mambas)		13; 14; 15
Pg. cocos		2	0	0	2*	0	0	0	1	1	?	1-5	Snakes e.g. forest cobras, boomslangs, green mambas), <i>Cercopithecus</i> monkeys		16; 17
Pg. zanzibaricus	60	2	0	2	2	2	0	0	1	1	?	1-5	Snakes e.g. forest cobras, boomslangs, green mambas)		13; 16; 17; 18
Galago senegalensis	52	2	0	1	2*	2	1	0	1	1	1	<6	Chimpanzees		19; 20; 21; 22; 23; 24; 25; 26; 27; 28; 29;30
G. gallarum		?	0	1	1	2	2	0	1	?	1	≤3	Snakes, raptors, jackals, mongooses, genets, wild cats	Live in the thorniest habitats of all galagos. Sleep and rest in the core of thorny vegetation.	17; 31; 32; 33; 34; 35
G. moholi	54	2	0	2	2*	1	1	0	1	0	?	1-8	Large snakes & monitor lizards, genets, Verreaux's eagle, small carnivores,		13; 36; 37; 38; 39; 40; 41
G. matschiei		2	0	?	2	?	?	?	1	0	?	?	Large snakes, viverrids, blue monkeys, baboons, chimpanzees		7; 42
Sciurocheirus alleni	65	2	0	1	2	2	1	?	1	1	1	1-6	Large snake, viverrids		1; 16; 43; 44
S. gabonensis		2	0	0	2	?	0	?	1	?	?	1-3	Large snakes, viverrids, leopard, African golden cat		16; 45
S. cameronensis		2	0	1	2	?	?	?	?	?	?	1-6			16; 44; 46
S. makandensis		?	0	?	?	?	?	?	1	1	?	1-4	Large snakes, viverrids, golden cats		47; 48
Euoticus elegantulus	64	0	2	0	0	1?	1?	0	1	1	?	3-4	Pythons, viverrids		1; 13; 16
E. pallidus		0	2	0	2	0	1	0	1	1	?	1-4	Central African linsang		13; 16; 49
Otolemur crassicaudatus	70	1	1	1	1*	2	0	?	1	0	?	1-4	Large snakes, raptors, leopards, chimpanzees		13; 16; 17; 50
O. garnettii	69	1	1	0	1	2	?	?	1	1	?	1-4	Large snakes, raptors		17; 51; 52; 53

Arctocebus calabarensis	89	0	2	0	0	2	1	0	0	0	?	1-2	Snakes, viverrids, monkeys		13
A. aureus		0	2	0	0	2	1	0	0	0	?	1-2	Large snakes, viverrids,		13
Perodicticus potto	88	0	2	0	0	2	1	0	0	0	?	?	Large snakes, African crowned eagles , civets, black-legged mongoose, leopards , African golden cats, <i>Cercopithecus</i> monkeys, mandrills, chimpanzees	Scapular neck shield, predator defense posture, drops to ground	6; 19; 54; 55; 56; 57
P. edwardsi		0	2	0	?	2	?	0	0	0	?	1-2		Scapular neck shield, predator defense posture, drops to ground	43; 58; 59
P. ibeanus		0	2	?	?	2	?	0	0	0	?	?	Leopard	Scapular neck shield, predator defense posture, drops to ground	54; 5
Nycticebus javanicus	93	0	2	0	0	2	1	2	0	0	1	4		Venomous, predator defense posture	60
N. bengalensis		0	2	0	0	2	1	2	0	0	1	?		Venomous, sleeps high in trees inaccessible positions or in dense thorny tangles	61
N. menagensis	91	0	2	0	0	2	1	0	0	0	1	1-3	Reticulated pythons, raptors	Venomous, predator defense posture	62
N. pygmaeus	91	0	2	0	0	2	1	1	1	1	1	1-5	Monitor lizards, raptor, small carnivores	Venomous, predator defense posture	63; 64
N. coucang	91	0	2	0	0	2	1	1	0	1	1	3	Reticulated python, monitor lizards	Venomous, predator defense posture	65; 66
N. hilleri	89	0	2	0	0	2	1	1	1	1	?	?	Changeable hawk eagle, Sumatran orangutans	Venomous, predator defense posture	67; 68; 69; 70
N. kayan		?	?	?	?	?	?	?	?	?	?	?		Venomous, predator defense posture	
N. bancanus		?	?	?	?	?	?	?	?	?	?	?	?	?	
N. borneanus		?	?	?	?	?	?	?	?	?	?	?	?	Venomous, predator defense posture	
Loris tardigradus	90	0	2	0	0	2	1	0	1	1	1	4		Sleeps in dense tangles, has cobra defense posture	71
L. lydekkerianus	92	0	2	0	0	2	1	0	1	1	1	4-5	Rusty spotted cat	Sleeps in dense tangles, has cobra defense posture	72; 73

1) Charles Dominique, 1977; 2) Bearder & Honess, 1992; 3) Hohmann & Fruth, 2008; 4) Ambrose & Butynski 2013a; 5) A. Luhrs, pers. obs.; 6) E. Pimley, pers. obs.; 7) Butynski, 1982; 8) Llorente et al., 2003; 9) Ambrose & Butynski 2013b; 10) Svensson et al., 2017; 11) Perkin, 2000; 12) Doody et al., 2001; 13) Nekaris & Bearder, 2011; 14) Butynski et al., 2006; 15) Génin et al., 2016; 16) Kingdon, 2015; 17) Y. De Jong & T. Butynski, pers. obs.; 18) Honess, Perkin & Butynski, 2013; 19) McGrew, Tutin & Baldwin, 1978; 20) Nishida et al., 1979; 21) Uehara, 1997; 22) Pruetz & Bertolani, 2007; 23) Off, Isbell & Young, 2008; 24) De Jong & Butynski 2009; 25) O'Malley, 2010; 26) Nash, Zimmermann & Butynski, 2013; 27) Svensson & Bearder, 2013; 28) Butynski & De Jong, 2014; 29) Butynski & De Jong, 2017; 30) Pruetz et al., 2015; 31) Butynski & De Jong & Butynski 2004a; 34) De Jong & Butynski, 2004b; 35) De Jong & Butynski, 2010; 36) Mzilikazi, Masters & Lovegrove, 2006; 37) Nowack, Mzilikazi & Dausmann, 2013; 34) De Jong & Butynski, 2004a; 34) De Jong & Butynski, 2004b; 35) De Jong & Butynski, 2010; 36) Mzilikazi, Masters & Lovegrove, 2006; 37) Nowack, Mzilikazi & Dausmann, 2013; 41) Pullen & Bearder, 2013; 42) Ambrose, 2002; 44) Ambrose & Pimley 2013; 45) Ambrose, 2013; 46) Nekaris, 2013b; 47) Ambrose, 2003; 48) Ambrose & Oates, 2013; 50) Rovero, Marshall, Jones & Perkin, 2009; 51) Lumsden & Masters, 2001; 52) De Jong & Butynski, 2004b; 35) Harcourt & Perkin, 2013; 54) Hart, Katembo & Punga, 1996; 55) Msuya, 2003; 56) Shultz, Noë, McGraw & Dunbar, 2004; 57) Pimley & Bearder, 2013; 58) Pimley, Bearder & Dixson, 2005b; 50) Nekaris et al., 2017; 61) N. Das, K.A.I. Nekaris & S.A. Poindexter, pers. obs.; 62) D.J. Stark, pers. obs.; 63) Kenyon et al., 2014; 64) K.A.I. Nekaris & Jayewardene, 2003; 72) Bearder, Nekaris & Buzyeli, 2002; 73) Nekaris, 2003b

305 Predation at sleep sites

Predation avoidance appears to be a main factor in sleep site choice. Benefits are associated with all the sleep site types regarding protection against predation. Known predators of lorisiforms include a wide range of species, including those that target lorisiforms at their sleep sites (Table 1). Snakes and monitor lizards can access tree holes and branches/forks, whereas monkeys and apes, and possibly also some snakes, can access tree holes and tangles. Among reptiles, monitor lizards *Varanus* spp. and reticulated python *Malayopython reticulatus* prey on *N. pygmaeus* and *N. coucang.* The smoothness of bamboo stems may provide protection for *Nycticebus* spp. and *Gd. demidovii*.

Diurnal raptors prey on lorisiforms, although recorded captures are scarce. Predators known to prey on lorisiforms are: crowned eagles *Stephanoaetus coronatus* on *P. potto* and *Galago* spp., Verreaux's eagle *Aquila verreauxii* on *G. moholi,* and changeable hawk-eagle *Nisaetus cirrhatus* on *N. coucang.* These captures likely took place during the day when the lorisiforms were at their sleep site.

317 Small mammalian carnivores, such as palm civets, linsangs and genets (Viverridae), may capture 318 lorisiforms when they enter or leave their sleep sites. Remains from *P. ibeanus* have been found in 319 leopard *Panthera pardus* scats and African palm civets *Nandina binotata* are known predators of *P.* 320 *edwardsi.*

321 Blue monkeys Cercopithecus mitis prey on Gd. thomasi and/or G. matschiei, with predation observed in 322 the afternoons. Sooty mangabeys Cercocebus atys have been observed poking Gd. demidovii out of their 323 nests with sticks. On Sumatra Nycticebus hilleri has been observed to be captured and killed during 324 daytime by P. abelii - we obtained confirmation from two sites-whereas there are no indications thus far 325 that further south on Sumatra that N. coucang is eaten by P. tapanuliensis (Table 2). Data from Borneo 326 indicate that predation on Nycticebus spp. by P. pygmaeus is less frequent, as we only were able to get 327 confirmation of one species, N. menagensis, being eaten at one site (Table 2). Pan troglodytes 328 occasionally prey on Galagidae (Fig. 3), mainly when sleeping in tree holes, sometimes using tools such 329 as sticks. Pan paniscus have been observed to force Gd. demidovii out of tree holes by inserting fingers 330 into the hole and then hitting the trunk (Table 2).

Humans are probably one of the main predators of lorisiforms. This relationship is especially true in Asia where *Nycticebus* and *Loris* are taken to meet the demand for the pet and traditional medicine trades, and where specialized hunters seek out sleep sites during the day.



FIGURE 3 Young chimpanzee Pan troglodytes in Guinea holding a dead northern lesser galago Galago
 senegalensis, having caught it in the daytime. Photo by: Chimpanzee Conservation Center / Charlotte
 Houpline.

TABLE 2 Day-time predation of lorisiforms by great apes

Species	Site, country	Predation	Reference
Pan paniscus	Lui Kotale, DR Congo	Galagoides demidovii forced out of tree hole	Hohmann & Fruth, 2008
P. troglodytes	Gombe Stream, Tanzania	Galago senegalensis retrieved from trunk and consumed	O'Malley, 2010
P. troglodytes	Fongoli, Senegal	<i>G. senegalensis</i> , frequently hunted, chimpanzees using tools	Pruetz & Bertolani, 2007; Pruetz et al., 2015
P. troglodytes	Mt. Assirik, Senegal	<i>G. senegalensis</i> and <i>Perodicticus potto,</i> remains found in fecal samples	McGrew et al., 1978
P. troglodytes	Mahale Mountains, Tanzania	<i>Otolemur crassicaudatus</i> and <i>Galago</i> spp. harassed and consumed by chimpanzees	Nishida et al., 1979; Uehara, 1997
P. troglodytes	Haute Niger, Guinea	Galagos . Chimpanzees observed killing, but not consuming, galagos.	C. Colin, pers. comm.
P. troglodytes	Ngogo / Kanyawara, Kibale, Uganda	Galago spp. and Perodicticus ibeanus. Interactions with galagos frequently observed. Female chimpanzee offspring carried dead galago, treated it like a baby	J. Negrey, pers. comm.; R. Wrangham, pers. comm.
P. troglodytes	Bossou forest, Guinea	Perodicticus potto harassed by chimpanzees	K. Hockings, pers. comm.
P. troglodytes	Tai, Ivory Coast	P.potto , females and offspring observed to hunt and feed on pottos	L. Luncz, pers. obs.; E. Pimley, pers. obs.
Pongo abelii	Ketambe, Indonesia	<i>Nycticebus hilleri</i> , observations of orangutans eating slow lorises	Utami & van Hooff, 1997; Hardus et al., 2012; S.S. Utami Atmoko, pers. comm., S. Rimba, pers. Comm.
P. abelii	Suaq, Indonesia	N. hilleri , observations of orangutans harassing and eating slow lorises	C. Schuppli, pers. comm.
P. pygmaeus	Tuanan, Indonesia	<i>N. menagensis,</i> observation of orangutan eating a slow loris	E. Vogel, pers. comm.

DISCUSSION

344 We show that lorisiforms use a wide range of sleep sites, with most taxa sleeping in dense tangles, 345 followed by holes and on branches/forks. Fewer species use nests and bamboo. It appears that the 346 ancestral lorisiform would have used dense tangles, and branches/forks as sleep sites. We suggest that 347 the use of tree holes and nests as sleep sites emerged ~30 (24-36) Mya in Africa, and the use of bamboo 348 as a sleep site emerged ~31 (23-26) Mya in Asia and later in Africa. The ability of infants to cling onto 349 their parents' fur appears to be the ancestral condition, and carrying infants in the mouth is a derived 350 condition and emerged in the African taxa. Our data provide support for Kappeler's (1998) hypothesis that 351 use of nests and tree holes is linked to having altricial infants that are not able to cling to fur, thus 352 providing them with a relatively safe location while adults forage.

353 Further understanding the comparative morphology of fur clingers may help us to infer nest using 354 behavior in the fossil record. We found a strong relationship between more generalized arboreal 355 lorisiforms with an IMI nearer to 100 with the absence of nest use and infants that fur cling. Tree hole use 356 was limited to animals with the lowest IMIs that are generally vertical clingers and leapers. Functionally, 357 animals with shorter arms, and hence lower IMI, might not be able to cling as well on tangles and 358 branches. Such morphological adaptations are further emphasized by the presence of a retia mirabilia 359 (where the arteries form vascular bundles that allows blood to flow even when the animal remains still) in 360 Loris, Nycticebus and Perodicticus allowing an enhanced grip (Ankel-Simons, 2007; Congdon and 361 Ravosa, 2016). In the fossil record IMI and the ability to engage in specialized grasping may help us to 362 interpret the sleeping patterns and sleep site selection of extinct species such as Carpolestes simpsoni, 363 that resemble the more generalized arboreal lorisiforms in this study that did not use tree holes (Bloch 364 and Boyer, 2002).

Sleep site selection can be seen partly as an anti-predation strategy, depending on predator species and densities (Charles-Dominique and Martin, 1972; Anderson, 2000). Factors important in reducing daytime predation appear to be: connectivity of sleep trees, use of thorny bushes, nest hole entrance size and selection of dense tangles of lianas and undergrowth and smooth-surfaced substrate such as bamboo.

The entrance size of sleep holes used by lorisiforms tends to be no larger than is necessary for the individual to enter. This minimizes the number of predator species that can enter or reach inside. Selection of tree holes with suitably small entrances that only enable the strepsirhine to enter/exit is not always possible, especially when the number of trees holes in an area is limited. As a result, holes with larger entrances are sometimes used. For example, *S. cameronensis* used tree holes with entrances of 20 cm diameter larger than necessary for it to access the hole (Pimley, 2002). More studies that systematically measure tree holes used by lorisiforms are needed.

376 Many species reuse sleep sites in an unpredictable order. This reuse allows them to become familiar with 377 the sleep site and facilitate escape during predation attempts (Di Bitetti, Vidal, Baldovino and 378 Benesovsky. 2000; Nekaris, 2003a; Qihai, Chengming, Ming and Fuwen, 2009; Svensson and Bearder, 379 2013). Rotation of sleep sites makes it more difficult for predators to ambush prey (Di Bitetti et al., 2000). 380 Other species, such as G. gallarum, frequently sleep on branches/forks in the center of trees or bushes 381 among a barrier of dense thorns, and use such areas on consecutive days (De Jong and Butynski, 382 2004a,b; Butynski and De Jong, 2013). Sleeping in tangles of dense vegetation reduces detection from 383 predators, provides protection from the elements and facilitates rapid escape, especially for smaller 384 lorisiforms such as Galagoides, Paragalago and Loris (Kappeler, 1998). Vegetation tangles and bamboo 385 have also been hypothesized as anti-predation strategy for Loris and Nytcicebus (Nekaris, 2014). The 386 slow and slender lorises, angwantibos and pottos are non-saltatory arboreal climbers, incapable of 387 leaping (Sellers, 1996). This locomotion demands constant connectivity to maintain substrate contact, as 388 well as an increased number of escape routes from predators (Voskamp, Rode, Coudrat, Wilson and 389 Nekaris, 2014).

Researchers have found animals exposed to high levels of predation to display less time spent in sleep, while those with less disturbances experience increased sleep quality (Samson and Shumaker, 2013, 2015a). This behavior is largely due to disturbances from predators during the sleeping period, as well as a need to be more alert (Zepelin, 2000; Lima, Rattenborg, Lesku and Amlaner, 2005). More field research on sleep quality is needed in primates, to determine if different sleep site types and predation pressures influence sleep patterns.

The use of different types of sleep sites within the same species of lorisiform suggests these species may be opportunistic generalists that are able to use the range of habitat features available to them or respond to varying sleep site selection pressures. Similar variability is seen in other primate species that have access to the same types of sleep sites in different environments, but do not select them based on differences in the site characteristics (Pontes and Soarse, 2005; Duarte and Young, 2011). Despite the range of sleep site types seen across the group however, sleep patterns are mostly consistent, with all species demonstrating nocturnal, not cathemeral behavior.

Given that most lorisiforms live in the tropics and only a few in the subtropics, with small amounts of variation in day length, most species have equal amounts of time available for sleep. The exceptions are *N. bengalensis*, and the southern-most populations of *G. moholi* and *O. crassicaudatus*, which have 4 hrs less available for sleep in winter than in summer (for *N. bengalensis* this is the boreal winter and for *G. moholi* and *O. crassicaudatus* it is the austral winter, when nights are significantly longer than in the summer). Several species of lorisiform are active for short periods during the day (Bearder, Nekaris and Curtis, 2006). These patterns are possibly linked to low temperatures or other adverse conditions.

410 Thermoregulation may be an important factor in sleep site selection in lorisiforms, especially for species 411 that live at high elevations and/or at high or low latitudes (Ruf, et al., 2015). Tree holes provide good 412 insulation against the cold (Schmid, 1998; Daussman, Glos, Ganzhorn and Helmaier, 2004), and buffer 413 against heat. Nests may also serve thermoregulatory requirements (Radespiel, Cepok, Zietemann and 414 Zimmermann, 1998; Lutermann, Verburgt and Rendigs, 2010; Nowack et al., 2013a). Galago moholi 415 return to their sleep site (usually a tree hole but also nests) earlier than usual during cold nights to huddle 416 with other individuals thus serving a thermoregulatory purpose (Bearder and Martin, 1980). The 417 requirement for thermoregulation may also explain why the smaller species of lorisiform such as Loris, 418 Galago, Galagoides, Paragalago and Sciurocheirus (which lose heat more quickly than their larger 419 relatives) tend to sleep together, thereby sharing body heat (Nowack, Wippich, Mzilikazi and Dausmann, 420 2013b; cf. Eppley, Watzek, Dausmann, Ganzhorn and Donati, 2017). Some of these smaller taxa, 421 including Pg. zanzibaricus and Gd. thomasi, bring in fresh plant material to line their sleep site (Bearder et 422 al., 2003).

423 Peckre et al. (2016) pointed out the need for more studies on infant carrying and its relevance to primate 424 evolution, in particular regarding the evolution of an enhanced grip. Based on nearly 20 years of new field 425 data, we help to confirm the view that fur clinging is an ancestral trait in lorisiforms, and that fur clinging 426 species rarely or never use tree holes (cf. Kappeler, 1998). Lorises and pottos have a shared derived trait 427 called the retia mirabilia, as well as a reduced second digit (Ankel-Simons, 2007). These morphological 428 traits produce an enhanced grip that has been suggested to be an anti-predation strategy (Charles-429 Dominique, 1977; Nekaris, 2014; Oates, 1984). Where in-depth studies were conducted on apes, the 430 importance of sleeping posture has proven to improve sleep quality (Samson and Shumaker, 2013; 431 Samson and Shumaker 2015a). Similarly, clinging to branches and a strong grip is also shown to be 432 related to continuous sleep during the diurnal period as well as a decreased frequency of measurable 433 fragmented sleep (KAIN and KDR, unpublished data). The confirmation of nest use as a derived state in 434 strepsirhines that evolved multiple times corresponds with the deep evolutionary divergence seen among 435 lemuriforms and lorisiforms. We provide a novel set of data that we hope will inform further studies 436 reconstructing aspects of primate evolution.

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