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Behavioral Ecology

Gregariousness, foraging effort, and social interactions in lactating bonobos and chimpanzees

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Manuscript Type:	Original article
Keywords:	sociality, fission-fusion, feeding competition, predation risk, bonobos, chimpanzees



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LAY SUMMARY

2 In some group-living species, individuals divide into subgroups to minimize feeding 3 competition, which can reduce social opportunities. Bonobos and chimpanzees exhibit such 4 dynamics, but chimpanzees face higher feeding competition and can spend substantial time 5 feeding alone. Despite this difference in grouping, females of the two species spend similar 6 amounts of time engaged in social interactions. 7 8 **FULL TITLE** 9 Gregariousness, foraging effort, and social interactions in lactating bonobos and chimpanzees 10 11 **ABBREVIATED TITLE** 12 Gregariousness and activity budgets in lactating females in Pan 13 14 ABSTRACT 15 Fission-fusion dynamics have evolved in a broad range of animal taxa and are thought to allow 16 individuals to mitigate feeding competition. While this is the principal benefit of fission-fusion, 17 few studies have evaluated its costs. We compared gregariousness, foraging budgets, and social 18 budgets between lactating bonobos and chimpanzees from wild populations to evaluate such 19 costs. Both species exhibit fission-fusion dynamics, but chimpanzees, particularly in East 20 African populations, appear to experience higher feeding competition than bonobos. We 21 expected lactating chimpanzees to be less gregarious than lactating bonobos; reduced

22 gregariousness should allow lactating chimpanzees to mitigate costs of higher feeding

23 competition without requiring more foraging effort. However, we expected the reduced

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24	gregariousness of lactating chimpanzees to limit their time available for affiliative social
25	interactions. Using long-term data from LuiKotale bonobos and Gombe chimpanzees, we found
26	that lactating chimpanzees were indeed less gregarious than lactating bonobos although feeding
27	and travel time did not differ between species. Contrary to our predictions, lactating females did
28	not differ in social interaction time, and lactating chimpanzees spent proportionately more time
29	interacting with individuals other than their immature offspring. Our results indicate that
30	lactating chimpanzees can maintain social budgets comparable to lactating bonobos despite
31	reduced gregariousness and without incurring additional foraging costs. We discuss explanations
32	for why lactating bonobos are more gregarious.
33	Keywords: sociality, fission-fusion, feeding competition, predation risk, bonobos, chimpanzees
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INTRODUCTION

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A major goal in the study of behavioral ecology is to understand the evolution of group living under different ecological conditions (Wilson 1975; Maynard Smith and Szathmary 1997). Extensive research demonstrates that one of the primary benefits of group living is enhanced predator detection, dilution, and/or defense capabilities, while one of the major costs of group living is increased competition for food resources (reviewed in Ward and Webster 2016). Costs associated with feeding competition are particularly salient to females, given that food generally limits female reproductive success more so than males' (Trivers 1972).

56 Due to intragroup feeding competition, foraging effort is a function of group size: individuals 57 living in permanently cohesive social groups experience reduced feeding efficiency as the 58 number of intragroup competitors increases (Beauchamp 2012; Markham et al. 2015). Therefore, 59 females in many social species cope with the high energetic requirements of lactation principally 60 by allocating more time to foraging effort (Lasiurus cinerus: Barclay 1989; Peromyscus 61 maniculatus: Hammond and Kristan 2000; Odocoileus virginianus: Therrien et al. 2008; Myotis 62 lucifugus: Dzal and Brigham 2013; Enhydra lutris nereis: Thometz et al. 2016). However, any 63 increase in time dedicated to foraging effort must come at the expense of time allocated to other 64 activities (Dunbar et al. 2009), as is the case in numerous vertebrate taxa (e.g., Octodon degus: 65 Ebensperger and Hurtado 2005; Oreamnos americanus: Hamel and Côté 2008; Rhinopithecus 66 *bieti*: Xiang et al. 2010; *Morus capensis*: Rishworth et al. 2014). One activity that may be 67 sacrificed to provide more time for foraging is affiliative social interactions; however, such 68 interactions play an important role in maintaining social bonds in many group living animals. 69 Indeed, a growing body of research highlights the positive relationship between social bond

strength and fitness across taxa (e.g., *Papio cynocephalus*: Silk et al. 2003; *Equus ferus caballus*:
Cameron et al. 2009; *Tursiops aduncus*: Stanton and Mann 2012; *Crotophaga major*: Riehl and
Strong 2018). Thus, sacrificing time for social interactions may carry costs in some taxa.

Fission-fusion social systems present additional means through which lactating females may mitigate feeding competition. Fission-fusion societies are characterized by fluid subgrouping patterns (Aureli et al. 2008) and have been described for diverse taxa (Couzin 2006), such as guppy shoals (*Poecilia reticulata*) (Kelley et al. 2011), sand tiger sharks (*Carcharias taurus*) (Haulsee et al. 2016), common ravens (Corvus corax) (Loretto et al., 2017), African lions (Panthera leo) (Mbizah et al. 2019), and Cape buffalo (Syncerus caffer caffer) (Wielgus et al. 2020). High fission-fusion dynamics are hypothesized to allow individuals to adjust subgroup size and composition in response to fluctuations in food availability and their own energetic requirements. Thus, females can mitigate the energetic costs of lactation by altering their grouping patterns to maintain energy balance without substantial increases in foraging effort, while also grouping when possible to maximize predator defense and social opportunities. However, this raises the question of whether fission-fusion dynamics impact the extent to which lactating females suffer from increased predation and/or reduced social interactions. By ranging alone or in smaller subgroups, lactating females may not need to increase foraging effort, but they may be more vulnerable to predators and/or their social activity may be constrained as a result of spending less time in the presence of group members.

91 The genus *Pan* provides a comparative framework through which to focus on the social costs of
92 fission-fusion dynamics while discounting potential effects of predation risk. The two great ape

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species that comprise *Pan*, bonobos (*P. paniscus*) and chimpanzees (*P. troglodytes*), share a recent phylogenetic history (Prüfer et al. 2012) and several core morphological and behavioral traits. In particular, both species are characterized by a relatively large body size relative to most other primate species, as well as a largely arboreal lifestyle (Fleagle 2013); these traits are hypothesized to reduce vulnerability to their most likely predator, African leopards (Panthera pardus pardus) (Isbell 1994; Janson and Goldsmith 1995; Zuberbühler and Jenny 2002). Indeed, evidence for leopard predation on bonobos and chimpanzees is rare (for all inferred cases of leopard predation in Pan, see: Boesch 1991; Furuichi 2000; Zuberbühler and Jenny 2002; D'Amour et al. 2006; Pierce 2009; Nakazawa et al. 2013), despite extensive evidence of leopard predation on monkeys (reviewed in Isbell 1994) including at study sites where leopard predation on chimpanzees has been documented and deemed rare (e.g., Nakazawa 2020). Despite broad similarities in morphology and social organization between bonobos and chimpanzees, they appear to face different levels of feeding competition and starkly different patterns of female social behavior (reviewed in Gruber and Clay 2016). Stable isotope analyses of hair samples from multiple *Pan* research sites across tropical Africa indicate clear species differences in the stability of food resources (Oelze, Fahy, et al. 2016). Stable isotope ratios provide a reliable proxy for diet because the isotopic characteristics of food components are incorporated into consumers' tissue in a predictable manner (Kohn 1999). Bonobos exhibit less variation in stable isotope ratios over time when compared to chimpanzees, indicating that

114 Douglas, et al. 2016; Oelze, Fahy, et al., 2016). Additionally, individual variation in females'

bonobo diet composition is more stable, i.e., less seasonal, than that of chimpanzees (Oelze,

115 stable isotope ratios did not vary based on dominance rank in bonobos (Oelze, Douglas, et al.

2016), whereas several studies have demonstrated variation in diet quality based on dominance

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rank in chimpanzees (e.g., Murray et al. 2006). These patterns strongly suggest that feeding competition is more intense in chimpanzees than in bonobos. Reduced feeding competition among female bonobos may facilitate what appears to be a general pattern of high female gregariousness across several long-term study sites (Wamba: Furuichi 2009; LuiKotale: Moscovice et al. 2017; Lomako: Hohmann and Fruth 2002; Waller 2011). This high gregariousness may in turn facilitate the high degrees of intrasexual affiliative social behavior characteristic of female bonobos (Tokuyama and Furuichi 2016; Moscovice et al. 2017, 2019). In contrast, female gregariousness varies across chimpanzee populations. Females in some East African chimpanzee populations (P. t. schweinfurthii) tend to be highly solitary, frequently ranging alone with their immature offspring in order to mitigate costs associated with exceptionally high feeding competition and seasonality (Wrangham and Smuts 1980). In other East African chimpanzee populations, females can be more gregarious (Wakefield 2008). Some female West African chimpanzee populations (P. t. verus) appear to experience reduced seasonality when compared to East African chimpanzees (Doran et al. 2002) and are more gregarious (Lehmann and Boesch 2008, 2009). What remains unclear is whether different patterns of fission-fusion dynamics differentially constrain the extent to which females can engage in affiliative social interactions because direct comparisons of gregariousness and social budgets have not been conducted in the two *Pan* species. While a study evaluating within-population variation in dyadic association strength across *Pan* populations found clear species differences that are in line with putative species differences in gregariousness, they did not make direct comparisons between the populations (Surbeck et al. 2017).

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140	In this study, we compared gregariousness, foraging budgets, and social budgets of lactating
141	females in wild populations of bonobos and chimpanzees. The lactation period represents the
142	female life history stage when energetic demands are highest and thus when constraints on social
143	interactions should be most pronounced. Here we compared the LuiKotale bonobo population to
144	the Gombe East African chimpanzee population. LuiKotale is characterized by low seasonality,
145	extensive primary forest, and modest resource competition (Hohmann et al. 2012; Oelze,
146	Douglas, et al. 2016; Oelze, Fahy, et al. 2016; Nurmi et al. 2018), while Gombe appears to be
147	characterized by high seasonality and heterogeneity in habitat structure (Wrangham and Smuts
148	1980; Williams et al. 2002; Murray et al. 2006). We hypothesized that lactating chimpanzees at
149	Gombe are less gregarious than lactating bonobos at LuiKotale due to higher feeding
150	competition at Gombe. We further predict that reduced gregariousness constrains the social
151	interaction budgets of lactating chimpanzees; while being less gregarious may allow lactating
152	chimpanzees to maximize foraging efficiency, they may then be limited in the extent to which
153	they can engage in social interactions. Based on this hypothesis, we predicted that the amount of
154	time that lactating females spend engaged in feeding and travel does not differ between species,
155	but that lactating chimpanzees spend less time in groups and less time engaged in affiliative
156	social interactions. We first compared gregariousness by evaluating the proportion of time that
157	lactating females spend ranging alone with their immature offspring. We then compared the
158	amount of time that lactating females spend engaged in feeding, travel, and affiliative social
159	interactions, and how lactating females allocate their affiliative social interactions.
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161	MATERIALS AND METHODS

162 Study site and subjects

Data were collected on the Bompusa West bonobo community at LuiKotale, Democratic Republic of the Congo, and on the Kasekela chimpanzee community at Gombe, Tanzania. All bonobos and chimpanzees included in our study were habituated to human observation. Maternal relatedness is known for all individuals from observations and genotyping. During the study periods, the Bompusa West community included up to 54 individuals and the Kasekela community included up to 68 individuals. We focused our analyses on lactating females whose youngest infants were less than 4.5 years of age as this is the average age by which infants in wild populations of both species are nutritionally weaned based on stable isotope analyses (~4 years based on δ^{15} N and ~5 years based on δ^{13} C in both species: Ngogo chimpanzees: Bădescu et al. 2017; LuiKotale bonobos: Oelze et al. 2020). This age range also overlaps with the average weaned age derived from data on suckling behavior in our chimpanzee study population (4.7 years: (Lonsdorf et al. 2020). We used approximate periods of lactation because precise ages of weaning are likely to vary (e.g., Borries et al. 2014) and are not known for the majority of individuals in our sample. We pooled data on lactating females into three age classes based on the age of their youngest infant ($0 \le 1.5, 1.5 \le 3$, and $3 \le 4.5$ years), given that lactating female energetic requirements may vary based on stage of infant development (see Emery Thompson et al. 2012).

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47 181 **Predictions**

182 We tested three predictions: 1) Lactating chimpanzees spend more time alone with their
183 immature offspring than do lactating bonobos; 2) Lactating females of the two species do not

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84 differ in feeding or travel time; 3) Lactating bonobos spend more time engaged in social 85 interactions, particularly with individuals other than their immature offspring.

87 **Time spent alone**

88 At both study sites, the total number of individuals ranging in subgroups (hereafter "parties") and 89 their identities are recorded systematically. Party scan data on lactating females are recorded 90 during focal follows (see next section for a description of focal follows) at regular intervals and 91 represent all individuals observed in the party during that interval; researchers record party scans 92 every hour at LuiKotale and every 15 minutes at Gombe. To make party scan data comparable 93 between sites, we aggregated all party scans over a given hour at Gombe and used the aggregated 94 on-the-hour party scan in our analyses. We only included lactating females for which at least 20 95 hours of party scans were available for a given infant age class (Table 1). We took several 96 additional steps to ensure that data from both study sites are comparable. First, we used 97 contemporaneous data from both sites, starting in July 2011, when two coauthors (CMM and 98 EVL) hired several new field staff to collect data at Gombe and conducted extensive training to 99 ensure that data collection remained consistent despite a change in field staff. Long-term party 200 scan data were available from LuiKotale for the same period. We thus included party scan data 201 from July 2011 through November 2016 for both study sites. Second, because the Gombe party 02 scan dataset is larger than the LuiKotale dataset due to more field researchers collecting data at 203 Gombe, we used the sample function in base R version 4.0.2 (R Core Team 2020) to randomly 04 subsample on-the-hour party scans from Gombe without replacement to match the LuiKotale .05 party scan sample size, based on number of lactating females, infant sex, and infant age class. 206 For example, if we had 10 total on-the-hour party scans from two lactating bonobos, each with

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207 one female infant in the 0 < 1.5 infant age class, we subsampled the Gombe data such that we 208 had approximately 10 total on-the-hour party scans from two lactating chimpanzees, each with 209 one female infant in the 0 < 1.5 infant age class.

211 We measured gregariousness of lactating females as the proportion of party scans in which the 212 lactating female was alone with her immature offspring, which we defined as offspring younger 213 than 12 years of age. This definition for immature offspring is consistent with previous research 214 on both of our study populations (e.g., Murray et al. 2006; Surbeck et al. 2011; Stanton et al. 215 2014; Markham et al. 2015) and a recent study indicating that chimpanzee offspring continue to 216 associate with their mothers until 12 years of age (Stanton et al. 2020). We do not claim that 12 217 years of age and older qualifies as adult; rather, individuals below this age are predominantly 218 immature. Thus, when a focal subject is alone with her immature offspring, i.e., not in a party 219 with other community members, we considered her to be "alone" in her own focal follow. 220 Researchers attempt to remain with the focal subject at both study sites, regardless of party size.

5 221

222 Feeding, travel, and social interactions

In addition to party scan data, researchers collect detailed behavioral data during focal follows of a lactating female and her immature offspring. At Gombe, a given focal follow focuses on a lactating female and her two youngest offspring simultaneously and lasts from several hours to a full day; the goal is to collect at least six hours of focal follow data on each focal subject during each month. However, focal follow lengths vary based on various uncontrollable factors such as losing sight of the focal subject during poor weather conditions. At LuiKotale, a given focal follow focuses on a lactating female and one of her immature offspring at a time and are

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generally conducted for one hour. Focal follows can be longer if the focal subject is alone with her immature offspring because researchers generally attempt to follow lone focal females continuously until she rejoins a larger party. These differences in focal follow duration and the number of immature offspring on which data are collected are due to practical constraints associated with focal following two immature bonobos simultaneously for extended durations, given that there are generally many immature bonobos present in parties and it can be very difficult to monitor multiple at once. However, to ensure consistency in our comparative analyses, the behavioral ethogram in place at LuiKotale was developed in collaboration with Gombe researchers and designed to be comparable by utilizing the same definitions for all behaviors of interest and by employing the same point sampling interval; behavioral data on the lactating female and her immature offspring are recorded during one-minute point samples and include the identity of social partners.

Despite using the Gombe protocol as a model to design the protocol at LuiKotale, we took additional steps to ensure that data are comparable between the two study sites. Focal follow data on bonobos were collected between July 2015 and July 2018; however, chimpanzee focal follow data were only available through November 2016. We thus utilized focal follow data on chimpanzees between November 2013 and November 2016 to match the number of years during which bonobo data were collected. Second, we again subsampled the larger Gombe dataset to approximately match the LuiKotale sample size, again using the sample function in base R; however, instead of subsampling one-minute point samples, we subsampled 60 consecutive point samples from a given focal female. We did this so the Gombe subsample more closely resembled the sample of predominantly one-hour focal follows from LuiKotale. Lastly, we again restricted

1 2		
2 3 4	253	our analyses to lactating females for which at least 20 hours of party scans were available for a
5 6	254	given infant age class (Table 2). Good observations included those one-minute point samples in
7 8	255	which the activity of the lactating female could be determined, as opposed to bad observations in
9 10 11	256	which the activity could not be determined due to poor visibility.
12 13	257	
14 15	258	We used focal follow data to compare foraging and social budgets by analyzing the following
16 17	259	behaviors (following Lonsdorf et al. 2014):
18 19 20	260	1. Feeding – Ingestion of solid food.
20 21 22	261	2. Travel – Continuous movement from one point to another.
23 24	262	3. Social Interactions – Engaging in either of the following behaviors:
25 26	263	a. Social Groom – Parting of another individual's hair with hands, fingers,
27 28 29	264	and/or lips and removal of debris or ectoparasites and/or receiving this
30 31	265	behavior from another individual.
32 33	266	b. Social Play – Non-aggressive interaction between two or more individuals that
34 35 36	267	include one or more of the following: tickling, wrestling, chasing, kicking,
36 37 38	268	rubbing, thrusting, biting, or pulling. May incorporate an object (e.g., tugging
39 40	269	of sticks back and forth).
41 42	270	
43 44	270	Statistical analyses
45 46 47	271	We conducted all analyses in R version 4.0.2 (R Core Team 2020) and RStudio version 1.3.1
48 49	272	(RStudio Team 2020) using the glmmTMB version 1.0.2 (Brooks et al. 2018), DHARMa version
50 51	273	0.3.2 (Hartig 2020), car version 3.0-9 (Fox et al. 2012), and emmeans version 1.5.0 (Lenth 2018)
52 53		
54 55 56	275	packages. To test our three predictions (described above), we fitted generalized linear mixed
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models (GLMMs) to each response variable (response variables for each prediction described below) using the glmmTMB function in the glmmTMB package with a beta-binomial error structure. We initially fitted GLMMs using binomial error structures but found that all models were overdispersed. Overdispersion occurs when variance is higher than predicted by the model because the model lacks an adjustable dispersion parameter (e.g., as in binomial and poisson models) (Bolker et al. 2009; Zuur et al. 2009). Beta-binomial models include an adjustable dispersion parameter that allows the model to predict variance appropriately for binomial proportion data (Harrison 2015). We reported results of nonparametric dispersion tests for all models using the testDispersion function (case sensitive) in the DHARMa package. None of our beta-binomial models exhibited overdispersion. We evaluated model assumptions by visually assessing quantile-quantile plots and the distribution of residuals plotted against fitted values using the simulateResiduals (case sensitive) function in the DHARMa package.

For all models, we first tested the interaction between species and infant age class. To determine the significance ($\alpha = 0.05$) of interaction effects, we conducted Wald Chi-Squared tests using the Anova (case sensitive) function in the car package (Kenward-Roger degrees of freedom approximation, type III sum of squares). If the interaction between species and infant age class was not significant, we removed it and refitted the model using species and infant age class as independent fixed effect predictors and conducted Wald Chi-Squared tests, again using the Anova function in the car package (Kenward-Roger degrees of freedom approximation, type II sum of squares). If the interaction between species and infant age class was significant, we conducted Tukey's pairwise post-hoc comparisons between species within each infant age class using the emmeans function in the emmeans package. For all models, we included lactating

female identity as a random effect because the same lactating female could be represented in multiple infant age classes.

To test our first prediction that lactating chimpanzees spend more time alone than lactating bonobos, we ran one set of models called Time Alone (here and below, we refer to one set of models as the interaction effect model followed by the refitted independent effects model if necessary). We calculated our response variable by dividing the number of party scans that a given lactating female was in a party alone with her immature offspring during each infant age class by the total number of party scans collected on that lactating female during that infant age class. We expected a significant interaction effect between species and infant age class or a significant effect of species, with lactating chimpanzees spending more time alone than lactating bonobos.

To test our second prediction that lactating females do not differ in feeding or travel time, we ran two sets of models called Feeding and Travel. We calculated our response variables by dividing the number of point samples that a given lactating female was engaged in feeding or travel, respectively, during each infant age class by the total number of good observations collected on that lactating female during that infant age class. We did not expect to find a significant interaction effect between species and infant age class nor a significant effect of species.

> To test our third prediction that lactating bonobos spend more time engaged in social interactions, we ran two sets of models called Social Interactions and Adjusted Social

Interactions. We calculated our response variable for Social Interactions by dividing the number

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of point samples that a given lactating female was engaged in social interactions during each infant age class by the total number of good observations collected on that lactating female during that infant age class. We calculated our response variable for Adjusted Social Interactions by dividing the number of point samples that a given lactating female was engaged in social interactions with individuals other than their immature offspring during each infant age class by the total number of social interaction point samples collected on that lactating female during that infant age class.

RESULTS

In support of our first prediction, we found that lactating chimpanzees spent more time alone with their immature offspring than did lactating bonobos (Figure 1; Table 3). The interaction between species and infant age class was not significant in the model for Time Alone ($X^2 =$ 1.510, df = 2, p = 0.470) (Table 4), but when we tested independent effects of species and infant age class, species had a significant effect ($X^2 = 26.321$, df = 1, p < 0.001), while infant age class did not have a significant effect ($X^2 = 0.414$, df = 2, p = 0.813). The nonparametric dispersion tests were not significant for either Time Alone model (interaction effect model: deviance ratio = 0.957, p = 0.960; independent effects model: deviance ratio = 1.002, p = 0.928).

In support of our second prediction, lactating females of the two species did not differ in feeding time (Figure 2) or travel time (Figure 3) (Table 3). The interaction between species and infant age class was not significant in the model for Feeding ($X^2 = 4.359$, df = 2, p = 0.113) or Travel $(X^2 = 0.850, df = 2, p = 0.654)$ (Table 4). When we tested independent effects of species and infant age class, species was not significant in either model (Feeding: $X^2 = 0.032$, df = 1, p =

345	0.857; Travel: $X^2 = 1.334$, df = 1, $p = 0.248$). However, infant age class had a significant effect
346	in both models (Feeding: $X^2 = 8.379$, df = 2, $p = 0.015$; Travel: $X^2 = 7.153$, df = 2, $p = 0.028$);
347	lactating females with older infants fed more (Figure 2) and traveled more (Figure 3) (Table 3).
348	The nonparametric dispersion tests were not significant for either Feeding model (interaction
349	effect model: deviance ratio = 1.066 , $p = 0.496$; independent effects model: deviance ratio =
350	1.146, $p = 0.216$) or for either Travel model (interaction effect model: deviance ratio = 0.859, $p =$
351	0.200; independent effects model: deviance ratio = 0.895 , $p = 0.328$).
352	

Against our third prediction, lactating females of the two species did not differ in time engaged in social interactions with any community member (Figure 4), and lactating chimpanzees spent proportionately more of their social interaction time interacting with individuals other than their immature offspring (Figure 5) (Table 3). The interaction between species and infant age class was not significant in our model for Social Interactions ($X^2 = 0.870$, df = 2, p = 0.647) or for Adjusted Social Interactions ($X^2 = 3.702$, df = 2, p = 0.157) (Table 4). When we tested independent effects of species and infant age class in the model for Social Interactions, neither species ($X^2 = 0.266$, df = 1, p = 0.606) nor infant age class ($X^2 = 2.745$, df = 2, p = 0.253) had significant effects. When we tested independent effects of species and infant age class in the model for Adjusted Social Interactions, species had a significant effect ($X^2 = 12.998$, df = 1, p < 0.001), but infant age class did not ($X^2 = 0.082$, df = 2, p = 0.960). The nonparametric dispersion tests were not significant for either Social Interactions model (interaction effect model: deviance ratio = 1.066, p = 0.608; independent effects model: deviance ratio = 1.043, p = 0.704) or for either Adjusted Social Interactions model (interaction effect model: deviance ratio = 0.988, p =1.000; independent effects model: deviance ratio = 0.977, p = 0.984).

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5 6	369	DISCUSSION
7 8 9	370	Many studies across animal taxa indicate that fission-fusion dynamics allow individuals to
9 10 11	371	reduce feeding competition by adjusting the size and composition of the subgroups that they
12 13	372	range in (e.g., Tursiops aduncus: Heithaus and Dill 2002; Ocaella heinsohni and Sousa
14 15	373	chinensis: Parra et al. 2011; Macropus giganteus: Favreau et al. 2018; Giraffa camelopardalis:
16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38	374	Bond et al. 2019; Neophron percnopterus majorensis: van Overveld et al. 2020). Given that
	375	feeding competition generally increases with increasing group size, females in fission-fusion
	376	societies can offset the high energetic costs of lactation by reducing their levels of
	377	gregariousness, thereby reducing feeding competition. We therefore hypothesized that lactating
	378	chimpanzees at Gombe mitigate the intense feeding competition that they face by being less
	379	gregarious than lactating bonobos, who are thought to facing less intense feeding competition. In
	380	support of our first two predictions, lactating chimpanzees spent more time alone than lactating
	381	bonobos, while feeding and travel time did not differ between the species. These result support
	382	the hypothesis that lactating chimpanzees mitigate high feeding competition by being less
	383	gregarious, given that in doing so they maintained foraging budgets comparable to their more
39 40 41	384	gregarious bonobo counterparts. Our results thus add evidence to the existing body of research
41 42 43	385	indicating that fission-fusion dynamics are a counterstrategy to feeding competition across taxa
44 45	386	(see above).
46 47	387	
48 49	388	We also hypothesized that the social budgets of lactating chimpanzees are constrained as a result

We also hypothesized that the social budgets of lactating chimpanzees are constrained as a result of being less gregarious. However, we did not find support for this hypothesis: against our third prediction, lactating females did not differ in total social interaction time, and lactating

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391 chimpanzees spent proportionately more of their social interaction time interacting with 392 individuals other than their immature offspring. These results suggest that despite being less 393 gregarious, lactating chimpanzees spend as much time engaged in direct social interactions as do 394 lactating bonobos. It is therefore unclear how lactating bonobos benefit from higher 395 gregariousness if they do not engage in more social interactions when compared to lactating 396 chimpanzees. One possibility is that grouping provides lactating bonobos with opportunities to 397 enhance social relationships in ways that do not require direct interactions. In some animal taxa, 398 spatial association with conspecifics is related to fitness, and not necessarily direct social 399 interaction. For example, in the greater ani (Crotophaga major), females that consistently nested 400 together were considered to have stable social relationships, and this stability increased fitness 401 (Riehl and Strong 2018). Similarly, in feral horses (*Equus ferus caballus*), composite social 402 integration scores were positively related to fitness, and these scores were based on measures of 403 spatial affinity between mares (Cameron et al. 2009). Thus, there are numerous ways in which 404 gregarious individuals could gain social benefits without necessarily engaging in direct social 405 interactions. In this view, gregariousness could be favored at LuiKotale simply because being in 406 relatively close spatial proximity to other group members confers social benefits. 407

On the other hand, grouping may also provide lactating bonobos with opportunities for modes of direct social interaction that we did not consider in our study. While our results indicate that lactating chimpanzees invest more time in grooming and playing with the broader social milieu, it may be that bonobos add to their social budget through other direct interactions. For example, female bonobos engage in genito-genital rubbing, a behavior that is thought to contribute to bond formation and maintenance (Furuichi 1989; Hohmann and Fruth 2000; Fruth and Hohmann

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2006). Genito-genital rubbing requires little time investment as it more closely resembles a behavioral event rather than a state. Female bonobos may thus have additional social currency at their disposal that does not require substantial time investments, but it is difficult to make direct comparisons with chimpanzees because chimpanzees do not habitually engage in genito-genital rubbing (but see Anestis 2004; Zamma and Fujita 2004). Similarly, another mode of social interaction not captured by our study is female-female coalitionary behavior. Female-female coalitions against males are prevalent in bonobos (Surbeck and Hohmann 2013; Tokuyama and Furuichi 2016; Nurmi et al. 2018) but not in chimpanzees (but see Newton-Fisher 2006), again restricting direct comparisons between the species. It may be the case that benefits accrued through modes of social interaction that we did not consider in this study are beneficial enough to favor grouping by lactating bonobos.

More gregarious bonobos could also gain benefits associated with predator defense. Evidence for variation in grouping patterns resulting from differences in predation risk is widespread across non-primate taxa (e.g., Suricata suricatta: Clutton-Brock et al. 1999; Junco hyemalis: Lima et al. 1999; Cervus elaphus: Childress and Lung 2003; Perdix perdix: Watson et al. 2007); however, such predator-prey systems are often characterized by relatively high rates of predation. In generating our hypothesis, we assumed that grouping patterns are primarily driven by feeding competition based on the limited empirical evidence for leopard predation on bonobos and chimpanzees (see Introduction); indeed the underlying assumption of most fission-fusion systems is that predation is sufficiently low enough to allow groups to fission (but see food-safety tradeoff in *Tursiops aduncus*: Heithaus and Dill 2002). However, predation pressure is not absent from LuiKotale. Leopards have not been observed at Gombe since roughly 1975 (see

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437	Pierce 2009) and are presumed to be locally extinct; at LuiKotale, bonobo hard tissue was found
438	in leopard scat (D'Amour et al. 2006) and researchers recently observed a non-lethal
439	confrontation between bonobos and a leopard (unpublished data, Fruth and Hohmann),
440	suggesting that leopards are indeed a threat to bonobos. Thus, we cannot rule out that predation
441	risk at LuiKotale can have major impacts on bonobo sociality. This would be broadly in line with
442	the influential Predator Risk Allocation Hypothesis (Lima and Bednekoff 1999), which posits
443	that the trade-off between foraging and vigilance is less a function of immediate
444	presence/absence of predators and more of the temporal pattern of predation risk over time.
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446	Irrespective of the costs and benefits of grouping, our finding that lactating chimpanzees can
447	maintain social budgets comparable to lactating bonobos despite reduced gregariousness
448	underscores the benefits associated with the flexibility in behavior that fission-fusion dynamics
449	provide. Bechstein's bats (Myotis bechsteinii) also illustrate this sort of flexibility: Kerth et al.
450	(2011) showed that individuals are able to maintain long-term social relationships despite high
451	fission-fusion dynamics. Similarly, we showed in a separate study that immature females from
452	LuiKotale and Gombe do not differ in time engaged in social play or social grooming, indicating
453	that the reduced gregariousness at Gombe does not constrain immature female social budgets
454	either (Lee et al. 2020). Results from our studies and the study by Kerth et al. (2011) suggest that
455	individuals in fission-fusion societies need not spend extensive time together in order to maintain
456	relationships with the broader social milieu. In those fission-fusion species for which social
457	relationships are likely critical components of fitness, selection may have favored social skills
458	that enable bond partners to maintain relationships even with limited association time. Future
459	research could evaluate this further by identifying the mechanisms by which individuals develop

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2 3 4	460	and maintain such relationships in species that appear to exhibit even less frequent encounters
5 6	461	with some associates, such as African forest elephants (Loxodonta cyclotis) (Fishlock and Lee
7 8 9	462	2013) or sperm whales (Physeter macrocephalus) (Whitehead et al. 1991). More specifically,
9 10 11	463	such research could focus on the role of different latencies between fusions within and between
12 13	464	species to determine whether it is the absolute amount of time that individuals associate and/or
14 15 16	465	the temporal patterning of fusions that influences bond formation and maintenance.
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FIGURE LEGENDS

Figure 1

Mean \pm *SE* percentage of time that lactating females spent ranging in parties with only their immature offspring. Note: This figure and all following figures represent raw data; asterisks indicate where the independent fixed effect of species was statistically significant.

Figure 2

Mean \pm SE percentage of time that lactating females spent feeding.

Figure 3

Mean \pm SE percentage of time that lactating females spent traveling.

Figure 4

Mean \pm *SE* percentage of time that lactating females spent engaged in social interactions with any community member.

Figure 5

Mean \pm SE percentage of social interactions in which lactating females spent engaged in social interactions with individuals other than their immature offspring.

TABLES

Table 1

Party scan sample size

Infort and along	Bonobo	Bonobo	Chimpanzee	Chimpanzee
Infant age class	female	male	female	male
0 < 1.5	2 78	4 238	2 78	4 238
1.5 < 3	3 318	1 79	4 318	1 79
3 < 4.5	3 243	1 25	4 243	1 25

Number of lactating females | number of on-the-hour party scans

Table 2

Focal follow sample size

Infant age class

0 < 1.5

1.5 < 3

3 < 4.5

1

Bonobo

male

4 | 177

1 | 23

2 | 63

Bonobo

female

4 | 125

3 | 119

3 | 99

Number of lactating females | total focal follow observation time

Chimpanzee

female

4 | 120

2 | 140

3 | 97

Chimpanzee

male

6 | 172

1 | 23

2 | 62

Table 3

GLMM parameter estimates for independent effects models

Model	Term	Estimate	SE	Z	р
	Intercept	-3.804	0.509	-7.481	-
Time alone	Chimpanzee	2.470	0.481	5.130	< 0.0
	Infant age class 1.5 < 3	-0.267	0.419	-0.637	0.5
	Infant age class 3 < 4.5	-0.140	0.405	-0.345	0.7
Feeding	Intercept	-0.507	0.109	-4.662	-
	Chimpanzee	-0.023	0.126	-0.180	0.8
	Infant age class 1.5 < 3	0.354	0.151	2.344	0.0
	Infant age class 3 < 4.5	0.319	0.132	2.411	0.0
	Intercept	-1.606	0.069	-23.338	-
	Chimpanzee	-0.093	0.080	-1.155	0.2
Travel	Infant age class 1.5 < 3	0.251	0.094	2.657	0.0
	Infant age class 3 < 4.5	0.105	0.085	1.242	0.2
	Intercept	-1.755	0.115	-15.224	-
	Chimpanzee	0.067	0.130	0.516	0.6
Social Interactions	Infant age class 1.5 < 3	-0.157	0.163	-0.960	0.3
	Infant age class 3 < 4.5	-0.249	0.162	-1.534	0.1
Adjusted Social Interactions	Intercept	-3.101	0.210	-14.802	-
	Chimpanzee	0.782	0.217	3.605	< 0.0
	Infant age class 1.5 < 3	-0.082	0.298	-0.276	0.7
	Infant age class $3 < 4.5$	-0.031	0.229	-0.135	0.8

Model	Term	Estimate	SE	Z
	Intercept	-3.900	0.719	-5.425
Time Alone	Chimpanzee * Age 1.5 < 3	0.680	1.302	0.523
	Chimpanzee * Age 3 < 4.5	-0.805	1.03	-0.779
Feeding	Intercept	-0.484	0.116	-4.013
	Chimpanzee * Age 1.5 < 3	0.531	0.284	1.871
	Chimpanzee * Age 3 < 4.5	-0.054	0.249	-0.215
	Intercept	-1.593	0.078	-20.415
Travel	Chimpanzee * Age 1.5 < 3	0.204	0.259	0.788
	Chimpanzee * Age 3 < 4.5	-0.069	0.164	-0.422
Social Interactions Adjusted Social Interactions	Intercept	-1.749	0.126	-13.840
	Chimpanzee * Age 1.5 < 3	-0.126	0.324	-0.390
	Chimpanzee * Age 3 < 4.5	0.206	0.292	0.705
	Intercept	-2.910	0.211	-13.799
	Chimpanzee * Age 1.5 < 3	0.506	0.495	1.023
	Chimpanzee * Age 3 < 4.5	0.878	0.470	1.869

FIGURES





















