

Birth timing generates reproductive trade-offs in a non-seasonal breeding primate

Jules Dezeure, Alice Baniel, Alecia J Carter, Guy Cowlishaw, Bernard

Godelle, Elise Huchard

► To cite this version:

Jules Dezeure, Alice Baniel, Alecia J Carter, Guy Cowlishaw, Bernard Godelle, et al.. Birth timing generates reproductive trade-offs in a non-seasonal breeding primate. Proceedings of the Royal Society B: Biological Sciences, 2021, 288 (1950), pp.20210286. 10.1098/rspb.2021.0286. hal-03254769

HAL Id: hal-03254769 https://hal.umontpellier.fr/hal-03254769

Submitted on 9 Jun2021

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

1 Birth timing generates reproductive trade-offs in a non-seasonal

2 breeding primate

3 Jules Dezeure^{1, 5,*}, Alice Baniel², Alecia J. Carter³, Guy Cowlishaw⁴, Bernard Godelle¹, Elise

4 Huchard¹

- ⁵ ¹Anthropologie Évolutive, Institut des Sciences de l'Évolution de Montpellier (ISEM),
- 6 Université de Montpellier, CNRS, IRD, EPHE, Montpellier, France.
- ⁷ ²Department of Anthropology, Stony Brook University, Stony Brook, NY, USA
- 8 ³Department of Anthropology, University College London, London, UK
- ⁹ ⁴Institute of Zoology, Zoological Society of London, Regent's Park, London NW1 4RY
- 10 ⁵Lead contact
- 11 *Corresponding author: Jules Dezeure
- 12 dezeurejules@gmail.com
- 13

14 ABSTRACT

15 The evolutionary benefits of reproductive seasonality are often measured by a single fitness 16 component, namely offspring survival. Yet different fitness components may be maximised by 17 dissimilar birth timings. This may generate fitness trade-offs that could be critical to 18 understanding variation in reproductive timing across individuals, populations and species. 19 Here, we use long-term demographic and behavioural data from wild chacma baboons (Papio 20 ursinus) living in a seasonal environment to test the adaptive significance of seasonal variation 21 in birth frequencies. We identify two distinct optimal birth timings in the annual cycle, located 22 4-months apart, which maximize offspring survival or minimize maternal interbirth intervals 23 (IBIs), by respectively matching the annual food peak with late or early weaning. Observed births are the most frequent between these optima, supporting an adaptive trade-off between 24 25 current and future reproduction. Furthermore, infants born closer to the optimal timing favouring maternal IBIs (instead of offspring survival) throw more tantrums, a typical manifestation of mother-offspring conflict. Maternal trade-offs over birth timing, which extend into mother-offspring conflict after birth, may commonly occur in long-lived species where development from birth to independence spans multiple seasons. Our findings therefore open new avenues to understanding the evolution of breeding phenology in long-lived animals, including humans.

32

33 MAIN TEXT

34 Introduction

35 Empirical studies investigating variation in reproductive timing have mostly focused on fast-36 lived seasonal breeders, whose development from birth to independence generally occurs within the most productive season [1]. In long-lived mammals, the reproductive cycle from 37 38 birth to weaning cannot similarly be squeezed into one annual food peak, and consequently, 39 females must choose which stage(s) of the reproductive cycle to synchronize with one or more 40 food peak(s). For example, female mammals could match the annual food peak to coincide with 41 either late-weaning or mid-lactation (two critical stages that require readily accessible food), 42 but usually not both. The reproductive timing strategy is likely to depend on how females trade-43 off the survival of their offspring (mortality risks tend to peak at the end of weaning) [2–4] with 44 their own reproductive costs (energetic demands tend to peak around mid-lactation) [5,6]. 45 Whether such reproductive timing strategies can vary within populations is largely unknown. 46 In addition, while evolutionary trade-offs between offspring quality and quantity have been 47 described both within and across species through associations between birth spacing and infant 48 growth and survival [7,8], the existence of maternal trade-offs over birth timing have only been 49 suggested theoretically [3] and never tested empirically in mammals (but for birds, see Fulica 50 *atra*: [9]).

51 Here, we investigate variation in maternal reproductive success and mother-offspring 52 relationships associated with variable birth timings in the annual cycle of wild chacma baboons 53 (Papio ursinus) living in a seasonal semi-arid savannah (Tsaobis, Namibia). Baboons are 54 African primates distributed across a wide latitudinal range and a classic model for 55 understanding how early humans adapted to seasonal savannahs [10,11]. In particular, baboons 56 typically breed year-round [12] and are therefore considered non-seasonal breeders, though the 57 distribution of births shows moderate seasonality (i.e. varies along the annual cycle) in some 58 species and populations [13–15]. In addition, infant baboons, like many young primates 59 including human toddlers, commonly perform tantrums, a manifestation of mother-offspring 60 conflict [16,17]. Using a combination of detailed long-term life-history and behavioural data 61 collected over 15 years (2005-2019), we first characterize the reproductive and environmental 62 seasonality of the Tsaobis baboons. Second, we quantify the consequences of birth timing on 63 two components of female fitness: offspring survival and maternal inter-birth intervals (IBIs) 64 and seek to identify two distinct birth timing optima. We further test whether individual females may vary in their birth timing strategies, and specifically individual traits predict whether a 65 66 female is more likely to give birth around one or the other optimum. In particular, dominance rank and parity can affect various aspects of individual reproductive performance, including 67 68 offspring survival and IBI [18-20], and may influence birth timing strategies accordingly. 69 Similarly, mothers conceiving close to the optimal timing that alleviates the energetic costs of 70 lactation may subsequently favour male over female embryos, which are more costly to produce 71 in sexually dimorphic mammals [18,21]. Third, we investigate if maternal care can mitigate the 72 costs of suboptimal birth timing for offspring, and whether infants born and weaned during 73 suboptimal periods had higher tantrum frequencies.

74

75 Materials and methods

76 Study population

77 Three habituated groups (named J, L and M) of wild chacma baboons were followed between 2005 and 2019: J and L since 2005, and M, a fission group from J, since 2016. They live in a 78 79 desert-edge population at Tsaobis Nature Park (22°23S, 15°44'50E) in Namibia, in a seasonal 80 and arid environment [22]. Water is always available through the presence of both natural seeps 81 and artificial water points for wildlife and livestock. A field team was present each year, mainly 82 during winter (between May to October), for a variable number of months (mean = 4.5, range: 83 1.9-7.9), that collected daily demographic and behavioural data, as well as GPS locations, while 84 following the groups on foot. All individuals, including infants, are individually recognizable 85 thanks to small ear markings performed during capture and/or other distinctive features.

86

Ethical Note. Our research procedures were evaluated and approved by the Ethics Committee
of the Zoological Society of London and the Ministry of Environment and Tourism, Namibia
(MET Research/Collecting Permits 886/2005, 1039/2006, 1186/2007, 1302/2008, 1379/2009,
1486/2010, 1486/2011, 1696/2012, 1786/2013, 1892/2014, 2009/2015, 2147/2016, 2303/2017,
RPIV00392018/2019), and adhered to the ASAB/ABS Guidelines for the Treatment of Animals
in Behavioural Research and Teaching.

93

94 Environmental data

In order to describe the relationship between reproductive and environmental seasonality, we
characterize two aspects of environmental seasonality at Tsaobis: rainfall and vegetation cover
(an index of food availability).

98 Daily rainfall in a 0.25×0.25 degree grid cell resolution (corresponding to 28×28 km 99 at this latitude) was extracted using satellite data sensors from the Giovanni NASA website 100 (product TRMM 3B42) [23]. We computed a rectangular geographic area that encompasses the

global ranging area of the Tsaobis baboons, using GPS locations collected by observers every
30 min when following the study groups. We used the minimal and maximal latitude and
longitude recorded between 2005 and 2019. Monthly cumulative rainfall (summed across daily
values) were computed between 2005 and 2019.

105 We used the Normalized Difference Vegetation Index (NDVI) as an index of food 106 availability. NDVI is obtained from the red:near-infrared reflectance ratio, with NDVI=(NIR-107 RED)/(NIR+RED), where NIR and RED are the amounts of respectively near-infrared and red 108 light, reflected by the vegetation and captured by satellites [24]. NDVI thus produces a 109 quantitative index of primary productivity with higher values corresponding to a higher degree 110 of vegetation cover [25]. It has previously been used as an indicator of habitat quality for the 111 Tsaobis baboons [26] and other baboon populations [27]. We further confirmed that temporal 112 variation in NDVI reflected temporal variation in rainfall: mean cumulative rainfall over the 113 past three months explained between 60-72% of the NDVI variation (Appendix 1). To index 114 food availability using NDVI, we first computed 100% isopleth home ranges for each group 115 using kernel density estimates with the adehabitatHR package ('kernelUD' function) [28], 116 based on the daily 30-min GPS locations from 2005-2019 (from 2016-2019 for M group). We 117 obtained one home range per group for the entire study period. We then extracted the mean 118 NDVI per 16 day-period on a 500 m \times 500 m resolution (these 16-days windows are imposed 119 by the resolution of the NASA datasets) across these periods using data provided by NASA 120 (MODIS13A1 product) [25] within these home ranges. Daily NDVI was computed by linear 121 interpolation and then averaged to obtain a monthly value. In contrast to rainfall, NDVI was 122 measured with greater resolution and for each group separately because baboons finely adjust 123 their ranging behaviour in relation to food availability [29].

124

125 Individual data

126 A female was considered adult when she reached menarche. The reproductive state of each 127 adult female was monitored daily. A female could be: (i) pregnant (assessed by the paracallosal 128 skin turning red and absence of cycles over the following months), with the exact start date of 129 pregnancy being determined post hoc following infant birth, and encompassing 190 days (mean 130 gestation length in this population, n = 13 pregnancies where both conception and birth were 131 observed, range: 181-200 days, SD = 5) between conception and birth; (ii) lactating, as long as 132 the female did not resume cycling after an infant birth; (iii) cycling, including both swollen 133 females in oestrus (i.e., sexually receptive with a perineal swelling) and non-swollen females 134 at other stages of their cycle. Conceptive cycles were established based on the beginning of a 135 pregnancy and were usually confirmed by a birth. The first post-partum cycle (i.e. cycle 136 resumption) is the first cycle following an infant's birth, when the female resumes cycling after 137 lactation. The exact date of the cycle resumption corresponds to the first day of oestrus of the 138 first post-partum cycle, i.e. the first day when a sexual swelling is recorded. The dates of these 139 reproductive events (conceptions, births and cycling resumptions) were either known with 140 accuracy when recorded by field observers, or estimated in the absence of observers using the 141 methods detailed in Appendix 2 and Table S1.

Female parity was known from life history records and defined as primiparous (between the birth of her first and second infant) or multiparous (after the birth of her second infant). Parity of adult females at the start of the study was assessed using both the presence of older offspring based on a combination of behavioural and genetic data [30], alongside female age estimated using teeth eruption patterns [31].

Female social rank was established annually for each group using *ad libitum* and focal
observations of agonistic interactions between adult females: supplants, displacements, attacks,
chases and threats (Huchard and Cowlishaw 2011). We computed a linear hierarchy using
Matman 1.1.4 (Noldus Information Technology, 2013) and then converted to a relative rank to

151 control for group size (i.e. the number of adult females in the group). Each female was thus152 assigned one rank per year, ranging from 0 (lowest ranking) to 1 (highest ranking).

153

154 Fitness data

155 We tested the influence of birth timing in the annual cycle on two fitness measures, namely 156 offspring mortality before weaning and the duration of the maternal interbirth interval. For each infant born between 1st January 2005 and 1st August 2018, we investigated whether it died 157 158 (yes/no) before weaning. The weaning age was identified as 550 days on the basis of the 159 maximum length of post-partum anoestrus (n = 33 cases for which both birth and cycle 160 resumption were known with accuracy, see also Appendix 3) and presumably reflected the 161 upper threshold of weaning age in our population [33,34]. Death was recorded when a corpse 162 was observed or when the infant had been missing in the group for five consecutive days. 163 Infants born later than August 2018 were not considered as their survival outcome was 164 unknown. Four infants that disappeared between consecutive field seasons were omitted 165 because we could not establish whether the age of death was before or after 550 days.

We defined interbirth intervals (IBI) as the number of days between two consecutive live births of the same female. We only considered IBIs for which the first infant reached weaning [18], i.e. survived until 550 days old. We discarded other IBIs as females resumed cycling rapidly after their infant's death when unweaned (median=21 days, range=9-51, n=9 observed death), and their IBIs would have been shortened regardless of environmental seasonality.

172

173 Behavioural observations

174 In order to characterize variation in maternal care and in mother-offspring conflict, we used 175 three behavioural indicators: suckling, infant carrying and tantrum frequencies. We also used

these behavioural data, along with life history data, to assign different developmental stages, including the different stages of weaning and the peak of lactation after an infant's birth (see Appendix 3). Field observers collected a total of 1185 hours of focal observation [35] of 20 (in 2017, 2018 and 2019) or 60 (in 2006) minutes long on 69 infants (mean \pm SD = 17.1 \pm 7.8 hours of observations per infants, range = 6.3–34.6) (see Appendix 4 for more details).

181

182 <u>Maternal care during weaning</u>

183 Maternal care was quantified through two measures: suckling frequency and infant carrying 184 frequency, which represent the two main energetic costs of maternal care before weaning 185 [5,36]. First, for each scan observation (taken every 5 min), we considered whether the infant 186 was suckling (1) or not (0) to investigate the effect of birth timing on variation in suckling 187 frequency. Here, suckling was recorded when the focal individual had its mouth on its mother 188 nipple, and we therefore could not distinguish comfort (when a juvenile suckles for reassurance, 189 without any milk transfer [37]) from nutritive suckling. We considered only infants aged 2- to 190 18-months-old for this analysis (Figure S1), using 11687 scans from 55 infants. The birth date 191 uncertainty for these 55 infants ranged from 0 to 130 days (with a median birth date uncertainty 192 of 16 days) and was taken into account in subsequent models (see Appendix 5).

193 Second, for each scan observation during which an infant was travelling, we determined 194 whether the infant was carried by its mother (1) or travelled on its own (0). This variable 195 allowed us to monitor the gradual decrease from full maternal dependence to full independence 196 during travelling. We considered infants aged from 2 to 12 months old for this analysis (Figure 197 S1), using 924 scans from 35 infants.

198

199 Mother-infant conflicts during weaning

200 We considered infant tantrums as a behavioural measure of mother-offspring conflict, reflecting 201 when an infant's request to access resources from its mother was not initially satisfied [16]. We 202 considered only infants aged 2 to 18 months old for this analysis (Figure S1), using 2221 focal 203 observations from 55 infants. During each focal observation, we determined if a tantrum 204 occurred (1) or not (0), based on a range of distinctive offspring vocalizations (gecks, moans 205 and loud screams) and behaviours (frenzied behaviour when infants hurl themselves to the 206 ground, sometimes accompanied by rapidly rotating their tail) that were recorded on a 207 continuous basis and are characteristic of baboon tantrums [38,39]. A tantrum was considered 208 to occur when at least two of these behaviours or vocalizations were recorded, separated by at 209 least 30 seconds (isolated complaints, and complaints that lasted fewer than 30 seconds, were 210 thus not considered as tantrums here). Tantrums were usually caused by maternal refusal of 211 access to the nipple or to carrying and more rarely by maternal absence.

212

213 Statistical analysis

214 <u>Characterization of the environmental and reproductive seasonality of the Tsaobis baboons</u>

215 First, to assess the strength and direction of reproductive seasonality, we used a Rayleigh test, 216 from circular statistics, which characterizes the deviation of circular data from a uniform 217 distribution, via the direction (μ) and length (R) of the mean vector summing all observed 218 events across the annual cycle (R=0 when the event is evenly distributed, and R=1 when all 219 events are synchronized, i.e. occurs the same day) [40]. Our sample comprised 241 conceptions, 220 215 births and 171 cycle resumptions which occurred between 2005 and 2019. Uncertainties in 221 those dates were taken into account in all subsequent analyses using 1000 randomized reproductive events for each variable (Appendix 5). 222

223

224 Birth timing effects on two fitness traits, and individual effects on birth timing

To quantify the effect of birth timing on the probability of offspring mortality before weaning (Model 1), we ran a generalized linear mixed model (GLMM) with a binomial error structure. We then ran a linear mixed model (LMM, Model 2), testing the effect of birth timing on IBIs.

228 In both models, we used a sine term to describe the timing of an infant's birth in the 229 annual cycle (see Appendix 6 for more details on this procedure) [26]. We included as random 230 effects year of infant birth and identity of the mother to control for repeated observations. We 231 also included maternal parity, rank (in the birth year of the focal infant) and infant sex as fixed 232 effects, because maternal parity and rank often affect reproductive traits in primates, including 233 baboons [18,41], while infant sex can affect both the mother's subsequent interbirth interval 234 [42] and the probability of infant survival in sexually dimorphic primates [43]. We also control 235 for group identity as a fixed effect in both models, as data were collected from only three groups 236 in this study [44].

237 We investigated the individual determinants of female reproductive decisions over birth 238 timing, based on 215 births from 62 females. Our two response variables were the deviations, in days, from the birth timing that minimises offspring survival (December 15th) in Model 3 239 240 and maternal IBIs (September 1st) in Model 4 (Table S2). For both Models 3 and 4, we tested 241 the influence of infant sex, female parity and rank (as fixed effects) on the proximity of birth to 242 the optimal timing for offspring survival (Model 3) or for maternal IBI (Model 4). We also 243 controlled for the identity of the mother and birth year as random effects, in order to take into 244 account the between-year environmental variation likely to affect birth timings. We included 245 group identity as fixed effects (as there was only three levels for this factor [44]). We tested the 246 significance of maternal identity using a likelihood-ratio test (LRT), comparing the model with 247 and without this random effect.

248

249 Birth timing effects on maternal care and tantrum probability

We ran three GLMMs with a binomial error structure to test the effect of birth timing on the probability of suckling (Model 5), infant carrying (Model 6), and tantrums (Model 7). Models 5 and 6 are scan-based data: during a scan observation, the infant is suckling (yes/no, Model 5), and during a travelling scan observation, the infant is carried by its mother (yes/no, Model 6). Model 7 is based on the entire focal observation as tantrum events are relatively rare: during a focal observation, the infant throws a tantrum (yes/no).

256 In order to investigate the potential effect of birth timing on maternal care and tantrum 257 probability, we used a sine wave term for infant birth date as a fixed effect (Appendix 6). We 258 included, as random effects, the identity of the infant (Models 5-7) to control for repeated 259 observations. We also added the focal observation as a random effect for Models 5-6. We 260 controlled for group identity and year of observation as fixed effects in all models, as there were 261 less than five levels for both factors [44]. In all models, we included maternal parity, rank (in 262 the year of birth of the focal infant) and infant sex as fixed effects. Such parameters are likely 263 to affect reproductive performances as well as the probabilities of maternal care and mother-264 offspring conflict [39,43]. For Model 7, we also controlled for the duration of focal observation 265 as a fixed effect.

266 For Models 5-7, we further controlled for the effects of infant age, which modulates the 267 amount of maternal care and probability of tantrums throughout early development [16,43]. We 268 considered four different possibilities for the form of the relationship between infant age and 269 the response variable, using a regression thin plate spline (general additive model), a simple 270 linear effect, and a polynomial regression (of 2 or 3 degrees), respectively [45]. To determine 271 the best fit, we ran these different preliminary models with no other fixed effect but including 272 all random effects (and the duration of focal observation for Model 7), and selected the model 273 minimizing the AIC. The age effect was linear for suckling and infant carrying probabilities 274 (Model 5 and 6), and a second-degree polynomial for tantrum probability (Model 7).

275 Lastly, mothers might be expected to invest more, and similarly infants might be 276 expected to have more requests for maternal care, during the lean season, irrespective of the 277 developmental trajectory of the infant, i.e. regardless of its age and birth timing (whether it was 278 born in the optimal period or not). Therefore, we also investigated the potential effect of 279 seasonality by assessing the influence of the observation date on suckling, infant carrying and 280 tantrum probabilities (see Appendix 7 for more details). We did not include in the same model 281 observation date and birth date, as they give redundant information (observation date is, by 282 definition, the sum of birth date and infant's age, and infant's age is already included as a fixed 283 effect). We present our models of birth date effects in the main text (Models 5-7, see also Table 284 S3), and our models of observation date effects in the Supporting Information (Models 5bis-285 7bis, Table S4).

The structure of each model, with the different fixed and random effects included,alongside sample size, is summarised in Table S5.

288

289 Statistical methods

290 All statistical analyses were conducted in R version 3.5.0 [46]. For the Rayleigh test, we used 291 the function 'r.test' from the R package 'CircStats' [47]. To run mixed models, we used 'lmer' 292 (for LMMs) or 'glmer' (for binomial GLMMs) function on the lme4 package [48]. To run 293 general additive mixed models (GAMMs) when investigating the best age effects on suckling, 294 infant carrying and tantrum probabilities, we used the 'gam' function of the 'mgcv' package 295 [45]. All quantitative fixed effects were z-transformed to facilitate model convergence. When 296 we obtained singular fits, we confirmed the results by running the same models with a Bayesian 297 approach, using the 'bglmer' and 'blmer' functions of the 'blme' package [49]. To diagnose the 298 presence of multicollinearity, we calculated the variance inflation factor for each predictor in 299 each full model using the 'vif 'function of the R 'car' package [50]. These were lower than 2.5 in all cases. To assess the strength of the fixed effects in each model, we used the Wald chisquare tests with associated P-values computed with the 'Anova' function of the R package (car' [50], and calculated the 95% Wald level confidence intervals. We further checked the distribution of residuals with 'qqPlot' function of the car package for LMMs [50], and with 'simulateResiduals' from DHARMa package for binomial GLMMs [51].

305

306 **Results**

Characterization of the environmental and reproductive seasonality of the Tsaobis baboons

Environmental seasonality was pronounced at Tsaobis (Figure 1A). Mean annual rainfall was low and variable (mean \pm SD = 192 \pm 143mm), falling mostly between January and April (Figure 1A). Seasonal variation in NDVI, a satellite-based proxy of primary productivity, followed a similar, but slightly lagged pattern, to rainfall (Figure 1A). The highest birth frequency occurred in October-November (i.e. 28.4% of annual births), preceding the peak in rainfall (February) and NDVI (March-April, Figure 1A).

315 Conceptions, births and cycle resumptions occurred throughout the year (Figure S2), 316 indicating an absence of a strict breeding season. We used circular statistics to test whether 317 moderate seasonality may still occur, computing respectively the mean annual angle (μ) and 318 Rayleigh tests (R and p-values) for the annual distribution of 241 conceptions, 215 births and 319 171 cycle resumptions recorded between 2005-2019. The frequency of conceptions and births 320 showed slight seasonal variations, which reached statistical significance for conceptions only 321 (conceptions: $\mu = May 12$, R=0.13, p=0.02; births: $\mu = November 18$, R=0.09, p=0.17; cycle 322 resumptions: μ = December 4, R=0.08, p=0.36, Figure S2).

323

324 2. Birth timing effects on two fitness traits, and individual effects on birth timing

We considered two indicators of maternal fitness. First, we assessed whether or not infants survived until weaning (550 days). In our sample, 39 infants out of 195 (i.e. 20%) died before weaning, at a median age of 74 days (range 1-284 days, n=17 known dates of death). Second, we gathered 120 IBIs from 43 adult females, ranging from 397 to 1132 days with a mean of 678 days (SD=128).

330 Birth timing influenced these two indicators of maternal fitness. First, birth timing affected offspring survival (Table S6): infants born between November 15th and January 1st 331 were the most likely to survive until weaning (Table S2), indicating an optimal birth timing for 332 offspring survival in the annual cycle (Figure 2A). Infants born in July 15th were 66% more 333 likely to die before being weaned than the infants born in December 15th (Table S6). Second, 334 335 the duration of maternal IBI was influenced by the timing of the birth opening the IBI (Table 336 S6): females giving birth between August 1st and September 15th had the shortest IBIs (Table 337 S2), indicating another different optimal birth timing for maternal reproductive pace in the 338 annual cycle (Figure 2B). Females giving birth in September 1st had IBIs 73 days shorter than 339 females giving birth in March 1st.

We then wondered whether some females might be more likely to time their births to maximise current over future reproduction, or vice versa. However, we failed to detect any significant variance associated with maternal identity on the deviation between observed birth and the optimal birth timing maximizing offspring survival (LRT=0.66, p=0.42) versus maternal IBI (LRT=0.00, p=0.98). This suggests that females did not consistently give birth in one timing over the other across successive births. Moreover, female parity, rank and infant sex did not influence the proximity of birth timing in relation to each optimum (Table 1).

347

348 3. Birth timing effects on maternal care and tantrum probability

In order to test whether maternal care may compensate for the costs of suboptimal birth timings in offspring, we investigated the effects of birth timing on the frequency of suckling and infant carrying. We did not find any effect of infant birth date on patterns of maternal care (Table S3).
Further analyses revealed that mothers increase maternal care in the dryer winter months, but such compensation occurs regardless of an infant's birth date (Appendix 7, Table S4).

We also investigated whether infants born in suboptimal timings may beg for maternal care more frequently, looking at tantrum frequencies. We found that infants born near the maternal IBI optimal timing, i.e. between August 1st and October 1st (Table S2), were more likely to exhibit tantrums than other infants (Table S3, Figure 3). Observation date did not affect tantrum frequencies, meaning that such an effect was independent of the season of observation (Table S4).

360

361 **Discussion**

362 We identify two distinct optimal birth timings in the annual cycle, respectively favouring 363 current reproduction (offspring survival) and future reproduction (maternal reproductive pace). 364 These are separated by four months, and the highest birth frequency occurs between these 365 optima, indicating that mothers balance current and future reproduction, though closer to the 366 optimal birth timing favouring offspring survival. Several reasons might explain why offspring 367 survival might be prioritized over maternal reproductive pace. First, lifespan and offspring 368 survival are the primary components of female lifetime reproductive success in long-lived 369 species such as baboons, while reproductive pace may be less important [52]. Second, shorter 370 IBIs might compromise infant survival independently of the effect of birth timing, and are thus not necessarily adaptive [7]. Finally, the effect size of birth timing is greater on offspring 371 372 survival than on maternal IBIs in our population.

373 More broadly, these results further our understanding of the evolution of vertebrate 374 reproductive timing in several ways. First, trade-offs over birth timing may be widespread in long-lived species with slow life histories, for which development from birth to independence 375 376 spans several months, therefore exceeding the length of the most productive season. In such 377 cases, different stage(s) of the reproductive cycle may be synchronized with one or more 378 seasonal food peaks, with the specific pattern dependent on the trade-offs females make among 379 different fitness components [53]. Such variation could account for empirical cases where the 380 observed birth peak fails to coincide with the birth timing expected on the basis of a single 381 fitness measure. For example, in humans from pre-industrial Finland, births did not concentrate 382 in the months with the highest infant survival expectations [54]. More generally, such trade-383 offs may contribute to explain the partial or total lack of breeding seasonality observed in some 384 large mammals [55], such as social primates including apes [15] and humans [56,57].

385 Second, while different species synchronize different stages of their reproductive cycle 386 with the seasonal food peak [1,2,58,59], this study reveals variations in breeding timing within 387 the same population. However, while mothers experience a trade-off between reproductive pace 388 and offspring survival in their birth timing, it is not clear if particular individuals consistently 389 favour certain strategies, as we did not detect any inter-individual effects of female identity, 390 parity or rank on parturition timing. Instead, intra-individual factors, such as maternal 391 reproductive history, may constrain the evolution of such individually-based specializations, if 392 only because the duration of IBIs - 22 months on average but with extensive variation - prevents 393 females from giving birth every two years at the same period of the year. In addition, birth 394 timings may be affected by many external factors beyond female control, such as male 395 reproductive strategies. In addition, the costs of waiting for the next optimal timing may often 396 outweigh the costs of giving birth at suboptimal timings.

397 Third, this study underlines the importance of weaning to understand the evolution of 398 mammalian reproductive schedules. Late-weaning is most critical for infants, as they must learn 399 to ensure their own provisioning. Matching that stage, which occurs between 12 and 18 months 400 of age in this population, with the most productive season may substantially enhance infant 401 survival (Figure 1B) [43,60,61]. Moreover, the peak of lactation, which is the most 402 energetically-costly reproductive stage for mothers [3,43], typically coincides with the onset of 403 weaning, occurring around 6 months after birth in this population. Matching lactation peak with 404 abundant resources can alleviate the costs of reproduction and help to accelerate the transition 405 to feeding independence by granting infants access to a wealth of weaning foods (Figure 1B) 406 [43]. It may contribute to explain the shorter interbirth intervals associated with this birth 407 timing. Such patterns may be very general. In the lemur radiation, for instance, despite a variety 408 of life-histories, ecologies and societies, and the fact that different species mate and give birth 409 at different times of year, all species synchronize weaning with the food peak [60]. Our 410 understanding of the ultimate causes of mammalian reproductive seasonality may gain from 411 granting more consideration to the dynamics and consequences of weaning, which may have 412 been underappreciated in comparison to the energetic costs of pregnancy and lactation [1,2,4].

413 Fourth, our results show that the trade-off over birth timing faced by mothers may 414 subsequently translate into mother-offspring conflict after birth. Although mothers adjust 415 maternal care seasonally, they do so regardless of the offspring's age. Offspring born at 416 suboptimal periods face the dry season in a critical developmental window (i.e., the end of 417 weaning), and maternal care is insufficient to buffer them entirely from the adverse 418 consequences that lead to higher mortality. Consequently, baboon infants respond by throwing 419 more tantrums, which may be an honest signal of need [38,62], just as children do in similar 420 situations [63]. More generally, these results shed light on the potential influence of 421 environmental fluctuations, and specifically seasonality, on mother-offspring conflicts over

maternal care. While the literature focusing on optimal birth spacing has mainly examined trade-offs between current and future reproduction for an implicitly stable level of resources [7,64], such a stability may rarely be encountered by mothers in the wild, who typically face extensive, but partly predictable, fluctuations in food availability. Taking into account the intensity and predictability of resource fluctuations may largely re-draw the landscape of strategic decisions available to mothers confronted with trade-offs between current and future reproduction in natural environments [65,66].

429 Our findings open new perspectives to understand the evolutionary drivers of vertebrate 430 reproductive seasonality, by revealing the occurrence of a maternal trade-off between current 431 and future reproduction over birth timing, extended by mother-offspring conflict during 432 weaning. Such a trade-off may commonly occur in organisms with a slow reproductive pace, 433 and future studies on such taxa should investigate the consequences of reproductive timing on 434 several fitness components. Indeed, multiple optimal birth timings in the annual cycle may 435 generate a bimodal birth peak or an extended birth season. Evolutionary trade-offs over birth 436 timing may therefore account for unexplained variation in the reproductive timing of long-lived 437 vertebrates, including the evolution of non-seasonal breeding in humans and other species.

438

439 ACKNOWLEDGMENTS

The authors are grateful to the Tsaobis Baboon Project volunteers from 2005 to 2019, and particularly to Harrison Anton, Charlotte Bright, Anna Cryer, Rémi Emeriau, Richard Gallagher, Chloe Hartland, Rachel Heaphy, Nick Matthews, Tess Nicholls, Vittoria Roatti and Ndapandula Shihepo for their dedicated effort at collecting focal observations on infant baboons. This research was carried out with the permission of the Ministry of Environment and Tourism, the Ministry of Land Reform, and the National Commission on Research, Science, and Technology. We further thank the Tsaobis beneficiaries for permission to work at Tsaobis, the Gobabeb Namib Research Institute and Training Centre for affiliation, and Johan Venter
and the Snyman and Wittreich families for permission to work on their land. We also thank
Jacinta Beehner and an anonymous reviewer for their constructive remarks on this manuscript.
Data used in this study are part of long-term data collected within the framework of the Tsaobis
Baboon Project, recently funded by a grant from the Agence Nationale de la Recherche (ANR
ERS-17-CE02-0008, 2018-2021) awarded to EH. This paper is a publication of the ZSL
Institute of Zoology's Tsaobis Baboon Project. Contribution ISEM n°XX.

454

455 **REFERENCES**

456 1. Bronson FH. 2009 Climate change and seasonal reproduction in mammals. *Philos*.

457 *Trans. R. Soc. B Biol. Sci.* **364**, 3331–3340. (doi:10.1098/rstb.2009.0140)

- 458 2. Brockman DK, van Schaik C. 2005 Seasonality and reproductive function. In
- 459 Seasonality in Primates: Studies of Living and Extinct Human and Non-Human
- 460 *Primates* (ed Cambridge University Press), pp. 269–305.
- 461 3. van Schaik CP, van Noordwijk MA. 1985 Interannual variability in fruit abundance
- 462 and the reproductive seasonality in Sumatran Long-tailed macaques (Macaca
- 463 fascicularis). J. Zool., Lond. **206**, 533–549. (doi:10.1111/j.1469-7998.1985.tb03557.x)
- 464 4. Janson C, Verdolin J. 2005 Seasonality of primate births in relation to climate. In
- 465 Seasonality in Primates: Studies of Living and Extinct Human and Non-Human
- 466 *Primates*, pp. 307–350. Cambridge: Cambridge University Press.
- 467 (doi:10.1017/cbo9780511542343.012)
- 468 5. Lee PC. 1996 The meanings of weaning: growth, lactation, and life history. *Evol*.
- 469 Anthropol. 5, 87–98. (doi:https://doi.org/10.1002/(SICI)1520-6505(1996)5:3<87::AID-

470 EVAN4>3.0.CO;2-T)

471 6. Emery Thompson M. 2013 Comparative reproductive energetics of human and

- 472 nonhuman primates. *Annu. Rev. Anthropol.* 42, 287–304. (doi:10.1146/annurev-anthro473 092412-155530)
- 474 7. Lee SD, Ruiz-Lambides A V., Higham JP. 2019 Higher offspring mortality with short
 475 interbirth intervals in free-ranging rhesus macaques. *Proc. Natl. Acad. Sci.* 116, 1–6.
- 476 (doi:10.1073/pnas.1817148116)
- 8. Bateson P. 1994 The dynamics of parent-offspring relationships in mammals. *Trends Ecol. Evol.* 10, 399–403. (doi:10.1016/0169-5347(94)90066-3)
- 479 9. Brinkhof MWG, Cavé AJ, Daan S, Perdeck AC. 2002 Timing of current reproduction
- 480 directly affects future reproductive output in european coots. *Evolution (N. Y).* 56, 400–
 481 411.
- 482 10. Fischer J *et al.* 2019 The Natural History of Model Organisms: Insights into the
 483 evolution of social systems and species from baboon studies. *Elife* 1960, 1–16.
- 484 11. Barrett L, Henzi SP. 2008 Baboons. Curr. Biol. 18, 404–406.
- 485 (doi:10.1016/j.cub.2008.02.074)
- 486 12. Swedell L. 2011 African papionins: diversity of social organization and ecological
- 487 flexibility. In *Primates in perspective*, pp. 241–277. Oxford: Oxford University Press.
- 488 13. Cheney DL *et al.* 2004 Factors affecting reproduction and mortality among baboons in
- the Okavango Delta, Botswana. Int. J. Primatol. 25, 401–428. (doi:0164-
- 490 0291/04/0400-0401/0)
- 491 14. Lycett JE, Weingrill T, Henzi SP. 1999 Birth patterns in the Drakensberg Mountain
 492 baboons (Papio cynocephalus ursinus). S. Afr. J. Sci. 95, 354–356.
- 493 15. Campos FA et al. 2017 Does climate variability influence the demography of wild
- 494 primates? Evidence from long-term life-history data in seven species. *Glob. Chang.*
- 495 *Biol.* **23**, 1–15. (doi:10.1111/gcb.13754)
- 496 16. Maestripieri D. 2002 Parent-offspring conflict in primates. Int. J. Primatol. 23, 923-

497

- 951. (doi:10.1023/A:1015537201184)
- 498 17. Trivers RL. 1972 Parental Investment and Sexual Selection. In *Sexual Selection and*499 *the Descent of Man 1871-1971* (ed Harvard University), pp. 136–207.
- 500 (doi:10.1002/ajpa.1330400226)
- 501 18. Gesquiere LR, Altmann J, Archie EA, Alberts SC. 2017 Interbirth intervals in wild
- 502 baboons: Environmental predictors and hormonal correlates. *Am. J. Phys. Anthropol.*

503 **166**, 107–126. (doi:10.1002/ajpa.23407)

- 504 19. Bulger J, Hamilton WJ. 1987 Rank and density correlates of inclusive fitness measures
- 505 in a natural chacma baboon (Papio ursinus) troop. *Int. J. Primatol.* **8**, 635–650.
- 506 (doi:10.1007/BF02735781)
- 507 20. Packer C, Collins DA, Sindimwo A, Goodall J. 1995 Reproductive constraints on
- aggressive competition in female baboons. *Nature* **373**, 60–63. (doi:10.1038/373060a0)
- 509 21. Lonsdorf E V. 2017 Sex differences in nonhuman primate behavioral development. J.

510 *Neurosci. Res.* **95**, 213–221. (doi:10.1002/jnr.23862)

- 511 22. Cowlishaw G, Davies JG. 1997 Flora of the Pro-Namib Desert Swakop River
- 512 catchment, Namibia: community classification and implications for desert vegetation

513 sampling. J. Arid Environ. **36**, 271–290. (doi:0140–1963/97/020271)

- 514 23. Huffman GJ, Bolvin DT, Nelkin EJ, Adler RF. 2016 TRMM (TMPA) Precipitation L3
- 515 1 day 0.25 degree x 0.25 degree V7.
- 516 24. Pettorelli N, Vik JO, Mysterud A, Gaillard JM, Tucker CJ, Stenseth NC. 2005 Using
- 517 the satellite-derived NDVI to assess ecological responses to environmental change.
- 518 *Trends Ecol. Evol.* (doi:10.1016/j.tree.2005.05.011)
- 519 25. Didan K, Barreto Munoz A, Solano R, Huete A. 2015 MOD13A1 MODIS/Terra
- 520 Vegetation Indices 16-Day L3 Global 500m SIN Grid V006 [Data set].
- 521 (doi:https://doi.org/10.5067/MODIS/MOD13A1.006)

- 522 26. Dezeure J, Dagorrette J, Baniel A, Carter AJ, Cowlishaw G, Marshall HH, Martina C,
- 523 Raby CL, Huchard E. 2021 Developmental transitions in body color in chacma baboon
- 524 infants: Implications to estimate age and developmental pace. *Am. J. Phys. Anthropol.*
- 525 **174**, 89–102. (doi:10.1002/ajpa.24118)
- 526 27. Zinner D, Peí Aez F, Torkler F. 2001 Distribution and habitat associations of baboons
 527 (Papio hamadryas) in central Eritrea. *Int. J. Primatol.* 22, 397–413.
- 528 28. Calenge C. 2006 The package 'adehabitat' for the R software: A tool for the analysis of
 529 space and habitat use by animals. *Ecol. Modell.* 197, 516–519.
- 530 (doi:10.1016/j.ecolmodel.2006.03.017)
- 531 29. Bronikowski AM, Altmann J. 1996 Foraging in a variable environment: weather
- 532 patterns and the behavioral ecology of baboons. *Behav Ecol Sociobiol* **39**, 11–25.
- 30. Huchard E, Knapp L, Wang J, Raymond M, Cowlishaw G. 2010 MHC, mate choice
 and heterozygote advantage in a wild social primate. *Mol. Ecol.*, 2545–2561.
- 535 (doi:10.1111/j.1365-294X.2010.04644.x)
- 536 31. Huchard E, Courtiol A, Benavides JA, Knapp LA, Raymond M, Cowlishaw G. 2009
- 537 Can fertility signals lead to quality signals? Insights from the evolution of primate
- 538 sexual swellings. *Proc. R. Soc. B Biol. Sci.* **276**, 1889–1897.
- 539 (doi:10.1098/rspb.2008.1923)
- 540 32. Huchard E, Cowlishaw G. 2011 Female female aggression around mating : an extra
 541 cost of sociality in a multimale primate society. *Behav. Ecol.* 22, 1003–1011.
- 542 (doi:10.1093/beheco/arr083)
- 543 33. Borries C, Lu A, Ossi-Lupo K, Larney E, Koenig A. 2014 The meaning of weaning in
- 544 wild Phayre's leaf monkeys: Last nipple contact, survival, and independence. Am. J.
- 545 Phys. Anthropol. 154, 291–301. (doi:10.1002/ajpa.22511)
- 546 34. Saltzman W, Tardif SD, Rutherford JN. 2011 Hormones and Reproductive Cycles in

547 Primates. In *Hormones and reproduction of vertebrates* (eds Norris DO, Lopez K), pp.

548 1–40. New York: NY Academic Press.

- 549 35. Altmann J. 1974 Observational Study of Behavior: Sampling Methods. *Behaviour* 49,
 550 227–267.
- 36. Altmann J, Samuels A. 1992 Costs of maternal care: infant-carrying in baboons. *Behav. Ecol. Sociobiol.* 29, 391–398. (doi:10.1007/BF00170168)
- 553 37. Bădescu I. 2018 The attainment of independence from the mother in primate infants
- and its implications for the evolution of cooperative breeding in hominins. In *The*
- 555 talking species : perspectives on the evolutionary, neuronal and cultural foundations of
- 556 *language* (eds EM Luef, MM Marin), pp. 169–194. Uni-Press Graz.
- 38. Barrett L, Henzi PS. 2000 Are baboon infants Sir Phillip Sydney's offspring? *Ethology*106, 645–658. (doi:10.1046/j.1439-0310.2000.00577.x)
- 559 39. Nicolson N. 1982 Weaning and the Development of Independence in Olive Baboons.
- 560 Harvard University, Cambridge, Massachusetts. (doi:10.13140/RG.2.1.4000.9761)
- 40. Batschelet E. 1981 *Circular Statistics in Biology (Mathematics in Biology)*. Academic
 562 Press.
- 41. Altmann J, Alberts SC. 2005 Growth rates in a wild primate population: ecological
 influences and maternal effects. *Behav. Ecol. Sociobiol.* 57, 490–501.
- 565 (doi:10.1007/s00265-004-0870-x)
- 566 42. Bercovitch FB, Berard JD. 1993 Behavioral Ecology and Sociobiology Life history
- 567 costs and consequences of rapid reproductive maturation in female rhesus macaques.
- 568 *Behav Ecol Sociobiol* **32**, 103–109.
- 569 43. Altmann J. 1980 Baboon Mothers and Infants. The University of Chicago Press.
- 570 44. Harrison XA, Donaldson L, Eugenia Correa-Cano M, Evans J, Fisher DN, Goodwin C,
- 571 Robinson B, Hodgson DJ, Inger R. 2017 Best practice in mixed effects modelling and

- 572 multi-model inference in ecology. *PeerJ Open Access*
- 573 (doi:10.7287/peerj.preprints.3113v1)
- 45. Wood SN. 2003 Thin plate regression splines. J. R. Stat. Soc. Stat. Methodol. Ser. B 65,
 95–114.
- 576 46. R Core Team. 2019 R: A language and environment for statistical computing.
- 47. Agostinelli C, Lund U. 2018 Package 'CircStats': Circular Statistics, from 'Topics in
 578 Circular Statistics'.
- 48. Bates D, Mächler M, Bolker BM, Walker SC. 2015 Fitting linear mixed-effects models
 using lme4. J. Stat. Softw. 67. (doi:10.18637/jss.v067.i01)
- 581 49. Dorie V. 2015 blme: Bayesian Linear Mixed-Effects models.
- 582 50. Fox J et al. 2019 Companion to Applied Regression.
- 583 51. Hartig F. 2020 Package 'DHARMa': Residual Diagnostics for Hierarchical (Multi584 Level / Mixed) Regression Models.
- 585 52. Weibel CJ, Tung J, Alberts SC, Archie EA. 2020 Accelerated reproduction is not an
- adaptive response to early-life adversity in wild baboons. *PNAS* **117**, 24909–24919.
- 587 (doi:10.1073/pnas.2004018117/-/DCSupplemental)
- 588 53. Jones TM *et al.* 2020 Parental benefits and offspring costs reflect parent-offspring
- 589 conflict over the age of fledging among songbirds. *PNAS* **117**, 30539–30546.
- 590 (doi:10.1073/pnas.2008955117)
- 591 54. Lummaa V, Lemmetyinen R, Haukioja E, Pikkola M. 1998 Seasonality of births in
- 592 Homo sapiens in pre-industrial Finland: maximisation of offspring survivorship? J.
- 593 *Evol. Biol.* **11**, 147–157. (doi:10.1046/j.1420-9101.1998.11020147.x)
- 594 55. Bronson FH. 1989 *Mammalian Reproductive Biology*. Chicago: The University of
 595 Chicago Press.
- 596 56. Bronson FH. 1995 Seasonal variations in human reproduction: environmental factors.

- *Q. Rev. Biol.* **70**, 55.
- 598 57. Ellison PT, Valeggia CR, Sherry DS. 2005 Human Birth Seasonality. In *Seasonality in*
- 599 Primates: Studies of Living and Extinct Human and Non-Human Primates, pp. 379–
- 600 399. Cambridge: Cambridge University Press.
- 601 58. Stouffer PC, Johnson EI, Bierregaard RO. 2013 Breeding seasonality in central
- 602 Amazonian rainforest birds. *Auk* **130**, 529–540. (doi:10.1525/auk.2013.12179)
- 603 59. Sinclair ARE, Mduma SAR, Arcese P. 2000 What determines phenology and

604 synchrony of ungulate breeding in Serengeti? *Ecology* **81**, 2100–2111.

- 605 (doi:10.1890/0012-9658(2000)081[2100:WDPASO]2.0.CO;2)
- 606 60. Wright PC. 1999 Lemur Traits and Madagascar Ecology: Coping With an Island
- 607 Environment. Yearb. Phys. Anthropol. 42, 31–72. (doi:10.1002/(sici)1096-
- 608 8644(1999)110:29+<31::aid-ajpa3>3.0.co;2-0)
- 609 61. Barrett L, Henzi SP, Lycett JE. 2006 Whose Life Is It Anyway? Maternal Investment,
- 610 Developmental Trajectories, and Life History Strategies in Baboons. In *Reproduction*
- 611 *and Fitness in Baboons: Behavioral, Ecological, and Life History Perspectives*, pp.
- 612 199–224.
- 613 62. Lummaa V, Vuorisalo T, Barr RG, Lehtonen L. 1998 Why Cry? Adaptive Significance
 614 of Intensive Crying in Human Infants. *Evol. Hum. Behav.* 19, 193–202.
- 615 63. Fouts HN, Hewlett BS, Lamb ME. 2005 Parent-Offspring weaning conflicts among the
 616 Bofi farmers and foragers of Central Africa. *Curr. Anthropol.* 46, 29–50.
- 617 64. Fotso JC, Cleland J, Mberu B, Mutua M, Elungata P. 2013 Birth spacing and child
- 618 mortality: An analysis of prospective data from the nairobi urban health and
- 619 demographic surveillance system. J. Biosoc. Sci. 45, 779–798.
- 620 (doi:10.1017/S0021932012000570)
- 621 65. Ljungström G, Francis TB, Mangel M, Jørgensen C. 2019 Parent-offspring conflict

- 622 over reproductive timing: Ecological dynamics far away and at other times may explain
- 623 spawning variability in Pacific herring. *ICES J. Mar. Sci.* **76**, 559–572.
- 624 (doi:10.1093/icesjms/fsy106)
- 625 66. de Villemereuil P et al. 2020 Fluctuating optimum and temporally variable selection on
- breeding date in birds and mammals. *PNAS*, 1–10. (doi:10.1073/pnas.2009003117/-
- 627 /DCSupplemental.y)
- 628
- 629

630 TABLES

631 **Table 1:** Predictors of female reproductive timing.

Estimates, confidence intervals, X² statistics and P-values of the predictors of the two linear mixed models (Models
3 and 4). The response variables are respectively the deviation from the offspring survival optimal birth timing,
i.e. from December 15th (Model 3), and the deviation from the maternal IBI optimal birth timing, i.e. from
September 1st (Model 4), in days, based on 215 births from 62 females. Female identity and year of infant birth are
included as random effects. For categorical predictors, the tested category is indicated between parentheses.

Fixed offect		Estimate	I	С	\mathbf{V}^2	D value
ГІХ			Lower	Upper	Λ^2	P-value
	Model 3: Deviat	l birth timi	ng			
Infant sex	(Male)	5.91	-7.42	19.23	0.76	0.385
Female parity	(Primiparous)	-12.77	-30.02	4.47	2.11	0.147
Female ranl	k	2.59	-4.82	10.00	0.47	0.493
Casua	(L)	5.16	-10.37	20.69	1 22	0.515
Group	(M)	-12.66	-44.70	19.39	1.55	0.313
	Model 4: Devi	iation from tl	he maternal [IBI optimal b	oirth timing	
Infant sex	(Male)	-3.19	-16.46	10.08	0.22	0.7637
Female parity	(Primiparous)	9.67	-7.49	26.82	1.22	0.269
Female ranl	k	-3.41	-10.75	3.92	0.83	0.362
Group	(L)	10.67	-4.70	26.04	1.02	0.282
Group	(M)	0.93	-30.91	32.78	1.72	0.362

638 **FIGURES**



639

640 **Figure 1:** Tsaobis baboons' reproductive timings in relation with environmental seasonality.

641 In Panel A, we plotted the proportion of conceptions (N=241, in light grey) and births (N=215, in dark grey) 642 recorded in 2005-2019 per month (Jan for January, Jul for July). We plotted the mean monthly cumulative rainfall 643 (in mm) per month in blue and the mean NDVI value per month in green between 2005 and 2019. We represented 644 the standard errors associated with vertical black segments. The pink and orange squares in the background 645 represent respectively the maternal IBI and the offspring survival optimal birth timings. In Panel B, we aimed to 646 represent the different phases of the female reproductive cycle, when the birth date occurs within the offspring 647 survival (December 15th) or maternal IBI (September 1st) optimal timing, according to seasonal variation of NDVI. 648 The green bar plot in the background indicates the mean NDVI per month (see y-axis). Pregnancy, indicated with 649 grey arrows, occurs the 6 months prior a birth. Early-weaning, indicated with salmon-colour arrows, occurs from 650 6 to 9 months after a birth. Lactation peak, indicated with black stars, occur around 6 months after a birth. Weaning 651 end, indicated with blue arrows, occurs from 12 to 18 months after a birth (see Appendix 3 for the characterization 652 of these different reproductive stages).



653



We plotted the predicted values of the full models (Model 1 looking at offspring mortality in panel A, and Model 2 looking at IBIs in panel B) according to the month of infant birth (Jan for January, Apr for April, etc.). The number of births observed for each month is indicated in blue below the bar. The dots represent the mean values,

658 while the vertical black bars represent its standard deviations. The offspring survival optimal birth timing is

identified as the period minimizing offspring mortality, i.e. from November to February, and indicated with orange

dots (Panel A). The maternal IBI optimal birth timing is identified as the period minimizing maternal interbirth

661 interval, i.e. from July to September, and indicated with pink dots (Panel B).



663 **Figure 3**: Influence of birth timing on tantrum probability.

Predicted values of tantrum probability (Model 7) at weaning (age 12 months), according to infants' birth month, based on 2221 focal observations from 55 infants. For graphical reasons, and given the low sample size of infants observed for some birth months, we pooled infants born in 2 consecutive months, so that Jan indicates infants born in both January and February, Mar in both March and April, etc. The brown horizontal bars indicate the median values of fitted values for each birth month category.

669

- 670
- 671
- 672

673

- 674 Supporting Information for:
- 675 Birth timing generates reproductive trade-offs in a non-seasonal

676 breeding primate

- 577 Jules Dezeure^{1, 5,*}, Alice Baniel², Alecia J. Carter³, Guy Cowlishaw⁴, Bernard Godelle¹, Elise
- 678 Huchard¹
- 679 ¹Anthropologie Évolutive, Institut des Sciences de l'Évolution de Montpellier (ISEM),
- 680 Université de Montpellier, CNRS, IRD, EPHE, Montpellier, France.
- ⁶⁸¹ ²Department of Anthropology, Stony Brook University, Stony Brook, NY, USA
- ³Department of Anthropology, University College London, London, UK
- ⁴Institute of Zoology, Zoological Society of London, Regent's Park, London NW1 4RY
- 684 ⁵Lead contact
- 685 *Corresponding author: Jules Dezeure
- 686 dezeurejules@gmail.com
- 687

689 CONTENTS

690 691	SUDDI EMENTADV TEVT	22
692	Annendix 1 Correlations between rainfall and NDVI at Tsaobis	22
602	Annendix 2 Estimations of the dates of concentions, births and evaluations	22
095	Appendix 2. Estimations of the dates of conceptions, on this and cycle resumptions	
694	Appendix 3. Characterization of developmental stages: weaning and lactation peak	35
695	Appendix 4. Methodology of infants' focal observations	37
696 697	Appendix 5. Controlling for uncertainties in the dates of conceptions, births and cycle resumptions in statistical analyses	s 37
698	Appendix 6. Method used to capture birth timing effects in Models 1-2 and 5-7	40
699	Appendix 7. Seasonal effects on maternal care and tantrum probability	41
700	TABLES	42
701 702	Table S1 : Different methods used to estimate the dates of births of the 215 baboon infants born at Tsaobis between 2005 and 2019.	42
703 704	Table S2: Identification of the best birth timing effect for Models 1-2 & 5-7: Δ AIC (AkaikeInformation Criterion) according to the phase of the sine wave term	43
705 706	Table S3: Birth timing and other predictors of the probability of suckling, infant carrying and tantrums	45
707 708	Table S4 : Seasonality and other predictors of the probability of suckling, infant carrying and tantrums	47
709	Table S5: Summary of the structure of all models included in the study.	49
710 711	Table S6 : Predictors of offspring mortality before weaning and maternal interbirth interval (IBI) duration.	50
712	FIGURES	51
713 714	Figure S1: Variation in the probabilities of suckling, infant carrying, and tantrums, according to infant age	51
715	Figure S2: Chacma baboons breed all year round	53
716	REFERENCES	54
717		

719 SUPPLEMENTARY TEXT

720 Appendix 1. Correlations between rainfall and NDVI at Tsaobis

721 In order to estimate the correlation between monthly rainfall and NDVI at Tsaobis, we used a 722 moving window approach. We expected that the cumulative rainfall over the preceding months, 723 rather than the rainfall during the current month, would be the best predictor of monthly NDVI. 724 First, we identified the time window maximizing the correlation between rainfall and NDVI 725 variation, testing periods covering 0 to 6 month(s) prior to the current month using an AIC-726 based selection procedure, and a univariate linear model containing only the fixed effect of 727 interest (cumulative rainfall over variable periods) and three response variables, namely the 728 monthly NDVI values associated with the home ranges of all three study groups. For these three 729 groups, the time window minimizing the model AIC was cumulative rainfall over the preceding three months. The adjusted value of the model R² measures the proportion of NDVI variance 730 731 explained by variation in cumulative rainfall over the past three months.

732

733 Appendix 2. Estimations of the dates of conceptions, births and cycle resumptions

We characterized the reproductive seasonality in our population considering three main reproductive events: conceptions, births, and cycle resumptions (i.e. the end of post-partum amenorrhea).

1. *Births.* The dates of births, conceptions, and cycle resumptions were directly observed where possible, but otherwise estimated for those periods when no observers were present. We observed a total of 84 births. Of those, 62 were seen by observers on the exact day, and 22 were witnessed after a short absence (leading to a small uncertainty in the actual date: median=17 days, range=1-30). When the conception only was observed (n=52 births), we estimated birth dates by adding the mean gestation length (n=190 days, range: 181-200 days, 743 SD=5, n=13 pregnancies where both conception and birth were observed) to the conception date. 744 Conception was considered to occur on the day of deturgescence (D-day) of the swelling during a conceptive cycle. This generated a total of 136 birth dates known with high accuracy. When 745 746 neither conception nor birth were observed (n=56 births), the birth date was estimated using infant coloration (based on the progressive loss of natal coat and skin coloration) following a 747 748 method recently described and validated in our population [1], with further refinement provided by the reproductive history of the mother (e.g., if the mother was pregnant during the last three 749 750 months of a field season, then the infant was necessarily born in the three months following the 751 end of this season given that a pregnancy lasts 190 days). Finally, when neither birth nor 752 conception was observed, and infant colour when first seen was unknown or uninformative (i.e., 753 the transition from natal to adult coat had already occurred) (n=23 births), we used the 754 reproductive state of females in the preceding field season to minimize uncertainty over birth 755 timing. For example, if a female was cycling the last day of the preceding season, the infant was necessarily born at least 190 days after this day. In total, our sample comprised 215 births 756 757 between 2005 and 2019, with a median uncertainty of 10 days (range: 0-153 days) (Table S1).

758 2. Conceptions. We observed 81 conceptions: 68 conceptions were witnessed 759 (observers were present during the conceptive cycle), and 13 occurred during a short absence 760 of observers (leading to a small uncertainty in the actual date: median=10 days, range=3-30). 761 The exact date of conceptions was the day of swelling detumescence of the conceptive cycle 762 (when witnessed) [2]. When the birth was observed but not the conception, the latter was 763 estimated to occur 190 days prior to birth (n=65 conceptions). When neither conception nor 764 birth were observed but birth occurred (i.e. no miscarriage), we estimated birth date as 765 explained above, and inferred conception from the birth date using the mean gestation period 766 (n=79). Finally, when pregnancy signs were seen (i.e. red paracallosal skin and an absence of 767 cycles) but conception was not observed and there was no birth due to a miscarriage or early

death (occurring before an infant was recorded by observers), we estimated the date of conception using female reproductive states (n=16 conceptions). For example, if a female was seen pregnant on the first day of a field season, we knew that the conception occurred from 0-190 days prior to this date (as gestation lasts 190 days in this population). All in all, we generated a sample of 241 conceptions, with a median uncertainty of 10 days (range: 0-164 days).

774 3. *Cycle resumptions*. Only cycle resumptions following a period of lactation were 775 included in our analyses. Cycle resumptions were observed in 64 cases. In 107 other cases, a 776 female was lactating at the end of a field season and then cycling at the beginning of the next 777 (median days between consecutive field seasons=225, range=83-584). To reduce the 778 uncertainty of the resumption date estimations in these cases, we calculated the minimum post-779 partum amenorrhea length (the time between birth and cycle resumption) (mean \pm SD = 353 \pm 780 89 days, range=223-550) based on the 33 cases for which both events were known, and used 781 this value as a minimum threshold in our estimations. We also excluded all cycle resumptions 782 for which the uncertainty exceeded one year. Our sample thus comprised a total of 171 cycle 783 resumptions, with a median uncertainty of 61 days (range= 0-272 days).

784

785 Appendix 3. Characterization of developmental stages: weaning and lactation peak

In order to understand which stage of the reproductive cycle might be timed with the seasonal food peak (Figure 1B), we needed to define the sequential behavioural stages of weaning, which is the infant's gradual transition to nutritional independence [3], along with the peak of lactation in our population. First, the onset of weaning can be defined as the period when solid foods start to account for an important part of an infant's energy intake, and is characterized by an increase in maternal refusals to accede to her offspring's suckling demands. The onset of 792 behavioural conflict between a mother and her offspring has therefore often been used as a 793 proximate measure of the early-weaning period [3]. In our population, tantrum probabilities 794 peak between 6 and 9 months of age (Figure S1), and we therefore used this age window to 795 characterize 'early-weaning'. The peak of lactation typically occurs just before the beginning 796 of weaning [3,4], when offspring have grown larger but are still fully dependent. So we can 797 consider that lactation peak occurs around 6 months after birth in our population. Similar ages 798 for early-weaning and lactation peak have been found in other baboon populations [5–7], albeit 799 weaning age and lactation durations can vary substantially between populations [8].

800 The end of weaning can be defined as the complete cessation of nursing, i.e. when 801 offspring feed exclusively on solid foods. Looking at behavioural data, suckling frequencies 802 decrease gradually from 2 to 18 months old, before stabilizing to ca. 2% of time from 18 to 24 803 months old (Figure S1). In addition, the maximum length of post-partum amenorrhea (PPA), 804 often used as a proxy for the end of weaning [3], was 550 days (i.e. 18.1 months) in this 805 population (based on n = 33 cases for which both birth and cycle resumption were known with 806 accuracy). We therefore considered the age of 18.1 months as an upper threshold marking the 807 end of weaning for all juveniles in our models on infant mortality and IBI (see main text). 808 However, age at the end of weaning is highly variable between individuals, as indicated by the 809 wide range of variation for PPA (8-18 months, mean = 12 months). To take this variation into 810 account, we defined the 'end of weaning' as the age window of 12-18 months after birth for 811 Figure 1B.

All in all, for Figure 1B, in order to better understand the relationship between reproductive phenology and environmental seasonality, we considered the lactation peak to occur around 6 months after birth, early-weaning between 6 and 9 months of age, and the end of weaning between 12 and 18 months of age. As a note of caution, these windows are strictly based on behavioural and life-history data, which show some limitations to evaluate the dynamics of lactation [9,10]. More objective measures, such as isotopic comparisons of motheroffspring hair or faecal samples [11–14], may help to refine these estimates.

819

820 Appendix 4. Methodology of infants' focal observations

821 Field observers collected behavioural data on infants aged between 2 and 24 months on a daily 822 basis from dawn until dusk over four periods: from October to December 2006, from July to 823 August 2017, from September to December 2018, and from April to July 2019. We collected a 824 total of 1185 hours of focal observation on 69 infants across four field seasons (mean \pm SD = 825 17.1 ± 7.8 hours of observations per infants, range = 6.3–34.6), with a mean of 40.7 focal 826 observations per individual (SD=29.4). Focal observations were spread equally across the day 827 (divided in four 3 h-long blocks) and focal individuals were chosen randomly, and never 828 sampled more than once within a block. Focal observations durations were 1 h in 2006 and 20 829 min in 2017-2019, with a minimum of 10 min in all cases. We recorded the following activities 830 on a continuous basis: suckling (when the focal individual had its mouth on its mother's nipple; 831 we could not distinguish comfort from nutritive suckling), travelling alone, infant carrying 832 (carried by the mother, either ventrally or dorsally) and other activities. We also collected 833 events related to mother-offspring conflicts (see in main text). In addition, we collected scan 834 observations every 5 minutes (n=16702 scans across 3081 focal observations), including the 835 activity of the focal individual.

836

837 Appendix 5. Controlling for uncertainties in the dates of conceptions, births and cycle

838 resumptions in statistical analyses

Bates of conceptions, births and cycle resumptions were estimated in many cases because the Tsaobis baboons are not followed year round (see Appendix 1). In addition, uncertainty in these estimations varied with the time of year, as we generally follow baboons during the cooler, 842 dryer months. In order to account for these uncertainties in our analyses, we ran a set of 843 randomizations to evaluate the robustness of the fixed effects found to be statistically 844 significant. For each reproductive event (conceptions, births, and cycle resumptions) for which 845 the date was associated to some uncertainty (i.e. exact date unknown), we created an extended 846 dataset including all possible dates of the full range of uncertainty (from the minimum to the 847 maximum date). For example if a baboon birth date was estimated to occur between October 2nd and December 23rd, we included all possible dates between October 2nd and December 23rd 848 849 in this extended dataset. Using this extended dataset, we generated 1000 simulations; in each iteration, a date was randomly drawn between the minimal and maximal estimate for each 850 851 reproductive event that was not known with certainty. Events known with certainty did not vary 852 throughout such simulations.

These simulations were integrated in different statistical analyses slightly differently. In our characterization of reproductive seasonality, we extracted the mean R, μ and p-value of the Rayleigh test for the 1000 simulated datasets of cycle resumptions, conceptions and births. We indicate these mean values in the main text. We also computed the 95% level confidence intervals of these p-values: for conceptions, p=0.019 – 0.021; for births, p=0.166 – 0.174; for cycle resumptions, p=0.328 – 0.358.

859 In our multivariate mixed models investigating the effect of seasonal birth timing on 860 offspring mortality before weaning (Model 1) and maternal interbirth intervals (Model 2), we 861 controlled for the uncertainties in dates of birth which could affect both our response variables 862 and our main fixed effect of interest (seasonal birth timing). For Model 1, we generated 1000 863 simulations with random birth dates drawn, for each birth, between minimal and maximal birth 864 date estimations for this particular birth, and subsequently ran 1000 mixed models, one for each 865 simulated value of the offspring's birth date and for each survival outcome (as birth date affects an offspring's age, and therefore its estimated age at death). For Model 2, we similarly 866

generated 1000 simulations with random birth dates drawn between minimum and maximum birth date estimations for the two births defining the IBI. We subsequently ran 1000 models with randomized values for both IBI (the response variable, i.e. number of days between the first and second birth) and the birth date fixed effect. For both models, we then extracted the 1000 p-values of our fixed effect 'seasonal birth timing' and computed the confidence intervals of these p-values (see the footnote of Table S6).

873 In our analysis of the individual determinants of birth timing (Models 3 and 4), we 874 similarly generated 1000 simulations of birth dates drawn between minimum and maximum 875 birth date estimations, and assessed for each of these randomly drawn births the deviation, in days, from December 15th for Model 3 and September 1st for Model 4 respectively. For both 876 877 models, we then ran 1000 LMMs using these deviations as the response variable (Models 3 and 878 4). We extracted 1000 p-values of our various fixed effects, and computed their 95% level 879 confidence intervals. No fixed effect was close to significance, and this information was thus 880 not added to the footnote of Table 1.

881 Finally, in our analysis investigating the effects of seasonal birth timing on maternal 882 care and mother-offspring conflict at the behavioural level, we similarly generated 1000 883 simulations of birth dates drawn between minimum and maximum birth date estimations. We 884 then ran 1000 GLMMs looking at the effect of seasonal birth timing, along with other 885 covariates, on the probabilities of suckling (Model 5), infant carrying (Model 6) and tantrums 886 (Model 7). We extracted the 1000 resulting p-values for our fixed effect 'seasonal birth timing', 887 computed their median and 95% level confidence interval, and added this information to the 888 footnote of Table S3.

Appendix 6. Method used to capture birth timing effects in Models 1-2 and 5-7

To investigate the influence of birth timings on offspring mortality (Model 1), female IBI (Model 2), suckling probability (Model 5), infant carrying probability (Model 6) and tantrum probability (Model 7), we used a sine term that captures the timing of an infant's birth in the annual cycle. Sine waves allow the introduction of a circular variable into a multivariate model as a fixed effect: the possible effects of the date of birth are circular with a period of one year, as January 1st is equally close to December 31st than to January 2nd. This sinusoidal term was as follows:

898

$sin(Date of Birth + \varphi)$

899 The date of birth in the formula above was converted in a radian measure, so that the period,

900 i.e. one year, equalled to $2^{*}\pi$, ranging from $2^{*}\pi/365$ for January 1^{st} to $2^{*}\pi$ for December 31^{st} .

901 We tested 12 different phase values φ (0, $\pi/12$, $2^*\pi/12$, $3^*\pi/12$, $4^*\pi/12$, $5^*\pi/12$, $6^*\pi/12$,

902 $7*\pi/12$, $8*\pi/12$, $9*\pi/12$, $10*\pi/12$, $11*\pi/12$), to account for different potential optimal periods

903 for the event of interest across the year [1], as our different response variables could be

904 minimized for different birth dates (and so tested all potential dates as minimal). For example,

a phase of 0 could maximize April 1st or October 1st depending on the sign of the estimate

906 (see Table S2). We ran sequentially these 12 multivariate models, containing all other fixed

907 and random effects (see below), and selected the best phase as the one minimizing the Akaike

908 Information Criterion (AIC) in this full model set: the phase of $7*\pi/12$ was retained for

- 909 offspring mortality probability, $2*\pi/12$ for IBI, $9*\pi/12$ for suckling, 0 for infant carrying, and
- 910 $2*\pi/12$ for tantrum probabilities (Table S2).

912 Appendix 7. Seasonal effects on maternal care and tantrum probability

When modelling suckling, infant carrying and tantrum probabilities (Models 5-7), we further tested for seasonal effects, i.e. effects of the date of observation, on the response variable. To do so, we applied the approach used to describe the effects of birth timings, i.e. a sine term of the date of observation (in radians) was entered as a fixed effect in the multivariate model. This sinusoidal term was as follows:

918

$sin(Date of observation + \varphi)$

The date of observation in the formula above was converted to a radian measure, so that the period, i.e. one year, equalled $2*\pi$, ranging from $2*\pi/365$ for the 1st of January to $2*\pi$ for the 31^{st} of December. We similarly tested 12 different phase values φ and selected the best phase as the one minimizing the AIC of the full models 5-7 (with all random and fixed effects, except the sine term of the date of birth). We found that $7*\pi/12$ was the best phase for suckling probability, $3*\pi/12$ for infant carrying probability, and $10*\pi/11$ for tantrum probability. The results of the models with observation date are presented in Table S4.

- 926
- 927
- 928
- 929
- 930
- 931
- 932
- 933

934 TABLES

- 935 Table S1: Different methods used to estimate the dates of births of the 215 baboon infants born936 at Tsaobis between 2005 and 2019.

Critoria used for estimation		N hirthe			
Citteria used for estimation			uncertainty	uncertainty	
	when first seen	estimated	(days)	(days)	
Birth observed in the field	Pink	62	0	0	
Birth occurred during a short	Pink	22	17	1-30	
field break	1 mix		17	100	
Conception date known	/	52	10	10-37	
Infant coloration & mother's	Pink or	56	61	6-151	
reproductive state [1]	transitional	50	01	0-131	
Mother's reproductive state	Grey or	23	67	21-153	
only	unknown	23	07	21-133	
Total	/	215	30	0-153	

940 **Table S2:** Identification of the best birth timing effect for Models 1-2 & 5-7: ΔAIC (Akaike

941 Information Criterion) according to the phase of the sine wave term

942 In order to identify the best birth timing effect on our various indicators of fitness and maternal care, we ran 12 943 different models, with 12 different phases φ for the sine wave term of the birth date (as a fixed effect), for each 944 full model (Models 1-2 & 5-7). If the estimate of the sine term fixed effect is positive, then the birth date maximised 945 is indicated in the 'Date maximised' column and the one minimised is indicated in the 'Date minimised' column. 946 On the contrary, if the estimate of the sine term fixed effect is negative, then the birth date maximised is indicated 947 in the 'Date minimised' column. ΔAIC of each model equals the AIC value of the considered model minus the 948 AIC value of the best model ($\Delta AIC=0$ for the best model, indicated in bold writing). We selected the best phase 949 as the one minimizing the AIC, i.e. for which $\Delta AIC=0$. For example, for Model 2, the best phase is $\varphi = 2 * \pi/12$, 950 and the estimate of the sine term fixed effect is positive (Table S6), indicating that IBIs are maximised in March 951 1st, and minimized in September 1st. Wherever the fixed effect 'birth date' was significant (Model 1, 2 and 7), we 952 considered all phases φ for which $\Delta AIC \le 2$ to define the optimal time window presented in the main text (see 953 greyer background), for instance between August 1st and September 15th for Model 2.

					Δ AIC		
Phase φ	Date maximised	Date minimised	Mortality (Model 1)	IBI (Model 2)	Suckling (Model 5)	Infant carrying (Model 6)	Tantrum (Model 7)
0	1st April	1st October	8.26	2.16	0.67	0	1.15
π/12	15th March	15th September	9.42	0.72	1.13	0.17	0.34
2 * π/12	1st March	1st September	9.45	0	1.62	0.65	0
3*π/12	14th February	15th August	8.12	0.33	1.96	1.29	0.39
4*π/12	1st February	1st August	5.66	1.61	1.92	1.85	1.49
5*π/12	15th January	15th July	2.91	3.36	1.46	2.16	2.94
6*π/12	1st January	1st July	0.84	5.02	0.86	2.18	4.20
7 * π/12	15th December	15th June	0	6.16	0.37	1.93	4.84
8*π/12	1st December	1th June	0.47	6.59	0.09	1.53	4.80
9 * π/12	15th November	15th May	1.99	6.29	0	1.05	4.20

	10 * π/12	1st November	1st May	4.10	5.31	0.08	0.56	3.27
	11 * π/12	15th October	15th April	6.34	3.84	0.32	0.17	2.19
954								
955								

957 Table S3: Birth timing and other predictors of the probability of suckling, infant carrying and

958 tantrums

959 Estimates, confidence intervals, X² statistics and P-values of the predictors of binomial generalized linear mixed 960 models of the probability of suckling (Model 5), infant carrying (Model 6) and tantrums (Model 7), including 961 infant's identity (and focal number for Models 5 and 6) as random effect, and focal observation time as an 'offset' 962 fixed effect for Model 7. These GLMMs are based on 11687 scan observations from 55 infants for Model 5, 924 963 scan observations from 35 infants for Model 6 and 2211 focal observations from 55 infants for Model 7. Significant 964 effects are indicated in bold. For relevant significant effect, we also indicated in the footnote the 95% level 965 confidence interval of the 1000 p-values taking into account birth date uncertainty. Infant birth date is fitted as a 966 sine term with a phase of $9^*\pi/12$ for suckling, 0 for infant carrying, and $2^*\pi/12$ for tantrum probabilities. For 967 categorical predictors, the tested category is indicated between parentheses.

Eined E	ffaat	Estimata	IC		V 2	D volue
FIXEd E	liect	Estimate -	Lower	Upper	Λ^2	P-value
		Model 5: Su	ıckling			
Infant birth date		-0.32	-0.76	0.13	1.97	0.16
Infant sex	(Male)	-0.01	-0.73	0.73	0.00	0.99
Female parity	(Primiparous)	-0.89	-2.20	0.42	1.76	0.18
Female rank		-0.09	-0.42	0.24	0.30	0.59
Infant age		-1.66	-1.97	-1.35	110	<10 ⁻⁴
Carona	(L)	0.18	-0.53	0.90	7 70	0.02
Group	(M)	1.51	0.36	2.66	1.19	0.02
	(2017)	0.21	-1.22	1.63		
Observation year	(2018)	1.73	0.84	2.62	40.45	<10 ⁻⁴
	(2019)	0.03	-0.82	0.88		
	Mo	del 6: Infan	t carrying			
Infant birth date		0.53	-0.18	1.24	2.13	0.14
Infant sex	(Male)	-0.94	-1.60	-0.28	7.78	0.005
Female parity	(Primiparous)	-1.08	-2.11	-0.05	4.21	0.040
Female rank		-0.51	-0.82	-0.19	10.0	0.002
Infant age		-1.94	-2.69	-1.20	26.2	<10-4
Group	(L)	0.01	-0.53	0.55	0.67	0.71
Group	(M)	-0.35	-1.34	0.64	0.07	0.71
	(2017)	-11.5	-262	239		
Observation year	(2018)	1.36	-0.36	3.09	3.69	0.30
	(2019)	0.86	-1.19	2.91		
]	Model 7: Ta	intrum			
Infant birth date		-0.32	-0.62	-0.03	4.53	0.033*
Infant sex	(Male)	-0.10	-0.45	0.25	0.33	0.57

Female parity	(Primiparous)	0.35	-0.32	1.02	1.02	0.31	
Female rank		0.03	-0.14	0.19	0.11	0.74	
T C 4	Age	-41.13	-54.06	-28.21	52 3 0	-10-4	
Infant age	Age ²	-19.07	-28.21	-9.92	53.28	<10-4	
9	(L)	-0.35	-0.68	-0.01	4.00	0.12	
Group	(M)	-0.21	-0.81	0.40	4.23	0.12	
	(2017)	0.12	-1.39	1.63			
Observation year	(2018)	0.30	-0.81	1.41	12.61	0.006	
	(2019)	-0.44	-1.57	0.68			
Offset		0.40	0.10	0.71	6.81	0.009	

* 95% CI: [0.04684 – 0.05234]

972 Table S4: Seasonality and other predictors of the probability of suckling, infant carrying and

973 tantrums

Estimates, confidence intervals, X² statistics and P-values of the predictors of the binomial GLMMs of the probability of suckling (Model 5bis), infant carrying (Model 6bis), and tantrums (Model 7bis). Each model includes infant's identity and year of infant's birth as random effects. Models 5bis and 6bis also included focal observation as random effects, whereas Model 7bis included focal observation duration as an offset fixed effect. Observation date is fitted as a sine term with a phase of $7*\pi/12$ for suckling probability, $3*\pi/12$ for infant carrying probability, and $10*\pi/12$ for tantrum probability. Significant effects are indicated in bold. For categorical predictors, the tested category is indicated between parentheses.

Fixed F	ffaat	Estimata	IC		\mathbf{V}^2	D voluo					
	llect	Estimate	Lower	Upper	Λ^2	P-value					
Model 5bis: Suckling											
Observation date		-1.66	-2.91	-0.40	6.70	0.0096					
Infant sex	(Male)	-0.03	-0.79	0.74	0.00	0.95					
Female parity	(Primiparous)	-0.89	-2.25	0.47	1.63	0.20					
Female rank		-0.11	-0.46	0.23	0.40	0.53					
Infant age		-1.62	-1.94	-1.30	97.47	<10-4					
Croup	(L)	0.30	-0.46	1.06	6 71	0.035					
Group	(M)	1.52	0.33	2.71	0.71	0.035					
	(2017)	-2.34	-4.88	0.20							
Observation year	(2018)	1.74	0.82	2.66	29.99	<10-4					
	(2019)	-2.64	-4.95	-0.33							
Model 6bis: Infant carrying											
Observation date		-1.14	-1.98	-0.30	7.12	0.0076					
Infant sex	(Male)	-0.90	-1.56	-0.24	7.20	0.0073					
Female parity	(Primiparous)	-0.78	-1.77	0.22	2.33	0.13					
Female rank		-0.45	-0.76	-0.15	8.31	0.0039					
Infant age		-2.40	-2.83	-1.97	120	<10-4					
Group	(L)	-0.003	-0.53	0.54	2.04	0.36					
Oloup	(M)	-0.63	-1.62	0.37	2.04	0.30					
	(2017)	-13.10	-2410	2384							
Observation year	(2018)	0.66	-1.19	2.52	12.2	0.007					
	(2019)	-0.73	-2.77	1.31							
Model 7bis: Tantrum											
Observation date		0.63	-0.16	1.42	2.44	0.12					
Infant sex	(Male)	-0.09	-0.45	0.26	0.27	0.60					
Female parity	(Primiparous)	0.07	-0.57	0.72	0.05	0.83					
Female rank		0.04	-0.13	0.21	0.23	0.63					
Infant aga	Age	-33.10	-43.45	-22.75	51.00	~10 /					
Infant age	Age ²	-20.84	-29.87	-11.81	51.09	~10-4					

Offset		0.40	0.10	0.71	6.73	0.0095
	(2019)	0.45	-1.18	2.09		
Observation year	(2018)	0.20	-0.91	1.30	0.45	0.93
	(2017)	0.57	-1.12	2.27		
Group	(M)	-0.05	-0.64	0.53	5.80	0.15
	(L)	-0.32	-0.66	0.02	3 86	0.15

Indicators	Fitnes	tness traits Birth timing			Maternal care		
Model number	1	2	3	4	5	6	7
Response variable	Offspring survival before weaning	Interbirth intervals (days)	Deviation from the offspring survival optimal birth timing	Deviation from the maternal IBI optimal birth timing	Suckling	Infant carrying	Tantrum
Model type	Binomial GLMM	LMM	LMM	LMM	Binomial GLMM	Binomial GLMM	Binomial GLMM
Number of observations	195	120	215	215	5089	924	2221
Number of individuals (juveniles / mothers)	57	43	62	62	55	35	55
Fixed effects	Infant birth date, infant sex, female parity, female rank, group	Infant birth date, infant sex, female parity, female rank, group	Infant sex, female parity, female rank, group	Infant sex, female parity, female rank, group	Infant birth date (or observation date, see Table S4), infant sex, female parity, female rank, infant age, group, observation year	Infant birth date (or observation date, see Table S4), infant sex, female parity, female rank, infant age, group, observation year	Infant birth date, (or observation date, see Table S4), infant sex, female parity, female rank, Infant age ² , group, observation year, focal duration
Random effects	Birth year, female identity	Birth year, female identity	Birth year, female identity	Birth year, female identity	Infant identity, focal number	Infant identity, focal number	Infant identity

Table S5: Summary of the structure of all models included in the study.

987 Table S6: Predictors of offspring mortality before weaning and maternal interbirth interval988 (IBI) duration.

989 Estimates, confidence intervals, X² statistics and P-values of the predictors of a binomial generalized mixed model 990 of the probability of offspring mortality before weaning (0/1: survived/died, Model 1) and a linear mixed model 991 of the duration of the maternal birth interval (IBI) (in days, Model 2), based on 195 observations from 57 females 992 for Model 1 and 120 observations from 43 females for Model 2. Female identity and year of infant's birth are 993 included as random effects in both models. Significant effects are indicated in bold. For the fixed effect 'birth 994 date', we also indicate in the footnote the 95% confidence interval of the average p-value of the simulated models 995 taking into account birth date uncertainty. Infant birth date is fitted using a sine term with a phase of $7*\pi/12$ for 996 infant mortality and of $2^*\pi/12$ for IBI, and. For categorical predictors, the tested category is indicated between 997 parentheses.

Eire d	Fixed Effect		IC	2	\mathbf{V}^2	D
Fixed	Effect	Estimate -	Lower	Upper	Λ^2	P-value
		Model 1: O	ffspring mor	rtality		
Infant birth d	late	-1.12	-1.84	-0.40	9.38	0.002*
Infant sex	(Male)	0.20	-0.76	1.15	0.16	0.685
Female parity	(Primiparous)	-0.83	-2.32	0.67	1.17	0.279
Female rank		-0.44	-0.95	0.07	2.87	0.090
C	(L)	-1.29	-2.39	-0.18	5.25	0.072
Group	(M)	-0.10	-4.08	3.88	5.25	0.072
		Model 2	: Maternal I	BI		
Infant birth d	late	36.84	7.59	66.09	6.10	0.014 †
Infant sex	(Male)	37.04	-0.36	74.45	3.77	0.052
Female parity	(Primiparous)	44.53	-3.24	92.29	3.34	0.068
Female rank		-25.73	-50.68	-0.77	4.08	0.043
Casara	(L)	-50.41	-105.19	4.38	2 21	0 101
Group	(M)	-31.09	-150.57	88.38	5.51	0.191

998 * 95% CI: [0.00967 – 0.01087]

999 † 95% CI: [0.02533 – 0.02821]

1000

1002 FIGURES



Figure S1: Variation in the probabilities of suckling, infant carrying, and tantrums, according
to infant age.



1008 panels, the size of black dots is proportional to the number of observations (see plot legends). The coloured curves 1009 show the predicted fit using a general additive function (method 'gam' of geom smooth function in 'ggplot2' R 1010 package). The darker area around each curve represents the confidence interval of the fitted curve. In order to 1011 determine the best infants' age window for each models (Models 5-7), we explored age-related variation in 1012 suckling (Panel A, Model 5), infant carrying (Panel B, Model 6) and tantrum (Panel C, Model 7) probabilities. We 1013 found that (A) suckling decreases gradually from 2 to 18 months old, before stabilizing to ca. 2% of the scans from 1014 18 to 24 months old, (B) the proportion of infant carrying gradually decreases during the first year of life in our 1015 population as in other baboon populations [6,15,16], and (C) tantrum occurrence started in early-life, peaked when 1016 infants were aged around 6-9 months, and then gradually decreased during the end of their first and second year

1017 of life.

1018





1020 Figure S2: Chacma baboons breed all year round.

1021 Number of conceptions (Panel A, N=241), births (Panel B, N=215) and cycle resumptions (Panel C, N=171) per

1022 month (from 1=January to 12=December) between 2005 and 2019. Births and cycle resumptions do not show

1023 significant seasonality, while conceptions significantly deviate from non-seasonality, with an average conception

1024 date in May. The black arrow length is the value of the Rayleigh statistic R, and its direction is μ . The numbers on

1025 the y-axis of each plot indicate the scale for the number of events on that plot.

1026

1027

1029 REFERENCES

- 1030 1. Dezeure J, Dagorrette J, Baniel A, Carter AJ, Cowlishaw G, Marshall HH, Martina C,
- 1031 Raby CL, Huchard E. 2021 Developmental transitions in body color in chacma baboon
- 1032 infants: Implications to estimate age and developmental pace. Am. J. Phys. Anthropol.
- 1033 **174**, 89–102. (doi:10.1002/ajpa.24118)
- 1034 2. Higham JP, Heistermann M, Ross C, Semple S, MacLarnon A. 2008 The timing of
- 1035 ovulation with respect to sexual swelling detumescence in wild olive baboons.

1036 *Primates* **49**, 295–299. (doi:10.1007/s10329-008-0099-9)

- 1037 3. Lee PC. 1996 The meanings of weaning: growth, lactation, and life history. *Evol*.
- 1038 Anthropol. 5, 87–98. (doi:https://doi.org/10.1002/(SICI)1520-6505(1996)5:3<87::AID-
- 1039 EVAN4>3.0.CO;2-T)
- 1040 4. Langer P. 2008 The phases of maternal investment in eutherian mammals. *Zoology*1041 111, 148–162. (doi:10.1016/j.zool.2007.06.007)
- 1042 5. Altmann J. 1980 Baboon Mothers and Infants. The University of Chicago Press.
- 1043 6. Nicolson N. 1982 Weaning and the Development of Independence in Olive Baboons.
- 1044 Harvard University, Cambridge, Massachusetts. (doi:10.13140/RG.2.1.4000.9761)
- 1045 7. Rhine RJ, Norton GW, Wynn GM, Wynn RD. 1989 Plant feeding of yellow baboons
- 1046 (Papio cynocephalus) in Mikumi national park, Tanzania, and the relationship between
- seasonal feeding and immature survival. *Int. J. Primatol.* **10**, 319–342.
- 1048 (doi:10.1007/BF02737420)
- 1049 8. Barrett L, Henzi SP, Lycett JE. 2006 Whose Life Is It Anyway? Maternal Investment,
- 1050 Developmental Trajectories, and Life History Strategies in Baboons. In *Reproduction*
- 1051 and Fitness in Baboons: Behavioral, Ecological, and Life History Perspectives, pp.
- 1052 199–224.
- 1053 9. Bădescu I. 2018 The attainment of independence from the mother in primate infants

- and its implications for the evolution of cooperative breeding in hominins. In *The*
- 1055 *talking species : perspectives on the evolutionary, neuronal and cultural foundations of*
- 1056 *language* (eds EM Luef, MM Marin), pp. 169–194. Uni-Press Graz.
- 1057 10. Borries C, Lu A, Ossi-Lupo K, Larney E, Koenig A. 2014 The meaning of weaning in
- 1058 wild Phayre's leaf monkeys: Last nipple contact, survival, and independence. Am. J.
- 1059 Phys. Anthropol. 154, 291–301. (doi:10.1002/ajpa.22511)
- 1060 11. Reitsema LJ. 2012 Introducing Fecal Stable Isotope Analysis in Primate Weaning
 1061 Studies. *Am. J. Primatol.* 74, 926–939. (doi:10.1002/ajp.22045)
- 1062 12. Bădescu I, Watts DP, Katzenberg MA, Sellen DW. 2016 Alloparenting is associated
- 1063 with reduced maternal lactation effort and faster weaning in wild chimpanzees. *R. Soc.*
- 1064 *Open Sci.* **3**. (doi:10.1098/rsos.160577)
- 1065 13. Crowley BE, Reitsema LJ, Oelze VM, Sponheimer M. 2015 Advances in Primate
- 1066 Stable Isotope Ecology Achievements and Future Prospects. Am. J. Primatol. 78,
- 1067 995–1003. (doi:10.1002/ajp.22510)
- 1068 14. Reitsema LJ, Partrick KA, Muir AB. 2016 Inter-individual variation in weaning among
- 1069 rhesus macaques (Macaca mulatta): Serum stable isotope indicators of suckling
- 1070 duration and lactation. Am. J. Primatol. 78, 1113–1134. (doi:10.1002/ajp.22456)
- 1071 15. Altmann J, Samuels A. 1992 Costs of maternal care: infant-carrying in baboons. *Behav.*1072 *Ecol. Sociobiol.* 29, 391–398. (doi:10.1007/BF00170168)
- 1073 16. Rhine RJ, Norton GW, Westlund BJ. 1984 The Waning of Dependence in Infant Free-
- 1074 Ranging Yellow Baboons (Papio cynocephalus) of Mikumi National Park. Am. J.
- 1075 *Primatol.* 7, 213–228.
- 1076
- 1077