



Cram, D. L., Monaghan, P., Gillespie, R. and Clutton-Brock, T. (2017) Effects of early-life competition and maternal nutrition on telomere lengths in wild meerkats. *Proceedings of the Royal Society of London Series B: Biological Sciences*, 284(1861), 20171383. (doi:[10.1098/rspb.2017.1383](https://doi.org/10.1098/rspb.2017.1383))

This is the author's final accepted version.

There may be differences between this version and the published version. You are advised to consult the publisher's version if you wish to cite from it.

<http://eprints.gla.ac.uk/147697/>

Deposited on: 20 September 2017

Enlighten – Research publications by members of the University of Glasgow  
<http://eprints.gla.ac.uk>

1           **Effects of early-life competition and maternal nutrition on**  
2                           **telomere lengths in wild meerkats**

3  
4           Dominic L. Cram<sup>1,2\*</sup>, Pat Monaghan<sup>3</sup>, Robert Gillespie<sup>3</sup> & Tim Clutton-Brock<sup>1,2</sup>

5  
6           <sup>1</sup> Department of Zoology, University of Cambridge, Downing Street, Cambridge CB2 3EJ,  
7           United Kingdom

8  
9           <sup>2</sup> Kalahari Meerkat Project, Kuruman River Reserve, P.O. Box 64, Van Zylsrus, Northern  
10          Cape 8467, South Africa

11  
12          <sup>3</sup> Institute of Biodiversity, Animal Health and Comparative Medicine, College of Medical,  
13          Veterinary and Life Sciences, University of Glasgow, Graham Kerr Building, Glasgow G12  
14          8QQ, United Kingdom

15  
16          \*Author for correspondence: dom.cram@gmail.com  
17

---

18  
19

20 **Abstract**

21 Early-life adversity can affect health, survival and fitness later in life, and recent evidence  
22 suggests that telomere attrition may link early conditions with their delayed consequences.  
23 Here, we investigate the link between early-life competition and telomere length in wild  
24 meerkats. Our results show that, when multiple females breed concurrently, increases in the  
25 number of pups in the group are associated with shorter telomeres in pups. Given that pups  
26 from different litters compete for access to milk, we tested whether this effect is due to  
27 nutritional constraints on maternal milk production, by experimentally supplementing  
28 females' diets during gestation and lactation. While control pups facing high competition had  
29 shorter telomeres, the negative effects of pup number on telomere lengths were absent when  
30 maternal nutrition was experimentally improved. Shortened pup telomeres were associated  
31 with reduced survival to adulthood, suggesting that early-life competition for nutrition has  
32 detrimental fitness consequences that are reflected in telomere lengths. Dominant females  
33 commonly kill pups born to subordinates, thereby reducing competition and increasing  
34 growth rates of their own pups. Our work suggests an additional benefit of infanticide may be  
35 that it also reduces telomere shortening caused by competition for resources, with associated  
36 benefits for offspring ageing profiles and longevity.

37

38 **KEY WORDS**

39 telomeres, early-life adversity, early-life stress, *Suricata suricatta*, meerkats, infanticide

40

41

42 **Introduction**

43 The early period of an animal's life can have a disproportionately influential role in  
44 determining health, survival and reproductive success later in life, even though it accounts for  
45 a relatively minor proportion of total lifespan [1]. Despite the importance of the early-life  
46 environment, our understanding of the physiological mechanisms underpinning its lasting and  
47 delayed consequences remains poor [2].

48

49 Telomere loss has recently been proposed as a potential molecular mechanism linking early-  
50 life adversity with later-life performance and ageing [3]. Telomeres are non-coding  
51 sequences at the ends of eukaryotic chromosomes that play a critical role in protecting  
52 genome integrity [4]. Telomeres shorten with each cell division, and this shortening is  
53 accelerated during early development and by stressors including oxidative damage and stress  
54 hormone exposure [5-7 but see 8]. When telomeres shorten beyond a critical point, the cell  
55 enters replicative senescence, and accumulation of senescent cells can impair tissue function  
56 in later life when cell renewal capacity is reduced [9]. A number of studies have shown that  
57 telomere length or rate of loss predicts survival and longevity in vertebrates [10-12] including  
58 humans [reviewed in 13], and short telomeres are associated with the systemic loss of  
59 function frequently observed in ageing individuals [14]. Accelerated telomere loss early in  
60 life may therefore advance the onset of senescence, thereby linking early-life conditions with  
61 later-life health and survival.

62

63 Early-life adversity promotes telomere shortening in a range of species, including salmon,  
64 humans and several birds [15-19]. In birds, offspring competing with more rivals, or rivals  
65 higher in the competitive hierarchy, exhibit accelerated telomere loss [20-25]. Studies of the  
66 consequences of offspring competition on telomere dynamics have thus far focussed almost  
67 exclusively on biparental species; the importance of early-life competition in species with  
68 other social systems therefore remains unclear.

69

70 In animal societies where multiple females breed the effects of early-life competition on  
71 telomere lengths are likely to be particularly pronounced, because the number of competing  
72 offspring is expected to be higher and competition therefore more intense. Where females  
73 breed asynchronously, greater age asymmetries between offspring will likely further  
74 exacerbate telomere loss for offspring that are younger or lower in the competitive hierarchy  
75 [23, 25]. Alternatively, sharing offspring care between multiple females may buffer offspring

76 against unpredictable environments [26] and improve growth and health [27], thus relaxing  
77 competition and slowing telomere attrition. Whether the effects of early-life competition on  
78 the rate of telomere attrition in animal societies are exacerbated by increased offspring  
79 number, or mitigated by cooperative care of young, remains unknown.

80

81 Where early-life adversity promotes the accumulation of ageing-related damage and poor  
82 telomere integrity, we would predict that selection would favour parental strategies that  
83 protect offspring, either by improving the environment or enhancing offspring resilience to  
84 adversity. Despite extensive evidence that early-life adversity is reflected in enduring  
85 deleterious effects on telomere lengths [15-20], and that short telomeres are linked with poor  
86 health and curtailed survival [12, 14, 28, 29], little is known about parental strategies  
87 associated with slowed offspring telomere attrition, and how effective they are [30].

88

89 Here, we investigate whether early-life adversity, in the form of intense pup competition, is  
90 associated with shortened telomeres in wild Kalahari meerkat pups at emergence from the  
91 natal burrow. Meerkats (*Suricata suricatta*) live in stable cooperatively breeding groups of up  
92 to 50 individuals. Reproduction is largely monopolised by a single dominant female, but  
93 older subordinate females also attempt to breed at a lower frequency [31]. Mean litter size is  
94 4.1 pups (range 1-8) [32]. Mixed litters are suckled indiscriminately by all lactating females  
95 [32], and pups therefore compete both with their littermates and with pups from other litters.  
96 Previous research suggests that pups compete for access to milk before emerging from the  
97 natal burrow, as experimental contraception of subordinate females leads to increased growth  
98 of the dominant's pups at emergence from the birth burrow [33]. Pups are also frequently  
99 observed aggressively competing for access to provisioning helpers after emergence [34].  
100 After investigating whether variation in the number of competing pups affects their telomere  
101 lengths, we test whether supplementing the mother's food intake during gestation and  
102 lactation mitigates the effects of competition on pup telomeres. We then investigate whether  
103 early-life telomere lengths predict survival into adulthood. Finally, we explore the extent to  
104 which mothers reduce pup competition by killing litters born to other females, and discuss  
105 how this strategy might impact telomere dynamics in her own pups. Such infanticide is  
106 common in meerkat groups, and is almost always perpetrated by heavily pregnant females  
107 [35].

108

109

## 110 **Methods**

### 111 *Study population*

112 Data collection was conducted in the context of a long-term study, monitoring a naturally  
113 regulated population of wild meerkats at the Kuruman River Reserve, South Africa (26° 58'S,  
114 21° 49'E), between 1994 and 2015. All meerkats were habituated to close observation (<1m)  
115 and individually recognizable using small dye-marks (ca. 2cm<sup>2</sup>, for adults and older pups) or  
116 trimming small patches of fur (ca. 0.5cm<sup>2</sup>, for newly-emerged pups) [36]. Virtually all  
117 (>95%) meerkats could be voluntarily weighed on electronic scales ( $\pm$  0.1g, Durascale, UK)  
118 before they commenced foraging in the morning, at midday and after sunset. Groups were  
119 visited 2-3 times per week to collect behavioural, life-history and body weight data.  
120 Observations of pregnancy, birth, infanticide, dominance, group size and rainfall were made  
121 using protocols detailed elsewhere [36, 37]. Mother and father identity were assigned  
122 genetically [38, 39].

123

### 124 *Pup tail tip sampling*

125 Meerkats are born in an underground burrow, emerging for the first time at age 3-4 weeks.  
126 Shortly after the litter's first emergence, a small biopsy of skin from the tail-tip was collected  
127 from each pup (age  $28.3 \pm 3.4$  days) for the determination of telomere length and parentage  
128 [39]. Skin samples were immediately transferred to 96% ethanol and stored at -20°C until  
129 DNA extraction.

130

### 131 *Supplementary feeding experiment*

132 To investigate the effects of early nutritional environment on telomere lengths, we fed  
133 pregnant females during gestation and lactation. In order to minimize inter-individual  
134 differences in body condition, our experimental procedure was limited to dominant females.  
135 The supplementary feeding protocol consisted of one hard-boiled egg per day (divided  
136 equally between the morning and afternoon observation sessions) commencing six weeks  
137 after the end of a dominant female's pregnancy, and continuing until the next parturition [40].  
138 Thereafter, fed dominant females received four eggs per week, until the pups were weaned.  
139 This feeding protocol occurred between August and November in 2011 and 2012. Control  
140 females were pregnant during the same period, and did not receive supplemental food.

141

142 *Observations of Infanticide*

143 We investigated how infanticide by dominant females affects the number of competing pups,  
144 and the likely consequences for telomere lengths in her own litter. While previous analyses of  
145 the distribution of infanticide have focussed on consequences for the victim mother (i.e.  
146 whether her litter survives or is killed [35, 37]), we quantified the *benefits* of infanticide for  
147 the perpetrator (i.e. how many competitor pups she removes). We identified periods when the  
148 dominant female is most likely to kill pups born to other females (the 30 days prior to her  
149 own parturition, hereafter termed ‘high infanticide period’) and least likely (the 30 days  
150 immediately after giving birth, hereafter termed ‘low infanticide period’) [27]. We then  
151 assessed subordinate litter survival probabilities and the total number of subordinate pups  
152 surviving to emergence during these two periods. Parturition for all females could be  
153 identified by sudden weight loss and change in body shape [36], and pup production for each  
154 period was measured as the number of pups born that survived to emergence from the birth  
155 burrow.

156

157 *qPCR determination of telomere lengths*

158 We used quantitative PCR (qPCR) analysis to measure telomere length in skin samples,  
159 based on published protocols with some modifications [41, 42]. This measure represents the  
160 average telomere length across cells in a sample, and is reported as the level of telomeric  
161 sequence abundance relative to a reference non-variable copy number gene (T/S ratio).  
162 Further details of DNA extraction and qPCR analysis can be found in the supplementary  
163 methods.

164

165 *Statistical analysis*

166 Statistical analyses were carried out in R version 3.2.3, using a step-wise model  
167 simplification approach [43, 44]. Initially all fixed terms of interest were fitted, followed by  
168 the stepwise removal of terms whose removal from the model resulted in a non-significant  
169 change in deviance (using maximum log-likelihood estimation), until the minimal adequate  
170 model (MAM) was obtained, in which only significant terms remained. Dropped terms were  
171 then added back in to the MAM to confirm their non-significance. The homoscedasticity and  
172 normality of residuals were confirmed by visual inspection, and all continuous predictors  
173 were scaled to a mean of 0 and standard deviation of 1. The significance of all terms was  
174 tested either by removing the terms from the MAM (if the term was in the MAM) or adding  
175 the terms to the MAM (if the term was not included in the MAM). Analysis using Akaike's

176 information criterion correcting for small sample size (AICc) and inspection of the top model  
177 set (for which AICc differed by  $< 2$ ) yielded qualitatively identical results [45]. We ran three  
178 sets of statistical models, first to investigate the determinants of pup telomere lengths in the  
179 large correlative dataset, second to investigate how experimental supplementary feeding of  
180 mothers impacted pup telomere lengths, and third to investigate the consequences of  
181 infanticide for pup competition.

182

### 183 1) What are the determinants of pup telomere lengths?

184 Our primary interest was the effect of the number of competing pups on telomere lengths at  
185 emergence from the natal burrow. For each sampled pup, we assessed the number of rival  
186 pups (aged under 90 days) present in the group, every day between the focal pup's birth and  
187 day of sampling for telomere length. The average of these daily rival counts represents our  
188 measure of overall competition experienced by the focal pup prior to sampling, hereafter  
189 termed 'pup number'. This estimate of pup competition includes littermates and pups from  
190 older and younger litters born to the dominant female and subordinate females.

191 We controlled for maternal factors that may influence offspring quality, including weight at  
192 conception, age (mean 4.9 years, range 1.2 - 8.0) and dominance status (dominant or  
193 subordinate) [46]. Social group size (average number of adult group members calculated as  
194 above for pup number) and rainfall (mm) in the month before birth can also both influence  
195 offspring quality [47]. Pup sex (male, female or unknown) and age at capture were also  
196 controlled for. We included these individual, maternal, environmental and social predictors,  
197 with our estimate of pup number, in a general linear mixed effects model (GLMM), with pup  
198 telomere length as the response. Cohort year, group ID, mother ID and litter ID were  
199 included as random terms, to account for the non-independence of pups within years, groups,  
200 mothers, and litters. Telomere lengths were available for 230 pups from 63 litters in 13  
201 groups, born between 2009 and 2012. We also tested the effect of paternal age (mean 4.1  
202 years, range 1.4 - 6.1) on pup telomere lengths in a reduced dataset for which the father's  
203 date of birth could be accurately determined (78 pups from 23 litters in 7 groups).

204

### 205 2) Does an experimentally improved nutritional environment mitigate the effects of pup 206 number?

207 To test the effect of supplementary feeding of the pregnant and lactating mother on pup  
208 telomere lengths, we included experimental treatment (fed/control) as a two-level factor in a  
209 GLMM, with pup telomere length as the response and litter ID as the random term. Given our

210 smaller sample size for the experimental dataset, only terms found to be significant in the  
211 larger correlative model were included, and two-way interactions between these and  
212 treatment. Telomere lengths were available for 25 pups from 8 litters in each treatment.

213

### 214 3) Do pup telomere lengths predict survival to adulthood?

215

216 We investigated whether pup telomere lengths predicted survival to adulthood (1 year old).  
217 Sub-adult meerkats do not disperse [31, 48], and any disappearance from the group before  
218 reaching adulthood is therefore likely to reflect mortality. We removed any individuals dying  
219 before reaching nutritional independence (90 days), as death at this early stage typically  
220 occurs due to starvation, predation, or becoming separated from the group, these sources of  
221 mortality are unlikely to reflect variation in telomere lengths. We used a binary term for  
222 survival to adulthood as the response in a binomial mixed effects model. We included pup  
223 telomere length as a predictor. We also controlled for other predictors known to influence  
224 telomere lengths and survival in young meerkats: sex, group size, rainfall, maternal  
225 dominance status, maternal age [47]. We controlled for the effects of pup body weight on  
226 survival, by including their bodyweight at age 40 days in the model. Group ID, mother ID  
227 and litter ID were included as random terms. This model was fitted to a dataset of 178  
228 individuals: 161 pups from 51 litters born to dominant females, and 17 pups from 7 litters  
229 born to subordinates. The maximum confirmed lifespan for meerkats in our population is  
230 12.2 and 12.4 years, for males and females respectively.

231

### 232 4) How does infanticide affect the number of competing pups?

233

234 We contrasted the fates of subordinate litters born in periods of high and low dominant  
235 female infanticide. First, for each dominant female parturition ( $n = 158$ ), we counted  
236 subordinate parturitions during the two periods (30 days before and after dominant  
237 parturition). Infanticide typically takes places shortly after birth, so we classed each  
238 subordinate parturition as a 'success' or 'infanticide' according to whether the litter survived  
239 its first two days (litter loss after this point is more likely to be due to starvation or predation  
240 [35, 37]). Although new-born litters remained in the burrow for up to four weeks, their  
241 survival could be recorded daily by observing whether the group continued to leave baby-  
242 sitters during foraging trips [35]. The number of successes and infanticides were then used as  
243 the response term in a binomial mixed effects model, with the high/low infanticide period

244 fitted as a two-level predictor. The random terms were dominant female pregnancy ID,  
245 dominant female ID and group ID. Second, for each dominant female parturition, we  
246 calculated the *total number* of emerging subordinate pups born during the two infanticide  
247 periods, and fitted this as the response term in a GLMM with a Poisson distribution. The  
248 main predictor of interest was the two-level high/low dominant female infanticide period, and  
249 we controlled for the number of subordinate females giving birth.

250

251 **Results**

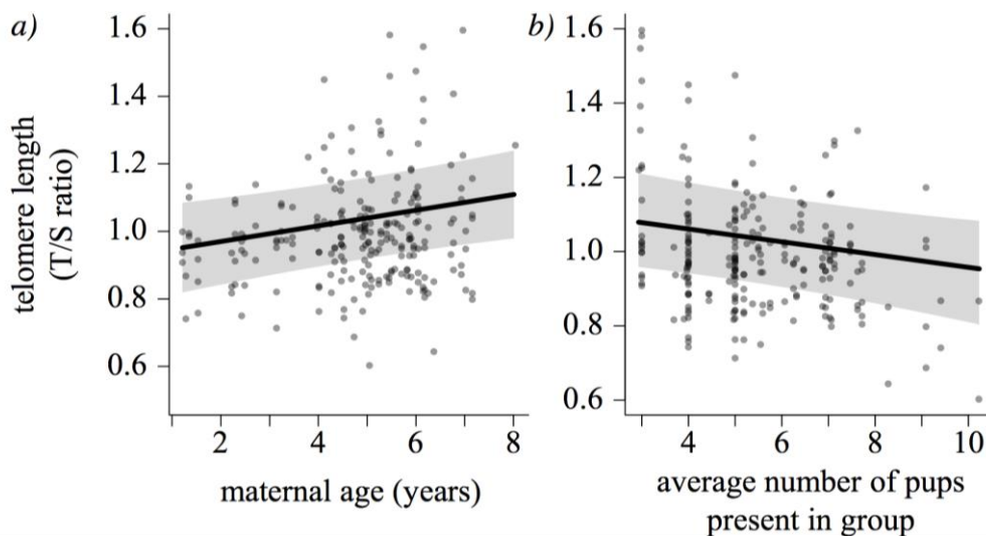
252 1) What are the determinants of pup telomere lengths?

253 Male and female pups had similar telomere lengths at four weeks ( $\chi^2_2 = 1.14$ ,  $p = 0.47$ ,  
254 supplementary table 1). Pup telomere lengths were not associated with their mother's  
255 dominance status, the number of helpers in the group, the pup's age or the amount of rainfall  
256 in the month before their birth (all  $\chi^2_1 < 1.53$ ,  $p > 0.23$ ). Older mothers produced pups with  
257 significantly longer telomeres ( $\chi^2_1 = 9.57$ ,  $p = 0.002$ , Fig 1 a). There was a trend for lighter  
258 mothers to produce pups with slightly longer telomeres, but this was not statistically  
259 significant ( $\chi^2_1 = 3.56$ ,  $p = 0.06$ ). In contrast to the positive effect of maternal age, in a  
260 reduced dataset for which father age was known, older fathers tended to produce pups with  
261 shorter telomeres, although this association was not significant ( $\chi^2_1 = 3.28$ ,  $p = 0.07$ ).

262

263 Controlling for the effect of mother's age, pup telomeres were significantly shorter when the  
264 number of competing pups was high ( $\chi^2_1 = 5.55$ ,  $p = 0.018$ , Fig 1b): telomeres were 13.3%  
265 longer when pup number was lowest compared to highest.

266



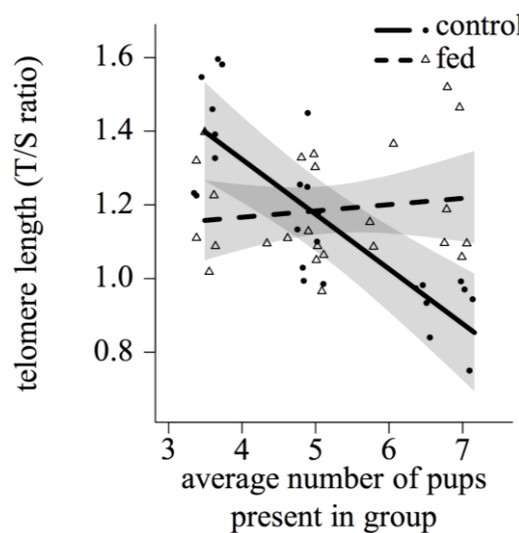
267

268 **Figure 1 a)** the positive association between maternal age and pup telomere length at  
269 emergence from the natal burrow. The line represents the model predictions from a GLMM,  
270 with an average pup number of 5.43. **b)** The negative association between the number of  
271 competitors a pup encounters in the first weeks of life, and its telomere length at emergence  
272 from the natal burrow. The line represents the model predictions from a GLMM, with an  
273 average maternal age of 4.86 years. In both figures, the points represent raw data, which are  
274 translucent for clarity. Shaded areas represent the 95% confidence intervals of each model  
275 prediction.

276  
277  
278  
279  
280  
281  
282  
283  
284  
285  
286  
287  
288

2) Does an experimental feeding of mothers mitigate the effects of pup number?

The effect of maternal supplementary feeding on pup telomere lengths was evident as a significant interaction between experimental treatment and pup number ( $\chi^2_1 = 16.47$ ,  $p < 0.001$ , Fig 2, Supplementary table 2). While control pups had shorter telomeres under greater pup competition, no similar pattern was observed in pups from fed mothers. In contrast to our larger, correlative dataset, in our restricted experimental dataset, pup telomere lengths were not significantly affected by maternal age, either as a single term or in the interaction with treatment (both  $\chi^2 < 2.23$ ,  $p > 0.14$ ). Retention of the non-significant maternal age in the model did not qualitatively change the results.



289  
290  
291  
292  
293  
294  
295  
296  
297  
298  
299  
300

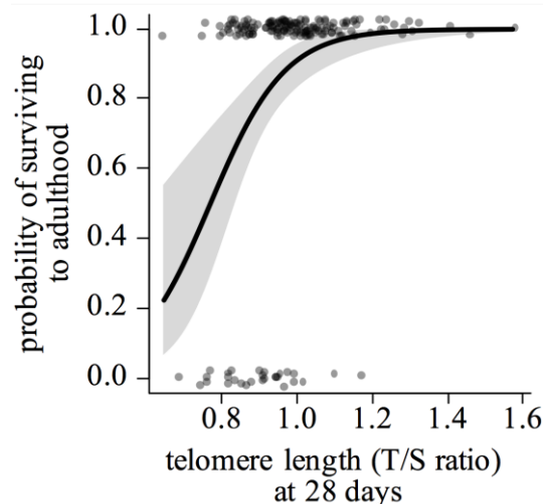
**Figure 2** the effect of experimental maternal feeding (during gestation and lactation) on pup telomere lengths is dependent on the number of competitor pups. In control litters (filled points and solid line) there is a negative relationship between the number of pups and telomere lengths, while in litters from mothers receiving supplementary feeding (open triangles and dashed line) this negative association disappears. Lines represent model predictions for a mean maternal age of 4.5 years, from a GLMM with telomere length as the response, and maternal age, experimental treatment, and the interaction between treatment and number of pups. Shaded areas represent the model's 95% confidence intervals for each model prediction line. Points represent raw data, and are jittered on the x-axis for clarity.

301

302 3) Do pup telomere lengths predict survival to adulthood?

303 A pup's probability of survival to adulthood was positively predicted by its weight ( $\chi^2_1=$   
304 16.24,  $p < 0.001$ , Supplementary table 3) and its mother's age ( $\chi^2_1= 4.88$ ,  $p = 0.027$ ). In this  
305 dataset, pups born to dominant females were less likely to survive to adulthood compared to  
306 those born to subordinates ( $\chi^2_1= 14.03$ ,  $p < 0.001$ ), however this may be driven by poor data  
307 availability for subordinates: only 17 pups (9% of this dataset) were born to subordinates.  
308 Pups were less likely to survive in larger groups ( $\chi^2_1= 4.15$ ,  $p = 0.042$ ). Controlling for these  
309 significant effects, pups with longer telomeres were significantly more likely to survive to  
310 adulthood ( $\chi^2_1= 17.93$ ,  $p < 0.001$ , Figure 3). Survival to adulthood was not significantly  
311 predicted by pup sex or rainfall (both  $\chi^2_1 < 0.82$ ,  $p > 0.36$ ).

312



313

314 **Figure 3** The positive association between pup telomere length and survival to adulthood.  
315 The line represents the model predictions from a GLMM, for a pup with a dominant mother  
316 and all other significant predictors at their mean (pup weight: 230g , maternal age: 4.8 years,  
317 group size: 19.2). The points represent raw data (jittered on the y-axis for clarity) and the  
318 shaded areas are 95% confidence intervals of the model predictions.

319

320 4) How does infanticide affect pup competition?

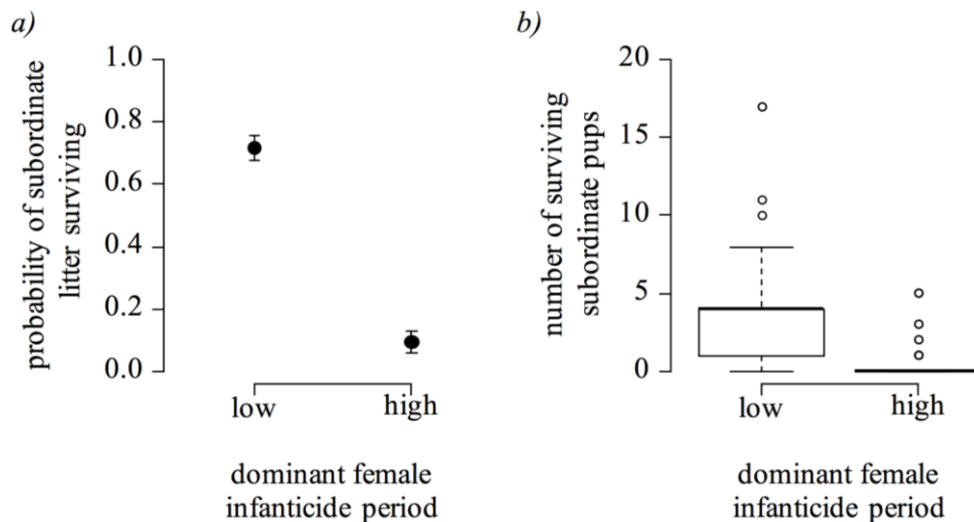
321

322 Pregnant dominant females commonly kill pups born to subordinate females shortly after  
323 they are born [35]. The probability of a subordinate litter surviving its first two days was  
324 9.4% if it was born during the high dominant female infanticide period (the 30 days before  
325 dominant female parturition), compared to 71.6% during the low dominant female infanticide

326 period (the 30 days after dominant female parturition) ( $\chi^2_1= 118.57$ ,  $p < 0.001$ , Fig. 4 a,  
327 Supplementary table 4i).

328

329 After controlling for the number of subordinate females giving birth ( $\chi^2_1= 41.38$ ,  $p < 0.0001$ ,  
330 Supplementary table 4ii), infanticide by the pregnant dominant female significantly reduced  
331 the number of subordinate pups surviving to emergence ( $\chi^2_1= 107.36$ ,  $p < 0.0001$ , Fig 4 b),  
332 and the number of surviving non-littermates that pups born to the dominant female had to  
333 compete with fell from a median value of 4 to zero. This suggests that the infanticidal  
334 behaviour of dominant females leads to a substantial reduction in pup competition for her  
335 own litter. From our analysis of the effects of pup number on telomere lengths, we estimate  
336 that the removal of 4 rival pups is likely to be associated with a 7.3% increase in telomere  
337 lengths for the dominant female's pups.



338

339 **Figure 4 a)** the effect of dominant female infanticidal behaviour on the probability of  
340 subordinate litters surviving their first two days of life. Points represent predicted means and  
341 standard errors from a binomial mixed model with dominant female infanticide risk as the  
342 only predictor. **b)** The effects of infanticide by dominant females on the total number of  
343 surviving pups produced by all subordinate females in their group.

344

### 345 Discussion

346 Our results show that early-life competition is associated with shortened telomeres in wild  
347 meerkats, and that this effect is evident by the time pups leave the natal burrow at  
348 approximately 28 days old. The detrimental effect of competition on pup telomere lengths  
349 disappears when mothers are given supplementary food during gestation and lactation,  
350 suggesting that the reduction in telomere lengths in pups facing more rivals is a consequence

351 of competition for food. Pups with short telomeres show a lower probability of survival to  
352 adulthood. Dominant females can reduce pup competition through killing other pups born in  
353 their group at times while they are pregnant, with likely benefits for their own pups' telomere  
354 lengths and survival.

355

356 Our findings from a wild, social mammal extend evidence that in several biparental bird  
357 species, offspring competing with more rivals, or rivals higher in the competitive hierarchy,  
358 exhibit accelerated telomere loss [20-25]. There are several non-mutually exclusive  
359 explanations for the association between high numbers of competing offspring and shortened  
360 telomeres in meerkats. Under greater competition, offspring typically expend more energy  
361 gaining access to food (either through elevated begging, or competing to access food [49-  
362 51]). Early-life adversity can also trigger physiological stress mechanisms, which confer  
363 short-term benefits, but are costly in the long-term, as they allocate resources to immediate  
364 survival at the expense of somatic maintenance [52]. Greater competition can also promote  
365 accelerated growth profiles, whereby offspring grow faster than their optimal developmental  
366 rate in order to out-compete rivals [53]. Aggressively competing to access food,  
367 physiological stress cascades and accelerated growth can all lead to elevated oxidative  
368 damage and accelerated telomere attrition [54-58 but see 8].

369

370 Our experimental results show that high numbers of competing pups are no longer associated  
371 with reduced telomere lengths when maternal food intake is increased during gestation and  
372 lactation. Previous evidence suggests that experimental food supplementation of weaned  
373 meerkat pups reduces aggressive pup competition [59], and that a relaxed early-life  
374 competitive environment slows telomere shortening in other species [23]. Increased maternal  
375 weight during gestation is positively associated with meerkat pup weight at weaning [46],  
376 suggesting that heavier females are better able to provision their young, leading to reduced  
377 early-life competition. Pups from experimentally fed mothers may therefore exhibit longer  
378 telomeres due to a relaxed early competitive environment, arising from improved pup quality  
379 at birth (thus enhancing pup competitive ability), or elevated maternal milk yield or  
380 micronutrient content (thus reducing the pups' need to compete for food) [60].

381

382 Our finding that variation in offspring telomere lengths is associated with maternal and  
383 paternal age should be interpreted with caution. Pups born to older mothers had longer  
384 telomeres, and while a female could provide better care for offspring as she grows older, it is

385 equally possible that early mortality of poor-quality females leads to disproportionate  
386 numbers of high-quality females in older cohorts [61, 62]. The positive effect of maternal age  
387 on pup telomere lengths may therefore be due to selective disappearance, rather than within-  
388 female change. Furthermore, maternal age at conception is unrelated to offspring telomere  
389 length in other mammals [63]; whether subsequent litters of pups have longer telomeres as  
390 the mother grows older is therefore unclear. We also find that paternal age has a weak  
391 negative effect on pup telomeres lengths. This result is surprising, given that paternal age at  
392 conception is typically positively associated with offspring telomeres (e.g. in humans [63]  
393 and chimpanzees (*Pan troglodytes*) [64]). It is possible that older male meerkats lose  
394 condition faster than humans or chimpanzees, with concomitant decreases in sperm and pup  
395 telomere lengths, but further work would be needed to clarify the role of paternal age at  
396 conception in meerkat telomere dynamics.

397

398 Shortened pup telomeres following early-life competition may be associated with significant  
399 fitness costs, given our finding that reduced telomere lengths predicted low survival to  
400 adulthood. Short telomeres and rapid telomere attrition are associated with reduced survival  
401 and curtailed longevity in a number of species, both in captivity and in the wild [10-12, 28].  
402 Given that we found short telomeres were linked to reduced survival during meerkats' first  
403 year of life, this likely does not reflect accelerated senescence, as senescence is typically only  
404 manifest after meerkats reach three years old [65]. Similarly, early-life telomere dynamics are  
405 linked with survival during the first years of life in other wild mammals [29], suggesting that  
406 telomeres are not only linked with ageing-related mortality, but provide an integrative bio-  
407 marker of somatic damage which can be associated with mortality at any age [66]. Whether  
408 telomere dynamics in adult meerkats are predictive of age-related mortality requires further  
409 investigation.

410

411 Our results suggest that infanticide by dominant females leads to marked reductions in the  
412 number of competitors faced by their own litters. Previous evidence suggests that  
413 experimental reductions of pup number, either by temporary pup removal or contraception of  
414 subordinate females, leads to increased weight gain in the remaining pups [33, 67]. Heavier  
415 pups are subsequently more likely to survive to adulthood and acquire dominance [67, 68],  
416 suggesting that this accelerated growth does not exceed the optimal growth rate, and  
417 therefore confers little costs. By eliminating rival offspring, dominant females are therefore

418 likely to improve the condition, survival and probability of dominance acquisition of their  
419 own litters.

420

421 Our findings highlight a further potential benefit of infanticide: removal of competitor pups  
422 may be associated with a significant increase in pup telomere lengths. Longer telomeres are  
423 associated with improved early-life survival in meerkats, and later-life benefits including  
424 delayed senescence and improved longevity in a number of other species [10-12, 14]. Such  
425 later-life benefits may be particularly important in meerkats: in dominants (who monopolise  
426 reproduction), the primary determinant of lifetime reproductive success is dominance tenure  
427 length [69]. Dominants of both sexes exhibit senescence and their tenure typically ends when  
428 they are unable to repel same-sex challengers [36, 39, 65, 70]. In addition to the above  
429 benefits for offspring condition and dominance acquisition, infanticide may therefore allow  
430 dominant females to improve pup telomere lengths, thus delaying their onset of senescence,  
431 extending their dominance tenures and increasing their lifetime reproductive success. While  
432 the level and type of parental care has been shown to influence offspring telomere lengths in  
433 humans and captive rhesus monkeys [30, 71, 72], to our knowledge this is the first evidence  
434 that a specific maternal strategy (killing competitor pups) has associated benefits for  
435 offspring telomere lengths.

436

### 437 **Conclusion**

438 Our results suggest that in a social species, where offspring competition may be particularly  
439 pronounced, an unfavourable early-life competitive environment accelerates telomere loss  
440 under natural conditions, with potentially lifelong consequences [12]. Despite the observed  
441 enduring detrimental effects of early-life adversity on telomere dynamics [20-25, 28, 58], and  
442 the clear selection pressure this places on parents, few studies have investigated whether  
443 parents are able to protect offspring telomeres by improving the early environment. In  
444 meerkats, dominant females kill rival litters to reduce competition for their own pups,  
445 resulting in improved pup condition and likely benefits for telomere lengths and longevity.  
446 Overall, our results highlight that both the early environment and protective parental  
447 strategies may affect offspring telomere lengths, and without detailed consideration of both  
448 we are likely to underestimate the role of telomere dynamics in shaping life-histories, ageing  
449 profiles and fitness.

450 **Ethics statement**

451 Our work was approved by the Animal Ethics Committee of the University of Pretoria, South  
452 Africa (no. EC010-13) and by the Northern Cape Department of Environment and Nature  
453 Conservation, South Africa (FAUNA 1020/2016).

454

455 **Data accessibility**

456 All data used in analyses and figures are included in the electronic supplementary material.

457

458 **Competing interests**

459 We declare we have no competing interests.

460

461 **Author contributions**

462 This study was designed by D.L.C. and T.H.C.-B.; P.M and R.G. planned and implemented  
463 the laboratory analyses and advised on interpretation of telomere data. D.L.C. planned and  
464 implemented the statistical analyses; D.L.C. and T.H.C.-B. wrote the paper, with extensive  
465 advice from P.M. and R.G.. All authors contributed to the manuscript, approved the final  
466 version and are accountable for the work.

467

468 **Funding**

469 The Kalahari Meerkat Project is supported by the Universities of Cambridge, Zurich and  
470 Pretoria. Components of this research were supported by the Natural Environment Research  
471 Council (grant no. NE/G006822/1) and the European Research Council (grant no. 294494).

472

473 **Acknowledgements**

474 We thank the Kalahari Research Trust for permission to work at the Kuruman River Reserve,  
475 and the neighbouring farmers for the use their land. Many thanks to Marta Manser, Dave  
476 Gaynor and Tim Vink for the organization of the field site and database. We are grateful to  
477 all the volunteers, students, and researchers who have assisted with data collection, especially  
478 Ben Dantzer, Sinead English, Stuart Sharp, Helen Spence-Jones, Chris Duncan and Nathan  
479 Thavarajah. Three anonymous reviewers provided helpful comments that improved an earlier  
480 draft of this work. We thank Iain Stevenson and Penny Roth for support and Markus Zöttl  
481 and Raff Mares for many useful discussions. We are especially grateful to Jenny York for  
482 advice throughout this work. Finally, we thank the Northern Cape Department of  
483 Environment and Nature Conservation for permission to conduct the research.

484

485 **References**

486

- 487 1. Lindström J. 1999 Early development and fitness in birds and mammals. *Trends Ecol*  
488 *Evol* **14**(9), 343-348. (doi:10.1016/S0169-5347(99)01639-0).
- 489 2. Tarry-Adkins J.L., Ozanne S.E. 2011 Mechanisms of early life programming: current  
490 knowledge and future directions. *Am J Clin Nutr* **94**(6 Suppl), 1765S-1771S.  
491 (doi:10.3945/ajcn.110.000620).
- 492 3. Nussey D.H., Baird D., Barrett E., Boner W., Fairlie J., Gemmell N., Hartmann N.,  
493 Horn T., Haussmann M., Olsson M. 2014 Measuring telomere length and telomere dynamics  
494 in evolutionary biology and ecology. *Methods Ecol Evol* **5**(4), 299-310. (doi:10.1111/2041-  
495 210X.12161).
- 496 4. Blackburn E.H. 1991 Structure and function of telomeres. *Nature* **350**(6319), 569-  
497 573. (doi:10.1038/350569a0).
- 498 5. Epel E.S., Blackburn E.H., Lin J., Dhabhar F.S., Adler N.E., Morrow J.D., Cawthon  
499 R.M. 2004 Accelerated telomere shortening in response to life stress. *Proc Natl Acad Sci*  
500 **101**(49), 17312-17315. (doi:10.1073/pnas.0407162101).
- 501 6. Badás E.P., Martínez J., Rivero de Aguilar Cachafeiro J., Miranda F., Figuerola J.,  
502 Merino S. 2015 Ageing and reproduction: antioxidant supplementation alleviates telomere  
503 loss in wild birds. *J Evol Biol* **28**(4), 896-905. (doi:10.1111/jeb.12615).
- 504 7. Salomons H.M., Mulder G.A., van de Zande L., Haussmann M.F., Linskens M.H.K.,  
505 Verhulst S. 2009 Telomere shortening and survival in free-living corvids. *Proc R Soc B*  
506 **276**(1670), 3157. (doi:10.1098/rspb.2009.0517).
- 507 8. Boonekamp J.J., Bauch C., Mulder E., Verhulst S. 2017 Does oxidative stress shorten  
508 telomeres? *Biol Lett* **13**(5). (doi:10.1098/rsbl.2017.0164).
- 509 9. Aubert G., Lansdorp P.M. 2008 Telomeres and Aging. *Physiol Rev* **88**(2), 557.  
510 (doi:10.1152/physrev.00026.2007).
- 511 10. Barrett E.L.B., Burke T.A., Hammers M., Komdeur J., Richardson D.S. 2013  
512 Telomere length and dynamics predict mortality in a wild longitudinal study. *Mol Ecol* **22**(1),  
513 249-259. (doi:10.1111/mec.12110).
- 514 11. Bize P., Criscuolo F., Metcalfe N.B., Nasir L., Monaghan P. 2009 Telomere dynamics  
515 rather than age predict life expectancy in the wild. *Proc R Soc B* **276**(1662), 1679.  
516 (doi:10.1098/rspb.2008.1817).

- 517 12. Heidinger B.J., Blount J.D., Boner W., Griffiths K., Metcalfe N.B., Monaghan P.  
518 2012 Telomere length in early life predicts lifespan. *Proc Natl Acad Sci* **109**(5), 1743-1748.  
519 (doi:10.1073/pnas.1113306109).
- 520 13. Boonekamp J.J., Simons M.J.P., Hemerik L., Verhulst S. 2013 Telomere length  
521 behaves as biomarker of somatic redundancy rather than biological age. *Aging Cell* **12**(2),  
522 330-332. (doi:10.1111/accel.12050).
- 523 14. Armanios M., Blackburn E.H. 2012 The telomere syndromes. *Nat Rev Genet* **13**(10),  
524 693-704. (doi:10.1038/nrg3246).
- 525 15. Ridout K.K., Levandowski M., Ridout S.J., Gantz L., Goonan K., Palermo D., Price  
526 L.H., Tyrka A.R. 2017 Early life adversity and telomere length: a meta-analysis. *Mol*  
527 *Psychiatry*. (doi:10.1038/mp.2017.26).
- 528 16. Herborn K.A., Heidinger B.J., Boner W., Noguera J.C., Adam A., Daunt F.,  
529 Monaghan P. 2014 Stress exposure in early post-natal life reduces telomere length: an  
530 experimental demonstration in a long-lived seabird. *Proc R Soc B* **281**(1782).  
531 (doi:10.1098/rspb.2013.3151).
- 532 17. McLennan D., Armstrong J.D., Stewart D.C., McKelvey S., Boner W., Monaghan P.,  
533 Metcalfe N.B. 2016 Interactions between parental traits, environmental harshness and growth  
534 rate in determining telomere length in wild juvenile salmon. *Mol Ecol* **25**(21), 5425-5438.  
535 (doi:10.1111/mec.13857).
- 536 18. Meillère A., Brischox F., Ribout C., Angelier F. 2015 Traffic noise exposure affects  
537 telomere length in nestling house sparrows. *Biol Lett* **11**(9). (doi:10.1098/rsbl.2015.0559).
- 538 19. Watson H., Bolton M., Monaghan P. 2015 Variation in early-life telomere dynamics  
539 in a long-lived bird: links to environmental conditions and survival. *J Exp Biol* **218**(5), 668.  
540 (doi:10.1242/jeb.104265).
- 541 20. Boonekamp J.J., Mulder G.A., Salomons H.M., Dijkstra C., Verhulst S. 2014 Nestling  
542 telomere shortening, but not telomere length, reflects developmental stress and predicts  
543 survival in wild birds. *Proc R Soc B* **281**(1785). (doi:10.1098/rspb.2013.3287).
- 544 21. Mizutani Y., Niizuma Y., Yoda K. 2016 How Do Growth and Sibling Competition  
545 Affect Telomere Dynamics in the First Month of Life of Long-Lived Seabird? *PLOS ONE*  
546 **11**(11), e0167261. (doi:10.1371/journal.pone.0167261).
- 547 22. Nettle D., Monaghan P., Boner W., Gillespie R., Bateson M. 2013 Bottom of the  
548 Heap: Having Heavier Competitors Accelerates Early-Life Telomere Loss in the European  
549 Starling, *Sturnus vulgaris*. *PLOS ONE* **8**(12), e83617. (doi:10.1371/journal.pone.0083617).

- 550 23. Nettle D., Monaghan P., Gillespie R., Brilot B., Bedford T., Bateson M. 2015 An  
551 experimental demonstration that early-life competitive disadvantage accelerates telomere  
552 loss. *Proc R Soc B* **282**(1798), 20141610. (doi:10.1098/rspb.2014.1610).
- 553 24. Reichert S., Criscuolo F., Zahn S., Arrivé M., Bize P., Massemin S. 2015 Immediate  
554 and delayed effects of growth conditions on ageing parameters in nestling zebra finches. *J*  
555 *Exp Biol* **218**(3), 491. (doi:10.1242/jeb.109942).
- 556 25. Stier A., Massemin S., Zahn S., Tissier M.L., Criscuolo F. 2015 Starting with a  
557 handicap: effects of asynchronous hatching on growth rate, oxidative stress and telomere  
558 dynamics in free-living great tits. *Oecologia* **179**(4), 999-1010. (doi:10.1007/s00442-015-  
559 3429-9).
- 560 26. Lukas D., Clutton-Brock T. 2017 Climate and the distribution of cooperative breeding  
561 in mammals. *R Soc open sci* **4**(1). (doi:10.1098/rsos.160897).
- 562 27. Clutton-Brock T.H., Brotherton P.N.M., Russell A.F., O'Riain M.J., Gaynor D.,  
563 Kansky R., Griffin A., Manser M., Sharpe L., McIlrath G.M., et al. 2001 Cooperation,  
564 control, and concession in meerkat groups. *Science* **291**(5503), 478-481.  
565 (doi:10.1126/science.291.5503.478).
- 566 28. Needham B.L., Rehkopf D., Adler N., Gregorich S., Lin J., Blackburn E.H., Epel E.S.  
567 2015 Leukocyte Telomere Length and Mortality in the National Health and Nutrition  
568 Examination Survey, 1999–2002. *Epidemiology (Cambridge, Mass)* **26**(4), 528-535.  
569 (doi:10.1097/EDE.0000000000000299).
- 570 29. Fairlie J., Holland R., Pilkington J.G., Pemberton J.M., Harrington L., Nussey D.H.  
571 2016 Lifelong leukocyte telomere dynamics and survival in a free-living mammal. *Aging Cell*  
572 **15**(1), 140-148. (doi:10.1111/accel.12417).
- 573 30. Asok A., Bernard K., Roth T.L., Rosen J.B., Dozier M. 2013 Parental responsiveness  
574 moderates the association between early-life stress and reduced telomere length.  
575 *Development and Psychopathology* **25**(3), 577-585. (doi:10.1017/S0954579413000011).
- 576 31. Clutton-Brock T.H., Hodge S.J., Flower T.P., Spong G.F., Young A.J. 2010 Adaptive  
577 suppression of subordinate reproduction in cooperative mammals. *Am Nat* **176**(5), 664-673.  
578 (doi:10.1086/656492).
- 579 32. Clutton-Brock T.H., Maccoll A., Chadwick P., Gaynor D., Kansky R., Skinner J.D.  
580 1999 Reproduction and survival of suricates (*Suricata suricatta*) in the southern Kalahari. *Afr*  
581 *J Ecol* **37**(1), 69-80. (doi:10.1046/j.1365-2028.1999.00160.x).

- 582 33. Bell M.B.V., Cant M.A., Borgeaud C., Thavarajah N., Samson J., Clutton-Brock T.H.  
583 2014 Suppressing subordinate reproduction provides benefits to dominants in cooperative  
584 societies of meerkats. *Nat Commun* **5**, 4499. (doi:10.1038/ncomms5499).
- 585 34. Hodge S.J., Flower T.P., Clutton-Brock T.H. 2007 Offspring competition and helper  
586 associations in cooperative meerkats. *Anim Behav* **74**(4), 957-964.  
587 (doi:10.1016/j.anbehav.2006.10.029).
- 588 35. Clutton-Brock T.H., P. N. M B., Smith R., McIlrath G.M., Kansky R., Gaynor D.,  
589 Riain M.J., Skinner J.D. 1998 Infanticide and expulsion of females in a cooperative mammal.  
590 *Proc R Soc B* **265**(1412), 2291. (doi:10.1098/rspb.1998.0573).
- 591 36. Hodge S.J., Manica A., Flower T.P., Clutton-Brock T.H. 2008 Determinants of  
592 reproductive success in dominant female meerkats. *J Anim Ecol* **77**(1), 92-102.  
593 (doi:10.1111/j.1365-2656.2007.01318.x).
- 594 37. Young A.J., Clutton-Brock T. 2006 Infanticide by subordinates influences  
595 reproductive sharing in cooperatively breeding meerkats. *Biol Lett* **2**(3), 385.  
596 (doi:10.1098/rsbl.2006.0463).
- 597 38. Griffin A.S., Pemberton J.M., Brotherton P.N.M., McIlrath G., Gaynor D., Kansky R.,  
598 O'Riain J., Clutton-Brock T.H. 2003 A genetic analysis of breeding success in the  
599 cooperative meerkat (*Suricata suricatta*). *Behav Ecol* **14**(4), 472-480.  
600 (doi:10.1093/beheco/arg040).
- 601 39. Spong G.F., Hodge S.J., Young A.J., Clutton-Brock T.H. 2008 Factors affecting the  
602 reproductive success of dominant male meerkats. *Mol Ecol* **17**(9), 2287-2299.  
603 (doi:10.1111/j.1365-294X.2008.03734.x).
- 604 40. Dubuc C., English S., Thavarajah N., Dantzer B., Sharp S.P., Spence-Jones H.C.,  
605 Gaynor D., Clutton-Brock T.H. 2017 Increased food availability raises eviction rate in a  
606 cooperative breeding mammal. *Biol Lett* **13**(4). (doi:10.1098/rsbl.2016.0961).
- 607 41. Cawthon R.M., Smith K.R., O'Brien E., Sivatchenko A., Kerber R.A. 2003  
608 Association between telomere length in blood and mortality in people aged 60 years or older.  
609 *Lancet* **361**. (doi:10.1016/s0140-6736(03)12384-7).
- 610 42. Criscuolo F., Bize P., Nasir L., Metcalfe N.B., Foote C.G., Griffiths K., Gault E.A.,  
611 Monaghan P. 2009 Real-time quantitative PCR assay for measurement of avian telomeres. *J*  
612 *Avian Biol* **40**(3), 342-347. (doi:10.1111/j.1600-048X.2008.04623.x).
- 613 43. Crawley M. 2007 *The R Book*. Chichester, UK, John Wiley and Sons; 527-528 p.
- 614 44. R Development Core Team. 2013 R: A Language and Environment for Statistical  
615 Computing. *R Foundation for Statistical Computing*.

- 616 45. Burnham K.P., Anderson D.R. 2003 *Model selection and multimodel inference: a*  
617 *practical information-theoretic approach*. New York, NY, Springer.
- 618 46. Russell A.F., Brotherton P.N.M., McIlrath G.M., Sharpe L.L., Clutton-Brock T.H.  
619 2003 Breeding success in cooperative meerkats: effects of helper number and maternal state.  
620 *Behav Ecol* **14**(4), 486-492. (doi:10.1093/beheco/arg022).
- 621 47. Russell A.F., Clutton-Brock T.H., Brotherton P.N.M., Sharpe L.L., McIlrath G.M.,  
622 Dalerum F.D., Cameron E.Z., Barnard J.A. 2002 Factors affecting pup growth and survival in  
623 co-operatively breeding meerkats *Suricata suricatta*. *J Anim Ecol* **71**(4), 700-709.  
624 (doi:10.1046/j.1365-2656.2002.00636.x).
- 625 48. Mares R., Bateman A.W., English S., Clutton-Brock T.H., Young A.J. 2014 Timing  
626 of predispersal prospecting is influenced by environmental, social and state-dependent factors  
627 in meerkats. *Anim Behav* **88**, 185-193. (doi:10.1016/j.anbehav.2013.11.025).
- 628 49. Drake A., Fraser D., Weary D.M. 2008 Parent-offspring resource allocation in  
629 domestic pigs. *Behav Ecol Sociobiol* **62**(3), 309-319. (doi:10.1007/s00265-007-0418-y).
- 630 50. Madden J.R., Kunc H.P., English S., Manser M.B., Clutton-Brock T.H. 2009 Calling  
631 in the gap: competition or cooperation in littermates' begging behaviour? *Proc R Soc B*  
632 **276**(1660), 1255. (doi:10.1098/rspb.2008.1660).
- 633 51. Kilner R.M. 2001 A growth cost of begging in captive canary chicks. *Proc Natl Acad*  
634 *Sci* **98**(20), 11394-11398. (doi:10.1073/pnas.191221798).
- 635 52. Monaghan P. 2013 Organismal stress, telomeres and life histories. *J Exp Biol* **217**(1),  
636 57. (doi:10.1242/jeb.090043).
- 637 53. Huchard E., English S., Bell M.B.V., Thavarajah N., Clutton-Brock T. 2016  
638 Competitive growth in a cooperative mammal. *Nature* **533**(7604), 532-534.  
639 (doi:10.1038/nature17986).
- 640 54. Costantini D., Marasco V., Møller A.P. 2011 A meta-analysis of glucocorticoids as  
641 modulators of oxidative stress in vertebrates. *J Comp Physiol [B]* **181**(4), 447-456.  
642 (doi:10.1007/s00360-011-0566-2).
- 643 55. De Block M., Stoks R. 2008 Compensatory growth and oxidative stress in a  
644 damselfly. *Proc R Soc B* **275**(1636), 781-785. (doi:10.1098/rspb.2007.1515).
- 645 56. Haussmann M.F., Longenecker A.S., Marchetto N.M., Juliano S.A., Bowden R.M.  
646 2012 Embryonic exposure to corticosterone modifies the juvenile stress response, oxidative  
647 stress and telomere length. *Proc R Soc B* **279**(1732), 1447. (doi:10.1098/rspb.2011.1913).

- 648 57. Moreno-Rueda G., Redondo T., Trenzado C.E., Sanz A., Zúñiga J.M. 2012 Oxidative  
649 Stress Mediates Physiological Costs of Begging in Magpie (*Pica pica*) Nestlings. *PLOS ONE*  
650 **7**(7), e40367. (doi:10.1371/journal.pone.0040367).
- 651 58. Nettle D., Andrews C., Reichert S., Bedford T., Kolenda C., Parker C., Martin-Ruiz  
652 C., Monaghan P., Bateson M. 2017 Early-life adversity accelerates cellular ageing and affects  
653 adult inflammation: Experimental evidence from the European starling. *Sci Rep* **7**, 40794.  
654 (doi:10.1038/srep40794).
- 655 59. Hodge S.J., Thornton A., Flower T.P., Clutton-Brock T.H. 2009 Food limitation  
656 increases aggression in juvenile meerkats. *Behav Ecol* **20**(5), 930-935.  
657 (doi:10.1093/beheco/arp071).
- 658 60. Landete-Castillejos T., García A., López-Serrano F.R., Gallego L. 2005 Maternal  
659 quality and differences in milk production and composition for male and female Iberian red  
660 deer calves (*Cervus elaphus hispanicus*). *Behav Ecol Sociobiol* **57**(3), 267-274.  
661 (doi:10.1007/s00265-004-0848-8).
- 662 61. Sanz-Aguilar A., Cortés-Avizanda A., Serrano D., Blanco G., Ceballos O., Grande  
663 J.M., Tella J.L., Donázar J.A. 2017 Sex- and age-dependent patterns of survival and breeding  
664 success in a long-lived endangered avian scavenger. *Sci Rep* **7**, 40204.  
665 (doi:10.1038/srep40204).
- 666 62. Vaupel J.W., Yashin A.I. 1985 Heterogeneity's Ruses: Some Surprising Effects of  
667 Selection on Population Dynamics. *Am Stat* **39**(3), 176-185.  
668 (doi:10.1080/00031305.1985.10479424).
- 669 63. Broer L., Codd V., Nyholt D.R., Deelen J., Mangino M., Willemsen G., Albrecht E.,  
670 Amin N., Beekman M., de Geus E.J.C., et al. 2013 Meta-analysis of telomere length in 19713  
671 subjects reveals high heritability, stronger maternal inheritance and a paternal age effect. *Eur*  
672 *J Hum Genet* **21**(10), 1163-1168. (doi:10.1038/ejhg.2012.303).
- 673 64. Eisenberg D.T.A., Tackney J., Cawthon R.M., Cloutier C.T., Hawkes K. 2017  
674 Paternal and grandpaternal ages at conception and descendant telomere lengths in  
675 chimpanzees and humans. *Am J Phys Anthropol* **162**(2), 201-207. (doi:10.1002/ajpa.23109).
- 676 65. Sharp S.P., Clutton-Brock T.H. 2010 Reproductive senescence in a cooperatively  
677 breeding mammal. *J Anim Ecol* **79**(1), 176-183. (doi:10.1111/j.1365-2656.2009.01616.x).
- 678 66. Simons M.J.P. 2015 Questioning causal involvement of telomeres in aging. *Ageing*  
679 *Research Reviews* **24, Part B**, 191-196. (doi:10.1016/j.arr.2015.08.002).

- 680 67. Clutton-Brock T., Russell A., Sharpe L., Brotherton P., McIlrath G., White S.,  
681 Cameron E. 2001 Effects of helpers on juvenile development and survival in meerkats.  
682 *Science* **293**(5539), 2446-2449. (doi:10.1126/science.1061274).
- 683 68. English S., Huchard E., Nielsen J.F., Clutton-Brock T.H. 2013 Early growth,  
684 dominance acquisition and lifetime reproductive success in male and female cooperative  
685 meerkats. *Ecol Evol* **3**(13), 4401-4407. (doi:10.1002/ece3.820).
- 686 69. Clutton-Brock T.H., Hodge S.J., Spong G., Russell A.F., Jordan N.R., Bennett N.C.,  
687 Sharpe L.L., Manser M.B. 2006 Intrasexual competition and sexual selection in cooperative  
688 mammals. *Nature* **444**(7122), 1065-1068. (doi:10.1038/nature05386).
- 689 70. Sharp S.P., Clutton-Brock T.H. 2011 Competition, breeding success and ageing rates  
690 in female meerkats. *J Evol Biol* **24**(8), 1756-1762. (doi:10.1111/j.1420-9101.2011.02304.x).
- 691 71. Enokido M., Suzuki A., Sadahiro R., Matsumoto Y., Kuwahata F., Takahashi N.,  
692 Goto K., Otani K. 2014 Parental care influences leukocyte telomere length with gender  
693 specificity in parents and offsprings. *BMC Psychiatry* **14**(1), 277. (doi:10.1186/s12888-014-  
694 0277-9).
- 695 72. Schneper L.M., Brooks-Gunn J., Notterman D.A., Suomi S.J. 2016 Early-Life  
696 Experiences and Telomere Length in Adult Rhesus Monkeys: An Exploratory Study.  
697 *Psychosom Med* **78**(9), 1066-1071. (doi:10.1097/psy.0000000000000402).  
698  
699

## 700 **Figure Legends**

701

702 **Figure 1 a)** the positive association between maternal age and pup telomere length at  
703 emergence from the natal burrow. The line represents the model predictions from a GLMM,  
704 with an average pup number of 5.43. **b)** The negative association between the number of  
705 competitors a pup encounters in the first weeks of life, and its telomere length at emergence  
706 from the natal burrow. The line represents the model predictions from a GLMM, with an  
707 average maternal age of 4.86 years. In both figures, the points represent raw data, which are  
708 translucent for clarity. Shaded areas represent the 95% confidence intervals of each model  
709 prediction.

710

711 **Figure 2** the effect of experimental maternal feeding (during gestation and lactation) on pup  
712 telomere lengths is dependent on the number of competitor pups. In control litters (filled  
713 points and solid line) there is a negative relationship between the number of pups and  
714 telomere lengths, while in litters from mothers receiving supplementary feeding (open  
715 triangles and dashed line) this negative association disappears. Lines represent model  
716 predictions for a mean maternal age of 4.5 years, from a GLMM with telomere length as the  
717 response, and maternal age, experimental treatment, and the interaction between treatment  
718 and number of pups. Shaded areas represent the model's 95% confidence intervals for each  
719 model prediction line. Points represent raw data, and are jittered on the x-axis for clarity.

720

721 **Figure 3** The positive association between pup telomere length and survival to adulthood.  
722 The line represents the model predictions from a GLMM, for a pup with a dominant mother  
723 and all other significant predictors at their mean (pup weight: 230g , maternal age: 4.8 years,  
724 group size: 19.2). The points represent raw data (jittered on the y-axis for clarity) and the  
725 shaded areas are 95% confidence intervals of the model predictions.

726

727 **Figure 4 a)** the effect of dominant female infanticidal behaviour on the probability of  
728 subordinate litters surviving their first two days of life. Points represent predicted means and  
729 standard errors from a binomial mixed model with dominant female infanticide risk as the  
730 only predictor. **b)** The effects of infanticide by dominant females on the total number of  
731 surviving pups produced by all subordinate females in their group.

732