The costs of competition: high social status males experience accelerated epigenetic aging in wild baboons

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25 Summary

Aging, for virtually all life, is inescapable. However, within species and populations, rates of biological aging (i.e., physical decline with age) vary across individuals.

- 28 Understanding sources of variation in biological aging is therefore central to understanding
- 29 the biodemography of natural populations. Here, we constructed a DNA methylation-based
- 30 predictor of chronological age for a population of wild baboons in which behavioral.
- 31 ecological, and life history data have been collected for almost 50 years (N = 277 blood
- 32 samples from 245 individuals, including 30 who were longitudinally sampled). Consistent
- 33 with findings in humans and model organisms [1-4], DNA methylation patterns exhibit a
- 34 strong, clock-like association with chronological age, but individuals are often predicted to
- 35 be somewhat older or younger than their known age. However, the two most robust
- 36 predictors of lifespan described for this population—cumulative early adversity and social
- 37 bond strength—do not explain this deviation. Instead, the single most predictive factor is
- 38 male dominance rank: high-ranking males are predicted to be biologically older than their
- 39 true chronological age, such that alpha males appear to be nearly a year older than their
- 40 known age. Longitudinal sampling indicates that males who climb the social hierarchy
- 41 subsequently look epigenetically "older," likely reflecting the high energetic costs of rank
- 42 attainment and maintenance in male baboons. Together, our results indicate that
- 43 environmental effects on survival and epigenetic age can be disjunct, and that achieving
- 44 high rank for male baboons—the best predictor of reproductive success—imposes
- 45 physiological costs consistent with a "live fast, die young" life history strategy.
- 46

47 Keywords: Epigenetic age, epigenetic clock, aging, DNA methylation, dominance rank,

48 baboons

49 **Results and discussion**

50 We used a combination of previously published [5] and newly generated reducedrepresentation bisulfite sequencing (RRBS) data from 245 wild baboons (N = 277 blood 51 52 samples) living in the Amboseli ecosystem of Kenya [6] to generate a DNA methylationbased age predictor (an "epigenetic clock:" [1, 2]). Starting with a data set of methylation 53 54 levels for 458,504 CpG sites genome-wide (Figure S1: Table S1), we used elastic net 55 regression to identify a set of 593 CpG sites that accurately predict baboon age to within a median absolute difference (MAD) of 1.1 years (Pearson's r = 0.762, p < 10⁻⁵³; Table S2; 56 57 median adult life expectancy in this population is 10.3 years for females and 7.94 for males 58 [7]). Because the clock was significantly more accurate in males (N = 135; MAD = 0.9 years; 59 Pearson's r = 0.86, p < 10⁻⁴⁰) than in females (N = 142; MAD = 1.6 years; r = 0.78, p < 10⁻²⁹; two-sided Wilcoxon test for differences in absolute error by sex: $p = 4.35 \times 10^{-9}$), we 60 separated males and females for all subsequent analyses (Figure 1A and 1B). 61

- 62 Overall, the clock performed favorably relative to other morphological or biomarker 63 predictors of age in this population. The epigenetic clock generally explained more
- 64 variance in true chronological age, resulted in lower median error, and exhibited less bias
- 65 than predictions based on body mass index (BMI) or blood cell composition data from flow
- 66 cytometry or blood smears (traits found to change with age in baboons [8, 9]). Its
- 67 performance was comparable to molar dentine exposure, a classical marker of age [10]
- 68 (Figure S2). For 16 males and 14 females, we had two samples collected at different points
- 69 in time. The predicted ages from these longitudinally collected samples were older for the
- 70 later-collected samples, as expected (Figure 1C-D; binomial test $p = 5.95 \times 10^{-5}$).
- 71 Furthermore, the change in epigenetic clock predictions between successive longitudinal
- samples positively predicted the actual change in age between sample dates (β = 0.312, p =
- 73 0.027).



- 74 **Figure 1. Epigenetic clock age predictions in the Amboseli baboons.** Predicted ages are shown relative to
- true chronological ages for **(A)** females (Pearson's r = 0.78, $p < 10^{-29}$, N = 142 samples) and **(B)** males (r =
- 76 0.86, $p < 10^{-40}$, N = 135 samples). Solid lines represent the best fit line; dashed lines show the line for y = x. (C)

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and (D) show predictions for individuals with at least two samples in the data set (14 females and 16 males,
respectively). In 26 of 30 cases (87%), samples collected later were correctly predicted to be from an older
animal.

81 In addition to differences in overall accuracy, sex differences were also apparent in 82 the slope of the relationship between predicted age and chronological age. Males show a 83 2.2-fold higher rate of change in predicted age, as a function of chronological age, compared to females (Figure 1A-B; chronological age by sex interaction in a linear model for 84 predicted age: $\beta = 0.448$, p < 10⁻¹⁸, N = 277). This result agrees with previous findings 85 showing that male baboons senesce more rapidly than females—a pattern shared with 86 87 most other primates investigated thus far, including humans [11]. Interestingly, sex 88 differences are not apparent in animals < 8 years, which roughly corresponds to the age at 89 which the majority of males have achieved adult dominance rank and dispersed from their 90 natal group [12-14] (N = 158, chronological age by sex interaction β = -0.038, p = 0.808). 91 Rather, sex difference becomes apparent after baboons have reached full physiological and 92 social adulthood (N = 119, chronological age by sex interaction β = 0.459, p < 10⁻⁶ in 93 animals \geq 8 years), when divergence between male and female life history strategies is 94 most marked [12-14] and when aging rates between the sexes are predicted to diverge [15-95 17]. This pattern suggests that within each sex, deviations between predicted age and 96 chronological age—commonly interpreted as a measure of "biological age" or accelerated 97 aging—may also be affected by environmental or life history variation, as has been 98 suggested in humans, lab mice, and captive rhesus macaques [3, 4, 18, 19].

99 To test this hypothesis, we focused on four factors of known importance to fertility 100 and/or survival components of fitness in the Amboseli baboon population. First, we investigated the effects of cumulative early adversity, which is a strong predictor of 101 102 shortened lifespan in female baboons: females who experience three or more major 103 sources of early adversity have expected adult lifespans that are a decade shorter than 104 those who experience none [20]. Additionally, those females are less capable of raising 105 their own juvenile offspring later in life, suggesting that early adversity compromises their 106 physical condition over the long-term [21]. Following [20, 21], we measured cumulative 107 adversity as a count of major adverse experiences suffered in early life, including low 108 maternal social status, early life drought, a competing younger sibling, maternal loss, and 109 high experienced population density. To maximize our sample size, we omitted early life 110 social connectedness (included in [20] but omitted for the same reason in [21]), because social connectedness data were missing for mothers born relatively early in the long-term 111 112 study. We predicted that high cumulative early adversity, which is linked to reduced 113 lifespan, would predict increased biological age.

Second, we considered social bond strength in adulthood, which is positively
associated with longer adult lifespan in female Amboseli baboons, human populations, and
several other wild social mammals [22-24]. We predicted that low social bond strength,
due to its relationship with decreased lifespan, would be associated with increased
biological age.

Third, we investigated the effects of dominance rank, which is a major determinant
of resource access in baboons. High-ranking males sire the most offspring, and highranking females experience shorter interbirth intervals, retain higher fertility during

122 droughts, and form stronger social bonds with males [22, 25-27].

Finally, we assessed the effect of BMI, which captures dimensions of both body condition and competitive advantage [8]. In this case, we calculated BMI relative to the expected value for each male's age, which eliminates the correlation between BMI and age and BMI and rank. The predictions for dominance rank and body mass index associations were less clear: improved resource access could conceivably slow biological aging, but increased investment in growth and reproduction (either through higher fertility in females or physical competition for rank in males) could also be energetically costly.

130 We tested these predictions by modeling the deviation between predicted age and 131 known chronological age (Δ_{age}) as a function of cumulative early adversity, ordinal 132 dominance rank, body mass index (controlling for age), and for females, social bond 133 strength to other females. Social bond strength was not included in the model for males, as 134 this measure was unavailable for a large proportion of males in this data set (53.8%). We 135 also included chronological age as a predictor in the model, as epigenetic age tends to be 136 systematically overpredicted for young individuals and underpredicted for old individuals 137 (Figure 1A-B); including chronological age in the model controls for this compression 138 effect. No predictor variables were strongly linearly correlated (all R² < 0.10; Table S3).

Surprisingly, despite being two of the strongest known predictors of lifespan in wild baboons, neither cumulative early life adversity nor social bond strength explain variation in Δ_{age} (Table 1). In contrast, high male dominance rank is strongly and significantly associated with larger values of Δ_{age} ($\beta = -0.785$, p = 7.0 x 10⁻⁴; Figure 2; Table 1). Alpha males are predicted to be an average of 10.95 months older than their true chronological age—a difference that translates to 11.5% of a male baboon's expected adult lifespan [7].

- 145 Dominance rank is strongly age structured in male baboons: males tend to attain their 146 highest rank between 7 and 12 years of age and fall in rank thereafter. Thus, nearly all males in the top four rank positions in our data set were between 7 and 12 years of age at 147 148 the time they were sampled (however, because not all 7 – 12 year-olds are high-ranking, 149 low rank positions include males of all age groups; Table S1, Figure S3). Our finding that 150 high rank predicts accelerated epigenetic aging therefore implies that males incur the costs 151 of high rank primarily in early to mid-adulthood, and only if they succeed in attaining high 152 rank. Accelerated epigenetic aging is thus a function of absolute rank values, regardless of 153 age, not deviations from the mean rank expected given a male's age (i.e., "rank-for-age," 154 which can be quantified as the residuals of male dominance rank modeled as a quadratic 155 function of chronological age: Figure S3). In support of this interpretation, a model that 156 includes rank-for-age as an additional covariate recapitulates the significant effect of 157 ordinal male rank (p=0.045), but finds no effect of rank-for-age (p=0.819; Table S4). In 158 contrast, we observed no evidence for rank effects on Δ_{age} in females, consistent with 159 overall sex differences in patterns of aging in primates and other mammals [15] and
- 160 marked sex differences in the effects of rank on other molecular phenotypes in the
- 161 Amboseli baboons specifically [11, 28].

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Table 1. Predictors of Δ_{age}^{1}				
Covariate	β	P-value	β	P-value
	(Female)	(Female)	(Male)	(Male)
Intercept	5.400	1.33 x 10 ⁻¹⁵	3.294	1.19 x 10 ⁻⁸
Cumulative early adversity	-0.050	0.807	-0.005	0.973
Social bond strength	0.382	0.164	_	—
Dominance rank	0.025	0.228	-0.078	7.39 x 10 ⁻⁴
Body mass index (age-adjusted)	0.026	0.682	0.111	6.33 x 10 ⁻³
Chronological age	-0.699	1.62 x 10 ⁻²⁸	-0.277	8.36 x 10 ⁻⁸

¹Separate linear models for Δ_{age} were fit for females (N = 66) and for males (N = 93) for whom no data values

163 were missing; social bond strength was not included in the model for males. Significant results are shown in

164 bold.



Figure 2. Dominance rank predicts epigenetic aging in male baboons. High rank is associated with elevated values of Δ_{age} ($\beta = -0.0785$, $p = 7.39 \times 10^{-4}$, N = 105). The y-axis shows relative epigenetic age, a measure of epigenetic aging similar to Δ_{age} that is based on the sample-specific residuals from the relationship between predicted age and true chronological age. Positive (negative) values correspond to predicted ages that are older (younger) than expected for that chronological age. Dominance rank is measured using ordinal values, such that smaller values indicate higher rank. Dots and error bars represent the means and standard errors, respectively. Gray values above the x-axis indicate sample sizes for each rank. 172

Previous work has shown that high-ranking male baboons, but not high-ranking
female baboons, up-regulate gene expression in inflammation-related and immune
response pathways [28]. Elevated or chronic inflammation is thought to be one of the
hallmarks of aging, and, in human populations, is one of the strongest predictors of
mortality risk [29-31]. Consistent with these observations, CpG sites in the epigenetic clock

- that increase in DNA methylation with age (N = 459 sites) are enriched in or near genes
- 179 that are up-regulated in the Amboseli baboons in response to the bacterial endotoxin
- 180 lipopolysaccharide (LPS), which is a strong driver of inflammation (Figure S3; Fisher's
- 181 exact test: log₂(odds ratio) = 1.88, p = 0.001; gene expression data from [28]). In contrast,
- 182 clock sites that decrease in DNA methylation with age (N = 134) are significantly enriched

183 in or near genes that are down-regulated after LPS exposure (Fisher's exact test: log₂(OR) =

3.28, p = 0.002). Clock sites, especially those for which DNA methylation is positively 184 185 correlated with age, are also enriched in genes, CpG islands, promoter regions, and putative enhancers, compared to the background set of sites we initially considered as candidates 186 187 for inclusion in the clock (Figure S4; Fisher's exact tests, all p < 0.005; similar functional 188 enrichment has been found in a human epigenetic clock [2]). Moreover, clock sites are 189 enriched in regions previously found to change in methylation levels with age [32] and in 190 regions showing regulatory activity in high-throughput reporter assays [33]. Together, 191 these results suggest that accelerated epigenetic aging in males reflects functionally 192 important changes in DNA methylation levels, concentrated in immune response and 193 inflammation-related pathways.

194 In baboon males, dominance rank reflects physical condition and fighting ability. 195 Rank is therefore dynamic across the life course, such that males in their prime (ages 7 – 196 12) are most likely to be high-ranking, and the same male is likely to occupy multiple 197 positions in the social hierarchy during his lifetime [14]. If accelerations in epigenetic age 198 are tightly coupled to rank, our results predict that across longitudinally collected samples, 199 a male who becomes higher ranking should look older for age in his second sample relative 200 to first, whereas a male who loses status should look younger for age in his second sample 201 relative to first. To assess this possibility, we calculated the residuals of the best fit line 202 relating chronological age and predicted age ("relative epigenetic age") for 14 males for 203 whom we had repeated samples collected when they occupied different dominance rank 204 positions (N = 28 samples, 2 per male). In this data set, the sample collected when males 205 were higher status typically predicted higher values of relative epigenetic age compared to 206 the sample collected when they were lower status, consistent with high rank-associated 207 accelerated biological aging (Figure 3; paired t-test, t = -2.31, p = 0.038). For example, in the 208 case of one young adult male who we sampled at rank 18 and then rank 1, as he was first 209 climbing the social hierarchy, the actual time that elapsed between samples was 0.79 years 210 but he exhibited an increase in *predicted* age of 2.6 years. Moreover, the two males that 211 showed a decrease in predicted age, despite increasing in chronological age (Figure 1D), 212 were among those that experienced the greatest drop in social status between samples. 213 Thus, change in rank between samples for the same male predicted change in Δ_{age} 214 controlling for chronological age ($R^2 = 0.24$, p=0.044). Consistent with our cross-sectional 215 results, we did not observe evidence for a relationship between change in Δ_{age} and rank-

216 for-age ($R^2 = 0.14$, p = 0.135).

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223 Together, our findings indicate that major environmental predictors of lifespan and 224 mortality risk (e.g., social bond strength and early life adversity in this population) do not 225 necessarily predict epigenetic measures of biological age. Although this assumption is 226 widespread in the literature, including for epigenetic clock analyses [34, 35], our results 227 are broadly consistent with empirical results in humans. Specifically, while studies of early 228 life adversity, which also predicts lifespan in human populations, find relatively consistent 229 support for a relationship between early adversity and accelerated epigenetic aging in 230 children and adolescents [36-41], there is little evidence for the long-term effects of early adversity on epigenetic age in adulthood [42-47]. Thus, while DNA methylation may make 231 232 an important contribution to the biological embedding of early adversity into adulthood 233 [48, 49], it does not seem to do so through affecting the epigenetic clock itself. Social and 234 environmental effects on the clock instead seem to be most influenced by concurrent 235 conditions, lending support to "recency" models for environmental effects on aging that posit that health is more affected by the current environment than past experience [50-52]. 236 237 Additional longitudinal sampling will be necessary to evaluate whether current conditions 238 alone can explain accelerated epigenetic aging, or whether it also requires integrating the 239 effects of exposures across the life course (the "accumulation" model: [50, 52]). Repeated 240 samples could also help exclude an alternative explanation for our findings: that viability 241 selection against individuals who experienced high early adversity attenuates the true 242 effect of cumulative early adversity on relative epigenetic age.

Finally, our analyses reveal that males who achieve high rank appear epigenetically older than expected given their known chronological age. Although high ranking males also tend to be larger due to increased muscle mass (Pearson's *r* between rank and BMI = 0.56, $p = 6.38 \times 10^{-9}$), we observed an additional, independent effect of age-adjusted BMI: males who had high BMI relative to their age also looked older for age, controlling for rank and age ($\beta = 0.1107$, p = 0.006) (Table 1). These two effects suggest that investment in body 249 condition, which is a crucial factor in male competitive ability, incurs other physiological

- costs that compound to influence biological age. Indeed, previous research on the Amboseli
- 251 baboons also points to costs of high rank, including high levels of glucocorticoids in alpha
- 252 males [53], increased expression of genes involved in innate immunity and inflammation
- 253 [28], and a trend towards elevated mortality risk [54]. These associations may arise
- because high-ranking males are both more likely to engage in physical conflict with other
- 255 males, and more likely to spend long periods of time in energetically costly mate-guarding
- [55, 56]. Alternatively, males who are able to make significant investments in bodycondition, while tolerating their accompanying costs, may be able to successfully maintain
- high rank. Importantly, since doing so is likely to contribute to higher lifetime reproductive
- success for male baboons, the fitness-associated benefits of high rank can exceed the costs,
- 260 even if accelerated biological aging increases the risk of mortality.
- 261

262 Acknowledgements

- We gratefully acknowledge the support provided by the National Science Foundation and 263 264 the National Institutes of Health for the majority of the data represented here, currently 265 through NSF IOS 1456832, NIH R01AG053308, R01AG053330, R01HD088558, and 266 P01AG031719. R.A.J. is supported by NIH F32HD095616 and J.A.A. by NSF #2018264636. 267 We thank the members of the Amboseli Baboon Research Project for collecting the data 268 presented here, especially J. Altmann for her foundational role in establishing the study 269 population and these data sets; J. Gordon, N. Learn, and K. Pinc for managing the database; 270 R.S. Mututua, S. Savialel, and I.K. Warutere for data collection in the field: and T. Wango and 271 V. Oudu for their assistance in Nairobi. We also thank the Kenya Wildlife Service, University 272 of Nairobi, the Institute of Primate Research, the National Museums of Kenva, the National 273 Council for Science, Technology, and Innovation, members of the Amboseli-Longido 274 pastoralist communities, the Enduimet Wildlife Management Area, Ker & Downey Safaris, 275 Air Kenya, and Safarilink for their assistance in Kenya. Finally, we thank current and past 276 members of the Tung, Alberts, Archie, and Altmann labs for their helpful feedback. This 277 research was approved by IACUCs at Duke University, University of Notre Dame, and
- 277 Princeton University and adhered to all the laws and guidelines of Kenva. For a complete
- set of acknowledgments of funding sources, logistical assistance, and data collection and
- 280 management, please visit http://amboselibaboons.nd.edu/acknowledgements/.
- 281
- 282 Author Contributions
- 283 Conceptualization, R.A.J., J.A.A., J.T., E.A.A., A.J.L.; Investigation, J.A.A., R.A.J., A.J.L., F.A.C.,
- 284 M.Y.A., T.N.V., and J.T.; Formal Analysis, J.A.A. and R.A.J.; Writing—Original Draft, R.A.J.,
- 285 J.A.A., and J.T.; Writing—Reviewing & Editing, R.A.J., J.A.A., A.J.L., T.N.V., M.Y.A., F.A.C., S.C.A.,
- E.A.A., and J.T. Funding Acquisition, J.T., S.C.A., and E.A.A. Supervision, J.T.
- 287
- 288 Declaration of Interests
- 289 The authors declare no competing interests.
- 290
- 291

292 STAR Methods

293 Study population and biological sample collection

This study focused on a longitudinally monitored population of wild baboons (*Papio cynocephalus*, the yellow baboon, with some admixture from the closely related anubis baboon *P. anubis* [57, 58]) in the Amboseli ecosystem of Kenya. This population has been continuously monitored by the Amboseli Baboon Research Project (ABRP) since 1971 [6]. For the majority of study subjects (N = 242 of 245 individuals), birth dates were therefore known to within a few days' error; for the remainder, birth dates were known within 3 months' error (Table S1).

301 All DNA methylation data were generated from blood-derived DNA obtained during 302 periodic darting efforts, as detailed in [28, 59, 60]. Samples were obtained under approval 303 from the Institutional Animal Care and Use Committee (IACUC) of Duke University and 304 adhered to all the laws and guidelines of Kenya. In brief, individually recognized study 305 subjects were temporarily anesthetized using a Telazol-loaded dart delivered through a 306 blow gun. Baboons were then safely moved to a new location where blood samples and 307 morphometric data, including body mass and crown-rump length, were collected. Baboons 308 were then allowed to recover from anesthesia in a covered holding cage and released to 309 their group within 2 – 4 hours. Blood samples were stored at -20 C in Kenya until export to 310 the United States.

311

312 DNA methylation data

DNA methylation data were generated from blood-extracted DNA collected from known individuals in the Amboseli study population (N = 277 samples from 245 animals; 14 females and 14 males were each sampled twice, and 1 female and 1 male were each sampled three times). Here, we analyzed a combined data set that included previously published reduced representation bisulfite sequencing [61] (RRBS) data from the same population (N = 36) [5] and new RRBS data from 241 additional individuals.

319 RRBS libraries were constructed following [62], using ~200 ng baboon DNA plus 0.2 320 ng unmethylated lambda phage DNA per sample as input. Samples were sequenced to a 321 mean depth of 17.8 (± 10.5 s.d.) million reads on either the Illumina HiSeq 2000 or HiSeq 322 4000 platform (Table S1), with an estimated mean bisulfite conversion efficiency (based on 323 the conversion rate of lambda phage DNA) of 99.8% (minimum = 98.1%). Sequence reads 324 were trimmed with Trim Galore! [63] to remove adapters and low quality sequence (Phred score < 20). Trimmed reads were mapped with BSMAP [64] to the baboon genome 325 326 (*Panu2.0*) allowing a 10% mismatch rate to account for the degenerate composition of 327 bisulfite-converted DNA. We used the mapped reads to count the number of methylated 328 and total reads per CpG site, per sample [64, 65]. Following [5, 32], CpG sites were filtered 329 to retain sites with a mean methylation level between 0.1 and 0.9 (i.e., to exclude 330 constitutively hyper- or hypo-methylated sites) and mean coverage $\geq 5x$. We also excluded 331 any CpG sites with missing data for $\geq 15\%$ of individuals in the sample. After filtering, we 332 retained N = 458,504 CpG sites for downstream analysis. For the remaining missing data 333 (mean number of missing sites per sample = $1.4\% \pm 3.5\%$ s.d., equivalent to $6,409 \pm 16,024$ 334 sites), we imputed methylation levels using a k-nearest neighbors approach in the R 335 package *impute*, using default parameters [66].

336

337 Building the epigenetic clock

338 We used the R package *glmnet* [67] to build a DNA methylation clock for baboons. 339 Specifically, we fit a linear model in which the predictor variables were normalized levels of 340 DNA methylation at 458,504 candidate clock CpG sites across the genome and the response 341 variable was chronological age. To account for the excess of CpG sites relative to samples, 342 *almnet* uses an elastic net penalty to shrink predictor coefficients toward 0 [68]. Optimal 343 alpha parameters were identified by grid searching across a range of alphas from 0 344 (equivalent to ridge regression) to 1 (equivalent to LASSO) by increments of 0.1. We 345 defined the optimal alpha as the value that maximized R² between predicted and true 346 chronological age across all samples. We set the regularization parameter lambda to the 347 value that minimized mean-squared error during n-fold internal cross-validation.

348 To generate predicted age estimates for a given sample, we used a leave-one-out 349 cross-validation approach in which all samples but the "test" sample were included for 350 model training, and the resulting model was used to predict age for the left-out test sample. Importantly, training samples were scaled independently of the test sample in each leave-351 352 one-out model to avoid bleed-through of information from the test data into the training 353 data. To do so, we first quantile normalized methylation ratios (the proportion of 354 methylated counts to total counts for each CpG site) within each sample to a standard 355 normal distribution. Training samples were then separated from the test sample and the 356 methylation levels for each CpG site in the training set were quantile normalized across 357 samples to a standard normal distribution. For predicting age in the test sample, we 358 compared the methylation value for each site in the test sample to the empirical cumulative 359 distribution function for the training samples (at the same site) to estimate the quantile in 360 which the training sample methylation ratio fell. The training sample was then assigned the same quantile value from the standard normal distribution using the function *qnorm* in R. 361

362

363 Comparisons to alternative predictors of aging

364 To assess the utility of the DNA methylation clock relative to other data types, we 365 compared its predictive accuracy to clocks based on three other age-related phenotypes: 366 tooth wear (percent molar dentine exposure [10]), body condition (body mass index: BMI 367 [8]), and blood cell type composition (blood smear counts and lymphocyte/monocyte 368 proportions from flow cytometry performed on peripheral blood mononuclear cells, as in 369 [28, 69]). Leave-one-out model training and prediction were performed for each data type 370 using linear modeling (i.e., not glmnet, since the number of features was much less than the 371 number of samples in this case). To compare the relative predictive accuracy of each data 372 type, we calculated the R² between predicted and chronological age, the median absolute 373 difference between predicted and chronological age, and the bias in age predictions (the 374 absolute value of 1- slope of the best fit line between predicted and chronological age) 375 (Figure S2).

Tooth wear. Molar enamel in baboons wears away with age to expose the underlying
dentine layer. Percent dentine exposure (PDE) on the molar occlusal surface has been
shown to be strongly age-correlated in previous work [10]. To assess its predictive power,
we obtained PDE data from tooth casts reported by Galbany and colleagues [10] for the left
upper molars (tooth positions M1, M2, M3) and left lower molars (tooth positions M1, M2,
M3) for 39 males and 34 females in our data set. For each molar position (M1, M2, M3)
within each individual, we calculated PDE as the mean for the upper and lower molars.

Because dentine exposure scales quadratically with respect to age [10], we fit age as a function of PDE using the following model: $age \sim \sqrt{PDE_{M1}} + \sqrt{PDE_{M2}} + \sqrt{PDE_{M3}}$.

Body mass index. For both male and female baboons in Amboseli, body mass 385 386 increases with age until individuals reach peak size, and then tends to decrease with age as 387 animals lose fat and/or muscle mass [8]. To quantify body condition using body mass, we 388 calculated body mass index (BMI) values for 139 males and 154 females for whom body 389 mass and crown-rump length data were available from periodic darting efforts. We 390 retained only measures taken from animals born into and sampled in wild-feeding study 391 groups, when sex-skin swellings (in females only) that could affect crown-rump length 392 measures were absent. BMI was calculated as mass (kilograms) divided by crown-rump 393 length (meters squared), following [70]. To assess the predictive power of BMI for age, we 394 built sex-specific piecewise-regression models. Breakpoints for the piecewise-regression 395 models (to separate "youthful" versus "aged" animals) were set at 8 years old for males and 396 10 years old for females, following findings from previous work on body mass in the 397 Amboseli population [8].

398 *Blood cell type composition.* The proportions of different cell types in blood change 399 across the life course, including in baboons [9]. We assessed the predictive power of blood 400 cell composition for age using two data sets. First, we used data collected from blood smear 401 counts (N = 134) for five major white blood cell types: basophils, eosinophils, monocytes, 402 lymphocytes, and neutrophils. Second, we used data on the proportional representation of 403 five peripheral blood mononuclear cell (PBMC) subsets: cytotoxic T cells, helper T cells, B 404 cells, monocytes, and natural killer cells, measured using flow cytometry as reported by Lea 405 and colleagues [28] (N = 53). Cell types were included as individual covariates for leave-406 one-out model training.

407

408 Sources of variance in predicted age

409 We asked whether factors known to be associated with inter-individual variation in 410 fertility or survival also predict inter-individual variation in Δ_{age} (predicted age from the epigenetic clock minus known chronological age). To do so, we fit linear models separately 411 412 for males and females, with Δ_{age} as the dependent variable and dominance rank at the time 413 of sampling, cumulative early adversity, relative BMI (corrected for age), and chronological 414 age as predictor variables [20]). For females, we also included a measure of social bond 415 strength to other females as a predictor variable, based on findings that show that socially 416 isolated females experience higher mortality rates in adulthood [22, 71]. Samples with 417 missing values for any of the predictor variables were excluded in the model, resulting in a 418 final analysis set of 66 female samples (from 59 females) and 93 male samples (from 84 419 males). The chronological ages of samples with complete data relative to samples with 420 missing data were equivalent for females (t-test, t = 1.95, p = 0.053) but were slighly lower 421 for males (t-test, t = -3.04, p = 0.003; mean chronological ages are 7.98 and 9.65 years for 422 complete and missing samples, respectively). Predictor variables were measured as 423 follows.

Dominance rank. Sex-specific dominance hierarchies were constructed monthly for
every social group in the study population based on the outcomes of dyadic aggressive
encounters. Ordinal dominance ranks were assigned to every adult based on these

427 hierarchies, such that low numbers represent high rank/social status and high numbers

represent low rank/social status [72]. Although most analyses of data from the Amboseli 428 429 baboons have used ordinal ranks as the primary measure of social status, in some analyses 430 proportional rank (i.e., the proportion of same-sex members of an individual's social group 431 that he or she dominates) has proven to be a stronger predictor of behavioral, life history, 432 or physiological outcomes [73]. In this study, we chose to use ordinal ranks, but proportional and ordinal dominance rank were highly correlated in our dataset (r^2 =.70. 433 434 p=1.13 x 10⁻⁵⁸). Using proportional rank rather than ordinal rank did not qualitatively 435 affect our analyses. Additionally, to investigate whether the patterns we observed are due 436 to a consistent effect of rank across all ages, or instead an effect of being high or low rank

relative to the expected (mean) value for a male's age, we also calculated a "rank-for-age"
value. Rank-for-age is defined as the residuals of a model with dominance rank as the
response variable and age and age² as the predictor variables (Fig S4).

440 *Cumulative early adversity.* Previous work in Amboseli defined a cumulative early 441 adversity score as the sum of 6 different adverse conditions that a baboon could experience 442 during early life [20]. This index strongly predicts adult lifespan in female baboons, and a 443 modified version of this index also predicts offspring survival [21]. To maximize the sample 444 size available for analysis, we excluded maternal social connectedness, the source of 445 adversity with the highest frequency of missing data, leaving us with a cumulative early 446 adversity score generated from 5 different binary-coded adverse experiences. These 447 experiences were: (i) early life drought (defined as ≤ 200 mm of rainfall in the first year of 448 life), which is linked to reduced fertility in females [27, 74]; (ii) having a low ranking 449 mother (defined as falling within the lowest quartile of ranks for individuals in the dataset). 450 which predicts rates of maturation [75-77]; (iii) having a close-in-age younger sibling (<1.5 451 years), which may redirect maternal investment to the sibling [78], (iv) being born into a 452 large social group, which may increase within-group competition for shared resources [27, 453 77, 79], and (v) maternal death before the age of 4, which results in a loss of both social and 454 nutritional resources [77, 80].

455 Body mass index. BMI was modeled as the residuals from sex-specific piecewise 456 regression models relating BMI to age. By taking this approach, we asked whether having 457 relatively high BMI for one's age and sex predicted higher (or lower) Δ_{age} .

458 Social bond strength. For this analysis, we measured female social bond strength to 459 other females using the dyadic sociality index (F-DSI) [54]. We did not include this 460 parameter (male's social bond strength to females) for the male model, because this 461 measure is unavailable for young males in this dataset. F-DSI was calculated as an 462 individual's average bond strength with her top three female social partners, in the 365 463 days prior to the day of sampling, controlling for observer effort. This approach is based on 464 representative interaction sampling of grooming interactions between females, in which 465 observers record all grooming interactions in their line of sight while moving through the 466 group conducting random-ordered, 10-minute long focal animal samples of pre-selected 467 individuals. Because smaller groups receive more observer effort per individual and per 468 dyad (and thus record more grooming interactions per individual or dyad), we estimated 469 observer effort for dyad *d* in year *y* as:

$$E_{d,y} = \frac{c_d(s_d)}{f_d}$$

471 where c_d is the number of days the two females in a dyad were coresident in the same

social group, s_d is the number of focal samples taken during the dyad's coresidence, and f_d 472 473 is the average number of females in the group during the dyad's coresidence.

F-DSI for each adult female dyad in each year is the z-scored residual, ε , from the 474 475 model:

- 476
- $\log(R_{d,y}) = \beta(\log(E_{d,y})) + \varepsilon$ where $R_{d,y}$ is the number of grooming interactions for dyad *d* in year *y* divided by the 477

number of days that the two individuals were coresident, and $E_{d,v}$ is observer effort. 478

479

480 Analysis of longitudinal samples

To test whether changes in rank predict changes in relative epigenetic age within 481 482 individuals, we used data from 5 males from the original dataset and generated additional 483 RRBS data for 9 samples, resulting in a final set of 14 males who each were sampled at least 484 twice in the data set, when they occupied different ordinal ranks (mean years elapsed 485 between samples = 3.92 ± 1.94 s.d.; mean absolute difference in dominance ranks = $6.86 \pm$ 486 5.07 s.d.). This effort increased our total sample size to N = 286 samples from 248 unique 487 individuals. To incorporate the new samples into our analysis, we reperformed leave-one-488 out age prediction with N-fold internal cross validation at the optimal alpha selected for the 489 original N = 277 samples (alpha = 0.1). For the 277 samples carried over from the original 490 analysis, we verified that age predictions were nearly identical between the previous 491 analysis and the expanded data set ($R^2 = 0.98$, $p = 2.21 \times 10^{-239}$).

492 Based on the new age predictions for males in the data set (N = 144), we again 493 calculated relative epigenetic age as the residual of the best fit line relating predicted age to 494 chronological age. We then used the 14 males with repeated DNA methylation profiles and 495 rank measures in this dataset to test whether, within individuals, changes in dominance 496 rank or rank-for-age explained changes in relative epigenetic age between samples.

497

498 Epigenetic clock enrichment analyses

To evaluate whether CpG sites included in the epigenetic clock were enriched in 499 500 functionally important regions of the baboon genome [32, 81], we used two-sided Fisher's 501 exact tests to investigate enrichment/depletion of the 593 epigenetic clock sites in (i) gene 502 bodies and exons, based on the Ensembl annotation Panu2.0.90; (ii) CpG islands annotated 503 in the UCSC Genome Browser; (iii) CpG shores, defined as the 2,000 basepairs flanking CpG 504 islands (following [32, 81, 82]); and (iv) promoter regions, defined as the 2,000 basepairs 505 upstream of the 5'-most annotated transcription start site for each protein-coding gene 506 (following [32, 81]). We also considered (v) putative enhancer regions, which have not 507 been annotated for the *Panu2.0* assembly. We therefore used ENCODE H3K4me1 ChIP-seq 508 data from humans [83] and the *liftOver* tool to define likely enhancer coordinates in 509 Panu2.0.

510 We also tested for enrichment of clock sites in regions of the genome that have been 511 identified by previous empirical studies to be of special interest. First, we considered 512 regions that likely have regulatory activity in blood cells, defined as all 200 base-pair 513 windows that showed evidence of enhancer activity in a recently performed massively 514 parallel reporter assay [33]. We used *liftOver* to identify the inferred homologous *Panu2.0* 515 coordinates for these windows, which were originally defined in the human genome. 516 Second, we defined age-related differentially methylated regions (age DMRs) in the

- 517 Amboseli baboons based on genomic intervals found, in previous analyses, to contain at
- 518 least three closely spaced age-associated CpG sites (inter-CpG distance ≤ 1kb), as described
- 519 in [32]. Third, we defined lipopolysaccharide (LPS) up-regulated and LPS down-regulated
- 520 genes as those genes that were significantly differentially expressed (1% false discovery
- rate) between unstimulated Amboseli baboon white blood cells and LPS-stimulated cells
- from the same individual, following 10 hours of culture in parallel [28].
- 523
- 524 Data and code availabity
- All sequencing data generated during this study are available in the NCBI Sequence Read
- 526 Archive (project accession PRJNA607996).

527	Refere	ences
528	1.	Hannum, G., Guinney, J., Zhao, L., Zhang, L., Hughes, G., Sadda, S., Klotzle, B., Bibikova,
529		M., Fan, JB., and Gao, Y. (2013). Genome-wide methylation profiles reveal
530		quantitative views of human aging rates. Molecular cell 49, 359-367.
531	2.	Horvath, S. (2013). DNA methylation age of human tissues and cell types. Genome
532		biology <i>14</i> , 3156.
533	3.	Petkovich, D.A., Podolskiy, D.I., Lobanov, A.V., Lee, SG., Miller, R.A., and Gladyshev,
534		V.N. (2017). Using DNA methylation profiling to evaluate biological age and
535		longevity interventions. Cell metabolism 25, 954-960. e956.
536	4.	Stubbs, T.M., Bonder, M.J., Stark, AK., Krueger, F., von Meyenn, F., Stegle, O., and
537		Reik, W. (2017). Multi-tissue DNA methylation age predictor in mouse. Genome
538		biology <i>18</i> , 68.
539	5.	Lea, A.J., Altmann, J., Alberts, S.C., and Tung, J. (2016). Resource base influences
540		genome - wide DNA methylation levels in wild baboons (Papio cynocephalus).
541		Molecular ecology 25, 1681-1696.
542	6.	Alberts, S.C., and Altmann, J. (2012). The Amboseli Baboon Research Project: 40
543		vears of continuity and change. In Long-term field studies of primates. (Springer),
544		pp. 261-287.
545	7.	Colchero, F., Rau, R., Jones, O.R., Barthold, J.A., Conde, D.A., Lenart, A., Nemeth, L.,
546		Scheuerlein, A., Schoeley, J., and Torres, C. (2016). The emergence of longevous
547		populations. Proceedings of the National Academy of Sciences <i>113</i> , E7681-E7690.
548	8.	Altmann, J., Gesquiere, L., Galbany, J., Onyango, P.O., and Alberts, S.C. (2010). Life
549		history context of reproductive aging in a wild primate model. Annals of the New
550		York Academy of Sciences 1204, 127-138.
551	9.	Jayashankar, L., Brasky, K.M., Ward, J.A., and Attanasio, R. (2003). Lymphocyte
552		modulation in a baboon model of immunosenescence. Clin. Diagn. Lab. Immunol. 10,
553		870-875.
554	10.	Galbany, J., Altmann, J., Pérez - Pérez, A., and Alberts, S.C. (2011). Age and individual
555		foraging behavior predict tooth wear in Amboseli baboons. American Journal of
556		Physical Anthropology 144, 51-59.
557	11.	Bronikowski, A.M., Altmann, J., Brockman, D.K., Cords, M., Fedigan, L.M., Pusey, A.,
558		Stoinski, T., Morris, W.F., Strier, K.B., and Alberts, S.C. (2011). Aging in the natural
559		world: comparative data reveal similar mortality patterns across primates. Science
560		<i>331</i> , 1325-1328.
561	12.	Alberts, S.C., and Altmann, J. (1995). Balancing costs and opportunities: dispersal in
562		male baboons. The American Naturalist 145, 279-306.
563	13.	Alberts, S.C., and Altmann, J. (1995). Preparation and activation: determinants of age
564		at reproductive maturity in male baboons. Behavioral Ecology and Sociobiology 36,
565		397-406.
566	14.	Alberts, S.C., Watts, H.E., and Altmann, J. (2003). Queuing and queue-jumping: long-
567		term patterns of reproductive skew in male savannah baboons, Papio cynocephalus.
568		Animal Behaviour 65, 821-840.
569	15.	Clutton-Brock, T.H., and Isvaran, K. (2007). Sex differences in ageing in natural
570		populations of vertebrates. Proceedings of the Royal Society B: Biological Sciences
571		274, 3097-3104.

572 573	16.	Kirkwood, T.B., and Rose, M.R. (1991). Evolution of senescence: late survival sacrificed for reproduction. Philosophical Transactions of the Royal Society of
574		London. Series B: Biological Sciences <i>332</i> , 15-24.
575	17.	Williams, G.C. (1957). Pleiotropy, natural selection, and the evolution of senescence.
576		evolution, 398-411.
577	18.	Maegawa, S., Lu, Y., Tahara, T., Lee, J.T., Madzo, J., Liang, S., Jelinek, J., Colman, R.J.,
578		and Issa, JP.J. (2017). Caloric restriction delays age-related methylation drift.
579		Nature communications <i>8</i> , 539.
580 581	19.	Ryan, J., Wrigglesworth, J., Loong, J., Fransquet, P.D., and Woods, R.L. (2019). A systematic review and meta-analysis of environmental, lifestyle and health factors
582		associated with DNA methylation age. The journals of gerontology. Series A,
583		Biological sciences and medical sciences.
584	20.	Tung, J., Archie, E.A., Altmann, J., and Alberts, S.C. (2016). Cumulative early life
585		adversity predicts longevity in wild baboons. Nature communications 7, 11181.
586	21.	Zipple, M.N., Archie, E.A., Tung, J., Altmann, J., and Alberts, S.C. (2019).
587		Intergenerational effects of early adversity on survival in wild baboons. eLife 8,
588		e47433.
589	22.	Archie, E.A., Tung, J., Clark, M., Altmann, J., and Alberts, S.C. (2014). Social affiliation
590		matters: both same-sex and opposite-sex relationships predict survival in wild
591		female baboons. Proceedings of the Royal Society B: Biological Sciences 281,
592		20141261.
593	23.	Holt-Lunstad, J., Smith, T.B., and Layton, J.B. (2010). Social relationships and
594		mortality risk: a meta-analytic review. PLoS Med 7, e1000316.
595	24.	Snyder-Mackler, N., Burger, J.R., Gaydosh, L., Belsky, D., Noppert, G.A., Campos, F.A.,
596		Bartolomucci, A., Yang, Y.C., Aiello, A.E., O'Rand, A., et al. Social determinants of
597		health and survival in humans and other animals. In review.
598	25.	Alberts, S.C., Buchan, J.C., and Altmann, J. (2006). Sexual selection in wild baboons:
599		from mating opportunities to paternity success. Animal Behaviour 72, 1177-1196.
600	26.	Gesquiere, L.R., Altmann, J., Archie, E.A., and Alberts, S.C. (2018). Interbirth intervals
601		in wild baboons: Environmental predictors and hormonal correlates. American
602		journal of physical anthropology <i>166</i> , 107-126.
603	27.	Lea, A.J., Altmann, J., Alberts, S.C., and Tung, J. (2015). Developmental constraints in
604		a wild primate. The American Naturalist 185, 809-821.
605	28.	Lea, A.J., Akinyi, M.Y., Nyakundi, R., Mareri, P., Nyundo, F., Kariuki, T., Alberts, S.C.,
606		Archie, E.A., and Tung, J. (2018). Dominance rank-associated gene expression is
607		widespread, sex-specific, and a precursor to high social status in wild male baboons.
608		Proceedings of the National Academy of Sciences <i>115</i> , E12163-E12171.
609	29.	Castagné, R., Garès, V., Karimi, M., Chadeau-Hyam, M., Vineis, P., Delpierre, C., Kelly-
610		Irving, M., and Consortium, L. (2018). Allostatic load and subsequent all-cause
611		mortality: which biological markers drive the relationship? Findings from a UK birth
612		cohort. European journal of epidemiology 33, 441-458.
613	30.	Levine, M.E. (2012). Modeling the rate of senescence: can estimated biological age
614		predict mortality more accurately than chronological age? Journals of Gerontology
615		Series A: Biomedical Sciences and Medical Sciences <i>68</i> . 667-674.
616	31.	López-Otín, C., Blasco, M.A., Partridge, L., Serrano, M., and Kroemer, G. (2013). The
617		hallmarks of aging. Cell <i>153</i> , 1194-1217.

618 32. Lea, A.J., Tung, J., and Zhou, X. (2015). A flexible, efficient binomial mixed model for
619 identifying differential DNA methylation in bisulfite sequencing data. PLoS genetics
620 11, e1005650.

- 621 33. Lea, A.J., Vockley, C.M., Johnston, R.A., Del Carpio, C.A., Barreiro, L.B., Reddy, T.E., and
 622 Tung, J. (2018). Genome-wide quantification of the effects of DNA methylation on
 623 human gene regulation. eLife 7, e37513.
- Liu, Z., Chen, B.H., Assimes, T.L., Ferrucci, L., Horvath, S., and Levine, M.E. (2019). The
 role of epigenetic aging in education and racial/ethnic mortality disparities among
 older US Women. Psychoneuroendocrinology *104*, 18-24.
- Shalev, I., and Belsky, J. (2016). Early-life stress and reproductive cost: A two-hit
 developmental model of accelerated aging? Medical Hypotheses *90*, 41-47.
- Brody, G.H., Miller, G.E., Yu, T., Beach, S.R., and Chen, E. (2016). Supportive family
 environments ameliorate the link between racial discrimination and epigenetic
 aging: A replication across two longitudinal cohorts. Psychological Science 27, 530541.
- Brody, G.H., Yu, T., Chen, E., Beach, S.R., and Miller, G.E. (2016). Family centered
 prevention ameliorates the longitudinal association between risky family processes
 and epigenetic aging. Journal of child psychology and psychiatry *57*, 566-574.
- Bavis, E., Humphreys, K., McEwen, L., Sacchet, M., Camacho, M., MacIsaac, J., Lin, D.,
 Kobor, M., and Gotlib, I. (2017). Accelerated DNA methylation age in adolescent
 girls: associations with elevated diurnal cortisol and reduced hippocampal volume.
 Translational psychiatry 7, e1223.
- Jovanovic, T., Vance, L.A., Cross, D., Knight, A.K., Kilaru, V., Michopoulos, V., Klengel,
 T., and Smith, A.K. (2017). Exposure to violence accelerates epigenetic aging in
 children. Scientific reports 7, 8962.
- Marini, S., Davis, K.A., Soare, T.W., Suderman, M.J., Simpkin, A.J., Smith, A.D., Wolf, E.J.,
 Relton, C.L., and Dunn, E.C. (2018). Predicting cellular aging following exposure to
 adversity: Does accumulation, recency, or developmental timing of exposure
 matter? BioRxiv, 355743.
- 647 41. Sumner, J.A., Colich, N.L., Uddin, M., Armstrong, D., and McLaughlin, K.A. (2019).
 648 Early experiences of threat, but not deprivation, are associated with accelerated
 649 biological aging in children and adolescents. Biological psychiatry *85*, 268-278.
- Austin, M.K., Chen, E., Ross, K.M., McEwen, L.M., Maclsaac, J.L., Kobor, M.S., and
 Miller, G.E. (2018). Early-life socioeconomic disadvantage, not current, predicts
 accelerated epigenetic aging of monocytes. Psychoneuroendocrinology *97*, 131-134.
- 43. Boks, M.P., van Mierlo, H.C., Rutten, B.P., Radstake, T.R., De Witte, L., Geuze, E.,
 Horvath, S., Schalkwyk, L.C., Vinkers, C.H., and Broen, J.C. (2015). Longitudinal
 changes of telomere length and epigenetic age related to traumatic stress and posttraumatic stress disorder. Psychoneuroendocrinology *51*, 506-512.
- Lawn, R.B., Anderson, E.L., Suderman, M., Simpkin, A.J., Gaunt, T.R., Teschendorff,
 A.E., Widschwendter, M., Hardy, R., Kuh, D., and Relton, C.L. (2018). Psychosocial
 adversity and socioeconomic position during childhood and epigenetic age: analysis
 of two prospective cohort studies. Human molecular genetics 27, 1301-1308.
- 45. Simons, R.L., Lei, M.K., Beach, S.R., Philibert, R.A., Cutrona, C.E., Gibbons, F.X., and
- Barr, A. (2016). Economic hardship and biological weathering: the epigenetics of

663		aging in a US sample of black women. Social Science & Medicine 150, 192-200.
664	46.	Wolf, E.J., Maniates, H., Nugent, N., Maihofer, A.X., Armstrong, D., Ratanatharathorn,
665		A., Ashley-Koch, A.E., Garrett, M., Kimbrel, N.A., and Lori, A. (2018). Traumatic stress
666		and accelerated DNA methylation age: a meta-analysis. Psychoneuroendocrinology
667		92, 123-134.
668	47.	Zannas, A.S., Arloth, J., Carrillo-Roa, T., Iurato, S., Röh, S., Ressler, K.J., Nemeroff, C.B.,
669		Smith, A.K., Bradley, B., and Heim, C. (2015), Lifetime stress accelerates epigenetic
670		aging in an urban. African American cohort: relevance of glucocorticoid signaling.
671		Genome biology 16. 266.
672	48.	Aristizabal, M.I., Anreiter, I., Halldorsdottir, T., Odgers, C.L., McDade, T.W.,
673		Goldenberg, A., Mostafavi, S., Kobor, M.S., Binder, E.B., and Sokolowski, M.B. (2019).
674		Biological embedding of experience: A primer on epigenetics. Proceedings of the
675		National Academy of Sciences 201820838
676	49	Hertzman ((2012) Putting the concent of biological embedding in historical
677	17.	nerspective Proceedings of the National Academy of Sciences 109 17160-17167
678	50	Ben-Shlomo V and Kub D (2002) A life course approach to chronic disease
679	50.	enidemiology: concentual models, empirical challenges and interdisciplinary
680		nerspectives (Oxford University Press)
681	51	Shanahan I. Coneland W.F. Costello F.I. and Angold A (2011) Child- adolescent-
682	51.	and young adult-onset depressions: differential risk factors in development?
683		Psychological medicine 41, 2265-2274
684	52	Shanahan M L and Hofer S M (2011) Molecular genetics aging and well-being
685	52.	Sensitive period accumulation and nathway models. In Handbook of aging and the
686		social sciences (Flsevier) nn 135-147
687	53	Cosquiere L.R. Learn N.H. Simoo M.C.M. Onvango P.O. Alberts S.C. and Altmann
688	55.	L (2011) Life at the top: rank and stress in wild male behoons. Science 222, 257-
680		360
690	54	Campos F A Villavicencio F Archie F A Colchero F and Alberts S C (2020)
691	54.	Social relationships social status and survival in wild baboons: A tale of two seves
602		Dhilosophical Transactions of the Doval Society Series B: Biological Sciences
603	55	Archia F A Altmann I and Alberts SC (2012) Social status products wound
604	55.	healing in wild behaving. Drogoodings of the National Academy of Sciences 100
605		0017 0022
695	56	Pasmusson K (1985) Changes in the activity hudgets of vellow haboons (Panio
607	50.	Rasinussen, R. (1965). Changes in the activity budgets of yenow baboons (Fapio
6097		cynocephaius) during sexual consortsnips. Benavioral Ecology and Sociobiology 17,
600	57	101-170. Alberta S.C. and Altmann I. (2001). Immigration and hybridization patterns of
700	57.	Alberts, S.C., and Alumann, J. (2001). Infining attoin and hyperbolic Variation patterns of
700		Drimatology, Official Journal of the American Society of Drimatologists 52, 120, 154
701	F 0	Tung L Charmontian ML Carfield D.A. Altmann L and Alberta S.C. (2000). Caractia
702	58.	rung, J., Charpentier, M.J., Garneid, D.A., Aitmann, J., and Alberts, S.C. (2008). Genetic
703		evidence reveals temporal change in hybridization patterns in a wild baboon
704	50	Alter and L Alberta C.C. Heiner C.A. Dubech L. Munuthi D. Costa T. Coffen E.
705	57.	Alumanni, J., Alberts, S.C., Halnes, S.A., Dubach, J., Muruthl, P., Coote, L., Genen, E.,
700		cheesman, D.J., Mututua, K.S., and Salyalei, S.N. (1996). Benavior predicts genes
/0/		structure in a wild primate group. Proceedings of the National Academy of Sciences
708		<i>73</i> , 5777-5801.

709	60.	Tung, J., Zhou, X., Alberts, S.C., Stephens, M., and Gilad, Y. (2015). The genetic
710		architecture of gene expression levels in wild baboons. Elife 4, e04729.
711	61.	Meissner, A., Gnirke, A., Bell, G.W., Ramsahoye, B., Lander, E.S., and Jaenisch, R.
712		(2005). Reduced representation bisulfite sequencing for comparative high-
713		resolution DNA methylation analysis. Nucleic acids research 33, 5868-5877.
714	62.	Boyle, P., Clement, K., Gu, H., Smith, Z.D., Ziller, M., Fostel, J.L., Holmes, L., Meldrim, J.,
715		Kelley, F., and Gnirke, A. (2012). Gel-free multiplexed reduced representation
716		bisulfite sequencing for large-scale DNA methylation profiling. Genome biology 13,
717		R92.
718	63.	Krueger, F. (2012). Trim Galore: a wrapper tool around Cutadapt and FastQC to
719		consistently apply quality and adapter trimming to FastO files, with some extra
720		functionality for MspI-digested RRBS-type (Reduced Representation Bisufite-Seq)
721		libraries. URL http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/.
722	64.	Xi, Y., and Li, W. (2009). BSMAP: whole genome bisulfite sequence MAPping
723		program. BMC bioinformatics 10, 232.
724	65.	Krueger, F. (2015). Trim Galore!: A wrapper tool around Cutadapt and FastQC to
725		consistently apply quality and adapter trimming to FastQ files. (0.4).
726	66.	Hastie, T., Tibshirani, R., Narasimhan, B., and Chu, G. (2001). impute: Imputation for
727		microarray data. Bioinformatics 17, 520-525.
728	67.	Friedman, J., Hastie, T., and Tibshirani, R. (2009). glmnet: Lasso and elastic-net
729		regularized generalized linear models. R package version 1.
730	68.	Friedman, J., Hastie, T., and Tibshirani, R. (2010). Regularization paths for
731		generalized linear models via coordinate descent. Journal of statistical software 33,
732		1.
733	69.	Snyder-Mackler, N., Sanz, J., Kohn, J.N., Brinkworth, J.F., Morrow, S., Shaver, A.O.,
734		Grenier, JC., Pique-Regi, R., Johnson, Z.P., and Wilson, M.E. (2016). Social status
735		alters immune regulation and response to infection in macaques. Science 354, 1041-
736		1045.
737	70.	Altmann, J., Schoeller, D., Altmann, S.A., Muruthi, P., and Sapolsky, R.M. (1993). Body
738		size and fatness of free - living baboons reflect food availability and activity levels.
739		American Journal of Primatology <i>30</i> , 149-161.
740	71.	Silk, J.B., Beehner, J.C., Bergman, T.J., Crockford, C., Engh, A.L., Moscovice, L.R., Wittig,
741		R.M., Seyfarth, R.M., and Cheney, D.L. (2010). Strong and consistent social bonds
742		enhance the longevity of female baboons. Current Biology 20, 1359-1361.
743	72.	Hausfater, G., Altmann, J., and Altmann, S. (1982). Long-term consistency of
744		dominance relations among female baboons (Papio cynocephalus). Science 217,
745		752-755.
746	73.	Archie, E.A., Altmann, J., and Alberts, S.C. (2014). Costs of reproduction in a long-
747		lived female primate: injury risk and wound healing. Behavioral ecology and
748		sociobiology <i>68</i> , 1183-1193.
749	74.	Beehner, J.C., Onderdonk, D.A., Alberts, S.C., and Altmann, J. (2006). The ecology of
750		conception and pregnancy failure in wild baboons. Behavioral Ecology 17, 741-750.
751	75.	Altmann, J., and Alberts, S.C. (2003). Intraspecific variability in fertility and offspring
752		survival in a nonhuman primate: behavioral control of ecological and social sources.
753		In Offspring: The Biodemography of Fertility and Family Behavior, K.W. Wachter

754		and R.A. Bulatao, eds. (Washington, DC: The National Academies Press).
755	76.	Altmann, J., Hausfater, G., and Altmann, S.A. (1988). Determinants of Reproductive
756		Success in Savannah Baboons, Papio cynocephalus. In Reproductive Success: Studies
757		of Individual Variation in Contrasting Breeding Systems, T.H. Clutton-Brock, ed.
758		(Chicago: The University of Chicago Press).
759	77.	Charpentier, M., Tung, J., Altmann, J., and Alberts, S. (2008). Age at maturity in wild
760		baboons: genetic, environmental and demographic influences. Molecular Ecology 17,
761		2026-2040.
762	78.	Altmann, J., Altmann, S.A., and Hausfater, G. (1978). Primate infant's effects on
763		mother's future reproduction. Science <i>201</i> , 1028-1030.
764	79.	Altmann, J., and Alberts, S.C. (2003). Variability in reproductive success viewed from
765		a life - history perspective in baboons. American Journal of Human Biology 15, 401-
766		409.
767	80.	Lea, A.J., Learn, N.H., Theus, M.J., Altmann, J., and Alberts, S.C. (2014). Complex
768		sources of variance in female dominance rank in a nepotistic society. Animal
769		behaviour <i>94</i> , 87-99.
770	81.	Vilgalys, T.P., Rogers, J., Jolly, C.J., Mukherjee, S., and Tung, J. (2018). Evolution of
771		DNA methylation in Papio baboons. Molecular biology and evolution <i>36</i> , 527-540.
772	82.	Irizarry, R.A., Ladd-Acosta, C., Wen, B., Wu, Z., Montano, C., Onyango, P., Cui, H., Gabo,
773		K., Rongione, M., and Webster, M. (2009). The human colon cancer methylome
774		shows similar hypo-and hypermethylation at conserved tissue-specific CpG island
775		shores. Nature genetics 41, 178.
776	83.	Consortium, E.P. (2012). An integrated encyclopedia of DNA elements in the human
777		genome. Nature 489, 57.
778		