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1 Heritable variation in telomere length predicts mortality in

2 Soay sheep

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24 Author Contributions

HF & DHN designed the study and wrote the first draft of the manuscript; JMP & JGP
conducted fieldwork and managed the long-term study; SLU, JD, LAS, KW & RVW
conducted the labwork; HF conducted the analyses; all authors contributed to the revised
manuscript.

- 29 This PDF file includes: Main Text; Figures 1 to 3.
- 30

31 Abstract

32 Telomere length (TL) is considered an important biomarker of whole-organism health and ageing. Across humans and other vertebrates, short telomeres are associated with 33 increased subsequent mortality risk, but the processes responsible for this correlation 34 remain uncertain. A key unanswered question is whether TL-mortality associations arise 35 36 due to positive effects of genes or early-life environment on both an individual's average 37 lifetime TL and their longevity, or due to more immediate effects of environmental stressors 38 on within-individual TL loss and increased mortality risk. Addressing this guestion requires 39 longitudinal TL and life-history data across the entire lifetimes of many individuals, which are difficult to obtain for long-lived species like humans. Using longitudinal data and samples 40 collected over nearly two decades, as part of a long-term study of wild Soay sheep, we 41 42 dissected an observed positive association between TL and subsequent survival using multivariate quantitative genetic models. We found no evidence that telomere attrition was 43 44 associated with increased mortality risk, suggesting that TL is not an important marker of 45 biological ageing or exposure to environmental stress in our study system. Instead, we find that among-individual differences in average TL are associated with increased lifespan. Our 46 analyses suggest that this correlation between an individual's average TL and lifespan has a 47 genetic basis. This demonstrates that TL has the potential to evolve under natural 48 conditions, and suggests an important role of genetics underlying the widespread 49 observation that short telomeres predict mortality. 50

51 Significance Statement

Telomeres play an important role in ageing and having relatively short telomeres is 52 associated with an increased risk of death in humans and other animals. Telomere length is 53 influenced by both genetic and environmental factors, both of which could potentially drive 54 the positive association with survival. We used lifelong telomere length measurements from 55 a population of wild sheep to disentangle this relationship. For the first time in a natural 56 population, our analyses reveal a genetic correlation between telomere length and longevity 57 58 but no association between telomere shortening and mortality risk. These findings have 59 important implications for our understanding of telomere dynamics and their role in health 60 and lifespan.

61

62 Main text

63 Introduction

Telomeres are repetitive sequences of non-coding DNA found at the terminal ends of linear 64 chromosomes, and they play an important role in maintaining DNA stability and integrity (1-65 3). Telomeres shorten during cell replication and in response to oxidative stress (4, 5), and 66 cellular senescence and apoptosis is triggered once telomeres reach a critically short 67 68 threshold (2). The important role of telomeres in cellular senescence has led to telomere shortening being considered as one of nine "hallmarks of ageing", and average telomere 69 length (TL) as an important biomarker of whole-organism health and biological ageing (6). In 70 71 humans, relatively short leukocyte telomeres have been linked to a range of age-related 72 diseases such as diabetes, cancer and cardiovascular disease (7-9) and increased subsequent mortality risk (10-12). A recent meta-analysis suggests this pattern may 73 generalise beyond humans: across studies from 20 non-model vertebrate species 74 (predominantly birds) there was an overall positive association between telomere length and 75 76 subsequent survival (13). Although evidence for a causal role for telomeres in whole-77 organism ageing and longevity remains weak (14), these findings highlight the potential 78 significance of TL as a biomarker of human and animal health (15, 16) and for our 79 understanding of life-history evolution (17, 18).

80 Studies in humans and other vertebrates have found evidence for consistent differences in 81 TL among individuals over multiple measurements (19, 20). Such repeatable amongindividual differences in any trait may result from the trait being under genetic influence, from 82 long-term effects of the early-life environment, and/or environmental conditions that persist 83 across the lifetime. There is good evidence that variation in average TL in blood cells has a 84 genetic basis in humans and other vertebrates, although estimates of the heritability (the 85 proportion of variation attributed to additive genetic effects) of TL are variable (21, 22). 86 Recent studies of wild vertebrates have also revealed considerable variation in adult TL 87 among birth cohorts, suggesting persistent impacts of early-life environment (23, 24). At the 88 89 same time, there is growing evidence that TL is highly dynamic across an individual's 90 lifetime, and meta-analyses of human and non-human animal studies show that experience 91 of diverse forms of environmental stress are predictive of shorter TL (25-27). Indeed, some 92 studies using longitudinal TL data have found that telomere shortening over successive 93 measurements rather than telomere length per se is predictive of mortality (28-30). Thus, the 94 emerging picture from studies in humans and other vertebrates is that shorter TL generally

predicts increased risk of subsequent mortality, and that variation in TL is under theinfluence of both genetics and environmental stressors.

The observation that shorter TL measurements predict increased mortality risk could be 97 underpinned by two, non-mutually exclusive processes operating across the lifetimes of 98 individuals. Firstly, individuals may differ in their average TL across life, and individuals with 99 100 shorter TL may be shorter lived. This pattern is referred to as the 'selective disappearance' 101 of individuals with shorter telomeres, and it implies that TL reflects constitutive differences among individuals (for example due to genetics or differences in early-life environment) 102 which shape their longevity (31, 32). Secondly, individuals may differ in their pattern of TL 103 104 change over time, and individuals showing the greatest telomere loss across successive measurements are more likely to die subsequently. This pattern is consistent with the idea 105 that within-individual telomere dynamics reflect recent and cumulative experiences of 106 environmental stress and physiological deterioration that also predict mortality. Neither 107 108 pattern necessarily implies a causal role for telomeres in driving the mortality risk of an 109 organism, because associations between TL and survival could result from both traits being correlated with underlying, unmeasured variables which causally impact survival (14, 18). 110 111 Nevertheless, unravelling the contribution of genetics, early-life environment and more immediate telomere shortening to the observed association between TL and survival is 112 essential for our understanding of TL as a biomarker of health and ageing (19). 113

To our knowledge, no study to date has assessed the relative importance of the different 114 processes underlying the relationship between TL and mortality risk across the entire 115 lifespan. To do so demands repeated measurements from across life to characterise among-116 and within-individual variation in telomere length, a population pedigree or genomic 117 118 information to separate genetic and environmental sources of variation, and detailed 119 information on individual health and fitness outcomes over the lifetime. Here, we use a multivariate mixed-effects modelling approach to analyse extensive, longitudinal data from a 120 long-term study of wild Soay sheep living on St Kilda, Scotland, to distinguish between 121 122 possible models of why shorter TL predicts increased mortality risk. We find that the 123 observed positive association between TL and mortality in this system is underpinned by 124 selective disappearance of individuals with shorter average TL. Importantly, our results 125 suggest this is largely driven by genetically-based differences in both TL and longevity.

126 **Results**

127 Soay sheep resident to our study area on St Kilda have been individually marked at birth and 128 closely monitored and repeatedly blood sampled across their lifetimes. Here, we measured 129 relative telomere length (RTL) in 3641 samples collected from 1586 individual sheep over a 19-year period (see Methods). We found that RTL declined with age in Soay sheep, with a 130 more rapid initial decline between measurements at around 4 and 16 months, followed by a 131 slower linear decline thereafter (Figure 1). The best fitting age function in our models of RTL 132 included a two-level factor for age class (lambs and adults aged ≥1 year) and a linear term 133 for age in years, which is equivalent to a segmented regression with a threshold at one year 134 of age (SI Appendix, Tables S1 & S2). There was considerable variation in RTL within age 135 groups (SI Appendix, Fig. S1). There was limited evidence for a sex difference in either 136 average RTL or the rate of change in RTL with age, although the significance of sex in our 137 model depended on model structure (see SI Appendix). The individual repeatability of RTL 138 over the lifespan was 0.214 (95%CI 0.169-0.252, SI Appendix, Table S2). Excluding the 139 variance attributed to qPCR plate and row (which represents measurement error) from the 140 141 denominator, the repeatability was 0.241 (95%CI 0.204-0.282, SI Appendix, Table S2). 142 Although RTL declined with age on average, we found evidence for consistent differences in 143 RTL among individuals.

There was detectable additive genetic variance for relative telomere length across all ages in 144 the population (Figure 2; SI Appendix, Table S3). The heritability of telomere length (the 145 proportion of variance explained by additive genetic effects) was 0.204 (95%CI: 0.158-146 0.252). The permanent environment effect was bound at zero (<0.001, 95%CI: <0.001-147 0.017), indicating that individual repeatability in telomere length could be almost entirely be 148 attributed to genetic rather than environmental effects. Maternal effects also explained a very 149 small proportion of the variance (<0.001, 95%CI <0.001–0.034, estimate bound at zero). The 150 year the sample was collected, qPCR plate and qPCR row each explained 3-4% of the 151 variance in telomere length (year: 0.031, 95%CI 0.014-0.084; gPCR plate: 0.043, 0.027-152 0.066; qPCR row: 0.038, 0.014–0.172). Excluding the measurement error terms of qPCR 153 plate and row from the total phenotypic variance, the heritability of telomere length was 154 0.233 (95%CI 0.189–0.279). This shows that variation in telomere length has a genetic basis 155 156 in wild Soay sheep.

We went on to estimate the genetic correlation between RTL expressed in lambs, which are still developing at the time of measurement (aged 4 months), and in older individuals, which are sexually mature and have largely completed growth. When estimated with a bivariate model of lamb and adult telomere length, the heritability of telomere length in lambs was estimated to be 0.285 (95%CI 0.206–0.369), and in adults 0.210 (95%CI: 0.156–0.255)

162 (Figure 2; SI Appendix, Table S4). The genetic correlation between lambs and adults was 163 close to 1 (0.916, 95%CI: 0.806-0.996, SI Appendix, Table S4), implying that largely the 164 same or linked genes influenced RTL across age groups. The residual correlation between lamb and adult telomere length was close to zero (0.036, 95% CI: -0.196-0.008, SI 165 166 Appendix, Table S4). The negligible residual correlation was expected given the lack of permanent environment effect underlying repeatable differences in individual RTL across all 167 ages (SI Appendix, Table S3). Repeatable among-individual differences across ages in RTL 168 were therefore driven predominantly by genetic rather than environmental effects, and a 169 similar set of genes influenced RTL across ages. 170

171 Using multivariate mixed-effects models, we next tested the strength and direction of correlations among RTL, body weight (both measured in August) and subsequent overwinter 172 173 survival at different hierarchical levels (among-individual, genetic, among-year, withinindividual). Although we were primarily interested in the RTL-survival association, the 174 175 inclusion of weight in our models allowed us to contrast the magnitude of the RTL-survival 176 association with a well-studied trait which is known to be linked to condition and fitness in our study system (33). As previously documented, August body weight was strongly 177 predictive of improved winter survival prospects ((33-35); Figure 3). Our multivariate models 178 revealed that this association was present at all hierarchical levels (Figure 3; SI Appendix, 179 Tables S5 & S6). Our initial phenotypic model indicated that heavier individuals on average 180 tended to have longer lifespans (Figure 3A; SI Appendix, Tables S5). Developing this model 181 into a pedigree-based quantitative genetic model revealed that this among-individual effect 182 was driven by similar sized contributions from genes and environment (Figure 3B; SI 183 Appendix, Table S6). Overwinter survival probabilities were higher in years where the 184 average body weight was higher (among-year effect), and individuals with relatively low 185 weight compared to their average weight were less likely to survive (residual effect; Figure 186 3). In our phenotypic model, there was little evidence for an association between RTL and 187 body weight at any hierarchical level (Figure 3A; SI Appendix, Table S5). However, the 188 189 quantitative genetic model revealed a negative genetic correlation (Figure 3B; SI Appendix, 190 Table S6). This was consistent in separate models of lambs and adults, although here credible intervals overlapped zero more widely (SI Appendix, Tables S7-S10). 191

We found a positive association between telomere length and overwinter survival probability at the among-individual level (Figure 3A; SI Appendix, Table S5). This suggests that individuals with longer telomeres on average across their lives tended to have longer lifespans. The 95% credible intervals overlapped zero for both the among-year and the residual covariance between telomere length and overwinter survival (Figure 3A; SI Appendix, Table S5). The absence of an association at the within-individual (or residual) level indicates that if an individual had a relatively short RTL measurement compared to their average in a particular year, this was not associated with an increased risk of mortality. Overall, these findings imply that average telomere length after weaning, rather than more immediate changes relative to an individual's average TL, predicted overwinter survival.

202 The quantitative genetic model revealed a positive association between telomere length and 203 overwinter survival probability at the genetic level, though the credible intervals narrowly 204 overlapped zero (Figure 3B; SI Appendix, Table S6). The covariance between TL and survival was very small (<0.001) at the permanent environment level (SI Appendix, Table 205 206 S6) and the correlation at this level had wide credible intervals which extensively crossed zero (Figure 3B). Our results were broadly consistent for lambs and adults across separate 207 208 models, although the credible intervals of the among-individual and genetic effects did narrowly cross zero within some age groups (Figures S2 & S3; SI Appendix, methods & 209 210 results, Tables S7-10). Overall, our results suggest that the among-individual positive 211 association between telomere length and survival identified in the phenotypic model was driven largely by genetic effects (Figure 3B; SI Appendix, Table S6). In other words, genes 212 213 conferring longer average telomeres after weaning also tended to be associated with longer lifespan. 214

215 **Discussion**

A growing number of studies across vertebrate species have shown that TL is heritable (21) 216 and that relatively short TL is predictive of increased mortality risk (13). We have dissected 217 the TL-mortality relationship across the natural lifespan of wild Soay sheep to identify the 218 processes that drive this association. We show that selective disappearance of individuals 219 220 with shorter average TL is the key process, rather than an association between the pattern of TL attrition and mortality. If TL was a marker of biological ageing, or it reflected variation in 221 222 condition resulting from environmental stress, we would expect telomere shortening to predict mortality, resulting in a positive residual correlation between TL and survival in our 223 224 models. We found no support for a correlation at the residual level, and only the amongindividual correlations between TL and survival were consistently different from zero. This 225 226 suggests that TL is not a useful marker of biological ageing in our system. Furthermore, it is 227 inconsistent with the hypothesis that TL reflects variation in physiological state or condition 228 resulting from recent or accumulating experience of environmental stress during adulthood, 229 which also influences mortality risk. However, our first TL measurement was taken at four 230 months of age – around the time of weaning for most lambs in our study population – and it is possible that associations between average TL and survival are driven by some combination of initial TL and the rate of telomere attrition prior to first measurement. An individual's average TL could therefore reflect its ability to maintain homeostasis and resist environmental stress during very early development, which in turn could predict its subsequent lifespan. Nonetheless, our results demonstrate that variation in TL is heritable and under directional selection, meaning it has the potential to evolve under natural conditions.

238 Our moderate estimate of the heritability of TL (around 20%, Figure 2) represents the first evidence for an appreciable genetic contribution to variation in TL in a wild mammal. Recent 239 240 studies in wild badgers and bats have used similar pedigree-based 'animal models' but found both the repeatability and heritability of TL to be negligible (36, 37). Our estimate was 241 242 lower than in a recent longitudinal study of farmed dairy cattle (32-38%; (38)), which is unsurprising given the expectation of reduced environmental variation in livestock compared 243 244 to wild systems. It is notable that both studies found negligible permanent environment 245 effects, implying that consistent differences in TL across life were largely the result of genetic rather than early environmental effects in both systems ((38); Figure 2). More broadly, while 246 247 human studies have tended to find moderate to high heritability of TL, the growing literature in birds presents a much more variable picture (21). Two laboratory studies of birds have 248 estimated the heritability of TL to be >100% (39, 40), whilst some studies in the wild suggest 249 little genetic influence on TL (heritability <5% (41, 42)). The reasons for the variation in 250 heritability estimates across studies remain an important area for future research. As well as 251 variation in the way that genes influence telomere dynamics, differences in the 252 environmental variation experienced by different populations or species, and the degree of 253 254 error associated with different telomere measurement methodologies are also likely to play a role (21). Our study used a qPCR method to measure TL, which is often described as having 255 greater measurement error than the 'gold standard' terminal restriction fragment (TRF) 256 approach (43). However, we note that the technical repeatability of our qPCR assay is high 257 and compares favorably to those reported for TRF studies (see Methods). Furthermore, 258 most previous studies in humans and birds have focused on specific age classes (e.g. 259 elderly humans or pre-fledging birds). Our data spans the entire lifetimes of individuals, and 260 261 we were able to demonstrate a very high genetic correlation between TL measured in lambs and adults. This provides clear evidence that the same or linked genes influence TL in early 262 and later life, an assumption that has rarely been tested in any study system (though see 263 264 (44)).

265 In a previous study, we identified an association between TL and mortality in a much smaller 266 sample of female Soay sheep (45). In that study, we reported a considerably lower 267 repeatability for TL (13%) than in the present study and were only able to detect a TLmortality relationship among younger females experiencing a high mortality winter. This 268 269 previous study was focused on a subset of females from four birth cohorts, and thus had relatively few samples from later adulthood (45). Our present study is distinguished by the 270 much larger data set and more complete population coverage, which has allowed us to 271 assess the repeatability of TL and determine that it is the among-individual differences in TL 272 across life that are predictive of mortality. We note that, when splitting our multivariate 273 models by age groups, the among-individual covariance between TL and survival was 274 positive and of a similar magnitude in adults and lambs, though the credible intervals 275 overlapped zero (SI Appendix, Figs. S2 & S3; Table S4). This indicates that the association 276 277 was not simply due to effects on lamb survival. Although some longitudinal vertebrate studies have found that telomere attrition predicted survival better than average or recent TL 278 (28, 29, 46), others have identified extremely high consistency in individual TL across 279 280 measurements (19, 20) and long-term associations between early-life TL and adult lifespan 281 (24, 47, 48). Most of these studies involved only one or two measurements of TL per 282 individual, and none used a multivariate mixed-effects modelling approach capable of fully 283 dissecting the contributions of genetic, individual, annual and residual sources to observed 284 TL-survival covariance.

Recent studies of elderly human cohorts have identified candidate SNP loci associated with 285 telomerase genes, which are involved in maintenance of telomeres and genomic integrity, 286 287 that not only predict average leukocyte TL but also subsequent morbidity and mortality (49-51). However, human twin studies also suggest that relatively short TL, independent of 288 genetic factors, predicts mortality (52). In support of a causal role for telomerase genetics in 289 mortality and ageing, studies of laboratory mice without telomerase appear to show early 290 291 onset of ageing phenotypes, while mice with genetically enlarged telomeres are longer lived 292 (53, 54). Our findings provide the first support for a role for genetics in observed TL-mortality 293 relationships from a non-human system outside the laboratory. Further work is required to determine whether specific genes, such as those involved in telomere maintenance, are 294 295 implicated in this relationship or whether the observed genetic correlations result from minor 296 effects of many different genes.

297 Our results imply that the observed relationship between TL and mortality is not causal. A 298 causal effect of short telomeres on survival would lead to the expectation that both 299 individuals with low average TL (among-individual covariance) and short TL at measurement 300 relative to their average (residual covariance) should be positively related to survival. But this 301 is not what we observed. Instead, it seems likely that the genes influencing some yet to be 302 determined aspect of an individual's overall frailty (or robustness) have a correlated influence on TL. These genes could be influencing TL determined during early embryonic 303 304 development and/or the rate of telomere attrition during gestation and neonatal life, as our 305 first TL measurement was not taken until around four months of age. However, comparing the magnitude of the TL-survival correlations with the correlations between body weight and 306 survival highlights that the association between TL and mortality is modest compared to a 307 measure that is more directly relatable to condition and health. It remains to be determined 308 whether our findings in Soay sheep will generalize to other species and systems, but our 309 work highlights the importance of using large data sets that span the entire lifetimes of many 310 individuals, in order to fully understand the drivers of associations between TL, health and 311 312 mortality risk. Overall, our results provide important insights into the genetics of lifespan in the wild and highlight the importance of long-term, longitudinal studies across different 313 species for our understanding of TL as a biomarker of health and fitness. 314

315 Materials and methods

316 Study system and sample collection

The Soay sheep (Ovis aries) is a primitive breed of domestic sheep that has been living on 317 the remote St Kilda archipelago with minimal human management for the last few millennia 318 (57°49'N, 8°34'W; (33)). Since 1985, the sheep resident within the Village Bay area of the 319 main island in the archipelago, Hirta, have been the subject of an individual-based study 320 321 (33). Individuals are caught and tagged within a few days of birth in the spring. Ten censuses are conducted on each of three annual field seasons, during spring (March -322 323 April), summer (July – August) and autumn (October – November), meaning the timing of 324 individual's disappearance is known with a high degree of accuracy (33). The vast majority 325 of sheep mortality occurs in late winter (85% of adult deaths occur January – April), and daily 326 carcass searches during this period mean that death dates are known to the nearest month 327 for most individuals. Each August, 50-60% of the resident population are caught in 328 temporary corral traps and blood sampled. Blood is collected from each individual into 9 ml lithium heparin Vacuettes and kept in a cool box or fridge from the point of sampling. The 329 330 blood is processed within 24 hours to separate the plasma and buffy coat fractions. The Vacuette is then spun at 1,008 x g for 10 minutes, the plasma layer drawn off and replaced 331 by the same quantity of 0.9% NaCl solution, gently mixed and spun again at 1,008 x g for 10 332 minutes. The intermediate buffy coat layer, comprising mainly white blood cells, is then 333

drawn off into a 1.5 ml Eppendorf tube and stored at -20 °C until used to assay leukocyte telomere length. All data collection was approved by the UK Home Office and carried out in accordance with the relevant guidelines.

337 Pedigree reconstruction

338 Parentage was inferred by genetic methods, except for some maternal links inferred by 339 observation (55). Multi-generational pedigree reconstruction was performed in the R package Sequoia (56), using 431 unlinked SNP markers. This likelihood-based approach 340 infers not only parent-offspring relationships, but also siblings and second-degree relatives. 341 In the resulting pedigree, a mother and/or father was assigned to 6082 individuals. After 342 343 pruning to only those individuals informative to the current analyses (using the R package MasterBayes (57)), the pedigree had a maximum depth of 13 generations and consisted of 344 2411 individuals, of which 2273 were non-founders and a total of 2050 maternities and 2172 345 346 paternities were assigned.

347 Sample selection and randomization

A total of 3891 August buffy coat samples from 1647 animals of known age collected 348 between 1998 and 2016 were selected for telomere length measurement. These samples 349 350 were selected from the total available buffy coat freezer archive (n=6775 from 3315 sheep) 351 based on the following exclusion criteria: samples collected before 1998 (n=1924); samples from individuals of unknown age (n=97); samples collected between 2013 and 2016 from 352 353 lambs or yearlings that were caught only once (n=454); samples collected from individuals that were only captured once but survived, and were therefore available for sampling in 354 355 subsequent years (n=409). These criteria were designed to maximise the longitudinality of the dataset while avoiding biasing the dataset against short-lived individuals (i.e. individuals 356 357 only sampled once because they died). Sample years were then randomly allocated to one 358 of four batches, each comprising 4–5 years (to reduce the number of samples that needed to 359 be removed from the freezer at any one time). Samples were then fully randomised within each batch, assigned a unique identifier from the start of batch 1 to the end of batch 4, and 360 361 processed from DNA extraction through to qPCR in this order.

362 **DNA extraction**

Genomic DNA was extracted from buffy coat on 96 well plates using the Macherey-Nagel Nucleospin 96 Blood kit (Cat# 740665). The samples were extracted on a liquid handling robot (Freedom Evo-2 150; Tecan) using a vacuum manifold. In order to facilitate passage of the sample through the DNA binding plate, the following step was included prior to automated extraction on the robot. 50 µl of buffy coat was mixed with 300 µl RBC lysis 368 solution (Qiagen; Cat# 158902) and incubated at room temperature for 5 minutes, before 369 centrifugation at 12200 rpm for 30 seconds. 250 µl of the supernatant was discarded and the 370 cell pellet was re-suspended in the residual supernatant, before it was transferred to a 96 well MN lysis block (Macherey-Nagel Nucleospin 96 Blood kit; Cat# 740665) and sealed. 371 The liquid handling robot was loaded with the MN lysis block with the seal removed, the 372 silica DNA binding plate, 100% Ethanol, lysis buffer BQ1, wash buffers B5 and BW and 373 finally a master mix containing PBS, proteinase K and RNAse A (binding plate, buffers and 374 proteinase K supplied with Macherey-Nagel Nucleospin 96 Blood kit, Cat# 740665; RNAse 375 A, Qiagen Cat# 158924; PBS, Sigma Cat# D1408). The extraction protocol followed 376 377 manufacturer's guidelines for use with a vacuum manifold with the following amendments. The robot added 96 µl 1X PBS, 25 µl Proteinase K and 4 µl RNAse A to each sample. Lysis 378 was performed on a shaker for 10 minutes. The vacuum steps for binding and the first two 379 washes were increased to 5 minutes and the final wash to 10 minutes with an additional 10-380 minute vacuum at the end to dry the membrane. If any lysate/wash failed to pass through the 381 382 silica membrane after a vacuum step the plate was removed from the robot and centrifuged 383 for 3 minutes at 4000 rpm. For the few samples that failed to pass though after centrifugation 384 the wash/lysate was removed by hand with a pipette. After dry centrifugation (3 minutes, 385 4000 rpm), DNA was eluted in a total of 150 µl elution buffer BE (Macherey-Nagel 386 Nucleospin 96 Blood kit; Cat# 740665) which was warmed to 60 °C prior to adding it onto the silica membrane. Elution was performed in two steps: first 100 µl buffer BE followed by 387 centrifugation (3 minutes, 4000 rpm), then 50 µl buffer BE followed by a final centrifugation 388 (3 minutes, 4000 rpm). 389

390 DNA extraction quality control

Following DNA extraction, a strict quality control protocol was implemented to measure DNA 391 concentration, integrity and purity (SI Appendix, Figure S4). Of the 3891 samples selected 392 393 for analysis, 42 samples were missing or accidentally omitted through human error. The 394 available samples were measured on a Nanodrop ND-1000 9 spectrophotometer (Thermo Scientific, Wilmington DE, USA). Samples yielding < 20 ng/µl were rejected, and re-395 396 extraction attempted for 491 of these samples. Overall, 87 samples failed to yield sufficient 397 DNA and were excluded from the analyses. Samples yielding $\geq 20 \text{ ng/}\mu\text{I}$ were checked for DNA purity. The acceptable range for absorption for 260/280 nm ratio (a measure of DNA vs 398 protein and RNA contamination) was 1.7-2.0. Samples falling outside of this range were 399 excluded from subsequent analyses (n=66, representing a failure rate of 1.76%). Two thirds 400 of samples were assayed for 260/230 nm ratio (a measure of salt and other impurities), and 401 samples with a ratio <1.8 were excluded from subsequent analyses (n=41, representing a 402 403 failure rate of 1.73%). Samples of sufficient yield and purity were standardized to 10 ng/ul

404 and DNA integrity was assessed by running 200 ng of DNA on a 0.5% agarose gel. Samples 405 were scored for integrity on a scale of 1 to 5 by visual examination of their DNA crowns, and 406 samples scoring 3 to 5 were excluded from further analyses (see (58)). Integrity was initially assayed for all of the first 1667 samples run. Only 16 of these samples were given a DNA 407 integrity score of 2, and 7 failed (failure rate of 0.42%). We randomly tested a guarter of all 408 subsequent samples for DNA integrity, with only 4 failing (representing a failure rate of 409 0.76%). In total, 11 samples were excluded due to poor DNA integrity. Overall, DNA was 410 successfully extracted and passed quality control requirements from 3644 buffy coat 411 samples (SI Appendix, Figure S4). 412

413 **Telomere length measurement**

414 Quantitative polymerase chain reaction

415 Relative leukocyte telomere length (RTL) was measured using real-time quantitative PCR (qPCR; (59)), using protocols we have previously developed and validated using blood 416 samples from sheep and cattle (45, 58). The qPCR method estimates the total amount of 417 telomeric sequence present in a sample relative to the amount of a non-variable copy 418 419 number reference gene. In this study we used the beta-2-microglobulin (B2M) as our reference gene (45, 58), with primers supplied by Primer Design (Cat# HK-SY-Sh-900, 420 Southampton, UK). For telomeric amplification tel1b (5'-CGG TTT GTT TGG GTT TGG GTT 421 TGG GTT TGG GTT TGG GTT-3') and tel 2b (5'-GGC TTG CCT TAC CCT TAC CCT TAC 422 CCT TAC CCT TAC CCT-3') primers were used. Telomere primers were manufactured, 423 HPLC purified and supplied by Integrated DNA Technologies (IDT, Glasgow, UK). Telomere 424 and reference gene reactions were run in separate wells of the same qPCR plate at a 425 concentration of 300 nM and 900 nM, respectively. Samples were diluted to 0.5 ng/µl with 426 buffer BE (Macherey-Nagel Nucleospin 96 Blood kit; Cat# 740665) immediately prior to 427 qPCR analysis. Each reaction was prepared using 5 µl of LightCycler 480SYBR Green I 428 Master Mix (Cat # 04887352001, Roche, West Sussex, UK) and 1 ng of sample DNA in a 429 total reaction volume of 10 µl. We used 384 well plates which were loaded with sample DNA 430 431 and master mix using an automated liquid handling robot (Freedom Evo-2 150; Tecan).

Each plate included 8 calibrator samples (1 ng/µl) to account for plate to plate variation and two non-template controls (NTC) consisting of nuclease free water. The calibrator sample was extracted from a large quantity of buffy coat prepared from blood supplied from a single domestic sheep (Cat# SHP-BUFCT-LIHP, Sera Laboratories International LTD, West Sussex, UK). We carried out a large number of extractions from this sample, applied the same quality control as above and then pooled the extracts and aliquoted them for subsequent use. A five-point standard curve, consisting of a four-fold serial dilution of the 439 calibrator sample (at concentrations: 20 ng/µl, 5ng/µl, 1.25 ng/µl, 0.3125 ng/µl and 0.078125 440 ng/µl) was included on the plate to provide a visual check that the samples amplified at the 441 correct cycle. Samples, calibrators, standard curve and NTCs were all run in triplicate. All qPCRs were performed using a Roche LC480 instrument using the following reaction 442 protocol; 10 min at 95 °C (enzyme activation), followed by 50 cycles of 15 s at 95 °C 443 (denaturation) and 30 s at 58 °C (primer annealing), then 30 s at 72 °C (signal acquisition). 444 Melting curve protocol was 1 min at 95 °C, followed by 30 s at 58 °C, then 0.11 °C/s 445 increase to 95 °C followed by 10 s at 40 °C (cool down). 446

447 Calculation of Relative Telomere Length

448 We used the LinRegPCR software package (version 2016.0; (60)) to correct our amplification curves for baseline fluorescence, and to calculate well-specific reaction 449 450 efficiencies and Cq values. A constant fluorescent threshold was set within the window of linearity for each amplicon group, calculated using the average Cq across all plates. The 451 threshold values used were 0.2 and 0.25, and the average PCR efficiency across all plates 452 was 1.876 and 1.881 for the B2M and telomere amplicon groups, respectively. Samples 453 454 were excluded from further analysis if the coefficient of variation (CV) across triplicate Cq values and triplicate PCR efficiency values for either amplicon was > 5% (n=1; note that for 455 > 95% of our samples the triplicate CV was < 2%), or if at least one of their triplicate 456 457 reactions had an efficiency that was 5% higher or lower than the mean efficiency across all 458 wells on that plate for the respective amplicon (n=2; > 95% of samples were within 2% of mean plate efficiencies; SI Appendix, Figure S4). 459

RLTL for each sample was calculated, following Pfaffl (61), using average reaction
 efficiencies for each plate and Cq for each sample determined by LinRegPCR as follows:

$$RTL = \frac{E_{TEL}(Cq_{TEL}[Calibrator] - Cq_{TEL}[Sample])}{E_{B2M}(Cq_{B2M}[Calibrator] - Cq_{B2M}[Sample])}$$

Where E_{TEL} and E_{B2M} are the mean reaction efficiencies for the respective amplicon group across all samples on a given plate; $Cq_{TEL}[Calibrator]$ and $Cq_{B2M}[Calibrator]$ are the average Cqs for the relevant amplicon across all calibrator samples on the plate; and $Cq_{TEL}[Sample]$ and $Cq_{B2M}[Sample]$ are the average of the triplicate Cqs for the sample for each amplicon.

467 Repeatability of telomere length

To assess the repeatability of our qPCR assay, the first qPCR plate (n=48 samples) was run 8 times over 4 consecutive days: four times over two days with samples in the same position 470 and then four times over two days with samples in an alternative row within the qPCR plate. 471 We calculated the overall repeatability of RTL as the proportion of variance explained by 472 sample identity over the total variance, in a linear mixed-effects model including only sample identity as a random intercept term (using restricted maximum likelihood estimation in 473 glmmTMB v.0.2.3 (62)). The overall repeatability of RTL was 0.866 (95% confidence 474 intervals 0.807-0.908). The repeatability of RTL measured in the same location across the 475 first four qPCR plates was 0.948 (95%CI 0.922-0.965), illustrating how repeatability can be 476 inflated if samples are consistently run in the same location. In a second model which 477 included sample identity, qPCR plate and qPCR row as random intercept terms, the 478 proportion of variance explained by qPCR plate and row was 0.005 and 0.063, respectively. 479 The repeatability of RTL in this model was 0.824 (95%CI 0.748-0.880), or 0.884 (95%CI 480 0.829–0.923) if the plate and row terms were excluded from the total variance (since they 481 represent measurement error). These repeatability estimates compare favourably with other 482 studies of telomere length measured by both qPCR (inter-assay repeatability: 0.85 (48); 0.82 483 484 (24)) and terminal restriction fragment (TRF repeatability: 0.86 (63)).

485 Data analysis

All analyses were conducted in the program R version 3.6.1 (64) using the package
 MCMCgImm v.2.29 (65) unless otherwise specified.

488 Relationship between telomere length, age and sex

489 Relative telomere length was approximately normally distributed in lambs, adults and overall (Figure S5). We ran a series of linear mixed-effects models to determine the function that 490 491 best described variation in telomere length with age. We included a two-level factor for age class (lamb: \sim 4 months of age; adult: \geq 1 year). To account for age-related variation within 492 493 the adult age class, we included age in years as a fixed covariate. We tested linear, 494 quadratic and cubic age terms, as well as threshold age functions with a range of 495 breakpoints (2-11 years). We also tested a four-level factor for age class (lamb: ~4 months of age; yearling: ~16 months; adult: 2-6 years; geriatric: >6 years. All models included 496 individual identity and sample year as random intercept terms to account for non-497 498 independence among observations. The qPCR plate and row for each sample were also included as crossed random intercept terms to account for variation associated with 499 500 measurement error. The models were run using maximum likelihood estimation in *glmmTMB* 501 v.0.2.3 (62) and AIC model selection was used to determine the best age function. We 502 selected the model with the fewest parameters within 2 ΔAIC of the model with the lowest AIC value. Once we determined the best fitting age function, we tested for a difference in 503 504 average telomere length between the sexes by including a two-level factor for sex in our

505 model. To test whether the ageing patterns differed between the sexes, we also tested for 506 interactions between sex and the selected age terms. The significance of additive and 507 interactive effects of sex was assessed using likelihood ratio tests (see SI Appendix). The repeatability of telomere length over the lifespan was estimated as the variance explained by 508 509 the random effect of individual over the total phenotypic variance (the sum of the random effects variance components plus the residual variance). There were 3641 observations of 510 telomere length from 1586 sheep available for this analysis. Of these individuals, 836 had 511 one RTL measurement available, 271 had two, 281 had three or four, and 198 had five our 512 513 more measurements.

514 Heritability of telomere length

A quantitative genetic animal model was used to estimate the additive genetic variance for 515 telomere length in Soay sheep. The model contained a two-level fixed factor for age class 516 (lamb: ~4 months; adult: \geq 1 year) and a linear covariate for age in years. We also included 517 sex as a two-level fixed factor to account for differences in average telomere length between 518 519 the sexes. The additive genetic effect was estimated using information on individual relatedness from the population pedigree. We included maternal identity in addition to the 520 additive genetic component to capture similarity among maternal siblings that is not 521 explained by the additive genetic effect (known as the maternal effect). We also included 522 523 individual identity as a random effect to capture consistent differences in measures from the 524 same individual that are not attributed to genetic effects, influenced for example by where 525 the individual lives or aspects of their early life environment (the permanent environment effect). Year of sample collection, qPCR plate and qPCR row were included as random 526 527 intercept terms. The heritability of telomere length was calculated as the variance explained by the additive genetic effect over the total phenotypic variance. This model was run for 528 1.1x10⁵ iterations, with 1x10⁴ burn-in and thinning interval of 50, resulting in 2000 stored 529 samples of the MCMC chain with minimal autocorrelation (<0.2). Parameter estimates are 530 presented as the posterior mode with 95% highest posterior density (HPD) intervals. 531 Parameter expanded priors were used for all variance components, and inverse-Wishart 532 533 priors for the residual variance. There were 3632 observations of telomere length from 1582 534 sheep available for this analysis (n=9 observations from 4 sheep were excluded because maternal identity was unknown). 535

We also ran a bivariate model to estimate the genetic correlation between telomere length in lambs (~4 months) and adults (≥ 1 year) which treated TL in lambs and adults as separate response variables. We included sex as a two-level fixed factor for both lambs and adults, with age in years included as a fixed covariate for adults only. qPCR plate and row were 540 included as random intercept terms across both models. Maternal identity was included as a 541 random effect in the lamb model and individual identity in the adult model to estimate 542 maternal and permanent environment effects, respectively. We estimated the unstructured variance-covariance matrix for the genetic, year and residual effects, which enabled us to 543 544 estimate the correlations across age classes at these different hierarchical levels. There is a possibility that the lamb residual variance could covary with the adult permanent 545 environment effect, but we did not attempt to model this covariance because we detected no 546 permanent environment effect in adults (see Results). The model was run 2.1x10⁵ iterations, 547 with 1x10⁴ burn-in and thinning interval of 200, resulting in 1000 stored samples of the 548 MCMC chain (autocorrelation <0.2). The genetic correlation was taken from the posterior 549 correlation of the stored samples with 95% HPD intervals. The bivariate model was run using 550 3632 measurements from 1582 individuals. Both lamb and adult telomere length 551 552 measurements were available for 424 individuals.

553 Associations between telomere length, August weight and overwinter survival

554 Phenotypic model

555 We used a multivariate mixed-modelling approach to examine the association between telomere length, weight and overwinter survival at different hierarchical levels. Relative 556 telomere length, body weight in kg (both measured August year t), and annual survival (to 1 557 May year t+1) were our response variables, modelled as Gaussian, Gaussian and threshold 558 distributions, respectively (corresponding to identity and probit link functions). Unstructured 559 variance-covariance matrices were estimated for some of the random effects, allowing us to 560 estimate the covariance among the three traits at different hierarchical levels. For each of 561 these random effects, we obtained a posterior distribution for the variance and covariance 562 563 between telomere length ("RTL"), weight ("Wt") and overwinter survival ("Surv"):

σ_{RTL}^2	$\sigma_{RTL,Wt}$	$\sigma_{RTL,Surv}$	
$\sigma_{RTL,Wt}$	σ_{Wt}^2	$\sigma_{Wt,Surv}$	
$\sigma_{RTL,Surv}$	$\sigma_{Wt,Surv}$	σ_{Surv}^2	

564 We estimated the covariance at the among-individual, among-year and residual levels. The among-individual level captures consistent differences between individuals (e.g. if individuals 565 566 that are consistently heavier tended to have longer lifespans). The covariance at the year level captures associations between average trait values among years (e.g. whether the 567 survival rate is higher in years when the average body weight is higher). The residual level 568 569 reflects covariance that is not captured by the among-individual and among-year levels (i.e. 570 if an individual has a relatively low body weight compared to its average has lower chances 571 of surviving).

The model was run for 5.3x10⁵ iterations, with 3x10⁴ burn-in and thinning interval of 500, 572 573 resulting in 1000 stored samples of the MCMC chain (autocorrelation <0.1). The residual 574 variance was fixed at 1 for survival (as is the standard recommendation for threshold models), and the latent variables were constrained to be between ±7 to avoid numerical 575 difficulties as the probabilities approached 0 and 1. In addition to the random effects of 576 identity and year, the telomere length model included random intercept terms for gPCR plate 577 and row, and the weight and survival models included random intercept terms for maternal 578 identity and birth year. We fitted age and sex in slightly different ways across the three 579 models, to reflect our understanding of differences in age-related variation in these three 580 581 traits. In the telomere model, we included age class (lamb vs adult) as a two-level fixed factor. No other fixed effects were included in the telomere model, since we were interested 582 in how telomere length covaried with the other traits across the lifespan. In the weight model, 583 we included age class (lamb vs adult) plus linear, quadratic and cubic age terms (capturing 584 variation in weight with age during adulthood) and their interactions with sex. In a previous 585 study, we detected a decline in weight across the year prior to death in this system using 586 587 univariate models of weight (35). We did not include a term of final year alive here, as this 588 effect is captured in the residual covariance between weight and survival in our multivariate 589 model. In the survival model, we included age class (lamb vs adult) plus sex and its 590 interaction with age class, as well as linear and quadratic age terms (capturing variation in 591 survival with age during adulthood). This model was run using 3569 measurements from 1574 individuals for which telomere length, August weight and overwinter survival 592 measurements were available. Telomere length and body weight were z-transformed prior to 593 inclusion in the model (mean=0, standard deviation=1). To check the consistency of our 594 results, we additionally ran separate models for lambs aged 4 months and adults aged ≥ 1 595 596 year (see Supplementary methods and results).

597 Genetic model

We extended our multivariate analysis to examine the association between telomere length, 598 August body weight and overwinter survival probability at the genetic level (the G matrix). As 599 600 with the univariate heritability analysis, we used information on individual relatedness from 601 the population pedigree to estimate the additive genetic variances and covariances. In addition to the additive genetic (co)variance, we estimated the covariance at the residual, 602 603 among-year and among-individual levels (the permanent environment covariance: the covariance among traits at the individual level that is not explained by genetic effects). The 604 model was run for 5.3×10^5 iterations, with 3×10^4 burn-in and thinning interval of 500, 605 resulting in 1000 stored samples of the MCMC chain (autocorrelation <0.1). Parameter 606 607 expanded priors were used for all variance components, and inverse-Wishart priors for the

residual variances. The residual variance was fixed at 1 for survival (as is the standard recommendation for threshold models), and the latent variables were constrained to be between ± 7 to avoid numerical difficulties as the probabilities approached 0 and 1. Other fixed and random effects were as stated above for the phenotypic models. We additionally ran separate animal models for lambs aged 4 months and adults aged \geq 1 year to check the consistency of our results (see Supplementary methods and results).

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773

774 Figure legends

775

Figure 1. Relative telomere length varied with age over the lifespan in Soay sheep (n=3641 776 777 observations of 1586 individuals). The points show raw data medians and standard errors for 778 each age, with females in black and males in grey (for clarity, n=3 observations of females 779 aged > 13 and n=6 observations of males aged > 8 are grouped with ages 13 and 8, respectively). The black lines show predictions from the best model (SI Appendix, Table S2), 780 781 grey shading represents 95% credible intervals around those predictions. The rug plot on the 782 inside of the x-axis shows the distribution of observations across the age range. Note that 783 the ageing pattern in the raw data and best-fitting age functions from mixed-effects models 784 are not expected to align perfectly when selective disappearance effects are present.

785

Figure 2. The proportion of variance in relative telomere length in Soay sheep explained by 786 different variance components. Estimates are based on the mode of the posterior distribution 787 from Bayesian quantitative genetic animal models. Estimates for age class 'all' came from a 788 789 univariate model incorporating all observations (n=3632 of 1582 individuals) that included sex, age class and age in years as fixed effects (SI Appendix, Table S3). The separate 790 estimates for adults (age \geq 1 year) and lambs (4 months old) came from a bivariate model 791 792 that accounted for the covariance between lamb and adult telomere length at the genetic, 793 year and residual levels (SI Appendix, Table S4).

794

795 Figure 3. The correlation between relative telomere length (RTL), August body weight and overwinter survival probability at different hierarchical levels in Soay sheep (n=3569 796 observations of 1574 individuals). Correlations were estimated as the mode of the posterior 797 798 distribution with 95% higher probability density intervals from multivariate Bayesian mixed-799 effects models. Panel A shows estimates for the among-individual (teal), among-year (black) 800 and residual correlations (grey) from a phenotypic model (SI Appendix, Table S5). Panel B shows estimates for the genetic (teal circles), permanent environment ("PermEnv", teal 801 triangles), among-year (black) and residual correlations (grey) from a quantitative genetic 802 803 animal model (SI Appendix, Table S6).



Age (years)



