High heritability of telomere length and low heritability of telomere shortening in wild birds

Christina Bauch1 | Jelle J. Boonekamp1 | Peter Korsten2 | Ellis Mulder1 | Simon Verhulst1

Abstract
Telomere length and telomere shortening predict survival in many organisms. This raises the question of the contribution of genetic and environmental effects to variation in these traits, which is still poorly known, particularly for telomere shortening. We used experimental (cross-fostering) and statistical (quantitative genetic “animal models”) means to disentangle and estimate genetic and environmental contributions to telomere length variation in pedigreed free-living jackdaws (Corvus monedula). Telomere length was measured twice in nestlings, at ages 4 (n = 715) and 29 days (n = 474), using telomere restriction fragment (TRF) analysis, adapted to exclude interstitial telomeric sequences. Telomere length shortened significantly over the nestling period (10.4 ± 0.3 bp day⁻¹) and was highly phenotypically (rP = 0.95 ± 0.01) and genetically (rG > 0.99 ± 0.01) correlated within individuals. Additive genetic effects explained a major part of telomere length variation among individuals, with its heritability estimated at h² = 0.74 on average. We note that TRF-based studies reported higher heritabilities than qPCR-based studies, and we discuss possible explanations. Parent–offspring regressions yielded similar heritability estimates for mothers and fathers when accounting for changes in paternal telomere length over life. Year effects explained a small but significant part of telomere length variation. Heritable variation for telomere shortening was low (h² = 0.09 ± 0.11). The difference in heritability between telomere length (high) and telomere shortening (low) agrees with evolutionary theory, in that telomere shortening has stronger fitness consequences in this population. Despite the high heritability of telomere length, its evolvability, which scales the additive genetic variance by mean telomere length, was on average 0.48%. Hence, evolutionary change of telomere length due to selection is likely to be slow.

KEYWORDS
early-life, inheritance, life-history, quantitative genetics, senescence, ageing
Individuals differ in lifespan and other fitness components, and identification of the molecular and physiological traits associated with fitness is a useful step in deciphering ecological and evolutionary dynamics of life histories. Telomere length and telomere shortening are such traits, because a shorter telomere length generally predicts a lower survival probability (meta-analyses in humans: Boonekamp et al., 2013; and wild vertebrates: Wilbourn et al., 2018) as does higher telomere shortening (in our study species: Salomons et al., 2009, Boonekamp et al., 2014; and other species: Barrett et al., 2013, Sheldon et al., 2021). Telomere length and telomere shortening are therefore increasingly being used as biomarkers of ageing and phenotypic quality across research areas such as epidemiology, ecology and evolutionary biology (Monaghan et al., 2018). However, to what extent both genetic and nongenetic factors contribute to among-individual variation in telomere length, and in particular to telomere shortening, remains poorly understood.

Telomeres consist of evolutionarily conserved, noncoding DNA sequence repeats (Blackburn, 1991) that together with the shelterin protein complex form the ends of chromosomes (de Lange, 2005) and contribute to genome stability (O’Sullivan & Karlseder, 2010). Telomeres are dynamic structures, in that their length shortens with age due to incomplete replication during cell division, which can be accelerated by DNA- and protein-damaging factors and attenuated or counteracted by maintenance processes (Chan & Blackburn, 2004). When telomeres reach a critically short length, the cell undergoes replicative senescence or apoptosis (Chan & Blackburn, 2004). In this way, telomere length and telomere shortening may play a causal role in senescence. Regardless of the question of causality, telomere length and telomere shortening are of interest as proxies for general health and senescence processes (Young, 2018).

To understand how telomere length and telomere shortening may play a role in shaping life histories from an evolutionary perspective, it is necessary to quantify their additive genetic variance on which natural selection can act. However, heritability estimates of telomere length differ markedly between studies, and this variation is not yet understood. To what extent variation in telomere length is due to inheritance or driven by the environment is therefore still under debate (Atema et al., 2015; Benetos et al., 2019; Broer et al., 2013; Dugdale & Richardson, 2018). Differences in the genetic background of populations, temporal–spatial variation of environmental influences, and methodological differences including limitations to separate genetic and environmental effects can all underlie variation in heritability estimates between studies (Becker et al., 2015; Dugdale & Richardson, 2018).

Telomere length at any given age is defined by the initial telomere length transferred through the parental gametes and the change in telomere length from the zygote stage onwards, which varies under the influence of both genes and the environment. For example, telomere shortening is generally accelerated with increased exposure to various (environmental) stressors; Angelier et al., 2018; Boonekamp et al., 2014; Kotsrach et al., 2007; McLennan et al., 2016; Seeker et al., 2021). Telomere shortening is typically faster early in life (Benetos et al., 2019; Fairlie et al., 2016; Salomons et al., 2009; Spurgin et al., 2018), potentially resulting from a greater susceptibility of telomeres to environmental and genetic effects in early life. Thus, exposure to environmental influences, age-dependent sensitivity to environmental effects and age-dependent gene expression all contribute to telomere length variability.

Most studies of telomere length heritability have measured telomere length at different ages in different individuals (reviewed in ref. Dugdale & Richardson, 2018), which affects the comparability of telomere length between individuals due to variation in exposure to environmental effects. In this way, sampling at variable ages will bias telomere length heritability estimates to an unknown extent, even when taking age variation into account in the estimation process. On the other hand, a shared environment among related individuals (e.g., due to parental care or location) can increase phenotypic similarity (e.g., in telomere length) and thereby inflate heritability estimates (Kruuk & Hadfield, 2007). Moreover, similarity in telomere length between parents and offspring can potentially arise in multiple ways, via additive genetic effects on gamete telomere length and telomere length regulation (Codd et al., 2013; Soerensen et al., 2012), epigenetic effects (e.g., mediated by paternal age; Bauch et al., 2019; Eisenberg, 2019; Noguera et al., 2018), and via early-life parental effects of genetic and nongenetic origin (Crisculo et al., 2017; Marasco et al., 2019). Quantifying telomere length heritability is therefore challenging.

Quantitative genetic “animal models” are a powerful statistical tool to disentangle genetic from environmental effects, which makes use of all relatedness information available in a pedigree (Wilson et al., 2010). As this is a “data-hungry” method, the number of studies using it to investigate causes of telomere length variation is currently limited, and results are highly variable (reviewed in refs. Dugdale & Richardson, 2018; Froy et al., 2021; Sparks et al., 2021). Moreover, all except one study on wild vertebrates (Vedder et al., 2021) measured telomeres using qPCR (quantitative polymerase chain reaction), which pools all telomeric sequences in the genome (located both terminally and interstitially; Nussey et al., 2014). It is not clear therefore how the currently available estimates relate to the heritability of the length of the terminal telomeres, which are subject to age- and stress-related shortening. Even less is known of genetic variation in telomere shortening, with one study on human twins reporting a heritability of 0.28 (95% confidence interval [CI] 0.16–0.44; Hjelmborg et al., 2015).

Heritability ($h^2$) is estimated as the ratio between the additive genetic variance ($V_A$) and the total phenotypic variance ($V_P$) $h^2 = V_A/V_P$ (Falconer & Mackay, 1996), and as such, low heritability can result from low additive genetic variance, but also from high total variance (e.g., because of large environmental effects). The evolutionary potential of a trait is dependent on its additive genetic variance, which scaled to the mean of the trait yields a metric known as the evolvability (Houle, 1992). Given the evidence for survival selection on both telomere length and telomere...
shortening (Barrett et al., 2013; Boonekamp et al., 2013, 2014; Sheldon et al., 2021; Wilbourn et al., 2018), their evolvability is of interest. However, we are aware of only one study estimating the evolvability of telomere length that was based on the absolute size of terminally located telomere length measured in base pairs. This study on captive field crickets found evolvability to be low despite high heritability (Boonekamp et al., 2021).

The aim of the present study was to quantify the additive genetic and environmental contributions to the variability of telomere length among and within individuals to estimate the heritabilities of telomere length and telomere shortening and the potential for telomere length to evolve over time in a population of free-living wild jackdaws (Corvus monedula). To this end, we measured telomere length in erythrocytes at the age of 4 days and for the majority of individuals again at 29 days (contingent primarily on their survival), using the golden-standard technique telomere restriction fragment (TRF) analysis adapted to measure terminally located telomeric sequences only (Nussey et al., 2014; Salomons et al., 2009). To disentangle genetic and environmental causes of variation in telomere length and shortening we (i) applied experimental cross-fostering and (ii) made use of the available multigenerational pedigree information by running uni- and bivariate “animal model” analyses (Wilson et al., 2010). Using these methods, we estimated the heritabilities of telomere length and telomere shortening, and—to test for potential age-dependent genetic effects—the genetic correlation of telomere lengths measured at different ages. We additionally performed parent–offspring regressions to assess the relative maternal and paternal contributions to the variation in offspring telomere length, while also taking into account effects of paternal age at conception (Bauch et al., 2019), and compared early-life telomere shortening between parents and offspring. Lastly, we estimated the additive genetic variance of telomere length in relation to absolute trait size (i.e., evolvability) to assess the potential for telomere length to evolve in our population. As telomere length has been shown to differ between the sexes in several species (Barrett & Richardson, 2011; Bauch et al., 2020), we also tested for sex differences in telomere length and telomere shortening in this study.

2 | MATERIAL AND METHODS

2.1 | Study population

We studied a jackdaw (Corvus monedula) population breeding in nestbox colonies located south of Groningen, The Netherlands (53°14′N, 6°64′E), which has been under investigation since 1996. Birds in our study population are ringed with numbered metal rings and colour rings shortly before fledging or as immigrants at first breeding in the study’s nestboxes. Breeders are highly site faithful and all breeding birds are identified by their unique colour ring combinations via telescope, photo or video camera. Jackdaws breed monogamously with low divorce rates and very rare extra-pair paternity (Henderson et al., 2000; Liebers & Peter, 1998). However, due to partner death, about 50% of the adults in our data set had two or more partners and the population comprises full as well as half-siblings produced over multiple years. From 1 day before the expected hatching date the nestboxes were checked daily for hatchlings. Freshly hatched chicks were marked by specific combinations of clippings of the tips of the toenails for identification until ringing. Exact ages were known for jackdaws native to the study’s nestboxes. Immigrants were assigned an age of 2 years when breeding for the first time, which is the modal age at recruitment in our population. The sex of all jackdaw parents and the majority of chicks has been identified either molecularly (Griffiths et al., 1998) or by behavioural observations and cross-reference with breeding partners.

2.2 | Pedigree

The pruned pedigree consists of 1007 relatedness-informative individuals, whereof 715 individuals held data on telomere length at the age of 4 days of which 474 individuals additionally held telomere length data when 29 days old. The pedigree spans six generations for known telomere length of either age. For details on relationships in the pedigrees see Table S1 and Figure S1. Summary statistics and pedigree images were generated using the R-package PEDANTS (Morrissey & Wilson, 2010).

2.3 | Cross-fostering

To experimentally disentangle additive genetic from early-life parental effects we performed two types of cross-fostering manipulations in our study population. In 2015 and 2016 in a sub-set of the nestboxes (n = 58 of 126 nests) complete clutches were exchanged between nests during mid-incubation. Nests were matched for clutch size and laying date (± 1 day), but in all other respects cross-fosters were performed randomly. In this way, foster parents already incubate and take care of the nestlings as soon as they hatch. Additionally, in all years, brood sizes were manipulated in most nests, with nestlings transferred between nests when the oldest nestling was 4 days old. Age-matched broods were reduced by removing three nestlings and adding one nestling and enlarged by adding three nestlings and removing one (for details, see Boonekamp et al., 2020). Consequently, manipulated broods were reduced by removing three nestlings and adding one nestling and enlarged by adding three nestlings and removing one (for details, see Boonekamp et al., 2020). Consequently, manipulated broods (i.e., between first and second telomere length measurement) comprised genetic and nongenetic offspring of parents and, thus, both genetic and nongenetic siblings. The multigenerational pedigree included individuals with various degrees of relatedness, which experienced the same or different rearing environments. The cross-fostering experiments add additional information to the data set by adding full siblings reared in different broods. For an overview of the study design and sample distribution over the different treatments see Figure 1.
2.4 | Blood sampling and measuring of telomere length

Telomere length was measured in blood samples taken between 2005 and 2016 from 715 individuals at the maximum age of 4 days (mean ± SD = 3.7 ± 0.6 days) and again 25 days later in a subset of 474 individuals (age 29 days; Figure 1). This interval between repeated telomere measurements of nestlings covers ~70% of their time in the nest (Röell, 1978). Fathers were caught and blood sampled when the brood was at least 14 days old, contributing telomere data of 82 fathers at conception across multiple years to the data set. Samples were stored in 2% EDTA buffer at 4–7°C and within 3 weeks snap frozen in a 40% glycerol buffer for permanent storage at −80°C. Telomere length of nucleated erythrocytes was measured performing TRF analysis under nondenaturing conditions using pulsed-field gel electrophoresis as in Salomons et al. (2009). See Text S1 for details, including a gel image (Figure S2). A sample of an individual consists of a characteristic telomere length distribution from erythrocytes of different ages and different chromosomes within cells. The individual average of this telomere length distribution was used in the present analyses (Salomons et al., 2009). Samples were distributed over 57 gels. Samples from nestlings of the same brood were partly spread over different gels. Repeated samples (two ages) from individuals were run on the same gel. As a quality index for intergel comparability, we calculated the coefficient of variation of one control sample from a 29-day-old jackdaw run on 26 gels, which was 6%, and of one control sample of a goose, with a similar telomere length distribution in a similar range, run on 31 other gels, which was 7%. Thus, intergel measurement accuracy remained similar over time. The high phenotypic correlation of telomere length within individuals measured at the two ages that we find in our data set (\( r_p = 0.95 \pm 0.01, n = 474, \text{Table 1} \)) reflects a high intragel repeatability for telomere length, since repeated samples were always on the same gel. This estimate has the advantage of covering the complete method procedure between blood sampling and final calculation of telomere length for a large data set.

2.5 | Statistical analyses

First, we investigated effects of age and sex on telomere length and telomere shortening. We ran linear mixed-effects models with telomere length as the dependent variable, age as a factor to investigate telomere shortening over the interval of 25 days and bird ID as a random effect. Gel ID was included as a random effect to control for methodologically induced variance. With a log-likelihood ratio test, comparing models with and without bird ID, we tested for significant between-individual variation of telomere length. Further, by including the factor sex and its interaction with age in the model, we tested for potential sex differences in telomere length and telomere shortening.

Second, we ran a series of univariate quantitative genetic “animal models” (Wilson et al., 2010) using early-life telomere length data and pedigree information holding various degrees of kinship to partition the total phenotypic variance in early-life telomere length (\( V_{p} \)) measured at the ages of 4 or 29 days into its additive genetic (\( V_{a} \)), residual (\( V_{r} \)) and varying other components of potential relevance: biological mother ID (\( V_{m} \)), estimates the variance in telomere length among nestlings from different mothers over and above the variance that is attributable to additive genetic effects,
or in other words it accounts for early-life maternal effects before eventual cross-fostering. The telomere length in offspring from the same biological mother may be more similar as compared to offspring from other mothers due to mother-specific effects, if, for example, egg content affects offspring telomere length (Haussmann et al., 2012). Similarly, biological father ID (V_{bf}), and mother or father ID after experimental cross-fostering could affect telomere length of nestlings by parental care effects (e.g., Vedder et al., 2018). Year (V_{Y}) was added to account for potential year effects on offspring telomere length. Brood ID (V_{B}) estimates the variance that can be attributed to the shared environment of brood mates. In the model with telomere length at age 4 days we included brood ID until the age of 4 days (only genetic siblings and natural brood size, B1; Figure 1). In the model with telomere length at age 29 days we additionally included brood ID of the following 25 days (genetic and nongenetic siblings, manipulated brood size, manipulated brood size,

Table 1 Results from a bivariate animal model analysis on telomere length (TL) measured at 4 days old in 715 individuals and again at 29 days old in a subset of 474 individuals. (a) Fixed effects, variance components and covariances with regard to telomere length. Brood ID changed due to experimental manipulation at age 4 days and accordingly the respective brood ID (B1, B2) was fitted for the specific telomere length measure (TL4, TL29). The variances for random effects were estimated for telomere lengths at each age and for the covariance between the two telomere lengths, except for biol. mother ID and gel ID, which each have a common variance for telomere length of both ages (see also Table S8). (b) Heritability estimates for telomere length and telomere shortening (ΔTL) derived from the bivariate model. (c) Correlations between TL at ages 4 and 29 days. For study design see Figure 1.

(a)

| Fixed effects                  | TL 4 days Estimate (SE) | TL 29 days Estimate (SE) | F  | df    | p      |
|--------------------------------|-------------------------|--------------------------|----|-------|--------|
| Intercept                      | 7,118.0 (75.0)          | 6,884.0 (79.9)           |    |       |        |
| Paternal age                   | −25.1 (13.4)            | −30.2 (14.2)             | 3.5| 1,269.7| .062   |
| Year                           |                         | 4.5                      | 1,286.8 | .034 |

(b)

| Heritability h^2 (SE)          | TL 4 days               | TL 29 days               | ΔTL |
|--------------------------------|-------------------------|--------------------------|-----|
| variance due to gel effect included | 0.603 (0.107)           | 0.660 (0.108)            | see Table S8 |
| biological variance only (gel effect excluded) | 0.709 (0.121)           | 0.765 (0.121)            | 0.088 (0.114) |

(c)

| Correlations between TL at ages 4 and 29 days |
|-----------------------------------------------|
| Phenotypic r (SE)                             | Additive genetic r (SE) |
| 0.954 (0.006)                                 | 0.999 (0.006) |
B2). To account for measurement differences between gels, we always included gel ID as a random effect \( (V_{\text{gel}}) \). As a fixed effect we included biological father age (Bauch et al., 2019). The fixed effect of sex was not significant and therefore excluded from the models. As we measured telomere length for the entire brood when the oldest nestling was 4 days old, while there is age variation within broods due asynchronous hatching, we tested whether the exact age of the nestling had an effect on telomere length. However, this was not the case and therefore this predictor was also excluded from the models. Statistical significance of fixed effects was determined using Wald tests and of random effects using log-likelihood ratio tests, comparing models with and without the specific random effect.

Third, we ran bivariate models with telomere length of both nestling ages. We included the same random effects as in the univariate model analysis. The additive genetic and year random effects were fitted with an unstructured variance–covariance matrix (i.e., for each random effect the variance was estimated for telomere length at each age as well as the covariance between the two telomere lengths). We initially included the random effect gel ID in the same manner, but this model structure prevented us from reliably estimating the variance for brood ID B1 (until age 4 days), possibly due to over-parameterization issues (Table S8). We therefore included gel ID as a common random effect estimating the common variance for the two ages in the final model. A comparison of the results from the two models (Table 1; Table S8) showed that the way the variance was fitted for gel ID had virtually no effect on the estimated genetic and phenotypic correlations, or the estimated heritability for telomere shortening. We also included biological mother ID estimating the common variance, because it did not significantly affect the variation in telomere length at any of the two ages. The statistical significance of fixed effects (paternal age) was determined using Wald tests.

Fourth, we ran a univariate model with telomere shortening as the dependent variable, including again the same random effects. With log-likelihood ratio tests, we determined the statistical significance of random effects on telomere shortening between the ages of 4 and 29 days.

We calculated narrow sense heritability \( h^2 \) estimates for telomere length as:

\[
h^2 = \frac{V_A}{(V_P - V_{\text{gel}})}
\]

excluding variation due to gel effects \( (V_{\text{gel}}) \) unless stated otherwise. For a more comprehensive understanding of the \( h^2 \)-estimate, which can be affected by fixed effects (Wilson, 2008), we reran the main univariate model with random effects only.

From the bivariate model results (Table 1) we additionally calculated the heritability estimate of telomere shortening, the phenotypic and additive genetic correlation between telomere lengths at the two ages, the coefficients of additive genetic variance of telomere length as well as the evolvabilities as follows.

To estimate the heritability \( h^2 \), see formula above) for telomere shortening \( (\Delta TL_{29} - \Delta TL_4) \), the additive genetic variance \( (V_A) \) for telomere shortening was derived from the variance–covariance matrix:

\[
V_{A29-4} = V_{A29} + V_{A4} - 2 \times \text{cov}_A
\]

where \( V_A \) is the additive genetic variance at age 4 or 29 days and \( \text{cov}_A \) is the additive genetic covariance between telomere length measured at age 4 and 29 days.

The genetic correlation \( r_G \) between telomere length at ages 4 and 29 days was calculated as:

\[
r_G = \frac{\text{cov}_A}{\sqrt{V_{A4} \times V_{A29}}}
\]

The phenotypic correlation \( r_P \) was calculated as:

\[
r_P = \frac{(\text{cov}_A + \text{cov}_R)}{\sqrt{(V_{A4} + V_{R4})(V_{A29} + V_{R29})}}
\]

with \( \text{cov}_R \) as the residual covariance between telomere length measured at age 4 and 29 days and \( V_R \) as the residual variance at age 4 or 29 days.

The coefficient of additive genetic variance of telomere length at both time points was calculated as:

\[
CV_A(x) = 100 \times \frac{\sqrt{V_A}}{x}
\]

where \( x \) is telomere length and \( x \) its mean (Houle, 1992). Whether additive genetic variances differ at the two ages was tested with a log-likelihood ratio test comparing bivariate models with constrained common variance and unconstrained variances.

The relative evolvability \( I_A \) of telomere length at both time points was calculated as:

\[
I_A = \frac{V_A}{x} = \left( \frac{CV_A}{100} \right)^2
\]

(Houle, 1992), multiplied by 100 to express it as a percentage.

Lastly, for comparison with the animal model results, we applied parent–offspring regressions using linear mixed-effects models of early-life telomere length (age 4 days) and telomere shortening during the nesting period (between age 4 and 29 days, \( \Delta TL \)) between parents and their offspring. We ran these analyses for mothers and fathers separately as data on both parents simultaneously were available in too few cases. When regressing offspring telomere length on father telomere length we ran an additional model that included father age as a fixed effect, as telomere length of consecutive offspring declines as fathers age (Bauch et al., 2019). Additionally, we calculated a father–offspring regression with father telomere length measured in the year of offspring conception. As the data set contains measures of telomere length or telomere shortening of
offspring of the same parents in multiple years and telomere length was analysed on different gels, we added parent ID, brood ID, year and gel ID as random effects.

All statistics were run in r (R Development Core Team, 2017) using ASReml-R software (versions 3 and 4; VSN International) and R-packages, lme4 (Bates et al., 2015) and lmerTest (Kuznetsova et al., 2017).

3 | RESULTS

3.1 | Descriptive statistics

Telomere length at the age of 4 days was on average 7,039 bp (SD = 591 bp, n = 715) and at the age of 29 days on average 6,729 bp (SD = 588 bp, n = 474; Figure 2). Telomeres shortened significantly within individuals over the measurement interval of 25 days by an average 260 bp (Table S2A). Telomere length differed significantly between individuals (χ² = 986.17, p < .001, Table S2A), and was highly phenotypically correlated within individuals between ages 4 and 29 days (r = .954 ± .006, n = 474, p < .001, Table 1, Figure 2). The sexes differed neither in telomere length (Table S2B) nor in telomere shortening (Table S2C).

3.2 | Animal model analyses

The uni- and bivariate animal model analyses revealed that the major part of the variation in telomere length at the age of 4 and 29 days was explained by additive genetic effects (Table 1; Tables S3-S8; Figure 3). The narrow sense heritability estimates for telomere length measured at the ages of 4 and 29 days, derived from the bivariate model, were h² ± SE = 0.71 ± 0.12 and 0.77 ± 0.12, respectively (Table 1). Heritability estimates based on respective univariate models, containing the same sample sizes, but not taking the relationship of repeatedly measured telomere length within individuals into account, were h² ± SE = 0.63 ± 0.13 (lower) for telomere length measured at the age of 4 days (Table S3) and h² ± SE = 0.88 ± 0.13 (higher) for telomere length measured at the age of 29 days (Table S6). Including paternal age as a fixed effect in the model did not significantly affect the heritability estimate for telomere length at age 4 days (h² ± SE = 0.63 ± 0.13, with paternal age, Table S3, vs. h² ± SE = 0.67 ± 0.13, without paternal age, Table S4).

In contrast to the high heritability of telomere length, the heritable component of the variation in telomere shortening was small with h² ± SE = 0.09 ± 0.11 (derived from the bivariate model, Table 1) and 0.08 ± 0.10 (derived from a univariate model with telomere shortening as dependent variable, Table S7).

Year was the only environmental factor that explained a significant part of the variance among individuals in early-life telomere length (age 4 days: 6%, Table 1, or 7%, χ² = 5.62, p = .018, Table S3; age 29 days: 7%, Table 1, or 6%, χ² = 2.29, p = .13, Table S6), indicating resemblance of telomere length between offspring within cohorts. Additional to the year effects present at the age of 4 days, there was a tendency for year effects leading to a significant variation in telomere shortening between the ages 4 and 29 days (ΔTL: 4%, χ² = 3.81, p = .05, Table S7).

Early-life maternal effects, represented by biological mother ID, explained small nonsignificant fractions of the variance in telomere length at both ages (age 4 days: 4%, Table 1, or 6%, χ² = 0.53, p = .47, Table S7).

**FIGURE 2** Repeated telomere length data within individuals (n = 474). (a) Telomere length at 29 days of age plotted against telomere length at 4 days of age in jackdaw nestlings. The dashed line represents x = y and hence the perpendicular distance below this line reflects the telomere shortening. (b) Telomere length at the ages 4 and 29 days, where lines connect repeated data of the same individual.
The mother that cared for the offspring from the age of 4 days onwards did not explain a noticeable amount of telomere length variation, and was bound to zero (Tables S6 and S7). The ID of the father that cared for the offspring was bound to zero in the model (Table S6) or explained a small, nonsignificant amount of variation in telomere shortening ($\Delta TL$: 2%, $\chi^2 = 0.15$, $p = .70$, Table S7). Thus, telomere length of offspring cared for by the same parent, whether biological or nonbiological, did not show increased similarity due to parental care, neither when measured at the age of 4 days nor at the age of 29 days, or in their telomere shortening over this period.

Brood ID (pre- and post age 4 days, B1 and B2, respectively) explained little of the variation in telomere length at both sampling ages (age 4 days: B1: 0.3%, Table 1, or 5%, $\chi^2 = 1.39$, $p = .24$, Table S3; age 29 days: B1: bound to 0, Table S6, and B2: 0.6%, Table 1, or 2%, $\chi^2 = 0.45$, $p = .50$, Table S6; $\Delta TL$: 2%, $\chi^2 = 0.09$, $p = .77$, Table S7). Thus, there was no significantly increased telomere length similarity of brood mates due to their shared brood environment, including brood-specific parental care effects.

### Table 2: Parent–offspring regressions. Comparison of telomere length (TL) measured at the age of 4 days (in bp) of (a) mothers and their offspring ($n = 113$ individual offspring of 31 mothers) and (b) fathers and their offspring ($n = 111$ individual offspring of 28 fathers)

| (a) Fixed effects | Estimate (SE) | $F$ | $df$ | $p$ |
|-------------------|---------------|-----|------|-----|
| Intercept         | 4,283.6 (781.6) |     |      |     |
| Mother’s TL (age 4 days) | 0.418 (0.106) | 15.44 | 1, 16.4 | .001 |

| (a) Random effects | Variance (SE) | Proportion of variance (SE) | $\chi^2$ | $df$ | $p$ |
|-------------------|---------------|----------------------------|---------|------|-----|
| Mother ID         | 71,974.6 (41,762.7) | 0.205 (0.110) | 3.940 | 1 | .047 |
| Brood ID (B1)     | –             | – | – | – | – |
| Year              | 92,126.7 (66,701.9) | 0.263 (0.148) | 4.763 | 1 | .029 |
| Gel ID            | 14,888.4 (23,115.2) | 0.042 (0.067) | 0.521 | 1 | .471 |
| Residual          | 171,639.6 (31,312.4) | 0.490 (0.125) |         |     |     |

| (b) Fixed effects | Estimate (SE) | $F$ | $df$ | $p$ |
|-------------------|---------------|-----|------|-----|
| Intercept         | 6,865.4 (830.0) |     |      |     |
| Father’s TL (age 4 days) | 0.102 (0.110) | 0.861 | 1, 25.6 | .362 |
| Father’s age at conception | −88.6 (38.2) | 5.386 | 1, 20.1 | .031 |

| (b) Random effects | Variance (SE) | Proportion of variance (SE) | $\chi^2$ | $df$ | $p$ |
|-------------------|---------------|----------------------------|---------|------|-----|
| Father ID         | 62,097.5 (46,496.8) | 0.200 (0.134) | 1.948 | 1 | .163 |
| Brood ID (B1)     | 27,415.8 (45,317.5) | 0.088 (0.147) | 0.297 | 1 | .586 |
| Year              | 11,885.5 (22,898.4) | 0.038 (0.072) | 0.446 | 1 | .504 |
| Gel ID            | 7,893.1 (23,205.1) | 0.025 (0.075) | 0.134 | 1 | .715 |
| Residual          | 200,755.8 (40,829.6) | 0.647 (0.137) |         |     |     |

*Brood ID was not included in the final model as it was bound to 0.*
Telomere lengths at the ages of 4 and 29 days were strongly genetically correlated ($r_g \pm SE = 0.999 \pm 0.006, p < .001$) and the additive genetic variances at the two ages (Table 1) did not differ ($p = .61$). Thus, there was no statistical support for a genotype by age interaction. The coefficients of additive genetic variance were 6.41% and 7.34% at ages of 4 and 29 days, respectively. Evolvability was 0.41% and 0.54% for telomere length at ages 4 and 29 days.

### 3.3 | Parent–offspring regressions

Telomere lengths of mothers and their offspring, all measured at the age of 4 days, were significantly positively related ($\beta = 0.42 \pm 0.11$), amounting to a telomere length heritability estimate of $h^2 = 0.84$ (Table 2a, Figure 4a). The same relationship for telomere lengths of fathers and their offspring was nonsignificant, whether paternal age was included in the model ($\beta = 0.10 \pm 0.11$, Table 2b) or not ($\beta = 0.18 \pm 0.10$, Table S9; Figure 4b). Older fathers produced nestlings with shorter telomere length, as reflected in a significant negative paternal age effect. The slope of the mother–offspring regression was more than twice as steep as the slope derived from the father–offspring regression (Table 2). In contrast, paternal telomere length measured at the age of offspring conception was significantly positively related to offspring early-life telomere length ($\beta = 0.31 \pm 0.07$; Table S10), amounting to a telomere length heritability estimate of $h^2 = 0.61$.

Telomere shortening from age 4 to 29 days was neither significantly correlated between mother and offspring ($\beta = 0.18 \pm 0.24$), nor between father and offspring ($\beta = 0.04 \pm 0.17$; Table S11, Figure 4c,d). Thus, while being far from statistically significant, parent–offspring regressions yielded low estimates of the heritability of telomere shortening as the animal models (Table 1; Table S7).
| Species                      | Age TL measured | Cross-foster | Controlled environment | Father age effect | Statistical analysis                                      | $h^2$ (95% CI)            | n        | Cell type | Laboratory method | References          |
|------------------------------|-----------------|--------------|------------------------|-------------------|----------------------------------------------------------|----------------------------|---------|------------|-------------------|----------------------|
| **Birds**                    |                 |              |                        |                   |                                                          |                            |         |            |                   |                     |
| King penguin                 | Offspring 10 days, parents at conception | No |                         |                   | Mid-parent-offspring mother–offspring                      | 0.2 (−0.02 to 0.42)        | 53      | Erythrocytes | qPCR              | Reichert et al. (2015) |
| White-throated dipper        | 7-17 days       | No           | Brood ID               | Cohort            | Mother–mid-offspring Animal model                         | 0.44 (0.048–0.83)          | 59      | Erythrocytes | qPCR              | Becker et al. (2015)  |
| Great reed warbler           | 8-10 days       | No           | Brood ID               | Mother ID         | No, but mother age effect Animal model                    | 1.08 (0.48–0.72)           | 17      | Erythrocytes | qPCR              | Asghar et al. (2015)  |
| Collared flycatcher          | 12 days         | Yes          | Brood triplets         |                   | Animal model (full sibling)                              | 0.09 (−0.04 to 0.15)       | 359     | Erythrocytes | qPCR              | Voillemot et al. (2012) |
| Seychelles warbler           | All ages        | No           | Mother ID              | Father ID         | Yes Mother–offspring Father–offspring Animal model        | 0.048 (<0.001–0.087)       | 284     | Erythrocytes | qPCR              | Sparks et al. (2021)  |
| Kakapo                      | 1–35 years      | No           |                         |                   | Mother–offspring Father–offspring                         | 0.84 (n.s.)                | 29      | Erythrocytes | TRF               | Horn et al. (2011)   |
| Tree swallow                 | Offspr. 12 days, parents at conception | Yes         | Brood size             |                   | Mid-parent–offspring                                     | 0.81 (n.s.)                | 122     | Erythrocytes | TRF               | Beelman et al. (2019) |
| Zebra finch                  | 9–636 days      | Yes          | Birth nest             | Rear nest         | Animal model                                             | 0.999 (0.87–1)             | 125     | Erythrocytes | TRF               | Atema et al. (2015)  |
| Common tern                  | 2–24 years      | No           | Permanent environment  |                   | Animal model                                             | 0.63 (0.50–0.73)           | 387     | Erythrocytes | TRF               | Vedder et al. (2021)  |
| Jackdaw                      | 4 days * 29 days * Fathers at conception (ac) | Yes         | Brood IDs              | Mother IDs        | Yes Mother–offspring Father–offspring Animal model        | 0.84 (0.42–1)              | 113     | Erythrocytes | TRF               | This study           |
| Reptiles                     |                  |              |                        |                   |                                                          |                            |         |            |                   |                     |
| Sand lizard                  | Hatchlings, known-age adults | No |                         |                   |                                                          |                            |         | Erythrocytes | TRF               | Olsson et al. (2011) |
| Fish                         | 122 days        | Yes          | Temperature Family ID  |                   | Animal models (full siblings of 9 families)               | 0.31 (0.05–0.97) to 0.47 (0.17–0.91) | 83, 213 | Brain tissue | qPCR              | Noreikiene et al. (2017) |
(Continues)
| Species                  | Age TL measured | Cross-foster | Controlled environment b | Father age effect | Statistical analysis | $h^2$ (95% CI)                  | n                  | Cell type | Laboratory method | References            |
|-------------------------|-----------------|--------------|--------------------------|------------------|---------------------|------------------------|---------------------|-----------|-------------------|----------------------|
| Mammals Greater mouse-eared bat Myotis myotis | 0–7+ years | No | Birth year Sample year Roost Temperature Rainfall | Animal models | 0.01 (0.00–0.04) to 0.06 (0.02–0.11) | 174 (504 samples) | Wing biopsy qPCR | Foley et al. (2020) |
| Holstein Friesian dairy cattle | All at birth +later in life | No | Birth year Genetic group Permanent env. | Random regression model incl. pedigree information | 0.36 to 0.47 (SE 0.05–0.08) | 308 females (1328 samples) | Leucocytes qPCR | Seeker et al. (2018) |
| Badger Meles meles ≤29 months * | All ages * | No | Mother ID Father ID Birth year Sample year Social group Permanent env. | Animal model | <0.001 (<0.001–0.043) * | 556 juveniles (837 samples) | Leucocytes qPCR | van Lieshout et al. (2020) |
| Soay sheep Ovis aries 4 months * > 1 year * | No | Mother ID Sample year Permanent env. | Animal model | 0.29 (0.21–0.37) * 0.21 (0.16–0.26) * | 1,582 (3,632 samples) | Leucocytes qPCR | Froy et al. (2021) |

*a Reared individually after hatching.
*b Refers to animal models in studies with multiple statistical approaches.
*c TRF including interstitial telomeric sequences (Southern blot).
*d No interstitial telomeric sequences in the genome.
*Heritability estimated for different age classes.
4 DISCUSSION

The main findings of our study were that telomere length was highly heritable in nestlings of free-living jackdaws, while heritable variation in telomere shortening was low. Telomeres shortened significantly and telomere lengths were phenotypically and genetically correlated within individuals. Nonetheless, as evidenced by the year effect, environmental influences can lead to significant variation in telomere length among individuals. Our finding that variation in telomere length and telomere shortening are primarily driven by additive genetic vs. environmental effects, respectively, suggests these traits differ in their potential as a fitness biomarker, and also in the mechanism through which this potential arises. Telomere length evolvability was low despite its high heritability.

Our estimate of telomere length heritability in jackdaws is similar to estimates reported by some other studies on free-living bird species such as terns and swallows (range $h^2 = 0.63–0.81$; Table 3), but substantially higher than estimates for penguins, dippers, warblers and flycatchers (range $h^2 = 0.0–0.48$; Table 3). One striking difference between those studies within the same taxonomic group (Aves) is that they used different telomere measurement techniques, TRF and qPCR, with higher and lower $h^2$ estimates respectively. Interestingly, TRF- and qPCR-based studies also report differences in telomere length repeatability, also being high and low, respectively (meta-analysis: Kärkkäinen et al., 2021). This is of relevance as phenotypic trait repeatability usually sets the upper limit to heritability (Dohm, 2002). The difference in heritability and repeatability estimates between TRF- and qPCR-based studies may be due to chance, given that the number of studies is still small, but may also be caused by real differences between the methods. With respect to the latter possibility, two main differences between TRF and qPCR measurements that may contribute to variation in repeatability and heritability estimates are that: (i) qPCR-based measurements include all telomeric sequences within the genome, and thus terminally and interstitially located telomorphic sequences, which are excluded when applying TRF to nondenatured DNA, whereas TRF-based measurements include a potentially variable subtelomeric region (Baird, 2005; Nussey et al., 2014); and (ii) measurement reliability tends to be lower when using qPCR (Horn et al., 2010; Morinha et al., 2020; Nussey et al., 2014). Regarding the potential influence of interstitial telomeric sequences on telomere length, their abundance in length and number varies between and within species (Delany et al., 2000; Foote et al., 2013; Meyne et al., 1990; Olsson et al., 2011). However, for the inclusion of interstitial telomeric sequences to explain the lower repeatability and heritability estimates when using qPCR, interstitial telomeric sequences would have to be more susceptible to environmental effects than terminal telomeres, resulting in a higher absolute change in the total number of telomere sequence repeats in the genome. There is at present little indication that this is a likely scenario. With respect to the subtelomeric region included in the TRF measurements, multiple tests on the extent of subtelomeric region measured, including a comparison between results from the two measurement techniques by Atema et al. (2019), suggest a minor influence on absolute telomere length, depending on the set of restriction enzymes used. We therefore consider, and also given the findings of Kärkkäinen et al. (2021), that measurement error, introduced at any point between sample collection and final estimation, is the most parsimonious explanation for the difference in repeatability and heritability estimates between qPCR- and TRF-based studies.

Heritability estimates for telomere length were high at both ages, and environmental effects had a relatively small influence on telomere length (Table 1; Tables S3–S8). Consequently, heritability estimates were similar whether derived from “animal model” analyses or parent–offspring regressions or whether shared environmental effects were controlled for in the models. The high phenotypic correlation of telomere length early and late in the nestling period, when telomere shortening is higher than later in life (Boonekamp et al., 2014; Salomons et al., 2009), and the consequently similarly high heritability estimates for both ages suggest that heritability of telomere length is high independent of age in our population, at least within the narrow age range over which we can make this comparison.

Among the tested environmental components, year effects explained a significant but small part of the variation in telomere length in our study population (Table 1; Tables S3 and S7). Birth cohort effects on telomere length have also been identified in other free-living vertebrates, including white-throated dippers (Cinclus cinclus) (Becker et al., 2015) and Soay sheep (Ovis aries) (Fairlie et al., 2016). However, the proportion of telomere length variance explained by year differed strongly between studies (6% in our study vs. 46% in dippers or nonsignificant in Seychelles warblers, Sparks et al., 2021). The cause of variation in telomere length between cohorts in our study population, already present at the age of 4 days, remains to be identified. Experimental studies found temperature conditions during incubation (Vedder et al., 2018) and glucocorticoid concentrations in the eggs (Haussmann et al., 2012) to affect offspring telomere length at this early stage in life, but whether these factors also modulate telomere length generally or depending on year remains to be established. The variation in telomere shortening between years that we found from the age of 4 days onwards can arise in combination with brood size, as shown previously in this population (Boonekamp et al., 2014).

To disentangle potential early-life parental effects on nestling telomere length, via egg content or parental care, we cross-fostered clutches and part of the offspring to nongenetic parents. Our results show that early-life parental effects did not increase telomere length similarity between nestlings (genetic and nongenetic siblings), as also reported for dippers (Becker et al., 2015) and Seychelles warblers (Sparks et al., 2021), but unlike reed warblers (Ashgar et al., 2015). Our results also do not support that a shared brood environment increased telomere length similarity of nest mates at any age, while this has been found, for example, in dippers (Becker et al., 2015).

Parent–offspring correlations of early-life telomere length were stronger for mothers than for fathers. This could result either from a stronger maternal inheritance of telomere length or from early-life effects affecting telomere length similarly in mother and offspring.
Alternatively, it may reflect that gamete telomere length changes more over life in fathers than in mothers, given that all oocytes are already present at birth while sperm is formed throughout life (Bauch et al., 2019). Early-life maternal effects were negligible according to our animal model results and are therefore unlikely to explain the higher similarity between mother and offspring early-life telomere length. However, we found a significantly positive correlation between father telomere length at the age of offspring conception and offspring telomere length at the age of 4 days. This suggests an epigenetic effect via declining telomere length in ageing fathers that is transferred to offspring via sperm (Bauch et al., 2019; Eisenberg, 2019; Sparks et al., 2021). Thus, taking into account paternal epigenetic inheritance, our results support quantitatively similar maternal and paternal contributions to telomere length inheritance.

We estimated the heritable component in the amount of telomere shortening early in life to be small (Table 1; Tables S7 and S11). The only other study of which we are aware that estimated heritability of telomere shortening, based on adult human twins, found a moderately low heritability ($h^2 = 0.28$; Hjelmborg et al., 2015). Telomere length early and late in the nesting period showed a strong genetic correlation between both ages ($r_c = .999$), thus supporting that telomere length at both ages is controlled by the same genes. This is in line with findings in other wild vertebrates later in life (Froy et al., 2021; Vedder et al., 2021). It should be noted, however, that the number of base pairs telomeres shortened over the nestling period (mean ± SD = $260 ± 171$) is small relative to the variation in telomere length at any age (SD = 590), potentially masking heritable variation in telomere shortening.

Fisher (1930) predicted that heritability of traits declines with their effect on fitness (a prediction that was empirically broadly confirmed; e.g., Mousseau & Roff, 1987). Our finding that additive genetic effects explained most variation in telomere length, but little variation in telomere shortening, fits this pattern, because previous investigations in our population showed that telomere shortening was a better predictor of survival than absolute telomere length (Boonekamp et al., 2014; Salomons et al., 2009).

It is tempting to suppose that a high heritability indicates that directional selection on telomere length could produce a fast evolutionary response. However, the high heritability estimate, representing the ratio between additive genetic and total phenotypic variance, results from the low environmental sensitivity that we find in our population in combination with a low amount of additive genetic variance (scaled to the mean: $CV = 6.41\%$ or 7.34%). The low additive genetic variance related to trait size resulted in a low evolvability estimate of telomere length. Telomere length evolvability was of similar magnitude in field crickets (Boonekamp et al., 2021). Whether this should be considered low or high is not obvious, but our evolvability estimate of 0.41% at age 4 days would allow an increase by 29 bp per generation, or close to 20 generations to increase population telomere length by 1 SD. More studies that provide evolvability estimates for comparison among populations and taxa and investigate the genetic link between telomere length and fitness are required to gain a more comprehensive picture.

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AUTHOR CONTRIBUTIONS

S.V. designed and organized the long-term project; J.J.B., S.V. and C.B. carried out fieldwork; E.M. and C.B. carried out laboratory work; C.B., J.J.B. and P.K. managed and analysed data; C.B. led the writing of the manuscript and all authors were involved in the reviewing and editing.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.k6djh9w74 (Bauch et al., 2021).

ORCID

Christina Bauch https://orcid.org/0000-0003-0218-5582

Jelle J. Boonekamp https://orcid.org/0000-0003-1900-627X

Peter Korsten https://orcid.org/0000-0003-0814-9099

Ellis Mulder https://orcid.org/0000-0002-1956-8734

Simon Verhulst https://orcid.org/0000-0002-1143-6868

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