Paternal Age and Transgenerational Telomere Length Maintenance: A Simulation Model

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Telomere length (TL) in offspring is positively correlated with paternal age at the time of the offspring conception. The paternal-age-at-conception (PAC) effect on TL is puzzling, and its biological implication at the population level is unknown. Using a probabilistic model of transgenerational TL and population dynamics, we simulated the effect of PAC on TL in individuals over the course of 1,000 years. Findings suggest a key role for an isometric PAC midpoint (PACmp) in modulating TL across generations, such that offspring conceived by males younger than the isometric PACmp have comparatively short telomeres, while offspring conceived by males older than the isometric PACmp have comparatively long telomeres. We further show that when cancer incidence escalates, the average PAC drops below the isometric PACmp and transgenerational adaptation to cancer ensues through TL shortening. We propose that PAC serves to maintain an optimal TL across generations.

Large, long-living mammals typically display short telomeres and repressed telomerase in somatic tissues1–4. Short telomeres can limit proliferative activity, diminishing the burden of cancer driver mutations and thereby attenuating cancer risk. However, short telomeres (and reduced telomerase activity) also increase predisposition to degenerative diseases, particularly in older animals. Converging evidence suggests that this cancer-degenerative disease trade-off operates not only across mammals but also within Homo sapiens1, as constitutively long telomeres5–8 and alleles that associate with long telomeres9–13 increase the risk of some cancers, while constitutively short telomeres14–16 and alleles that associate with short telomeres13,17,18 increase the risk of adult-onset cardiovascular disease (CVD), the main degenerative disease afflicting contemporary humans19. At the population level, across generations and in the face of changing environments (e.g., due to human migratory tendencies), this telomere length (TL)-mediated cancer-degenerative disease trade-off might require meticulous maintenance.

Most of the empirical data on human TL is based on leukocyte TL (LTL), which reflects TL in other somatic cells20. LTL is a highly heritable21–23 complex genetic trait that exhibits wide inter-individual variation from birth onwards24,25. In previous work, we proposed that polygenic adaptation is one of the mechanisms that maintain the TL-mediated cancer-degenerative disease trade-off26. Here we propose, and illustrate by simulations, that the effect of paternal age at conception (PAC) on the offspring's TL might be another mechanism that contributes to the maintenance of this trade-off.

The PAC effect, expressed as a longer LTL in offspring conceived by older men23,27–31, apparently stems from longer telomeres in sperm of older men31,32. This contrasts with the general phenomenon of telomere shortening with age in somatic tissues. Telomerase, the reverse transcriptase that elongates telomeres33, and sperm selection are potential explanations for the longer TL in sperm of older men. Telomerase, which is largely silent in somatic tissues, is active in the testes and presumably in the male germ stem cells34. Analysis of the PAC effect in twins also suggests the possibility of an age-dependent selection of sperm with longer telomeres29. The longer telomeres in the male germline are apparently inherited by the offspring according to Mendelian principles35,36.

While the effects of PAC on offspring’s TL across one24,27–29,31 and two30 generations have been studied, little is known about how PAC impacts TL dynamics across multiple generations. To gain insight into potential

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implications of the PAC effect for future TL trajectories, we incorporate the PAC effect into a simulation model of transgenerational TL that integrates the effects of heritability, PAC and cancer on TL.

Results

Model basic parameters. The model comprises a founding population (n = 1,000), male-female mating, fertility based on female and male ages, birth and death (Materials and Methods; Supplementary Figs S1 and S2). As TL is highly heritable21–24, and as studies showed that maternal inheritance is stronger than paternal inheritance23,24, the model applies a set of assumptions, including: (a) a mode of TL inheritance, whereby each parent's genetic makeup contributes directly to the offspring's TL using an inheritance factor that weights the maternal and paternal contribution to the offspring's TL based on empirical observations (Materials and Methods)21–24; (b) the PAC effect of 15 base pairs (bp) per each additional year of paternal age24,27–29,31; (c) age-dependent TL shortening16,20,21,28,31,32,37; and (d) a stochastic factor based on a standard deviation of TL observed across newborns24,25 in which the TL-outcome is normally distributed (mean = 0 bp; SD = 700 bp).

We used Monte Carlo simulation to drive the model and track TL of the population with/without the PAC effect and with/without TL-dependent cancer. Vital parameters, e.g., mortality, fertility, etc., were derived from CDC data (Supplementary Table S1)38,39. To make the simulations computationally tractable in the face of exponential growth, we restricted the transgenerational timespan to 1,000 years and imposed a constraint on the maximum number of lifetime births per individual (Materials and Methods).

The initial series of simulations covered a range of birth TLs (7,000–11,000 bp) in the founding population, but key simulations focused on birth TL of 9,500 bp, the approximate birth TL empirically observed in humans24,25.

A key parameter in the simulations was the ‘telomeric brink’ (TB), i.e., TL = 5,000 bp, which denotes a 50% probability of imminent death because of compromised ability to maintain homeostasis through cell replication (Supplementary Fig. S3)40,41. The TB (Materials and Methods) is based on the average TL observed in dyskeratosis congenita, a monogenic disease with critically short TL due detrimental mutation in telomere maintenance genes41. Another key parameter was TL-associated cancer (and its consequential mortality). This parameter is based on formulation of TL-dependent cancer incidence in which the probability of acquiring cancer, presumably due to the accumulation of driver mutations, in the course of a given year is a function of an individual's age and TL40.

Simulations without cancer. Our first series of simulations focused on TL dynamics in the presence of the TB, but without cancer or the PAC effect, over the course of 1,000 years for birth TLs of 7,000–11,000 bp in the founding population. The selective pressure exerted by the TB, together with the mixing TLs through inheritance, plus the stochastic effect across the population, resulted in upward birth TL trajectories whose steepness was inversely related to mean birth TL in the founding population (Fig. 1).

We then repeated these simulations with the PAC effect functioning in two independent modes: (i) offspring conceived by older fathers having a longer TL (a PAC+ mode), presumably because TL is longer in sperm of older men (Fig. 2A), and (ii) offspring conceived by younger fathers having a shorter TL (a PAC− mode), presumably because TL is shorter in sperm of younger men (Fig. 2B). The PAC+ mode further increased the upward transgenerational TL trajectories without bound, while the PAC− mode caused a reversal of the TL trajectories downward to the TB. For mean TL at birth = 7,000 bp in the PAC− mode, the initial transgenerational trajectory was slightly upward, then assuming a downward course with wide oscillation (i.e. year-to-year variance). For all other mean TLs at birth in the PAC− mode, the trajectories first descended towards ~7,000 bp with minimal TL oscillations and then plunged downward with wide TL oscillations. These configurations reflect the independent and opposing pressures of the TB (upward) and the PAC− effect (downward) on the transgenerational TL trajectories. The widening TL oscillations with the passage of time stems from the contraction of sample size as the population moves towards extinction due to critically short TL.

Figure 1. Transgenerational telomere length (TL) dynamics in the presence of the telomeric brink (TB) over the course of 1,000 years. Telomere length at birth for the founding population ranges between 7,000–11,000 bp.
Clearly, whether in the positive or negative mode, a unidirectional PAC effect exerts a ‘destabilizing’ influence on the transgenerational TL trajectory. For this reason, focusing on mean birth TL of 9,500 bp, we tested a simultaneously positive and negative PAC effect, i.e., a bidirectional (PAC±) mode in which the relative change in the offspring TL is determined with regard to a fixed age about which the PAC effect is centered, i.e., a PAC midpoint (PACmp). In this mode, when paternal age was older than the PACmp, it acted in a PAC+ mode, lengthening TL in the offspring by 15 bp/year for each paternal year greater than PACmp; when PAC age was younger than the PACmp, it acted in a PAC− mode, shortening TL in the offspring by 15 bp/year for each paternal year less than PACmp. Thus, when the mean PAC for a population is above the PACmp, there is an overall transgenerational lengthening of population TL; in contrast, when the mean PAC is below the PACmp, there is an overall transgenerational shortening of population TL (Fig. 3A). As per the PAC− model (Fig. 2B), the wider TL oscillations for PACmp = 50 or 55 years reflects the contraction of sample size as the population moves towards extinction due to critically short TL.

Through simulations, we then found the PAC age for a founding population with mean birth TL = 9,500 bp, where the transgenerational TL trajectory was flat (i.e., at equilibrium). We refer to this PAC as the isometric PACmp (34.2 years; Fig. 3A). Next, we examined the effect of this specific isometric PACmp on mean birth TLs ranging between 7,000–11,000 bp in a founding population. At this PACmp, the transgenerational TL trajectories converged toward 9,500 bp (Fig. 3B). These findings underscore a key outcome of the model, namely, PACmp and mean population TL are inter-connected traits. In this way, isometric PACmp drives convergence of the mean population TL toward the initial TL from which the isometric PACmp is derived.

Simulations with cancer. To illustrate the PAC-environment interaction, our next series of simulations were based on theoretical scenarios of a changing environment wherein the cancer incidence escalates, e.g. due to an increase in the cumulative mutation load. Focusing on mean birth TL of 9,500 bp (year 0), we simulated the transgenerational TL trajectories, based on the following settings: baseline cancer (scale 1), and increased
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short telomeres in offspring of younger fathers, i.e., PAC
considerably between 1876 and 1979 (from 36.9 years to 27.9 years) and only started to increase after 197948.
In human evolution. Comprehensive historical analyses show, for instance, that in the US mean PAC declined
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Discussion
The PAC effect may apply across the primate lineage, since it is also observed in chimpanzees47. This begs the
question why the PAC effect has evolved. Our simulations show an upward transgenerational TL trajectory when
the PAC effect is expressed in the inheritance of comparatively long telomeres by offspring of older fathers, i.e.,
PAC+ (Fig. 2A), and a downward trajectory when the PAC effect is expressed in the inheritance of comparatively
short telomeres in offspring of younger fathers, i.e., PAC− (Fig. 2B). A bidirectional (PAC+) and isometric PACmp
effect (Fig. 3) engender stability of TLs across generations and are thus more evolutionarily likely. The biology
of the male germline affords the evolution of this effect, since testes are formed in utero and sperm production
commences at puberty, forging functional sperm shortly thereafter and throughout the remaining life course43.
Given the numerous replications of stem cells, which continue unabated in the male germline, sperm TL may be
amenable to modification29,43,44.
Over the course of human evolution, the isometric PACmp has probably fluctuated near the mean age of male
reproduction. In light of increasing mean PAC in recent decades in many western countries45–47, we might expect
that TL has been progressively increasing across generations. However, several decades represent a tiny period
in human evolution. Comprehensive historical analyses show, for instance, that in the US mean PAC declined
considerably between 1876 and 1979 (from 36.9 years to 27.9 years) and only started to increase after 197948.
Estimates of historic and pre-historic paternal age, which are likely more evolutionarily informative, are difficult
to come by. In contemporary hunter gatherer populations, mean PAC is approximately 40 years49, suggesting that
TL may have been getting successively shorter in western countries over the past century and that the recent rise
in PAC has reversed this trend.

Figure 4. Transgenerational telomere length (TL) dynamics over the course of 1,000 years in the presence of
cancer incidence (scales 2 and 4) in which the cancer incidence was scaled upward 2 and 4 fold, respectively. For
each cancer incidence scale, we simulated the transgenerational TL in response to increased cancer based on two
sub-scenarios: (a) without the PAC effect, and (b) with the PAC effect. To this end, we allowed the model to oper-
ate without PAC for the first 500 years, and then introduced the PAC effect into the model beginning at year 500.
In the absence of the PAC effect, increased cancer attenuated the upward transgenerational TL trajectories due
to the upward pressure imposed by the TB (Fig. A,B). However, even in the face of a four-fold increase in cancer
incidence, the transgenerational TL trajectory remained upward, suggesting that the TB continued to drive up TL
and thereby increase cancer incidence. In contrast, introduction of the PAC+ effect (at isometric PACmp) not
only stabilized the transgenerational TL in the absence of cancer, but also shifted the direction of the transgen-
erational TL trajectory downward in the presence of cancer. This PAC effect was proportional to the magnitude
of the increase in cancer incidence (Fig. 4C,D), presumably because older individuals were likely to die from the
disease, which thereby decreased the mean PAC of the population.
To further understand the relation between PAC and the transgenerational TL dynamics in the face of can-
cer, we expanded the scope of the simulations to include founding populations with mean birth TL ranging
from 6,000 to 15,000 bp (year 0). For each mean birth TL, we identified a unique isometric PACmp that stabilized
the transgenerational TL for the period during which the PAC effect was included in the simulation (i.e. years
500–1000), denoted as iso-PAC500. The relationship between each iso-PAC500 and this TL was then plotted for
TL-dependent cancers at scales 1, 2, and 4-fold. Results were fitted to logistic curves and exhibited two key fea-
tures (Fig. 5): First, a younger iso-PAC500 maintained a comparatively long equilibrium TL. However, the relation
between the two variables was nonlinear and complex. This likely reflects the opposing selective pressures of the
TB, pushing TL upward from the bottom of the TL distribution, and cancer, pushing TL downward from the top
of the distribution. Second, as the cancer scale factor increased, the iso-PACmp decreased. Taken together, findings
suggest that the iso-PACmp and equilibrium TL are related within a given environment, whereby a change in one
determines a change in the other.
Another major finding of our simulations is that the PAC effect and cancer can work interactively because older men are more likely to die from cancer, thereby shifting downward the PAC distribution and thus TL. It follows that the reverse might also be true, i.e., if later life survival increased, the distribution of PAC and thus TL would be shifted upward. Such an effect on TL might allow relatively rapid changes towards an optimal TL in response to environmental and demographic conditions. We note in this regard that in contemporary humans, both cancer and CVD primarily occur during the post-reproductive years. Natural selection, mediated via factors such as polygenetic adaptation and PAC, may have adjusted in relatively short time periods the optimal TL to balance cancer against degenerative diseases and lessens their impact during the reproductive years.

Since the cancer-degenerative disease trade-off is a prominent explanation for TL evolution and there are clear empirical precedents to build on, our simulation model focuses on these factors. Another key driver of the optimal TL might be exposure to infection and thus TL-dependent immune function during early life and thereafter. In present societies, drivers of the PAC effect on TL may also act through demographic pathways. For instance, the age of conception is positively correlated with social-economic status and inversely with early-life poor health. Life expectancy and TL are correlated; thus, correlations of TL with health prior to and during reproductive ages may affect PAC. Moreover, cancer in older family members may affect the PAC of younger members since cancer impacts both family economic resources and family member health. Such PAC drivers are expected to be complex but may be significant and fast acting mechanisms for population adaptation in a changing environment.

Finally, we hope that our model will be further refined and serve as a catalyst for testing through simulations the impact of diverse environmental and demographic settings on the optimal TL in the human population.

Materials and Methods

General approach. Our model is based on empirical observations from epidemiological studies and theoretical considerations, as described under Results. It comprises population-level processes that are incorporated into four modules (Supplementary Figs S1,S2). The first module characterizes each individual (virtual human) by sex, date of birth, date of death, birth TL, and the rate of age-dependent TL attrition; together, birth TL and TL attrition define each individual’s TL throughout his/her life course. Based on cross-sectional and longitudinal studies of age-dependent TL shortening, the attrition in TL during the first two decades is taken as 1,400 bp and at 25 bp per year thereafter.

The second is a module of TL inheritance, whereby a newborn’s birth TL is specified by a combination of independent processes. One process specifies TL at birth as a weighted average of the maternal and paternal birth TLs, respectively. The weights are given by a maternal inheritance factor of 57.5%. Next, to account for natural TL variance, a stochastic process is used to adjust the birth TL by a random amount taken from a normal distribution with standard deviation 700 bp. The PAC effect is taken as 15 bp/yr. For PAC+ models, the isometric PAC midpoint (PACmp) is given as 15 years (the minimum PAC in the fecundity module below). Thus, all fathers older than 15 years will endow their offspring with increasingly longer TLs. For PAC− models, the PACmp is given as 55 years (the maximum PAC in the fecundity module). Thus, all fathers younger than 55 years will endow their offspring with shorter TLs. Intermediate models for the PAC effect are similarly parameterized by a PACmp between 15 and 55 years (Supplementary Fig. S4).

The third module, fecundity, accounts for the distribution of maternal and paternal ages. This model assumes that all living females become pregnant with a probability distribution parameterized solely by age.
(Supplementary Table S1). Male fertility is similarly modelled in an all-or-none fashion, such that fertile males have uniform probability of a successful mating during their reproductive years and are infertile otherwise. Due to computational limitations imposed in proportion to population size, rapid population expansion was attenuated by arbitrarily restricting each female to a maximum of 3 lifetime births to facilitate a simulated period of 1,000 years.

The fourth module, mortality, determines the individual’s death year, which may be ascribed to one of three independent processes. A baseline mortality rate is taken as the probability that an individual dies in a given year. Baseline mortality is parameterized only by age (Supplementary Table S1), and is therefore independent of TL-related causes of mortality. TL-dependent causes of death are modeled independently based on whether the effect is proportional or inversely proportional to TL (i.e. effects that are more likely to occur in individuals with relatively longer telomeres vs. effects that are more likely to arise in individuals with relatively shorter telomeres). Because TL must be always positive, the probability of death must approach 100% as TL approaches zero. Causes of death linked to short TL, therefore, fall under a TB Model, whereby TL is inversely correlated with probability of death. This is characterized as a logistic function over TL centered at 5,000 bp (Supplementary Fig. S3). Conversely, death related to long TL is determined by a Cancer Model, which is parameterized by both age and birth TL. (This model is based on equation 8 in supplement40. These potential causes of death are independently simulated for each year of the individual’s life and the earliest cause of death determines the age at death of the individual. Because birth TL is always positive and finite, the TB model guarantees that all individuals will eventually die.

Together, these four modules were run through Monte Carlo simulation (Supplementary Fig. S2) written in the Scala programming language, with an initial population consisting of 500 males and 500 females and allowed to proceed for a period of 1,000 cycles (years). The state of the population was tracked year-over-year in the simulator and summary statistics generated corresponding to mean newborn TL. The simulation itself was run for multiple module variants and parameterized by trial number, presence or absence of a PAC effect with corresponding $PAC_{\text{cancer}}$, TL-dependent cancer module (including a scaling factor for arbitrariness increasing cancer risk), maternal inheritance, fecundity module, mortality module, and a constant birth TL for the founding population. Each module was then run in parallel on the Rutgers University Harpertown High Performance Computing Cluster at Rutgers University, consisting of approximately 300 compute nodes each outfitted with an 8-core Xeon 2.83 GHz (Intel E5440) Processor, and 16 GB memory. Finally, the outputs from each of these simulations (50 runs per scenario) were combined for analysis.

Early simulations of the PAC effect displayed an abrupt initial decline in trangenerational TL with wide variance which stabilized over the course of several hundred cycles. This decline most likely arose from the model starting with only young individuals (since the model had to run longer for individuals to age) and only young men thus being able to reproduce initially. To overcome this artifact, we constructed a founding population with a heterogeneous age distribution as follows: 100 individuals were generated each year over the course of 100 years (total = 10,000 individuals) prior to initiating the simulation. Of the surviving population, 1,000 individuals were randomly chosen to establish the founding population. In this way, the founding population reflects the wide age distribution arising from the mortality module.

Data and Materials Availability

The code of our model is available at http://njms.rutgers.edu/centers_institutes/humandevelopmentandaging/.

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**Author Contributions**

K.H. and A.A. conceptualized the design of the study. K.H. and R.S. collected the data used for the final analysis. K.H. performed the data analysis (simulation) and interpreted the findings. K.H., R.S., J.A., J.K., D.E. and A.A. drafted, reviewed, and made critical revisions of the manuscript. The final approval of the version to be submitted for publication was conducted by K.H., R.S., J.A., J.K., D.E. and A.A.

**Additional Information**

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