In a Stationary Population, the Average Lifespan of the Living Is a Length-biased Life Expectancy, Version 2

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In a stationary population, the average lifespan of the living is a length-biased life expectancy

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ABSTRACT:
What is the average lifespan in a stationary population viewed at a single moment in time? Even though periods and cohorts are identical in a stationary population, we show that the answer is not life expectancy, but a length-biased version of life expectancy. That is, the distribution of lifespans of the people alive at a single moment is a self-weighted distribution of cohort lifespans, such that longer lifespans have proportionally greater representation. One implication is that, if death rates are unchanging, the average lifespan of the current population always exceeds period life expectancy. This result connects stationary population lifespan measures to a well-developed body of statistical results; provides new intuition for established demographic results; generates new insights into the relationship between periods, cohorts, and prevalent cohorts; and offers a framework for thinking about mortality selection more broadly than the concept of demographic frailty.
How long, on average, will the people who are currently alive live? It turns out that, in a stationary population, this “average lifespan of the living” (ALL) can be interpreted as a length-biased life expectancy—meaning, a transformation of life expectancy in which lifespans are re-weighted by their own values. We will show that this has several important implications—such as that period life expectancy is always less than the average lifespan of the people alive in that period—and provides intuition for a number of previous results about stationary populations.

Length bias, also known as size bias or a self-weighted distribution, occurs when units are observed in proportion to their size (e.g., Correa and Wolfson 1999, Patil 2005, Lann and Falk 2005). Demographers are likely to be most familiar with length-biased sampling in the context of Preston’s (1976) analysis of the family sizes of children vs. women. A survey that asks gestational parents how many children they have will receive one report per parent, while a survey that asks children how many children are in their family will receive one report from each one-child family, but six reports from each six-child family, and so forth (Bytheway 1974, Preston 1976, Patil and Rao 1978, Ruggles 2012, Song and Mare 2014, Song 2020). Similarly, a length-biased—or self-weighted—lifespan distribution is a distribution of lifespans that is weighted by lifespan, such that the longer-lived are more likely to be observed, in proportion to their longer lifespans. We will show that, in a stationary population, the average lifespan of the living (ALL) is such a length-biased lifespan. In what follows, we formally prove this result; provide an intuitive explanation that also clarifies relationships between the lifespans of periods, cohorts, and prevalent cohorts; offer several empirical illustrations; and discuss how this result provides intuitions and implications for other demographic results and frameworks.

1. Proof
   a. Background: Length Bias
   Consider a non-negative real-valued random variable \( Y \geq 0 \)—lifespans or family sizes, for example—with probability density function \( f(y) \) and mean \( E[Y] > 0 \). The length-biased distribution arising from \( f(y) \) has the density:

   \[
   f_{lb}(y) = \frac{yf(y)}{E[Y]}
   \]
and the mean of this length-biased distribution is:

\[ E_{lb}[Y] = \frac{E[Y^2]}{E[Y]} . \]  \hspace{1cm} (1)

This length-biased mean can be re-expressed, by using the definition of the variance

\[ \sigma_Y^2 = E[Y^2] - E[Y]^2, \]

as a function of the squared coefficient of variation, \( cv_Y^2 = \frac{\sigma_Y^2}{E[Y]^2} \):

\[
E_{lb}[Y] = \frac{\sigma_Y^2 + E[Y]^2}{E[Y]}
\]

\[ = E[Y] + \frac{\sigma_Y^2}{E[Y]}
\]

\[ = E[Y] (1 + cv_Y^2) \]  \hspace{1cm} (2)

These equations describing length-biased samples are well-known in the sampling statistics literatures (e.g., Cochran 1977: 249-255, deCarvalho 2016).

b. Proof: Length-Biased Life Expectancy

We will show that, in a stationary population—one with constant birth and death rates, zero net migration at any age, and zero population growth— the living population’s distribution of age at death, denoted \( X \), is a length-biased sample of the cohort distribution of \( X \). Denote the number of survivors at any age \( x \) \( l(x) \), such that the cohort survivorship is \( l(x) / l(0) \) and the population at any moment in time has size

\[ \int_0^\omega l(x) \, dx \] where \( \omega \) is the oldest attainable age. Let the number of deaths at each age be denoted \( d(x) \). In a stationary population, at each instant the number of births equals the number of deaths, so \( l(0) = \int_0^\omega d(x) \, dx \). Thus, the density of ages at death is given by \( f(x) = d(x) / l(0) \). The average lifespan (life expectancy at birth) of this cohort, \( e(0) \), is:

\[
e(0) = \frac{\int_0^\omega l(x) \, dx}{l(0)} = \frac{\int_0^\omega xd(x) \, dx}{\int_0^\omega d(x) \, dx}
\]  \hspace{1cm} (3)
(where \( \int_0^\infty l(x)dx = \int_0^\infty xd(x)dx \) because both express the total lifespan of the cohort by alternatively summing the incremental lifetime lived at each age, or the full lifespan of those dying at each age).

In this population, the lifespan distribution of the living, \( f_t(x) \), is a length-biased transformation of the lifespan distribution of each cohort (or period) \( f(x) \):

\[
f_t(x) = \frac{xd(x)}{\int_0^\infty l(x)dx} = \frac{xf(x)/l(0)}{e(0)/l(0)} = \frac{xf(x)}{E_{f(x)}[X]}.
\]

The first line of Eq. (5) reflects that, among the people currently alive, the number of people who will die at exact age \( x \) is the product of the per-cohort deaths at age \( x \), \( d(x) \), and the \( x \) cohorts that have future deaths at age \( x \); and the density that will die at exact age \( x \) is the ratio of those \( xd(x) \) deaths to the size of the full living population, which is \( \int_0^\infty l(x)dx \). Life expectancy, \( e(0) \), is the expected age at death in the cohort, denoted \( E_{f(x)}[X] \) (with the \( f(x) \) subscript to underscore that this expectation applies to the cohort, not the living population). Since Eq. (5) instantiates Eq. (1), Eq. (5) establishes that the distribution of lifespans in the living population is a length-biased sample of the lifespans in the stationary population's cohort or period.

Similarly, Eq. (5), along with Eq. (2) and (3), implies that the average lifespan of the living is:

\[
ALL = \frac{\int_0^\infty x^2d(x)dx}{\int_0^\infty xd(x)dx} = \frac{E_{f(x)}[X^2]}{E_{f(x)}[X]} = e(0) + \frac{\sigma_x^2}{e(0)}
\]

(6)
An alternative derivation of these expressions for the ALL, not relying on Eq. (2) and (3) but derived directly from the life table functions, is given in Appendix 1.¹

Length bias is not the only perspective in which to view these results, but in the remainder of this article, we use the length-biased interpretation of the ALL as a length-biased sample to develop new insights and connect these results to other concepts in demography and other fields.

2. Intuition
The results in Eq. 5 and 6 connect lifespan demography to well-developed areas of statistics and help to clarify other results about stationary populations, discussed below. The intuition for this result is simple. By definition, the living population dies at an age older than its current age. This has the consequence that older ages of death are disproportionately likely among the living population, compared to its original birth cohorts. This selective sample of lifespans is illustrated in Figure 1, which shows a Lexis diagram in which the living population is defined by a vertical line and their future deaths are only those deaths included in the triangle on the right of this line. In epidemiology, the upper triangle in Figure 1—representing the deaths that contribute to the lifespan of the living—is called a prevalent cohort (e.g., Törner et al. 2011). The two horizontal bands reflect two example ages at death, illustrating that older ages at death will be experienced by more currently-living cohorts than younger ages at death will. This concordance between age at death and the number of cohorts that can still reach that age at death produces length bias.

The results illustrated here offer new perspective on the relationship between cohorts, periods, and prevalent cohorts. Cohort life expectancy takes everyone who

¹ Other derivations are also possible. For example, our Equation (6) can be recovered from a decomposition of life expectancy provided by Cohen (2015), which divides life expectancy into three components defined by any arbitrary age \( x \): the lifespan lived by those who die before age \( x \), the first \( x \) years lived by those living beyond \( x \), and the remaining life years lived beyond \( x \), each weighted by the proportion of the original birth cohort that experiences them. The ALL in a stationary population can then be understood as the survivorship-weighted sum, over all ages, of Cohen’s second and third decomposition components, representing the past and future lifespans of those who survive to each successive age. Here, our goal is simply to provide a single lens through which some key demographic lifespan measures can be related to one another.
starts their lives at a single moment in time and follows that unselected sample across time. Period perspectives take everyone alive in a moment, regardless of when they began, but stay within that single moment in time. Prevalent cohorts, whose lifespan is the ALL, are distinct from both the cohort and the period perspectives. Like periods, they first select people on their longevity but then, like cohorts, follow them through their own lifespans. Thus, the length bias arises because the ALL is asking a cohort question about a period slice of the data.

An alternative formulation underscores what is unique about prevalent cohorts. Cohort life expectancy and period mean age at death are the average lifespans of a birth cohort or “death cohort” (e.g., Riffe, Schöley, and Villavicencio 2017), respectively. Thus, cohort and period lifespans each constrain birth or death time (respectively) to occur at a single instant, neither earlier nor later, while leaving the time of the other vital event (death or birth) unconstrained. In a stationary population, these alternative constraints generate the same distribution. Prevalent cohorts, in contrast, impose a one-sided constraint on each of birth time and death time: the ALL summarizes the lifespans of people who were born before time $t$ and die after $t$. This produces the triangle shape in the Lexis diagram because, for young ages at death, there are very few combinations of birth and death cohort that meet this constraint, but for old ages at death, there are very many combinations of birth and death cohort that meet this constraint.

A practical application of this result is that, if age-specific death rates are unchanging over time, the lifespan of the population alive in a period will necessarily be greater than the period life expectancy.\(^2\) More broadly, other contexts that define a ‘cohort’ through cross-sectional membership in some category will produce similarly length-biased estimates of the duration of category membership. For example, epidemiological research designs that begin by selecting people with some incurable medical condition at a particular time point, and then follow those people as a cohort to see how long they live—a prevalent cohort study design—generate length-biased lifespan estimates analogous to the bias in the ALL (Zelen and Feinleib 1969). The results here underscore that demographers can conceptualize living populations and their relationship to underlying birth cohorts with the same analytical tools that epidemiologists use to represent prevalent cohort samples and their relationship to

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\(^2\) This implication assumes that everybody does not die at the same age, as we explain below.
disease incidence (e.g., Alho 1992, Addona et al. 2009, Carone et al. 2012, Keiding et al. 2019). Finally, a conceptual implication—recalling the familiar family size example—is that lifespans cluster time the way that families cluster children.

3. Empirical illustration: Life expectancy, variation in the age at death, and the ALL

The expression for the ALL given in the last line of Eq. 6 generates useful insights into the relationship between the ALL, life expectancy, and variance in the age at death. It implies that the ALL is always greater than life expectancy (except in the special case that everyone dies at the same age, in which case the ALL equals life expectancy). More generally, Eq. 6 shows that, holding life expectancy fixed, a population with more variable ages at death will have an older ALL that is more divergent from life expectancy. Conversely, holding the variation in the ages at death fixed, a population with a larger life expectancy will have a smaller ALL that is more similar to life expectancy.

The relationships between life expectancy, variation in the age at death, and the ALL are illustrated by calculating the ALL implied by model life tables. As a simple illustration, Table 1 summarizes four such life tables, which form two pairs. The first pair are two life tables with identical life expectancy (60 years) but divergent variance (represented in the table as the standard deviation, for ease of interpretation). The one with the smaller variance (the UN’s male Far East Asian life table) has an ALL that is 8 years above its life expectancy, while the one with the larger variance (the Coale-Demeny female South life table) has an ALL that is more than 13 years above its life expectancy. The second pair are two life tables with nearly-identical variance but radically different life expectancy. In this case, the life table with the larger life expectancy (the UN’s female Latin American life table at life expectancy 67) generates an ALL that is less than 10 years above life expectancy, while the low-lifespan table (Coale and Demeny’s male North life table at life expectancy 21) generates an ALL that is more than 30 years above life expectancy. In this last population, the combination of

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3 UN Extended Model Life Tables were downloaded from https://www.un.org/en/development/desa/population/publications/mortality/model-life-tables.asp on October 22, 2020. Specifically, we used the complete (non-abridged) life tables in one-year age increments.
relatively large variance and very small life expectancy generate an extreme
discrepancy between the lifespan of the population alive in cross-section and the
lifespan of a cohort.

Figure 2 illustrates how this divergence between life expectancy and the ALL occurs
by showing the age at death distribution for the living population (hollow bars) and for
birth cohorts (filled bars) implied by these four model life tables. As is well-known,
deaths at young ages have an outsized effect on life expectancy. Figure 2 shows that
these deaths are dramatically underrepresented in the living population’s lifespans. For
example, someone who dies at 1 month old would need to appear in the living
population at age less than 1 month, and the population contains few people in this
narrow age band. More generally, the distribution of ages at death is systematically
pulled upward among the living, compared to the cohorts from which they spring.

Figure 3 takes a more global view of these relationships across the full set of UN
Extended Model Life Tables. (The life tables marked with Xs are the four example life
tables just discussed.) Panels A and B show that, at low levels of life expectancy, the
ALL diverges widely from life expectancy, while at high levels, it diverges very little.
This relationship between the ALL and life expectancy is not only driven directly by the
level of life expectancy, but also by life expectancy’s historical relationship with
variance in the age at death. Panel C illustrates that historical relationship for these
model life tables. This relationship between life expectancy and variation in the age at
death can be roughly periodized into three phases. First, the earliest increases in life
expectancy (away from the levels experienced by early human societies) may also have
increased variation in the age at death, relative to a context in which most deaths were
during childhood. Second, in the past few centuries, large increases in life expectancy
were driven by declining mortality at young ages when such deaths were less common
than in early human societies, and these changes tended to substantially reduce
variation in the age at death. Third, in high-longevity populations, more recent
increases in longevity are concentrated at older ages and may tend to reduce variation
in the age at death only a small amount, and sometimes increase it (Engelman et. al.
2010, Tuljapurkar 2010). Since the ALL will diverge from life expectancy the most when
the variance is high and life expectancy is low (Eq. 6), these relationships help to explain
why, as Panels A and B of Fig. 3 showed, the ALL is maximally divergent from life
expectancy when life expectancy is roughly in the age range 20-40, and converges
rapidly toward life expectancy once life expectancy reaches roughly age 50. Panel D of Fig. 3 shows the resulting relationship between the variance in the age at death and the ALL. Even though, fixing life expectancy, a larger variance in the age at death means a larger divergence of the ALL from life expectancy, the historical relationship between life expectancy and variance complicates the overall relationship between variance and the ALL. As life expectancy increases across these model life tables, the ALL’s divergence from life expectancy travels along the arcs shown in Panel D, from the uppermost values (representing low life expectancy contexts) then out to the right, then down toward the bottom left.

A few examples help to calibrate expectations about how much longer the living population will live, compared to birth cohorts. As Figure 3 implies, even in stationary populations, this additional lifespan can differ dramatically across populations. In a comprehensive review of hunter-gatherer societies, Gurven and Kaplan (2007) suggest that life expectancy at birth in such societies ranges from roughly 21-37; the model life tables in that range of life expectancy estimate average lifespans of the living that are 18-35 years older than life expectancy. On the other hand, for low-mortality populations, the divergence is much smaller. Recent U.S. life expectancy values are 76 for men and 81 for women. Model life tables at those values generate ALLs that are, respectively, 2.9-5.7 years and 2.1-4.3 years older than life expectancy. For hypothetical populations with life expectancy of 95 or older, in these models, the ALL diverges from life expectancy by less than one year.

4. The ALL, the average age, and the average remaining life

The last line of Eq. 6, which represented the ALL as a function of life expectancy and the variance in the cohort age at death, also provides a core intuition for a well-known, but somewhat inscrutable, formula for the mean age of a stationary population, denoted \( A \). The formula, given by Ryder (1975: 8) and Preston et al. (2001: 112), is:

\[
A = \frac{e(0) + \frac{\sigma_x^2}{e(0)}}{2}
\]  

(7)
Thus, from Eq. (6) and Eq. (7), ALL=2A: in a stationary population, the average lifespan of the living is twice the average age.

One way to understand this formula is to see that it follows from two properties of stationary populations. One is the property shown here: that the average lifespan of a stationary population’s members alive in a snapshot is a length-biased life expectancy. The second distinctive feature of stationary populations is that the distribution of lifetimes lived so far—that is, age—equals the distribution of lifetimes still to come—that is, remaining life expectancy (Müller et al. 2004, 2007; Vaupel 2009; Villavicencio and Riffe 2016). This property can be considered a form of time symmetry in stationary populations, implying that in a stationary population observed at a specific moment in time, a randomly selected individual is equally likely to be observed at any point in their lives, and on average is observed halfway through their lifespan (Kim and Aron 1989, Goldstein 2009). Thus, the average age of the population and the average remaining lifespan are each half of the ALL. In combination, these two properties imply that the average age and average remaining lifespan in a stationary population each equal one half of the length-biased life expectancy, generating Eq. 7.5

This perspective similarly provides intuition for a proof given by Finkelstein (2008: 268) that a stationary population with a larger life expectancy than a second stationary population need not also have a larger average remaining lifespan: thus, it is possible for $e_1(0) > e_2(0)$ while $A_1 < A_2$ (where $A$, which we used to denote mean age, also

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4 Specifically, Eq. (7) draws on the fact that the average age in a stationary population equals the average expected lifespan. This result was shown in a demographic context by Kim and Aron (1989) and Goldstein (2009), with elaborated historical context in Goldstein (2012). It was broadened to the more general claim about stationary populations referenced here—that the distribution of ages equals the distribution of past lifespans—in Vaupel (2009) and is generally considered to have been independently discovered in work by Brouard (1989) [which the authors are unable to read in its original French] and Carey (Müller et al. 2004, 2007) and subsequently (Villavicencio and Riffe 2016) dubbed the Carey-Brouard inequality. An alternative proof of the broader claim is presented in Rao and Carey (2015). The details of the Rao and Carey proof are controversial (Villavicencio and Riffe 2016, Rao 2021) but the claim is not. Finally, exploration of these time symmetries has been extended to other measures in stationary populations (Riffe 2015) and modified for stable populations (Vaupel and Villavicencio 2018). The intuition we provide here, based on length bias, is only one way of gaining insight into the mean age of a stationary population based on the symmetry of past and future lifespans.

5 Ryder (1975: 8-11) discusses implications of Eq. (7) for the relationship between life expectancy, the coefficient in the variation in the age at death, and the mean age of a stationary population. His discussion is an intellectual forerunner to our discussion in Section 3, above.
equals mean remaining lifespan). Finkelstein’s clever proof, which involves making a succession of changes to a survival curve, does not (to our mind) provide a clear intuition as to the mechanism by which a cohort with a larger life expectancy can have a smaller remaining lifespan. Given that average remaining lifespan equals average age in a stationary population, Eq. 6 and Eq. 7 clarify that this possibility will come to pass if the variation in the age at death in the second cohort is large enough to offset its smaller life expectancy. In that circumstance (and only that circumstance), a randomly-selected member of this second cohort at a single moment in time will have a longer lifespan than a randomly-selected member of the reference cohort, even though a randomly-selected member at birth will have a shorter lifespan.

5. Connections to other demographic concepts
a. Mortality Selection Without Frailty

By highlighting the inherent bias in lifespan measures that begin with a population defined by a period, the length-biased lifespan in Eq. (5) also gives a somewhat different perspective on mortality selection than demographers’ and biostatisticians’ traditional perspective based on frailty modeling. Frailty is a construct representing systematic and sustained individual differences in longevity (Manton et al. 1979, Vaupel and Yashin 1985, Vaupel and Missov 2014). It implies that, if the frailer people who die young did instead live to old age, their old-age mortality would be higher than that of the robust people who actually do survive to old age. In other words, the concept of frailty implies that there are stable differences in individuals’ expected longevity, and, therefore, differences in actual longevity select the population by continually removing those with the lowest expected longevity (Feehan and Wrigley-Field 2021: 369-371.).

Yet recent theoretical and empirical investigations (Steiner and Tuljapurkar 2012; Caswell 2014; Hartemink, Missov, and Caswell 2017) suggest that the vast bulk of heterogeneity in lifespans may be due to stochastic differences in individual luck, rather than stable frailty. The results here highlight the importance of lifespan variability in generating mortality selection, whether or not there is any such frailty in the population. Indeed, no concept of frailty is needed to generate any of the results in this article. Imagine that everyone in the population has exactly the same age-specific hazards and lifespans differ purely by chance. Even in this situation, the ALL reflects a
length-biasing selection on individual differences in actual longevity—even though those differences in actual longevity do not reflect differences in individuals’ expected longevity. Viewing the population in a cross-section intrinsically means viewing individuals in proportion to their lifespan. The cross-section selects cohort members, not on frailty (expected longevity), but on actual longevity.

b. Longevity of the living population vs. longevity of living cohorts: Comparison with the Cross-Sectional Average Lifespan (CAL)

The ALL is part of “a family of mortality indicators that make use of cohort information but refer to only one period” (Guillot and Payne 2019: 418). One member of that family deserves special mention because, like the ALL, it uses cohort mortality to describe the longevity of cohorts alive during a particular period. That measure is the cross-sectional average lifespan (CAL), proposed by Brouard (1986) and developed by Guillot (2003). A comparison between the ALL and CAL is informative about both measures.

The CAL at time $t$ is defined in terms of the cohort-specific survivorship $p_c(x, t - x) = l_c(x) / l_c(0)$, the survivorship of cohort $c$ at age $x$, as:

$$CAL(t) = \int_0^\infty p_c(x, t - x)dx$$

Thus, “$CAL(t)$ is the sum of proportions of survivors among the various cohorts present in the population at time $t$” (Guillot 2003: 42). We note that this implies that the CAL summarizes lifespans using only the deaths included in the left-hand triangle of our Fig. 1 (past deaths of living cohorts), in contrast to the ALL, which uses only the deaths included in the right-hand triangle (future deaths of living people).^6

This key contrast between the CAL and the ALL—whether past or future deaths of living cohorts comprise the lifespans that the measure summarizes—also means that the two measures describe different population units: the ALL describes the (future) longevity of living people, while the CAL describes the (past) longevity of living cohorts. That relationship, in the CAL, of living people to their current cohort survivorships—

^6 This restriction to past deaths of living cohorts generates one of the key practical advantages of the CAL as a period longevity indicator: while it is based on genuine cohort, rather than synthetic cohort, mortality (unlike period life expectancy), it requires no data on the future (unlike cohort life expectancy or the ALL).
reflecting the mortality that such people did not succumb to—may or may not proxy meaningful aspects of their own lives. In many contexts, being a rare survivor of a cohort with high past mortality, compared to a survivor of a cohort with lesser mortality, might capture something quite meaningful about the survivor (such as the intensity of the disease exposures each may have endured). On the other hand, in highly unequal and segregated contexts, high past mortality concentrated in a particular (disadvantaged) subpopulation might not be a meaningful descriptor of the experiences of survivors drawn largely from a different (advantaged) subpopulation within the same national population. In contrast to most other longevity measures, the ALL describes living populations directly, rather than the cohorts into which they were born.

6. Conclusion
We have shown that, in a stationary population, the lifespans of the living population form a length-biased sample of the cohort (and period) lifespan distribution. This insight connects demographic lifespan measures with a well-developed body of statistics (e.g., Cochran 1977: 249-255, deCarvalho 2016), including many epidemiological applications (e.g., Hill et al. 2003, Asgharian et al. 2006, Törner et al. 2011), and provides new intuition for existing demographic results. It implies that, when death rates are unchanging, the average lifespan of the living always exceeds period life expectancy, and exceeds it by more years when variation in the age at death is large and life expectancy is low.

These results are relevant whenever the population of interest is the population that currently experiences a state, rather than the population that ever experiences a state. Here, we considered the state of being alive, comparing the lifespan of the living to the lifespan of a cohort. The same results would apply when characterizing the expected length of current marriages (Alho 2016), disease durations among those who currently have a disease (Zelen and Feinleib 1969), or spells of incarceration (Patterson and Preston 2008) or unemployment (Beach and Kaliski 1983) among those currently incarcerated or unemployed. Each of these applications involves what we termed “asking a cohort question about a period slice of the data,” introducing length bias.
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REFERENCES


Figure 1. The average lifespan of the living (ALL) is the average of the lifespans that end in the upper right triangle. These deaths represent the future deaths of the living population. The deaths in the left triangle reflect the past deaths of living cohorts. The horizontal bands illustrate the unequal number of cohorts that contribute deaths at different ages at death (e.g., 40 cohorts for age 40 and 60 cohorts for age 60).
<table>
<thead>
<tr>
<th>Life Table Family &amp; Gender</th>
<th>Life Expectancy</th>
<th>Standard Deviation of the Age at Death</th>
<th>Average Lifespan of the Living (ALL)</th>
<th>ALL – life expectancy divergence</th>
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<td>8.17</td>
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<tr>
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<td>21</td>
<td>25.43</td>
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<td>30.74</td>
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</tbody>
</table>

Table 1. Average lifespan of the living and its divergence from life expectancy for four model life tables. The first two life tables share a life expectancy; the second two share a variance (reported here in terms of the standard deviation) in the age at death.
Figure 2. Distribution of lifespans in a life table (solid bars) and its associated living population (hollow bars), corresponding to the life tables summarized in Table 1. The living population’s lifespans are older than the life table’s due to the selection intrinsic to the prevalent cohort. The distributions are displayed to age 110 (the model life tables extend to age 130). Note that, in the upper left panel, the living population appears visually to have a younger distribution of ages of death than the life table; this is a consequence of the relative absence of very young deaths among the living population creating proportionately more deaths at middle ages. Note also that the visually small peaks in the lower right panel result from the extremely high infant mortality of the life table, which distorts the graph scale.
Figure 3. ALL, life expectancy, and variance in the age at death in the full set of UN Extended Model Life Tables. The Xs distinguish the four life tables portrayed in Table 1 and Figure 2. (A) The average lifespan of the living (ALL) as a function of life expectancy. (B) The divergence of the ALL from life expectancy as a function of life expectancy. (C) Variance in the age at death as a function of life expectancy. (D) The divergence of the ALL from life expectancy as a function of variance in the age at death.
Appendix 1. Alternative proof that the average lifespan of the living (ALL) is a length-biased life expectancy.

Here we present an alternative proof that the ALL is a length-biased life expectancy, to complement the proof given in Equation 6 and provide additional intuition for the result. The proof begins by representing the ALL as the total lifespan of everyone alive, divided by the number of people alive:

\[
ALL = \frac{\int_0^\omega \int_y^x xd(x)dx dy}{\int_0^\omega l(x)dx} \\
= \frac{\int_0^\omega xd(x)\int_y^x dydx}{\int_0^\omega l(x)dx} \\
= \frac{\int_0^\omega x^2d(x)dx}{\int_0^\omega l(x)dx} \\
= \frac{\int_0^\omega \int_0^\omega x^2 d(x)dx}{\int_0^\omega \int_0^\omega d(y)dy dx} \\
= \frac{\int_0^\omega x^2d(x)dx}{\int_0^\omega xd(x)dx} \\
= \frac{l(0)\int_0^\omega x^2f(x)dx}{l(0)\int_0^\omega xf(x)dx} \\
= \frac{E_{f(x)}[X^2]}{E_{f(x)}[X]}. 
\]

(S1)

The expression for the total lifespan of the living, in the numerator of the right-hand side of the first line of Eq. S1, may appear inscrutable. It represents the total lifespan \(x\) of the \(d(x)\) people who die at age \(x\) within each cohort whose current age is \(y \leq x\). The total lifespan of the living therefore equals \(\int_0^\omega \int_y^x xd(x)dx dy\). From there, the proof employs “two standard strategems in formal demography” (Vaupel 2010: 1144), translating the number of people currently living \(\int_0^\omega l(x)dx\) to the number of future deaths \(\int_0^\omega \int_x^\omega d(y)dy dx\), and switching the order of integration.
Reference to Appendix 1