

Healthy life expectancy, mortality, and age prevalence of morbidity

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Abstract

In calculating period healthy life expectancy, the use of age-specific morbidity prevalence patterns assumes that age captures the important time-variation in the given health condition, i.e. that the disabling process is related to how long an individual has lived. However, many morbidity patterns are better classified by time-to-death. At advanced ages the conflation of an increasing chronological-age mortality pattern and a time-to-death morbidity pattern produces an apparent morbidity pattern that increases with advancing age. Differences in period healthy life expectancy over time or between populations cannot easily be partitioned into morbidity and mortality components because the period morbidity pattern may depend on an unknown future time-to-death process not captured by period mortality. We illustrate these concepts formally and empirically, using morbidity data from the U.S. Health and Retirement Study. While holding the time-to-death morbidity pattern fixed, we show that mortality reduction alone reduces the total life years with disability. We estimate the magnitude of this bias for different realistic morbidity patterns. This has implications for any between- or within-population comparisons of period healthy life expectancy conditioned on different age patterns of mortality.

1 Introduction

Healthy life expectancy (HLE) is among the most widely used metrics of population health. It combines information on mortality and morbidity to summarize the expected years of life lived in good health, however measured. If healthy life expectancy increases faster than life expectancy, morbidity is compressed into a smaller proportion of life. HLE can change because of changes in mortality, morbidity, or both.

Most often, HLE is calculated on the basis of morbidity prevalence and a life table. In calculating HLE, the use of age-specific morbidity prevalence

data implicitly assumes that a chronological age pattern best characterizes variation in the health characteristic over the lifespan. However, Riffe et al. (2016) show that the prevalence of many health characteristics in old ages is better measured by a pattern over time-to-death (TTD) or by both age and TTD. Most morbidity patterns increase with age, and can claim empirical regularity in this regard. However, we explain how the observed shape of a morbidity age-curve can change due to changes in mortality even with no change in underlying morbidity. At advanced ages the conflation of an increasing chronological-age mortality pattern and a TTD morbidity pattern produces an apparent morbidity pattern that also increases with advancing age.

In the cohort perspective the true and apparent morbidity patterns imply the same HLE. Problems arise in the period perspective. Differences in period HLE over time or between two populations cannot easily be partitioned into morbidity and mortality components, because the period morbidity component will depend on some unknown future TTD process. For the same reason, comparisons of disability prevalence rates by age are not recommended between populations with different underlying age schedules of mortality.

In this paper we illustrate these concepts formally and empirically, using morbidity data from the U.S. Health and Retirement Study (RAND 2017, HRS 2017). Assuming a fixed TTD morbidity function, we show that mortality reduction alone can lead to a lower fraction of years lived in poor health, and consequently morbidity compression. We estimate the magnitude of potential biases for different disabling processes, given different levels of mortality extracted from the Human Mortality Database (HMD 2015). We first explain how the age pattern of morbidity may partly be a function of mortality using both formulas and a schematic illustration.

2 Morbidity as a function of time to death

A joint pattern of morbidity prevalence by age and time to death could in principle obtain any shape. Empirically, such joint patterns appear to fall into a small set of basic types, and typically these patterns slope monotonically in a single direction over these two dimensions. For example, Figure 1 shows US female poor self-reported health (SRH) prevalence after age 70 broken down by age and TTD under three views. Panel 1a shows these data as a surface, whose more horizontally running prevalence contour lines indicate that time-to-death is a more important time measure to describe prevalence of poor SRH than age. Figures 1b and 1c, show the same prevalence data plotted with respect to the respective TTD and age margins. Of these, the TTD lines in 1b yield a more compact set of patterns, which one could imaginably summarize with a single line without much loss of

precision.

Thus our convenience assumption thus far of a single fixed morbidity prevalence over time-to-death is not overly unrealistic, at least for characteristics with this joint prevalence pattern.

This pattern holds for many, but not all, indicators of health and disability (Riffe et al. 2016). There is not a broad evidence base on such joint age and TTD patterns. While our simplifying assumption of fixed TTD patterns is an acceptable abstraction of the particular joint patterns from which they were derived, these patterns only serve to demonstrate an arithmetic point in the extreme. The degree to which prevalence is better measured by age or time-to-death for different disabling conditions remains an open research question.

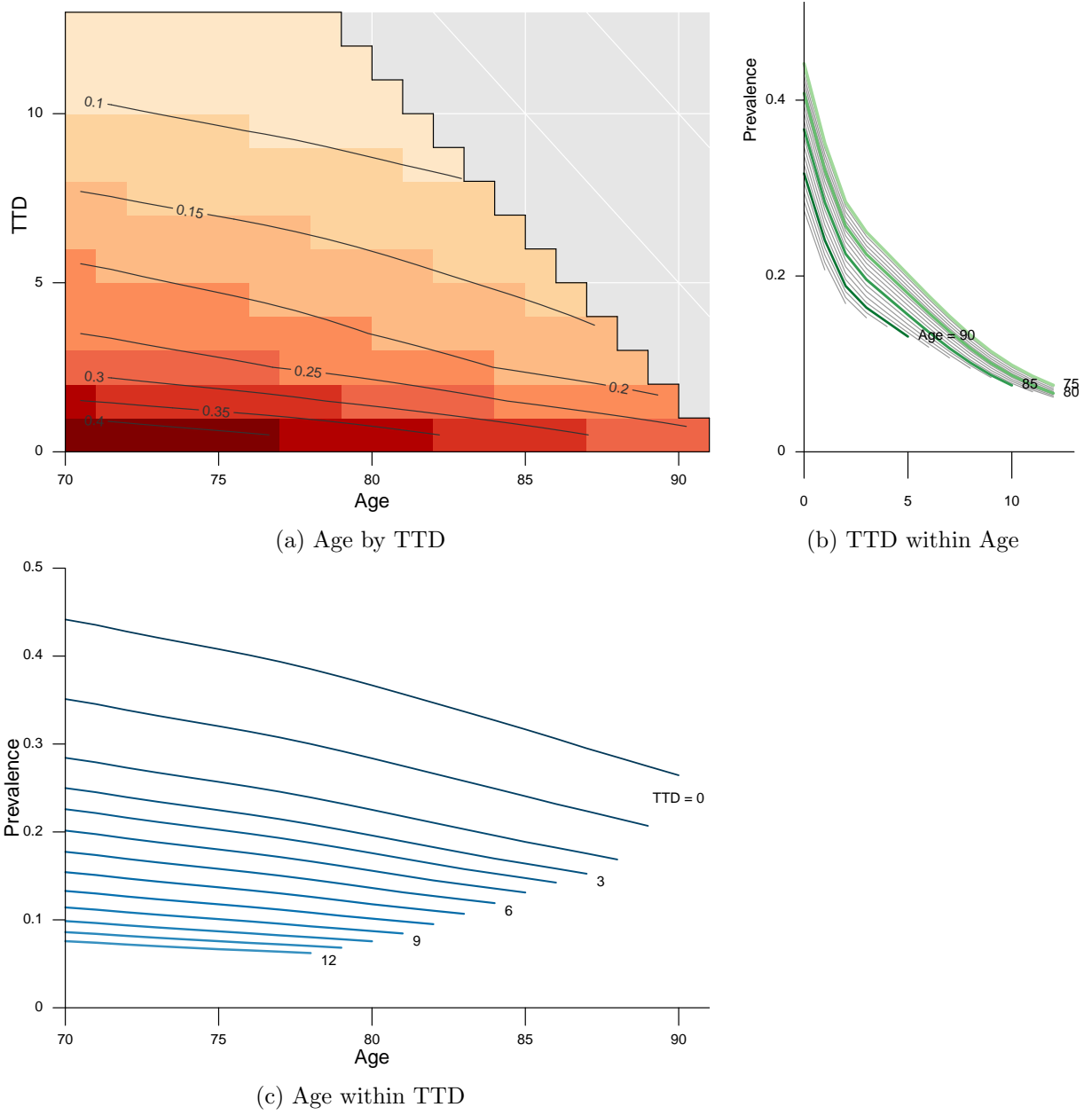


Figure 1: Prevalence of females from the 1920-1924 cohort with self-reported poor health, by time-to-death and age. The same data are shown by age over the time-to-death margin in (b) and by time-to-death over the age margin in (c). Comparing (b) and (c) we conclude that a single time-to-death pattern would better approximate the surface (a) than would a single age pattern.

Imagine a bad health condition, G , with prevalence that varies only as a function of time to death, y , and not as a function of chronological age,

a . Since the TTD prevalence distribution is very closely concentrated at the end of life, there will still be an apparent age function, $g^*(a)$. In this case $g^*(a)$ is a heterogeneous aggregate based on both mortality and the underlying TTD process:

$$g^*(a) = \frac{\int_0^\omega g(y)N(a, y) \, dy}{N(a)} \quad (1)$$

$$= \frac{\int_0^\omega g(y)N(a)\mu(a+y)\frac{\ell(a+y)}{\ell(a)} \, dy}{N(a)} \quad (2)$$

$$= \int_0^\omega g(y)f(y|a) \, dy \quad , \quad (3)$$

where $N(a)$ is the population aged a , $\ell(a)$ is life-table survivorship, and $\mu(a)$ is the force of mortality. $f(y|a)$ is the conditional remaining-years distribution, which gives the probability of dying in y years given survival to age a . The expression (3) says that the proportion of those in age a that has condition G does not depend on population structure at all, but only on future mortality rates and the TTD pattern of G , $g(y)$.

A function such as $g(y)$ would have implications for the interpretation of period age patterns of morbidity, and by extension, HLE. If a function such as $g(y)$ holds, it is tautologically true that the measurement of HLE in completed cohorts (or stationary populations) will be identical whether calculated on the basis of $g^*(a)$ or the underlying $g(y)$ pattern. Distortions only arise in the interpretation of period HLE under changing mortality, or with period HLE comparisons between populations with different mortality.

Since morbidity prevalence in this scenario is partly a function of mortality, the age patterns of morbidity for populations with different mortality levels or patterns cannot be compared without additional information. Under these circumstances, it is also deceptively tricky to partition period HLE differences into underlying morbidity and mortality components, because the morbidity component is (arithmetically) a function of an uncertain future mortality pattern that accompanies the apparent age pattern of morbidity. Although cohort HLE (a gold standard) is theoretically unbiased (Imai and Soneji 2007)¹, and therefore comparable, this quantity cannot be faithfully decomposed into morbidity and mortality components based on age patterns of morbidity and mortality alone if the underlying morbidity pattern is a function of time-to-death.

A toy example serves to illustrate these concepts. Figure 2 provides a schematic overview of two stationary populations. The underlying survival pattern of these two populations is based on period survival curves

¹We confirm that this remains so even if morbidity prevalence is strongly patterned by time-to-death.

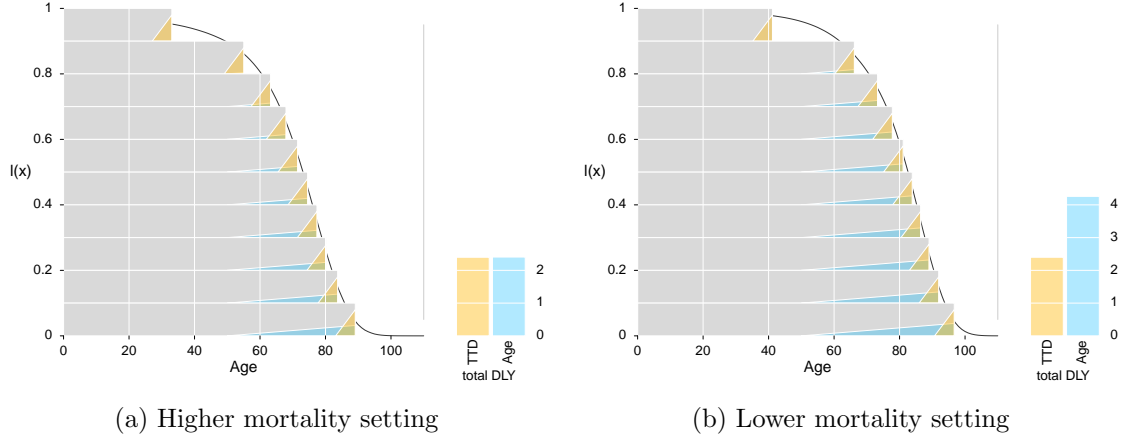


Figure 2: Schematic survival curves from higher (2a) and lower (2b) mortality populations. Each population is subjected to the same chronological age (blue) and time-to-death (yellow) morbidity prevalence patterns. The total prevalence of each sums to disability life years (DLY), drawn on the right of each survival curve. In 2a the age and time-to-death prevalences imply the same DLY. In 2b the time-to-death DLY is identical to 2a, but the age DLY is two years higher, due entirely to improved longevity.

from Japanese males in 1970 (a) and 2010 (b) (HMD), but the reader may imagine these as two hypothetical populations. Population (a) has a life expectancy of 69.3, while population (b) has a life expectancy of 79.5, slightly more than 10 years higher. For demonstration, we partition each survival curve into 10 lifespan quantiles, represented with horizontal bars. Our simple TTD prevalence, $g(y)$, is drawn with identical yellow triangles at the end of each lifespan bar. Onset begins 5 years before death and culminates with 80% prevalence. The chronological prevalence function is drawn with blue triangles, with onset at age 50 reaching a maximum prevalence of 50% at hypothetical age 111.5. Both prevalence functions are identical for populations (a) and (b).

The resulting disability life years (DLY) is shown with barplots next to each stationary population. In population (a) the age and TTD prevalence functions yield the same DLY (and HLE). In population (b) the time-to-death DLY is identical to population (a), but the age DLY is nearly twice as high. For the age prevalence function, it is correct to conclude that increased longevity leads to increases in prevalence, but for the TTD prevalence function there is no morbidity-mortality trade-off. Instead, improved longevity leads to increased proportions of life lived disability-free, albeit with no change in the absolute concentration of morbidity in the final years of life. Analyses based on the standard Sullivan method (Sullivan 1971) are only capable of predicting increased DLY when projecting from the mortal-

ity of (a) to (b). This is so for both kinds of morbidity because the TTD prevalence pattern is erased when the same condition is measured over age. The same Sullivan method can also only conclude that the morbidity of the TTD process is more compressed in (b) than in (a), even though its essential character is unchanged. Prevalence functions are in fact more nuanced than those presented here, often varying by both age and TTD, but our example provides a useful heuristic to understand a previously-undescribed source of bias in common applications of the Sullivan method.

3 Data and methods

Estimating morbidity prevalence by time-to-death is straightforward for health surveys with mortality follow-up modules. We use the RAND version P of the U.S. Health and Retirement Study (HRS),² which consists of 12 survey waves from 1992 through 2014 and an extensive set of measures for various aspects of morbidity. Time-to-death is calculated directly for deceased respondents as the difference between the date of death and interview date. Within-individual time-to-death trajectories of health measures are therefore directly available as such in the data. For a given binary health measure on an age by TTD grid, the 1-year age by 1-year TTD specific prevalence ($g_{a,y}$) is defined as the average value of that respective health measure in that age by TTD interval.

To ease interpretation, we produced age by TTD prevalence surfaces in which random variation is filtered out by applying logistic regression models with a given binary health measure as the outcome and natural splines for birth cohort, TTD, and age as covariates. We used the predictions from these regression models to produce a prevalence surface for each variable. To account for the longitudinal nature of the HRS, potentially resulting in correlated error terms, we reproduced the HRS sampling structure in a bootstrap procedure (Efron and Tibshirani 1994). Annotated R-code for the procedure and the generation of all empirical results for this analysis is available in an online repository. The bootstrap procedure is described in a technical appendix. In a sensitivity analysis, we investigated the influence on our results of the underlying assumption that morbidity prevalence is a smooth function over birth cohort, TTD, and age; both natural splines and LOESS curves over the other time measures (Riffe et al. 2016) produced highly similar results. We produced these surfaces for each health measure and by sex separately.

²The HRS is sponsored by the National Institute on Aging (grant number NIA U01AG009740) and is conducted by the University of Michigan

4 Results

4.1 How reasonable is the assumption of a time-to-death prevalence pattern?

4.2 A time-to-death morbidity pattern interacted with mortality

The age pattern of morbidity prevalence observed in the cross-section (in older ages) depends on the extent to which prevalence is principally described by age versus time-to-death and on the underlying mortality level. If the prevalence is principally a function of TTD, the specifics of its shape are also important. In Figure 3 we show the age-translations of different schematic TTD prevalence patterns when interacted with fictitious stationary populations under different mortality levels.

The first column contains three different types of disability, all of which are experienced by half of the population at the time of death, but which differ in the timing of onset prior to death and in the steepness of the curve with the approach to death. The first type of disability is virtually nonexistent 5 years prior to death, but then increases very rapidly as death approaches. The middle variant of disability is rare 15 years before death, but increases to about 20 percent of the population 5 years before death and rises sharply thereafter. The bottom figure depicts a disabling process that although still strictly determined by time-to-death, is common and accumulates very slowly starting from about 50 years before death.

The second column translates the time-to-death disability prevalence curves into the “apparent” chronological age prevalence of disability for different mortality levels depicted by the death density curves above. These mortality levels roughly correspond to USA males in 2002 $e_{60} = 20.0$ years, Canadian females in 2004 $e_{60} = 25.0$ years, and projected Japanese females a decade or so from now $e_{60} = 30.0$ years (latest observed level in 2012 was $e_{60} = 28.3$ years (HMD 2015)). In all cases increasing remaining life expectancy results in decreasing age-specific disability prevalence by chronological age. With steeply increasing disability prior to death (first row), the differences in disability prevalence are largest above age 80, where the bulk of mortality occurs, while with more gently increasing disability (second and third rows) the greatest differences in disability age-prevalence curves appear at younger ages.

These differences induced by mortality alone are not trivial. In the middle variant, which closely resembles the TTD prevalence of disability in bathing, a 10-year increase in e_{60} results in a 50 percent drop in disability prevalence at age 80 from around 20 to 10 percent. Meanwhile, the age at which a quarter of the population were considered disabled in this scenario differed by about 5 years with a 5-year improvement in e_{60} from 20 to 25.

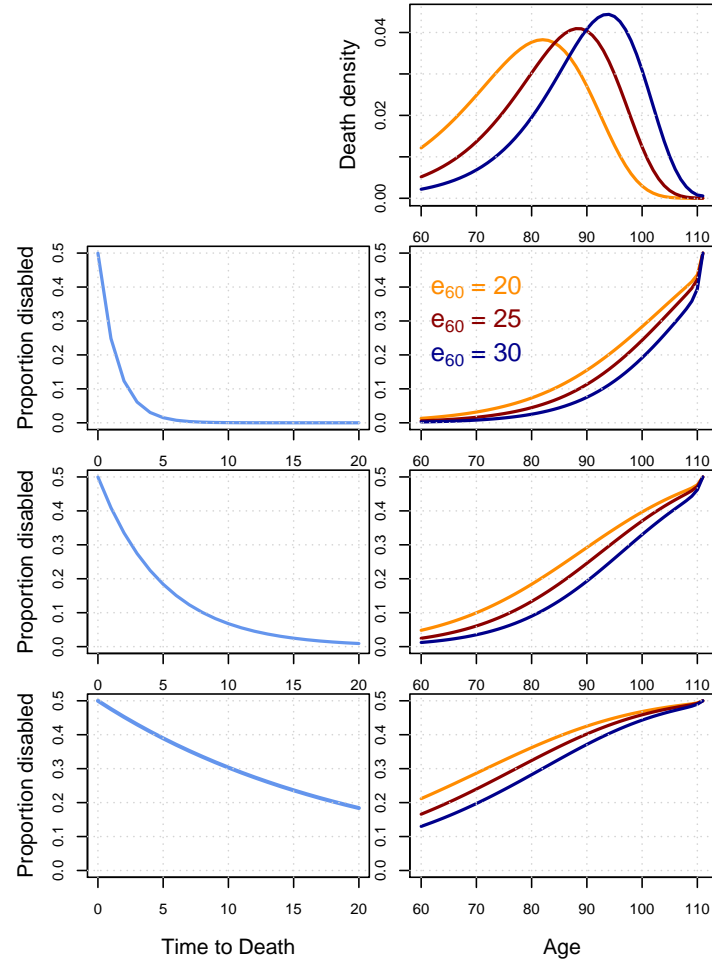


Figure 3: The age pattern of morbidity prevalence in ages 60+ derived from interacting different TTD prevalence patterns (left column) with fictitious stationary populations subject to the color-coded death distributions depicted in the top right figure.

4.3 Estimating bounds for the impact of mortality differences on estimates of disabled life years

The Sullivan method is the most commonly used method to partition life expectancy into estimates of the total life years lived in a state of good health (HLY) or disability (DLY). Its popularity owes to its minimal data requirements. Only current age-specific disability prevalence rates are needed in addition to a life table or age-specific mortality rates. Specifically, the number of person-years with disability ${}_n\pi_x \cdot {}_nL_x$ in age-group x to $x + n$ are the product of the person-years lived from the life table ${}_nL_x$ and the proportion disabled ${}_n\pi_x$. The total DLY is the sum of ${}_n\pi_x \cdot {}_nL_x$ over all age groups (Sullivan 1971).

It has long been recognized that the Sullivan method does not produce a pure synthetic measure of health expectancy, since it combines flow data of current mortality incidence with stock data of morbidity prevalence. While the mortality flow responds immediately to period change, for instance from medical innovations, the morbidity stock is slower to change because it reflects past cohort experiences with disability incidence and recovery (Mathers and Robine 1997, Barendregt et al. 1994). Moreover, as was illustrated in Figure 3, the prevalence at any age may not only depend (mathematically)) on past transitions into and out of disability but also on future deaths.

Nevertheless, comparing populations on the basis of life years lived in a state of good health (HLE) or disability (DLY) is standard practice in population health. The difference in either metric, either a within-population difference over two time periods or a between-population difference in the same time period, is often decomposed into mortality and morbidity components on the basis of differences in survivorship and morbidity age-prevalence respectively (Nusselder and Looman 2004, Andreev et al. 2002). According to Nusselder and Looman (2004), the corresponding mortality and disability effects at each age group for a within-population decomposition are:

$${}_nMOR_x = \left[\frac{{}_n\pi_{x(t)} + {}_n\pi_{x(t+y)}}{2} \right] \cdot \Delta {}_nL_x \quad (4)$$

$${}_nDIS_x = \left[\frac{{}_nL_{x(t)} + {}_nL_{x(t+y)}}{2} \right] \cdot \Delta {}_n\pi_x \quad , \quad (5)$$

where Δ refers to the change in the variable from time t to $t + y$, ${}_n\pi_x$ is prevalence in the discrete age group $[x, x + n)$, and L_x is life table exposure. Discrete prevalence ${}_n\pi_x$ is essentially our $g^*(a)$ from equation (3). The sum of the two components over all ages is equal to the total change in DLY. These formulas are essentially an application of standard Kitagawa (1955) decomposition.

Although it is true that these two components arise from changes in survivorship and morbidity prevalence respectively, difficulties arise in the

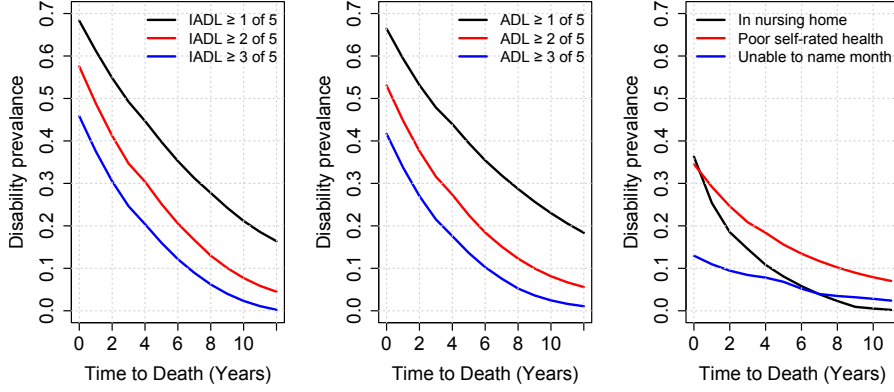


Figure 4: The disability prevalence by time to death for various disability types, based on U.S. HRS data (females).

interpretation of the components as pure mortality and disability effects. If the time-to-death profile of disability prevalence does not change over different mortality regimes, inducing mortality decline alone will result in declines in the age-pattern of prevalence component, as we demonstrated. This method of decomposition will therefore still attribute a portion of the change in DLY to morbidity, and we consider this to be a source of bias.

To get a sense of the upper magnitude of this bias, we test how the disability component of this decomposition method would change when a fixed time-to-death prevalence of morbidity is applied to different mortality regimes using empirical data. Morbidity data come from the US Health and Retirement Study while mortality and exposure data come from the Human Mortality Database (HMD 2015).

Specifically, we consider the age prevalence of difficulties in carrying out at least 1, 2, or 3 (out of 5) functional Activities of Daily Living (ADL), difficulties in carrying out at least 1, 2, or 3 (out of 5) instrumental Activities of Daily Living (IADL), living in a nursing home, having poor self-rated health, and being unable to name the month of the year. We estimate these time-to-death prevalences for quinquennial cohorts from 1905 to 1930, separately for males and females. We calculate the time-to-death $g(y)$ profiles of disability prevalence for all extinct lifespans above age 65 for each cohort. Estimation is done via a loess smooth over age, time-to-death, and birth cohorts. We then average the time-to-death profiles of disability over lifespans and over the 6 cohorts. The time-to-death prevalence of morbidity for each disability type is shown for females in Figure 4.

We calculate the apparent period chronological age prevalence of morbidity, $g^*(a)$, for all medium to large populations of the Human Mortality

Database, had they experienced the US time-to-death profile of morbidity. Eastern European countries are excluded from this exercise due to widely varying age patterns of mortality, particularly in the years surrounding political transition. To calculate the age-pattern of morbidity, we assume the survival pattern of each lifetable to be a stationary population and apply a discretization of equation (3).

We then make pair-wise comparisons of DLY between each population in the same year for the years 1980, 1990, and 2000. For within-country comparisons, we compare each population in 10-year jumps, for all years starting from 1950. Altogether this leads to 187 within-population comparisons and 1785 between-population comparisons for each sex. Finally we decompose the change or difference in DLY between the population pairs into mortality and morbidity components using the Nusselder and Looman (2004) method described above and in Equations (4) and (5). The true value of the change in DLY and the true value of the disability component are both zero by design. Thus, the estimated disability component from this decomposition gives a gauge of bias.

We compare the association between the change in the disability component (Equation 5) and the increase or difference in remaining life expectancy at age 60 for each population pair in Figure 5. By doing so we aim to provide a rough empirically-based estimate of the upper bound of the change in the disability component that is attributable to the different underlying mortality levels of any two populations being compared. Thus if female e_{60} increases in a country by 5 years, up to about 1 year of the reduction in DLY that is attributed to the disability component could be solely arising from the decrease in mortality, in the case where disability is measured as having difficulty in at least three ADLs. Departure from this upper bound in real life depends on the degree to which the disability prevalence changes on a time-to-death versus chronological age axis, the extent to which the US average time-to-death prevalence is representative, and the departure from the stationary population assumption.

Overall, the relationship between the change in disability component and the increase in e_{60} is strikingly linear although the slopes differ for males and females and by disability type. To some extent this is because the final level of disability prevalence (i.e. in the final year of life) differs by disability type. In Appendix Figure 6 we standardize for the maximum disability prevalence to give a clearer comparison between disability types. Even after standardization, the change in the disability component is greater with larger e_{60} differences for disability types with more gradually changing prevalence levels by time-to-death and for females who likewise have more gradually changing prevalence levels by time-to-death. This is because steep time-to-death profiles of disability concentrate disability prevalence near the death distribution, whereas gradually changing (but still entirely time-to-death) disability profiles spread out disability prevalence over a wider range

of ages before each age at death.

If projecting morbidity prevalence, one can also refer to these results as a heuristic on the bias inherent in assuming a fixed age-pattern of morbidity.

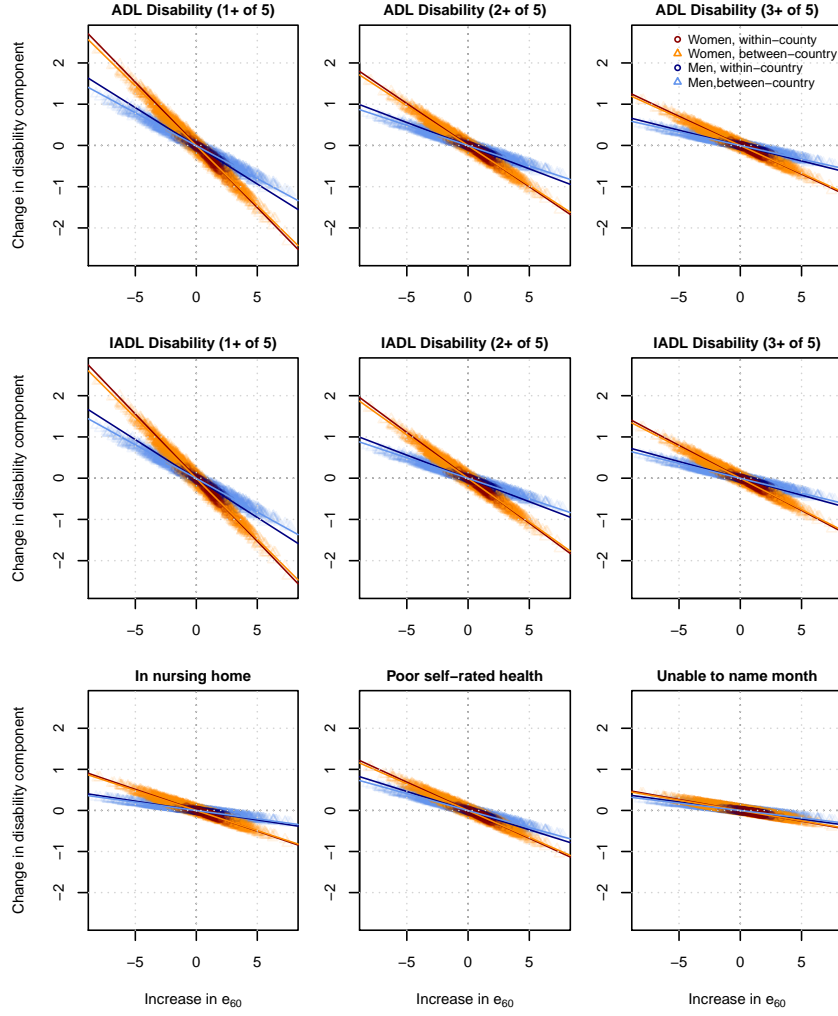


Figure 5: Results of the hypothetical decomposition exercise. The size of the morbidity component using a standard decomposition method is plotted against the difference in remaining life expectancy at age 60 (e_{60}) in each pair of populations. Linear trend lines are also provided for each sex and decomposition type.

5 Discussion

Healthy life expectancy is a popular tool for analyzing population health, forming a snapshot of the hypothetical life years lived in good or poor health. This information is well-summarized in both the period and cohort perspectives, irrespective of whether morbidity prevalence is a function of chronological age or time until death. However, difficulties arise in the interpretation of period differences in this quantity. The chronological age pattern of disability can increase or decrease solely as a function of mortality change, even when the underlying morbidity function is held constant. Thus, for instance, observed widening ratios in the age profiles of disability prevalence between subgroups (Crimmins and Saito 2001) cannot be attributed to changes in the disabling process without taking into account changing mortality profiles. This observation calls into question the practice of forecasting observed age-specific rates of decline in disability (Manton et al. 2006, Khaw 1999), and especially the more common practice of holding age-patterns of disability fixed in morbidity projections. Health economists refer to a similar ‘red herring’ argument, namely that medical costs are more closely associated with time-to-death than with chronological age. As a result, health care cost projections based on a chronological age rather than time-to-death pattern of expenditure are artificially inflated (Zweifel et al. 1999, Geue et al. 2014).

We argue that the better way to measure changes in health or disability is from a cohort perspective (Manton and Land 2000, Manton et al. 2008, Christensen et al. 2013). Manton and Land (2000), for instance, found large differences between period and cohort estimates of active life expectancy (ALE). ALE at ages 65 and 85 was between 1.6 and 2.6 times larger in the cohort perspective than for similar period estimates, and the expected years of life disabled were smaller in the cohort perspective. Additionally, they uncovered larger differences between the cohort and period perspectives for men than women, which they attribute to differences in disability transition rates between the sexes. The cohort-period direction of difference in ALE is consistent with a fixed TTD pattern combined with improving mortality, as of our toy example in Section 2. We hypothesize that some of these larger differences might also be attributable to larger mortality reduction among men. Further, while cohort HLE estimates are unproblematic as an index, decompositions of HLE differences between cohorts (with different mortality) into age-patterned morbidity and mortality components would still be biased. This is because the age pattern of morbidity is itself decomposable into morbidity and mortality components.

As our example illustrates, differences in underlying mortality can lead to differences in the age profile of disability. That mortality levels and disability prevalence are related is perhaps not surprising. Several studies have looked at the macro relationship between overall mortality levels and sex differences in HLE. At higher levels of life expectancy, female advantage

in healthy life expectancy diminishes, or even reverses into male advantage (Van Oyen et al. 2013). As well, the larger the proportional female advantage in longevity, the larger the female excess in the proportion of life in poor health (Luy and Minagawa 2014). Additionally, although the association between the severity of chronic conditions and poor health was found to be similar for men and women, morbidity prevalence rates are generally higher among women (Case and Paxson 2005). It would be worthwhile to investigate whether there might not only be differences in the composition of chronic conditions between the sexes, but whether the underlying morbidity pattern itself might differ between the sexes in its joint age and time-to-death pattern (Riffe et al. 2016).

Empirically, not all end-of-life health conditions are exclusive functions of time to death, but morbidity often varies as a function of both time measures, and expressing morbidity prevalence as a function of both age and time-to-death, $g(a, y)$, can increase precision (Wolf et al. 2015, Riffe et al. 2016). There is great variety in the temporal variation of the prevalence of late-life health conditions. There is also great variety in individual trajectories with the approach to death (Lunney et al. 2003).

That morbidity prevalence may for certain health conditions be patterned by time to death does not imply that morbidity incidence is necessarily a function of time to death. First, a sequence of health states wherein mortality risk increases in each successive state could produce a time-to-death prevalence pattern.³ Second, it is also plausible that some morbidity conditions are linked to a more general process of dying, thereby linking morbidity to a process that ends with death and consequently producing a time-to-death prevalence pattern. For example, certain conditions may manifest themselves that are not (primary) causes of the impending death but consequences of nearness to death caused by some other (primary) factor. Neither of these explanations conflicts with the reality that causes must precede effects, and that therefore death cannot cause the morbidity that precedes it (Lynch 2015).

Increasingly, data on mortality incidence and recovery are available from multiple waves of survey panels such as the HRS, allowing researchers to calculate healthy life expectancy using sophisticated multistate models. Unfortunately such data is not available in all countries, or if it is, time trends are limited to the recent past. We are not arguing that a TTD approach should replace multistate models when such data is available. Our aim is rather to expose the implications of comparing healthy life expectancy from age-structured prevalence-based models with different underlying mortality regimes.

³Such a minimal state model with no age patterns for state transitions or mortality risk within states can produce both an overall age pattern of mortality and a time-to-death pattern of morbidity prevalence.

To model prevalence as a function of time-to-death requires no surreal understanding of how things work, but is rather a modelling choice (Wolf et al. 2015). When modeling for descriptive or exploratory purposes (as we have done to produce e.g. Figure 3, and possibly for projections, one can safely use TTD as a predictive variable. However, using time-to-death in models intended for causal interpretation is more perilous; Some argue that TTD may function as a proxy for unobserved variables such as biomarkers for impending mortality (Wolf et al. 2015). Others argue that including this variable as a proxy in models will introduce omitted variable bias (Lynch 2015). Regardless, even in the descriptive case, the reader may understand a simple TTD morbidity prevalence pattern as generally hypothetical, since the current evidence base is limited in scope. Much more empirical work is needed in order to determine whether modeling morbidity prevalence as a function of time to death is more widely applicable to other health conditions, younger ages, more recent birth cohorts, and other populations in different stages of epidemiological transition. Further, even joint age-TTD prevalence patterns, such as that of Figure ?? may change over time, and are not fixed as in our examples. Nevertheless, the distortions demonstrated are likely to arise in everyday practice when comparing health trends over age between populations or over time, since many health conditions appear to show strong time-to-death components. To further test the bias we hypothesize and to design more robust synthetic measures of life lived in good or poor health, more effort should be given to data collection, mortality followup, and measurement practices.

6 Conclusions

We describe a likely source of bias in comparisons of healthy life expectancy or life years lived in poor health. This bias derives from a failure of age structure to capture the principal pattern of time variation in morbidity. In short, age standardization does not guarantee comparability of morbidity levels. Comparisons of period healthy life expectancy calculated from marginal age patterns of morbidity prevalence cannot be directly partitioned into morbidity and mortality effects if the joint age by time-to-death morbidity prevalence pattern varies at all by time-to-death.

In Section 2 we show how simplistic time-to-death and age-patterns of morbidity interact with lifetable survivorship to yield opposite conclusions on morbidity compression under improved mortality. A fixed increasing age pattern of morbidity predicts a greater burden of morbidity under improved survival, but a fixed time-to-death pattern of morbidity predicts no change in the average time spent in poor health, and therefore morbidity compression. In Section 4.2 we describe in greater detail how a range of schematic time-to-death morbidity patterns interact with different stationary mortality levels

to produce age patterns of morbidity. In Section 4.1 we provide an example of a variable whose joint prevalence pattern over age and time-to-death is better summarized by a single time-to-death pattern than by an age pattern. We demonstrate in Section 4.3 that the degree of bias in partitioning effects depends on the shape and level of time-to-death variation.

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A Decomposition after standardization

In this appendix we standardize the decomposition exercise of section 4.3 for the maximum disability prevalence to give a clearer comparison between disability types.

B Description of bootstrapping technique

The Health and Retirement Survey (HRS) data is a longitudinal dataset, and hence observations are not independent but correlated within individuals. To account for the influence of the correlated error structure on the estimation of Time to Death (TTD) surfaces, we applied a bootstrap (Efron Tibshirani, 1994) procedure in which we approximated the sampling structure of the HRS. In our application, bootstrap sizes of 999 provided the best approximation of the distribution of our estimator; if our method is adapted to other datasets, we recommend the use of bootstrap diagnostics applied to

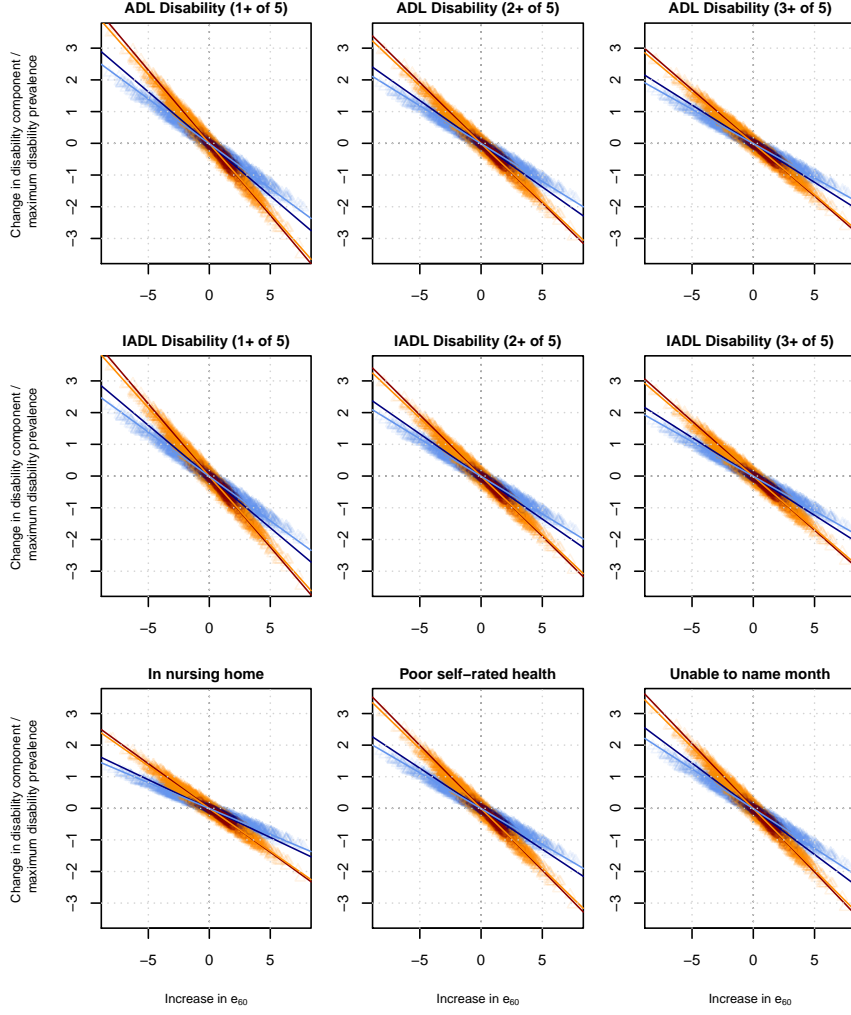


Figure 6: Results of the hypothetical decomposition exercise standardized to the maximum disability prevalence for each disability type. The size of the morbidity component using a standard decomposition method is plotted against the difference in remaining life expectancy at age 60 (e_{60}) in each pair of populations. Linear trend lines are also provided for each sex and decomposition type.

various locations on the TTD surface (with special attention to the “edges” of the TTD surface) to determine bootstrap size.

B.1 Bootstrap steps

An R-code application of our method is available as Supplemental Material. The method, and therefore the code, works through the following 9 steps:

1. Choose an outcome variable (i.e. BMI, hospital stay duration, etc.).
2. Create an empty dataset with time dimensions of interest (in our application: birth year, time to death, and chronological age, but this could be more dimensions if desired), which we call the Lexis space
3. Resample individuals with replacement.
4. Create natural cubic splines for the dimensions of the Lexis space.
5. Fit regression model of a type appropriate to the outcome variable to the resampled data with the natural cubic splines (step 4) as covariates.
6. Use the regression fit to predict the Lexis space and save these predictions.
7. Repeat steps 3 to 6 until desired bootstrap size is achieved.
8. Take the median or mean over the bootstrap iterations for each Lexis cell to create a Lexis space with median or mean estimates.
 - Optional: take $\alpha/2$ and $1 - \alpha/2$ quantiles over the bootstrap iterations for each Lexis cell in order to create Lexis spaces containing the lower and upper $1 - \alpha$ confidence bounds of the estimates.
9. Take a cross-section of the Lexis space with median or mean predictions in order to plot the Lexis surface.

The rest of this appendix sheds more light on these steps. See also the supplementary R-code for the implementation of the method.

B.2 Resampling with replacement and probability weights

The HRS samples individuals each year. Sampled individuals stay in the sample until either death or drop-out. To account for the changing population structure of the United States, probability weights for an individual can differ between observations (i.e. they change over time).

In order to account for the longitudinal nature of the data, we resample individuals following the sampling structure of the HRS. A (re)sampled individual contributes all of his or her observations to that respective (re)sample.

This type of resampling is also known as blockwise bootstrapping in a time series context (Bühlmann and Künsch 1995). The probability for an individual to be sampled is equal to the HRS probability weight of the first observation of that individual (RAND variable `wtresp`). We resample individuals with replacement. The number of individuals resampled is equal to the number of individuals in the original sample.

The information contained in the probability weights of subsequent observations of individuals is also used: We create new weights termed rescale weights, which are each individuals probability weights divided by that individuals respective sampling probability (and hence, the rescale weight on the first observation for each individual is always 1). These rescaled weights are used in the regression procedure (see section B.3), where the sampling probability represents the representativeness of the individual in their respective demographic category, the rescale weights capture to what extent this weight changes as a consequence of changes in population structure, and therefore captures to what extent an individuals representativeness changes. In our application of the method, the vast majority of resampling weights did not exceed the 0.90 to 1.10 range, i.e. the sampling probability was never inflated or deflated by more than 10 percent.

B.3 Regression natural cubic splines

To produce smooth age by TTD surfaces we applied regression models to the outcome of interest (BMI, hospital stays, etc.). Covariates in these regression models were natural cubic splines of the dimensions age, birth year, and time to death. The cubic spline is defined as follows:

$$S(x) = \begin{cases} S_0(x) = a_0x^3 + b_0x^2 + c_0x + d_0 & t_0 \leq x \leq t_1 \\ \vdots & \\ S_{k-1}(x) = a_{k-1}x^3 + b_{k-1}x^2 + c_{k-1}x + d_{k-1} & t_{k-1} \leq x \leq t_k \end{cases}, \quad (6)$$

where x represents points on the dimension (in our application, a time dimension) of interest, and t_0 and t_k denote the two extremes (endpoints) of that part of the dimension of interest for which we wish to create this spline. Furthermore, the constraints

$$S_{k-1}(x_i) = S_k(x_i) \quad (7)$$

$$S'_{k-1}(x_i) = S'_k(x_i) \quad (8)$$

$$S''_{k-1}(x_i) = S''_k(x_i) \quad (9)$$

with $i = 1, 2, \dots, k - 1$ must be satisfied. This ensures that the segments of the spline meet at the boundaries (first constraint) in a smooth manner

(second and third constraint). The additional constraint

$$S''_{k-1}(x_i) = S''_k(x_i) \quad (10)$$

makes it into a natural cubic spline, i.e the second derivative of each polynomial is set to 0 at the endpoints of our part of the time dimension of interest.

The part of the dimension of interest that can be found between t_0 and t_k is subdivided into $k - 1$ segments, the connection of these segments are known as knots. In our application, knots for birth year were placed at the year 1902.5 to 1925.5 at 5-year intervals. Knots for time to death were placed at 0.5, 1, 2, 4, 7.5 and 10 years to death. Knots for chronological age were placed at 72.5 to 97.5 years of age at 5 year intervals. We denote the natural cubic spline for age as $A(x)$, for birth cohort as $C(x)$ and for time to death as $T(x)$.

For continuous outcomes, we used these splines in linear regression models, i.e.

$$\mathbb{E}[Y|x_a, x_c, x_t] = A(x_a) + C(x_c) + T(x_t) \quad , \quad (11)$$

where Y is the outcome variable of interest, and x_a, x_c, x_t represent (observed) points on the age, birth year and time to death axes, respectively. For binary outcomes, we used logistic regression, i.e.

$$\text{logit}\{\mathbb{E}[Y|x_a, x_c, x_t]\} = A(x_a) + C(x_c) + T(x_t) \quad . \quad (12)$$

Ordinal outcomes were dichotomized during data handling and therefore also modeled using logistic regression.

For count outcomes we used zero-inflated Poisson regression, which is a mixture model, combining logistic and Poisson regression. This type of model separates modeling the probability that an observation is an “excess” 0 (or not) from the “count” component (which is modeled if the observation is not an excess 0). The count component was therefore modeled as:

$$\ln\{\mathbb{E}[Y|x_a, x_c, x_t]\} = A(x_a) + C(x_c) + T(x_t) \quad . \quad (13)$$

The 0 or not component was modeled with logistic regression as denoted above. We used this type of model because most of our observations contained a very large number of zeros relative to observed counts and therefore appeared to come from a mixture distribution. Ordinary Poisson models can be compared with zero-inflated Poisson models using the Vuong test, but this was omitted in our application. Most count variables in our dataset were bounded, e.g. the number of nights that can be spent in the hospital is bounded by 365 (maximum number of days in a year). In those instances, we included those bounds in our Poisson model through an offset term:

$$\ln\left\{\mathbb{E}\left[\frac{Y}{\text{max bound}}|x_a, x_c, x_t\right]\right\} = A(x_a) + C(x_c) + T(x_t) \quad (14)$$

\implies

$$\ln\{\mathbb{E}[Y|x_a, x_c, x_t]\} = A(x_a) + C(x_c) + T(x_t) + \ln(\max \text{ bound}) \quad .$$

Since the functions $A(x)$, $C(x)$ and $T(x)$ are cubic natural splines, they represent a vector of coefficients corresponding to $S(x)$ as defined earlier. Specification of our models in this manner with natural splines was inspired by work of Carstensen (2007).

Because the three time dimensions included in our models are not linearly dependent, additional constraints on one dimension (such as required in APC modeling, see e.g. Clayton and Schifflers (1987)) were not required. The application of splines along different time dimensions, or such splines in addition to splines of the already time dimensions, did not improve our smoothed TTD surfaces. Similarly, a higher number of knots had only minor effects on the smoothed TTD surfaces. Finally, we also compared our natural cubic splines to other flexible curve fitting methods (such as LOESS and GAM), and found natural cubic splines to provide the best fit overall to the empirical data, as expressed in distance to squared and absolute residuals and propensity for systematic under- or over-estimation of the data along relevant time dimensions (including edge effects).

Splines are known to function less well near the endpoints (t_0 and t_k); to determine if this was present in our data, we investigated the residuals of regression models. We found only weak evidence for the existence of edge effects, and for most outcomes edge effects were absent. For those outcomes where edge effects were potentially present, as a sensitivity analysis, in our resampling procedure we purposely oversampled observations near the edges and estimated TTD surfaces under this regime. We did not find this to substantially affect our conclusions.

B.4 Predicting the age by TTD surface

To produce age by TTD surfaces, we created a dataset with birth year, time to death, and chronological age as columns. The minimum and maximum values of birth year, time to death and chronological age were equal to those found in the empirical data, and intermediate values were spaced at 1-year intervals. This dataset therefore represents an empty Lexis space. Using the fitted regression models, we then predicted outcomes for this Lexis space; i.e. we created a column with an expected value for each birth year, time to death and chronological age combination (henceforth: Lexis cell) in the Lexis space. Each iteration of the bootstrap added a new column of such predictions. To produce the final Lexis surface, we then plot the median prediction for each Lexis cell combination. Since three dimensions are more difficult to visualize due to overplotting, in our paper we chose the predictions of the 1915-1919 birth cohort and plotted along the time to death and age dimensions.

We found that, overall, predictions for each Lexis cell were normally distributed, and TTD surfaces based on Lexis cell averages were nearly identical to those for TTD surfaces based on median values. However, since some cells had more skewed predictions, namely when the outcome was binomial and predictions were close to 0 or 1 (the edge of the parameter space), we chose the median as the appropriate measure of centrality. Additionally, the median, being the 50% quantile, is also consistent with 95% quantile bootstrap confidence intervals. As their name implies, 95% quantile bootstrap confidence intervals can be calculated by taking the 2.5% and 97.5% quantile for each Lexis cell and plotting those surfaces. The value of the median will, by definition, not exceed the 2.5% and 97.5% quantiles, whereas in theory the mean could exceed these confidence bounds in highly skewed distributions.