Association between telomere length in blood and mortality in people aged 60 years or older

Richard M Cawthon, Ken R Smith, Elizabeth O'Brien, Anna Skvatchenko, Richard A Kerber

During normal ageing, the gradual loss of telomeric DNA in dividing somatic cells can contribute to replicative senescence, apoptosis, or neoplastic transformation. In the genetic disorder dyskeratosis congenita, telomere shortening is accelerated, and patients have premature onset of many age-related diseases and early death. We aimed to assess an association between telomere length and mortality in 143 normal unrelated individuals over the age of 60 years. Those with shorter telomeres in blood DNA had poorer survival, attributable in part to a 3.18-fold higher mortality rate from heart disease (95% CI 1.36–7.45, p = 0.0079), and an 8.54-fold higher mortality rate from infectious disease (1.52–47.9, p = 0.05). These results lend support to the hypothesis that telomere shortening in human beings contributes to mortality in many age-related diseases.

Lancet 2003; 361: 393–95

Telomere length declines with age in all mitotic tissues apart from germline tissue, in which it is fully maintained by the enzyme telomerase. Patients with autosomal dominant dyskeratosis congenita carry mutations in the gene encoding the RNA component of telomerase,1 have short telomeres and other indications of accelerated ageing, and die prematurely, usually from severe infections secondary to bone-marrow failure.

The shortest average telomere lengths in blood DNA from normal elderly individuals overlap with the highest average lengths in blood from patients with dyskeratosis congenita.1 Furthermore, telomere lengths in blood accord with those in other tissues.2 This raises the question of whether shorter telomeres in blood DNA are associated with increased mortality rates from multiple age-related diseases. We aimed to assess telomere length versus overall survival and cause-specific mortality in a sample of individuals not selected for the presence or absence of any specific disease or clinical disorder.

We studied unrelated Utah residents aged 60–97 years who donated blood between 1982 and 1986,3 and for whom follow-up survival data were available. We obtained birth and death dates from the Utah Population Database (UPDB), and from the Social Security death index. Survival analysis by cause of death (from Utah death certificates, coded to the International Classification of Diseases, 9th and 10th revisions) was restricted to individuals with UPDB identification numbers. The University of Utah's Institutional Review Board approved the study.

The relative ratio of telomere repeat copy number to single copy gene copy number (T/S ratio) in experimental samples compared with a reference DNA sample was established by quantitative PCR of original blood DNA. In another set of DNA samples, we measured T/S ratios relative to the same reference DNA sample, and mean terminal restriction fragment (TRF) lengths were ascertained.4 The slope of the plot of mean TRF length versus T/S for these samples served as the conversion factor for calculation of approximate telomere lengths in bp for each T/S ratio in this survival study.

We used Cox’s proportional-hazards regression models to test whether differences in telomere length (analysed as a continuous variable) were associated with differences in survival. In all other analyses, telomere length was treated as a dichotomous trait (shorter vs longer), using all available samples in each comparison (ie, bottom half of the telomere length distribution vs top half, and bottom 25% vs top 75%).

Because older individuals tend to have shorter telomeres than those who are younger, use of one telomere length distribution for the entire sample of people would result in a higher proportion of older than younger participants being scored as shorter for telomere length. To achieve more balanced proportions of participants with shorter versus longer telomere lengths at every age, we stratified the sample into six categories of age at blood draw (60–64 years, 65–69 years, 70–74 years, 75–79 years, 80–84 years, and 85 years and older), and telomere length distribution was determined independently within each category. Individuals in the bottom half for telomere length in each age group were pooled together, and their survival was compared with that of the pooled top half individuals. Similarly, individuals in the bottom quartile for telomere length in each age group were pooled, and their survival was compared with that of the pooled top 75% in individuals. No significant difference in mean

### All-cause mortality

<table>
<thead>
<tr>
<th>Number of individuals/number of deceased</th>
<th>Mortality rate ratio* (95% CI)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Both sexes 143/101</td>
<td>1.86 (1.22–2.83)</td>
<td>0.004</td>
</tr>
<tr>
<td>Women 71/64</td>
<td>2.16 (1.07–4.39)</td>
<td>0.033</td>
</tr>
<tr>
<td>Men 72/55</td>
<td>1.94 (1.01–3.74)</td>
<td>0.047</td>
</tr>
</tbody>
</table>

### Cause-specific mortality:

<table>
<thead>
<tr>
<th>Cause</th>
<th>Number of individuals/number of deceased</th>
<th>Mortality rate ratio* (95% CI)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart †</td>
<td>124/30</td>
<td>3.18 (1.36–7.45)</td>
<td>0.008</td>
</tr>
<tr>
<td>Cerebrovascular §</td>
<td>124/15</td>
<td>1.35 (0.36–5.13)</td>
<td>0.660</td>
</tr>
<tr>
<td>Cancer ‡</td>
<td>124/12</td>
<td>1.43 (0.34–6.03)</td>
<td>0.625</td>
</tr>
<tr>
<td>Infectious §</td>
<td>124/8</td>
<td>8.54 (1.52–47.9)</td>
<td>0.015</td>
</tr>
<tr>
<td>Other known ‡</td>
<td>124/16</td>
<td>2.15 (0.71–6.50)</td>
<td>0.174</td>
</tr>
<tr>
<td>All known causes apart from heart and infectious §</td>
<td>124/43</td>
<td>1.70 (0.82–3.53)</td>
<td>0.156</td>
</tr>
</tbody>
</table>

*Ratio of the death rate for participants with shorter telomeres to the death rate for those with longer telomeres. †Ratio is for individuals from the bottom half of the telomere length distribution versus those from the top half of the distribution. §Only assessed in individuals with UPDB identification numbers. ‡Ratio is for individuals from the bottom 25% of the telomere length distribution versus those from the top 75% of the distribution.

Mortality rate ratios associated with short versus long telomeres in whole-blood DNA.

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THE LANCET • Vol 361 • February 1, 2003 • www.thelancet.com

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age at blood draw between compared groups was noted (bottom vs top half p=0·894, bottom 25% vs top 75% p=0·469).

Survival was assessed beginning with the time at blood draw, except as noted. We used Cox models to control for variation in mortality rate due to age differences, both between the age at blood draw categories, and within each age group.

We studied 143 individuals (age 60–97 years). 101 deaths were known by mid-2002. For the remaining 42 participants, we could establish a date at which they were last known to be alive, post-blood draw.

Telomere length ranged from 1930 to 4310 bp. Every 1-year increase in age at blood draw was associated with a 0·0048 decrease in the relative T/S ratio (95% CI 0·00137–0·00823, p=0·0074), corresponding to about 14 bp of telomere sequence lost per year. Women and men did not differ significantly in the rate of telomere shortening estimated from these cross-sectional data (p=0·645). Women’s telomeres were 3·5% longer than were men’s after adjustment for age (p=0·157).

Individuals with shorter telomeres had a mortality rate nearly twice that of those with longer telomeres (table). The loss in median survival associated with shorter telomeres was 4·8 years for women, and 4·0 years for men (averaged across all categories of age at blood draw; figure). Telomere length was a significant predictor of mortality for people aged 60–74 years (p=0·021), and a moderate predictor for those aged 75 years and older (p=0·086) (table, figure). Telomere length, when analysed as a continuous variable, was inversely associated with the age-adjusted mortality rate (r=−1·87, 95% CI −3·35 to −0·392, p=0·013).

Excess mortality risks associated with short versus long telomeres did not vary by sex (p=0·878), age at blood draw (p=0·946), or time since blood draw (p=0·851). The excess mortality rates of those in the bottom half of the telomere length distribution remained significant even when only individuals surviving at least 5 years after the blood draw (n=112) were included in the analysis (p=0·0063).

Individuals from the bottom half of the telomere length distribution had a heart disease mortality rate that was over three times that of those from the top half of the distribution (table). This increased risk of dying from heart disease remained significant even when the analysis was restricted to individuals who survived at least 5 years after the blood draw (number of heart disease deaths 21; mortality rate ratio 4·87 [95% CI 1·59–14·9], p=0·006). Mortality rates for cerebro-
vascular disease and cancer were not significantly higher in individuals with shorter telomeres (Table).

The mortality rate from infectious disease was eight times higher for individuals in the bottom 25% of the telomere length distribution than for individuals in the top 75% (p=0.015). Of the deaths from infectious disease, the shortest time between blood draw and death was 1.5 years.

Our results lend support to the hypothesis that telomere shortening contributes to the rise in mortality rates from multiple diseases typically seen with ageing. Alternatively, telomere shortening might not affect mortality, but might be controlled by (and so serve as a useful indicator of) progression of a process of senescence that raises mortality rates by other mechanisms.

Short telomeres in blood could indicate the presence of an age-related disease that has, perhaps, triggered a shift in the proportions of white-blood-cell subsets, thereby reducing average telomere length. This scenario, however, predicts that the strength of the association between telomeres with higher mortality should decline as the number of years that individuals needed to survive after blood draw increases, and we did not see such a decline.

Telomere length in age-matched people can be affected by many factors, including telomerase activity, rate of cell division, and amounts of oxidative stress, which in turn might be determined by genetic and environmental factors.

Effectiveness of vaccination for Haemophilus influenzae type b

David Garner, Vivienne Weston

Several cases of invasive Haemophilus influenzae type b (Hib) infection have been identified in children previously vaccinated against this disease. We have reviewed all cases of Hib in Nottingham, UK, since the conjugate vaccine was introduced in the primary course of childhood immunisations in 1992. Our results show a worrying increase in the frequency of Hib in Nottingham, which is much the same as the national trends. We believe that the medical profession should be aware of the return of Hib, and that we now need to review the need for a booster vaccination for what was thought to be a disease of the past.

Lancet 2003; 361: 395–96

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Acute epiglottitis used to frequently strike fear into the hearts of general practitioners and paediatricians in the UK. On Oct 1, 1992, Haemophilus influenzae type b (Hib) conjugate vaccine became part of the primary course of childhood immunisations at 2, 3, and 4 months of age and during 1993, all children younger than 4 years were offered “catch-up” vaccinations with a single dose of vaccine. This programme greatly reduced the frequency of acute epiglottitis and invasive Hib disease throughout the UK. However, over the past 3 years the frequency of invasive Hib in the UK has more than doubled and this disorder should still be considered a cause of acute laryngitis.

In Nottingham, from October to December, 2001, we recorded four cases of invasive Hib disease in children younger than 4 years of age who had previously been vaccinated. Thus, the microbiology consultant alerted the consultant in Paediatric Intensive Care to the presence of Hib within vaccinated children—a timely intervention. The following week a child, previously vaccinated against Hib, was referred to Paediatric Intensive Care Unit from Accident and Emergency with severe group. The consultant in Paediatric Intensive Care was immediately alerted to the presence of Hib epiglottitis and proceeded to manage this child appropriately. The diagnosis was later confirmed by culture of Haemophilus influenzae type b from blood. Without the warning from the microbiology department, instigation of appropriate antimicrobial treatment might have been unnecessarily delayed in this child.

There were 238 cases of invasive Hib per 100 000 children in England and Wales in 1991, the year before the childhood immunisation programme was introduced; by 1996, prevalence had dropped to 0·92 per 100 000; and in

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Cases of Invasive Haemophilus influenzae type B (Hib) in Nottingham Public Health Laboratory 1992–2001