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# Genome-wide meta-analysis points to CTC1 and ZNF676 as genes regulating telomere homeostasis in humans

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Leukocyte telomere length (LTL) is associated with a number of common age-related diseases and is a heritable trait. Previous genome-wide association studies (GWASs) identified two loci on chromosomes 3q26.2 (*TERC*) and 10q24.33 (*OBFC1*) that are associated with the inter-individual LTL variation. We performed a meta-analysis of 9190 individuals from six independent GWAS and validated our findings in 2226 individuals from four additional studies. We confirmed previously reported associations with *OBFC1* (rs9419958  $P = 9.1 \times 10^{-11}$ ) and with the telomerase RNA component *TERC* (rs1317082,  $P = 1.1 \times 10^{-8}$ ). We also identified two novel genomic regions associated with LTL variation that map near a conserved telomere maintenance complex component 1 (*CTC1*; rs3027234,  $P = 3.6 \times 10^{-8}$ ) on chromosome17p13.1 and zinc finger protein 676 (*ZNF676*; rs412658,  $P = 3.3 \times 10^{-8}$ ) on 19p12. The minor allele of rs3027234 was associated with both shorter LTL and lower expression of *CTC1*. Our findings are consistent with the recent observations that point mutations in *CTC1* cause short telomeres in both *Arabidopsis* and humans affected by a rare Mendelian syndrome. Overall, our results provide novel insights into the genetic architecture of interindividual LTL variation in the general population.

## INTRODUCTION

Rare mutations in telomere maintenance genes—resulting in critically short leukocyte telomere length (LTL)—cause monogenic diseases with a spectrum of clinical manifestations. These include diseases such as dyskeratosis congenita, idiopathic pulmonary fibrosis and liver cirrhosis, the common manifestation of which is often aplastic anemia (1-4). In contrast, little is known about the consequences of common genetic variants that contribute to the considerable inter-individual variation in LTL in the general population (5). Patients with clinical manifestations of atherosclerosis and those at risk for this aging-related disease often display short LTL (6-9). Moreover, recent studies have reported LTL to be inversely related to survival in the elderly (10-12). Therefore, genetic variants that affect LTL dynamics (LTL at birth and age-dependent attrition) might play a role in the aging of the cardiovascular system and in longevity.

To date, genome-wide association studies (GWASs) have identified two main genetic variants associated with LTL variation (rs3772190 at 3q26.2 and rs4387287 at 10q24.33) (13–15). However, these findings explained only a very small part of the genetic variation in LTL (<1 and 2.26% for rs3772190 and rs4387287, respectively) (13,14). As has been demonstrated for other complex traits, the larger the sample size, the more likely it is to unveil additional LTL-regulating genes. This would bolster understanding of the role of LTL dynamics in aging, aging related-disease and longevity.

Here, we report the results of a GWAS meta-analysis, performed on a total of 9190 individuals from six independent observational studies. We have identified two novel LTL loci (on chromosome 17p13.1 and 19p12), highlighted a number of genes for which verification in larger studies is warranted and reaffirmed that LTL in humans is fashioned by a large number of common genetic variants with small individual effects.

# **RESULTS**

We carried out a meta-analysis of GWAS of LTL in 9190 individuals of European ancestry from six collaborating studies: TwinsUK (n = 3222), the Family Heart Study (FamHS) (n = 2508), the Framingham Heart Study (FHS) (n = 1146),

the Cardiovascular Health Study (CHS) (n = 1061), Hypertension Genetic Epidemiology Network Study (HyperGEN) (n = 920) and the Bogalusa Heart Study (BHS) (n = 333). Blood samples were collected in each cohort and LTL was measured by Southern blot (Table 1). Mean age ranged across the cohorts from 35 to 75 years and the mean LTL varied from 6.33 to 7.22 kb (Table 1).

Each cohort underwent GWA for LTL on 2.5 million imputed single-nucleotide polymorphisms (SNPs) with adjustment for age, sex, BMI and smoking status (pack years). In the individual cohorts participating to this study, the GWAS inflation ( $\lambda_{GC}$ ) ranged from 0.995 to 1.076, indicating that there was no significant population stratification or that it was very minor. In addition, there was no between-study heterogeneity as indicated by the *P*-values of the heterogeneity test ranging from 0.15 to 0.68 (Table 2). Finally, the quantile–quantile plot demonstrated that the statistical association for all SNPs included in the meta-analysis follows the distribution expected under the null hypothesis (Supplementary Material, Fig. S1).

We calculated that our study (n = 9190) had 80% power to detect a variant (minor allele frequency 10%) which has an effect on LTL of  $\pm 0.12$  kb at a statistical threshold of  $P < 5 \times 10^{-8}$  (Supplementary Material, Fig. S2).

Our analysis confirmed the previously reported associations (13,14) of LTL with the oligonucleotide/oligosaccharide-binding fold containing one gene (*OBFC1*; rs9419958  $P=9.1\times10^{-11}$ ) and the telomerase RNA component (*TERC*) gene (rs1317082,  $P=1.1\times10^{-8}$ ) (Fig. 1). We also identified two novel loci associated with LTL at the conventional genome-wide significance level ( $P<5\times10^{-8}$ ). These newly identified loci include variants in a conserved telomere maintenance complex component 1 (*CTC1*; rs3027234,  $P=3.6\times10^{-8}$ ) and in the zinc finger protein 676 (*ZNF676*; rs412658,  $P=3.3\times10^{-8}$ ) (Fig. 2).

A total of 2226 individuals from four independent studies were included in replication analyses of the novel two genome-wide significant SNPs (rs3027234 and rs412658) (Table 1). There was no evidence for heterogeneity of effect size among the replication studies (Table 2). In the replication sample set, the effect of the minor allele was in the same direction as in the discovery sample for both SNPs. In the

Table 1. Cohort characteristics for the telomere length GWAS analysis

	N Mean, age (year range)		Women (%)	Body mass index (mean $\pm$ STD)	Telomere length (mean $\pm$ STD)	
Cohort						
Framingham Heart Study	1146	59 (33-86)	51	$28.0 \pm 5.0$	$6.96 \pm 0.58$ (Kb)	
Family Heart Study	2508	57 (30-93)	54	$28.9 \pm 5.7$	$6.78 \pm 0.67$ (Kb)	
Cardiovascular Health Study	1061	75 (67–95)	62	$26.6 \pm 4.4$	$6.33 \pm 0.61$ (Kb)	
Bogalusa Heart Study	333	$35(20-48)^a$	42	$28.0 \pm 6.7^{a}$	$7.22 \pm 0.70 \text{ (Kb)}$	
HyperGEN	920	50 (18-87)	50	$29.5 \pm 6.3$	$6.78 \pm 0.61$ (Kb)	
TwinsUK	3222	48 (18-82)	92	$26.1 \pm 4.9$	$6.97 \pm 0.68$ (Kb)	
Replication					· · ·	
Health ABC	337	73.6 (69-80)	49	$27.7 \pm 4.8$	$6.13 \pm 0.51$ (Kb)	
Jerusalem LRC	620	43.2 (41–46)	33	$27.2 \pm 4.5$	$7.00 \pm 0.63$ (Kb)	
ADELAHYDE/Nancy/ERA-France	316	61.7 (25-84)	51	$27.4 \pm 4.6$	$6.28 \pm 0.74 \text{ (Kb)}$	
Danish collection	964	80.7 (58–101)	77	$24.3 \pm 4.0$	$5.74 \pm 0.67 \text{ (Kb)}$	

<sup>&</sup>lt;sup>a</sup>These values include two time points for each individual.

**Table 2.** Meta-analysis results for the most significant SNPs  $(P < 5 \times 10^{-8})$  at each locus

Most significant SNP	Chromosome	Location	minor/major allele	MAF	β <sup>a</sup> (SE)	P-value	Het P <sup>b</sup>	Analysis	closest gene
rs9419958 rs1317082 rs412658	10 3 19	105665936 170980279 22151280	t/c g/a t/c t/c t/c	0.1353 0.2875 0.3534 0.3697 0.3564	0.0829 (0.013) 0.0679 (0.011) 0.0559 (0.010) 0.0241 (0.021) <b>0.0497 (0.009)</b>	9.13E-11 1.14E-08 3.32E-08 1.49E-01 9.75E-09	0.15 0.26 0.65 0.69 <b>0.68</b>	Discovery Discovery Discovery Replication Combined	OBFC1 TERC ZNF676
rs3027234	17	8076817	t/c t/c t/c	0.1678 0.2304 0.1794	-0.0669 (0.012) -0.0226 (0.022) - <b>0.0573 (0.011)</b>	3.58E - 08 2.80E - 01 <b>2.29E - 08</b>	0.45 0.78 <b>0.40</b>	Discovery Replication Combined	CTC1

<sup>&</sup>lt;sup>a</sup>Effect reported in Kbp relative to the minor allele.

combined meta-analysis of the discovery and replication samples, both rs3027234 and rs412658 (Table 2) showed a smaller *P*-value (rs412658,  $P = 9.7 \times 10^{-9}$ ; rs3027234,  $P = 2.3 \times 10^{-8}$ ) than in discovery.

Furthermore, we examined whether the effect of the top four SNPs associated with LTL variation identified in the discovery stage was modified by gender. Although we observed higher *P*-values for the female-specific analyses compared with those of the males, the effect size was not significantly different between the genders (*P*-values ranging from 0.07 to 0.89) (Supplementary Material, Table S2). However, in our study, the female to male ratio was nearly 2 to 1 (Supplementary Material, Table S2), which may compromise the power to find a significant gender interaction for *ZNF676* and *TERC* SNPs.

To explore the potentially functional impacts of these two top SNPs, we used genome-wide expression data from the Multiple Tissue Human Expression Resource (MuTHER) (16) (http://www.muther.ac.uk/) based on  $\sim$ 170 unselected twins sampled for skin, adipose tissue and lymphoblastoid cell lines (LCLs). We focused our analysis only on LCLs and found that the minor allele (T) of rs3027234 is highly associated with lower expression of *CTC1* ( $P = 4.8 \times 10^{-6}$ ). No expression data were present in the analyzed array for *ZNF676*.

Further, we used two independent pathway-based methods [STRING (17) and GRAIL (18)] to identify connections

among our single-marker associations and link them with broader biological processes. Although both approaches are based on published data linking the gene products of our top hits in functional annotated pathways, they use substantially different methods.

With GRAIL, we investigated an excess of connectivity between the four established/novel loci (seed regions) and those loci that did not reach genome-wide significance, but showed suggestive associations ( $P < 5 \times 10^{-5}$ ) with LTL variation (query regions). Among the 75 suggestively associated loci (Supplementary Material, Table S1), we observed five genes to be connected with those at the genome-wide significant loci at  $P_{\rm grail} < 0.01$ . These included *TERT* ( $P_{\rm grail}$  3.2  $\times$  10<sup>-5</sup>), *ZNF256* ( $P_{\rm grail}$  1.4  $\times$  10<sup>-3</sup>), *ZNF587* ( $P_{\rm grail}$  4.5  $\times$  10<sup>-3</sup>), *WDR21A* ( $P_{\rm grail}$  7.3  $\times$  10<sup>-3</sup>) and *RPS29* ( $P_{\rm grail}$  7.6  $\times$  10<sup>-3</sup>).

We then used STRING to uncover additional protein interaction pathways among the genes within 500 Kb of the 79 independent SNPs with  $P < 5 \times 10^{-5}$ .

We observed two clusters (Supplementary Material, Fig. S3). The major cluster included *TERT* (rs10069690,  $P = 3.6 \times 10^{-6}$ ), *SMAD3* (rs17208967,  $P = 3.5 \times 10^{-6}$ ), *YWHAZ* (rs884488,  $P = 3.6 \times 10^{-5}$ ), *PDK1* (rs4972842,  $P = 2.0 \times 10^{-5}$ ), *PIK3R1* (rs16897810,  $P = 1.3 \times 10^{-5}$ ), *GNG2* (rs7154422,  $P = 4.1 \times 10^{-5}$ ), *ADRA1B* (rs10515803,  $P = 3.7 \times 10^{-5}$ ), *PROK2* (rs6790891,  $P = 3.2 \times 10^{-5}$ ), *CHRNA7* (rs11637923,  $P = 4.0 \times 10^{-5}$ ), *DLC1* (rs17254431,

<sup>&</sup>lt;sup>b</sup>P-value of the heterogeneity test.

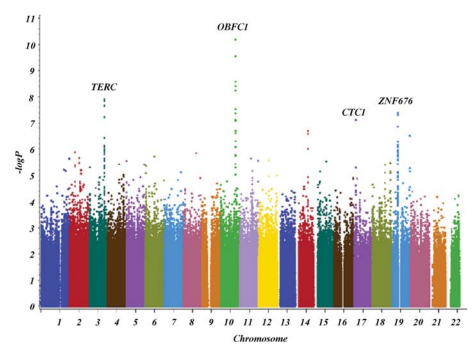


Figure 1. Manhattan plot of discovery meta-analysis.

 $P=2.3\times10^{-5}$ ), LRP1B (rs11886422,  $P=1.8\times10^{-5}$ ) and GRM7 (rs9843107,  $P=9.9\times10^{-7}$ ). Of particular interest for LTL homeostasis was the interaction between TERT and YWHAZ (19), TERT and SMAD3 (20) and the interaction between CTC1 and OBFC1 (21), which formed the second protein–protein cluster.

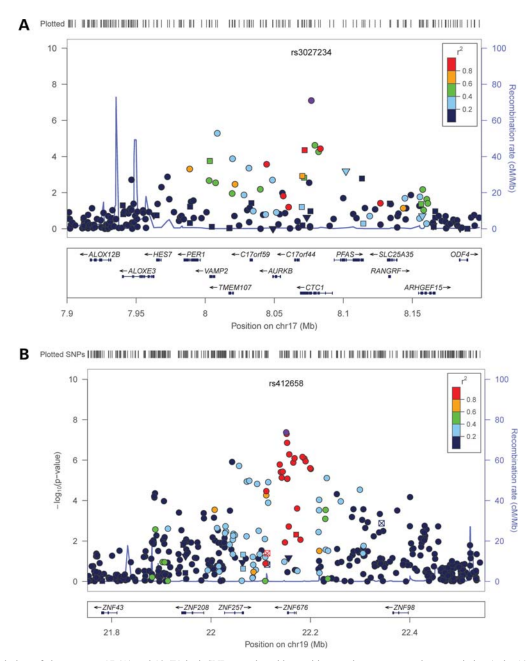
#### DISCUSSION

In this meta-analysis of GWAS results from six observational studies, we confirmed associations with LTL at loci harboring *OBFC1* and *TERC*, two intrinsic telomere maintenance genes. We, also, identified genome-wide significant associations of two new loci [a conserved telomere maintenance *CTC1* and a zinc finger protein 676(*ZNF676*)] with LTL variation. Expression analysis showed that the rs3027234 minor allele was associated with both short LTL and reduced expression of the *CTC1* gene. Based on its position on the sequence (rs3027234 is in introns 11–12, at 132 bp from exon 11) and the LD patterns of the region, we hypothesize that this variation (or another SNP in LD) may disrupt (or create a new) splicing site which results in the reduced gene expression

The variance in telomere length explained by all four identified LTL loci was  $\sim 1.6\%$ . The minor allele of rs3027234 and the major allele of rs412658 were associated with a mean LTL that was, respectively,  $\sim 57$  and 49 bp shorter. To put this in context, in TwinsUK, LTL decreased by 27 bp on average per year (22); therefore, carriage of rs3027234 minor allele was associated with shorter LTL equivalent to 2.1 years of average age-related LTL attrition rate.

The genetic influence of telomere maintenance genes on telomere dynamics in the hematopoietic systems is clearly

displayed in rare Mendelian diseases that result from major mutations in genes that encode the catalytic subunit of telomerase reverse transcriptase (TERT), TERC and other proteins that regulate telomerase and protect telomeres (1-4). The cardinal manifestation of these diseases is aplastic anemia, presumably because very short telomeres affect replicative capacity within the hematopoietic system. In addition, common variants near the TERT affect susceptibility to a variety of cancer subtypes (23-28), suggesting that more subtle variation in telomere maintenance increases cancer risk. At the same time, recent GWASs have also found that loci that contain *OBFC1* and *TERC* explain some of the interindividual variation in LTL in the general population (13,14). The protein encoded by OBFC1 is a member of the heterotrimer CST complex that was originally discovered in budding yeast (Cdc13, Stn1 and Ten1); this complex is conserved in plants and vertebrates (21,29). In human cells, CTC1 carries out the functions of cdc13 (30), while OBFC1 is equivalent to Stn1. Thus, the genes that encode the human heterotrimer CST complex are CTC1, OBFC1 and TEN1. The CST complex is distinguished from the shelterin complex, which consist of six proteins (TRF1, TRF2, TIN2, TPP1, RAP1 and POT1) (31). While the shelterin complex is distinctly involved in telomere maintenance through telomere protection, the CST complex is also involved in other tasks that are not entirely understood. Based on knock-downs of CTC1 and OBFC1, it appears that the complex is not only involved in telomere maintenance, perhaps by negatively regulating telomerase (30,32), but is also engaged in the functioning of the DNA replicative fork, which is not strictly related to telomere protection (29,33). In a recent study, point mutations in CTC1 were identified in Coats plus disorder, a rare recessive Mendelian syndrome (34). Individuals carrying mutations in CTC1 had a short LTL.



**Figure 2.** Regional plots of chromosome 17 (**A**) and 19 (**B**) loci. SNPs are plotted by position on chromosome against association ( $-\log 10 P$ -value) with LTL using discovery GWAS meta-analysis data. In each panel, the SNP with the strongest association is denoted with a purple dot: the P-value attached represents the final P-value obtained in the meta-analysis (Table 2). Estimated recombination rates (from HapMap-CEU) are plotted in blue to reflect the local LD structure on a secondary Y-axis. The SNPs surrounding the most significant SNP (purple dot) are color-coded (see legend) to reflect their LD with the lead SNP (using pairwise  $r^2$ -values from HapMap CEU). Genes position, as well as transcription direction, is shown below the plots (using data from the UCSC genome browser, genome.ucsc.edu).

The findings that *OBFC1* (14) and now *CTC1* are associated with LTL in the general population underscore that the CST complex is indeed engaged in telomere maintenance in humans.

The mechanisms that link *CTC1* to LTL evidently stem from the gene being a member of the CST complex, but the role of *ZNF676* in LTL regulation is unknown. Theoretically, ZNF676 can modify LTL in two broad ways. First, by direct binding to DNA it might alter the expression (repression/

activation) of genes engaged in telomere maintenance and through its interaction with RNAs/proteins alter the post-translational signaling of these genes (35,36). Second, there is evidence that the single-stranded telomeric DNA can fold into a structure known as a G-quartet or G-quadruplex (37,38), which at the 3' end of the telomeres might inhibit telomere elongation by telomerase (37). Zinc finger proteins can bind specifically to and stabilize G-quadruplex DNA (39–41), including telomeric DNA (39,41).

Interestingly, we found that two SNPs that did not reach genome-wide significance, but showed suggestive levels of association ( $P < 5 \times 10^{-5}$ ) with LTL variation, are in or near genes (TERT and SMAD3) pivotal for telomere regulation. Indeed, our results are consistent with findings that the expression of the human telomerase reverse transcriptase (hTERT) (a key element in telomerase activation, telomere maintenance and tumor development) is mediated by SMAD3 (20,42–44).

Leukocyte telomeres may become critically short during the long human lifespan because the hematopoietic system is the most proliferative tissue in the body. Thus, the premise that GWAS of LTL might ultimately provide a mechanistic insight into the role of genes in the telomere dynamics of hematopoietic stem cell has been borne out by this and recent investigations (13,14). That is because LTL dynamics principally reflect hematopoietic stem cell telomere dynamics (45,46). Based on the existing body of research, it is now clear that a large subset of genes contain common variants with very small effects on human LTL maintenance. These genes, as shown recently (47), might play a role in human longevity.

Finally, as women have longer LTL than men (48,49) and African Americans have longer LTL than whites of European ancestry (11,50,51), it will be important to test in future studies whether the functions of the LTL genes identified thus far by GWAS (and other LTL-regulating genes) are modified by sex or ethnicity and whether these genes interact with the environment. In this study, we examined the possibility that the loci identified may have a gender effect. Although we observed that gender had no effect on the function of the identified genes, we could not rule out any possible gender effect because the sample size of the males included in our analysis did not have enough power compared with females.

## **MATERIALS AND METHODS**

#### Cohorts

A detailed description of demographic of the six cohorts participating in this GWAS consortium (the FHS, the FamHS, the CHS, the BHS, the HyperGEN and TwinsUK) can be found in Table 1. All studies received institutional review-board approvals and all participants provided written informed consent for participation in the parent study, including genetic analysis and the use of DNA for the measurement of LTL. Further details related to the discovery cohorts can be found elsewhere (50,52–57). The same covariates (age, age<sup>2</sup> sex, smoking history) were included GWA analysis by all members of the consortium. Evidence of non-European ancestry was assessed in all cohorts by PCA comparison with HapMap populations.

## LTL assay

LTL measurement was performed by Southern blot analysis of the terminal restriction fragments, generated by the restriction enzymes *HinfI* and *RsaI* after verification of DNA integrity (58). The overlay method was used for the FHS and the FamHS (49) and the standard method was used for the other cohorts. These two methods use the same measurement

principles and are highly correlated. The mean length of terminal restriction fragments (expressed in kilobases) was used as a measure of LTL. Coefficients of variation ranged from 0.9 to 2.4% in replicate samples.

#### Genome-wide genotyping

For the FHS, genotyping was conducted on 9274 participants using the Affymetrix 500K mapping array and the Affymetrix 50K gene-focused molecular imprinted polymer array (SNP Health Association Resource at http://www.ncbi.nlm.nih. gov/projects/gap/cgi-bin/study.cgi?study\_id=phs000007.v2.p1). Participants were excluded from association analyses if the call rate across all genotyped SNPs was <97%. A total of 503 551 SNPs available for analysis after exclusions and 1146 individuals met the eligibility criteria, based on LTL measurements and covariate information. The genotype data for 2.5 million SNPs were imputed using MACH (version 1.0.15) (59) software for autosomal SNPs and the reference panel for the imputation was the publicly available phased haplotypes from HapMap (release 22, build 36, CEU population). The final genotyping data for analysis were composed of 2 543 887 SNPs in HapMap using allele dosage (0-2) values. Family members were included. Thus, significance of the association between SNPs and LTL was tested using the linear mixed-effects regression model, which considers the random individual effects correlated within pedigree according to kinship relationships.

For the FamHS, participants were genotyped in three phases using the Illumina HumMap 550, Human 6100-Quadv1 or Human 1M-Duov3 arrays. Because telomere lengths were only measured on blood samples obtained at the second exam of FamHS, all of the subjects of European descent in this study were genotyped on the HumMap550 array. For the subjects of European descent, there were 530 092 SNP markers available for analysis in the FamHS. After exclusions for deviations from the Hardy-Weinberg equilibrium  $(P < 10^{-6})$ , minor allele frequency < 1% and markers not in the HapMap, 499 558 markers were used for imputation. MACH (version 1.0.15) (59) was used to impute up to 2 543 887 SNPs using the publicly available phased haplotypes from HapMap (release 22, build 26, CEU population) as a reference population. A random sample of unrelated 100 men and 100 women, excluding individuals with the highest rates of missing genotypes, was used to estimate parameters that were then applied to the remaining subjects. The GenAbel (60) package in R was used for analysis to allow for the pedigree structures to be modeled appropriately.

For the CHS, genotyping was performed using the Illumina 370CNV BeadChip system on 1080 samples from individuals of European ancestry were available with both LTL measurements and GWA analysis. Because all other cohorts included only Caucasian individuals, the African-American participants were excluded from this analysis (n = 19). Participants with a call rate  $\leq 95\%$  were excluded. The following exclusions were applied to identify a final set of 306 655 autosomal SNPs: call rate  $\leq 97\%$ , Hardy–Weinberg equilibrium  $P < 10^{-5}$ , more than two duplicate errors or Mendelian inconsistencies (for reference Centre d'Etude du Polymorphisme Humain trios), heterozygote frequency = 0 or SNP not found in HapMap.

Imputation was performed using BIMBAM (v0.99) (61) with reference to HapMap CEU using release 22, build 36 with one round of imputations and the default expectation—maximization warm-ups and runs. During the analysis, SNPs were excluded for variance on the allele dosage  $\leq 0.01$ .

For the BHS, participants were genotyped on the Illumina Human610 Genotyping Beadchip. Genotypes were called using the BeadStudio clustering algorithm. Samples were filtered based on the 10th and 50th percentile GenCall score and overall call rates (<99%). Related individuals were identified using Identity-by-descent measures in PLINK (62). Individuals were filtered such that there were no pairs with pi-hat >0.10. This resulted in a total of 333 individuals. SNPs were filtered based on call rates (<90%). Cluster plots for SNPs with call rates between 90 and 95% or with cluster separation scores <0.30 were manually inspected. By genotyping 29 samples in duplicate (18 known replicates and 11 blind replicates), we observed reproducibility >99.99%. There were 626 145 genotyped SNPs used in the reproducibility analysis. Imputation was performed with MACH (v. 1.0.16) (59) using phased CEU haplotypes from HapMap Phase 2 (release 22) and 550 798 genotyped SNPs. A random sample of 200 individuals was used to estimate recombination and error rates before imputing the entire sample. This resulted in imputing 2 543 887 SNPs. By masking 0.1% of the genotypes before imputation, we observed allelic error rates of 1.6% and genotypic error rates of 3.1%. As LTL for BHS participants was measured at two time points, we modeled and tested associations between SNPs and the two LTL measures using a linear mixed model as implemented in the nlme R package (48). A covariance structure for the two LTL measurements was chosen by testing all spatial structures with the model, and the structure that resulted in the lowest Akaike Information Criterion score was used (in this study, the best covariance was exponential). If an SNP had been genotyped, the genotype value (0, 1 or 2) was used as a predictor, whereas if the SNP was imputed, the estimated dosage was used.

For HyperGEN, a subset of 1319 subjects of European descent (none from Framingham) were genotyped on an Affymetrix 5.0 array. After removing the HyperGEN subject overlap with the Family Heart Study, those without telomere lengths or those who did not meet quality control standards, 920 Caucasian subjects had complete data for inclusion in the meta-analysis. The total 358 327 SNPs were used for imputation using MACH (59) after excluding SNPs with Mendelian errors ( $P < 10^{-6}$ ) or minor allele frequencies <1%. The final number of imputed SNPs was 2 355 427. The GenABEL package (60) in R was used for analysis to allow for the pedigree structures to be modeled appropriately.

TwinsUK samples were typed with the Infinium 317K and 610K assay (Illumina, San Diego, USA) at two different centers, the Centre for Inherited Diseases Research (USA) and the Wellcome Trust Sanger Institute. We pooled the normalized intensity data and called genotypes on the basis of the Illluminus algorithm. No calls were assigned if the most likely call was less than a posterior probability of 0.95. Validation of pooling was done by visual inspection of 100 random, shared SNPs for overt batch effects; none was observed. We excluded SNPs that had a call rate <97% (SNPs with MAF  $\geq 5\%$ ) or <99% (for 1%  $\leq$ MAF <5%), Hardy–

Weinberg P-values  $< 10^{-6}$  and minor allele frequencies < 1%. We also removed subjects where genotyping failed for > 2% of SNPs. The overall genotyping efficiency of the GWA was 98.7%. Imputation of genotypes was carried out using the software IMPUTE (63).

Because of the relatedness in the TwinsUK cohort, we utilized the GenABEL software package (60) which is designed for GWAS analysis of family-based data by incorporating a pair-wise kinship matrix calculated using genotyping data in the polygenic model to correct relatedness and hidden population stratification. The score test implemented in the software was used to test the association between a given SNP and LTL.

#### **GWA** meta-analysis

The meta-analysis was carried out independently by two analysts using METAL (64). The results were then compared for consistency.

In the main meta-analysis, we combined the individual P-values obtained for each cohort using Fisher's method (65), which allows P-values and direction of effect in each cohort to be combined taking cohort size into account, independently of  $\beta$ -estimates. Individual cohort results were corrected for residual inflation of the test statistics using lambda of genomic control (GC) estimates (66). The GC values were estimated for each study using all analyzed SNPs.

The level of conventional genome-wide significance level was established at  $P < 5 \times 10^{-8}$ . A *P*-value between  $5 \times 10^{-8}$  and  $5 \times 10^{-7}$  was considered strongly suggestive of association.

The inverse variance weighted method was used to combine the cohort-specific  $\beta$ -estimates to evaluate the effect size only for the markers that reached a genome-wide significance level.

The estimate of the variance explained by the top four SNPs at each locus were calculated in a two-stage approach using GCTA (v 1.0) (67). TwinsUK (n = 3222) genotypes and LTL information were used for this analysis. In the first stage, we estimated the genetic relationship matrix for all the genotyped SNPs in order to account for family structure in the second stage. In the second stage, we performed a restricted maximum likelihood analysis to estimate the variance explained by the top four SNPs. In this stage, age and sex were also included as covariates.

#### **Replication cohorts**

Samples from the Jerusalem LRC Longitudinal Study (N = 620) (68), Health ABC study (N = 337) (69), from ADELAHYDE-Nancy study and ERA-France study (N = 316) (70,71) and from a Danish collection (N = 964) (47) were included in the replication stage. A detailed description for each cohort can be found elsewhere (47,68,69). All studies received institutional review-board approvals and all participants provided written informed consent, including genetic analysis and the use of DNA for the measurement of LTL.

LTL for all the samples was measured by Southern blot analysis of the terminal restriction fragments, as in the discovery data set (47,58).

A linear regression analysis adjusting for same covariates (age, age<sup>2</sup> sex, smoking history) included in the discovery analysis was performed in the replication sample to test association between the two novel SNPs and LTL variation.

#### Power calculations

Power calculations were performed using QUANTO (v 1.2.4) (72). We calculated the power in our combined discovery cohorts (n = 9190) to detect effects of various sizes at a genome-wide significance level ( $P < 5 \times 10^{-8}$ ) for variants with allele frequencies ranging from 0.1 to 0.5.

## Pathway analysis

Grail

We used the GRAIL (18) online tool to evaluate whether genome-wide loci associated with LTL were enriched for connectivity between genes representing particular pathways. The method performs a text-based analysis looking at abstracts in PubMed (we used abstracts prior to December 2006 to avoid confounding from GWAS results). We investigated connectivity between our genome-wide significant signals (seed regions) and those which did not reach genome-wide significance but were suggestively associated with LTL ( $P < 5 \times 10^{-5}$ ) (query regions) as previously described (73).

We identified 75 query regions defined by all SNPs with a  $P < 5 \times 10^{-5}$ . We also pruned SNPs with  $r^2 > 0.05$  in each region represented by more than one SNP using SNAP pairwise LD online tool (74).

#### String

We used the Search Tool for the Retrieval of Interacting Genes/Proteins (STRING) (www.string-db.org) (17) to perform a more in-depth pathway analysis. The interactions explored by STRING include direct (physical) and indirect (functional) associations. We examined potential interactions among all the gene proteins in a range of 500 kb from the 79 independent SNPs with a  $P < 5 \times 10^{-5}$  utilized in the GRAIL analysis.

#### **Expression analysis**

We used the genome-wide expression data from the LCLs from the Multiple Tissue Human Expression Resource (MuTHER) (16). The expression values were derived from a subset of twins from TwinsUK, which were also included in the association analysis. The analysis was performed only on rs3027234 and the expression levels of CTC1 because there was no expression data for ZNF676. The analysis was performed using MERLIN (75), taking in account the family structure. For this analysis, the significance was defined as P < 0.05, as only one independent test was performed.

## SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

Conflict of Interest statement. None declared.

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