









Natural variation in plant telomere length is associated with flowering time

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Abstract

Telomeres are highly repetitive DNA sequences found at the ends of chromosomes that protect the chromosomes from deterioration during cell division. Here, using whole-genome re-sequencing and terminal restriction fragment assays, we found substantial natural intraspecific variation in telomere length in *Arabidopsis thaliana*, rice (*Oryza sativa*), and maize (*Zea mays*). Genome-wide association study (GWAS) mapping in *A. thaliana* identified 13 regions with GWAS-significant associations underlying telomere length variation, including a region that harbors the telomerase reverse transcriptase (*TERT*) gene. Population genomic analysis provided evidence for a selective sweep at the *TERT* region associated with longer telomeres. We found that telomere length is negatively correlated with flowering time variation not only in *A. thaliana*, but also in maize and rice, indicating a link between life-history traits and chromosome integrity. Our results point to several possible reasons for this correlation, including the possibility that longer telomeres may be more adaptive in plants that have faster developmental rates (and therefore flower earlier). Our work suggests that chromosomal structure itself might be an adaptive trait associated with plant life-history strategies.

Introduction

Telomeres are regions of repetitive sequences that cap the ends of eukaryotic chromosomes to protect them from deterioration and from eliciting a DNA damage response

(Shay and Wright, 2019). During DNA replication, failure to fill in terminal base pairs at the lagging strand leads to the end-replication problem (Olovnikov, 1971, 1973; Watson, 1972), resulting in the shortening of chromosome ends

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Background: Telomeres are DNA-protein complexes found at the ends of eukaryotic chromosomes that protect the genome from instability and damage. In humans, telomeres shorten each time a cell divides, and accelerated telomere shortening leads to aging-associated diseases. In both plants and animals, telomere lengths often vary not only between but also within species. Such telomere length variation is associated with differences in life span or aging in animals, but the functional importance of natural telomere length variation in plants remains unclear.

Question: What is the biological significance of carrying different telomere lengths for natural plant populations? We asked this question in three model plant species: *Arabidopsis thaliana*, (Asian rice) *Oryza sativa*, and maize (*Zea mays*).

Findings: In all three plant species, we discovered that individuals with longer telomeres flowered significantly faster than those with shorter telomeres. Through genome-wide association mapping (GWAS), we identified several candidate genomic regions underlying natural telomere length variation, including a region harboring the telomerase reverse transcriptase (*TERT*) gene. We also discovered associations of telomere length with the genotypes' geographic origin, suggesting that plants from temperate climates have the longest telomeres. Our data indicate that longer telomeres may be more adaptive in plants that have faster developmental rates and therefore flower earlier. Overall, our discovery of the association of flowering time with telomere length across the three evolutionarily different plant lineages suggests that this response may be a universal phenomenon across the plant kingdom and implies that telomere length variation in plants correlates with different life strategies and adaptations in response to the environment.

Next steps: The main open question is how telomere length variation mechanistically leads to changes in plant flowering time. Future genetic experiments will test whether telomere length affects flowering time directly or indirectly.

during each cell division and the eventual loss of replicative capacity (Hayflick and Moorhead, 1961; van Deursen, 2014). To prevent this loss of chromosomal DNA at the termini, the ribonucleoprotein enzyme complex telomerase, whose core components consist of a telomerase reverse transcriptase (*TERT*) and RNA template (*TR*; Osterhage and Friedman, 2009; Song et al., 2019), binds to single-stranded telomeric DNA at the 3'-end and processively extends the telomere sequence (Wu et al., 2017). Other specialized telomere binding proteins are also recruited to prevent the telomere from being detected as damaged DNA (Fulcher et al., 2014).

Eukaryotic telomeres consist of a tandem repeat of TG-rich microsatellite sequences (Podlevsky and Chen, 2016). The core telomeric repeat sequence is conserved among species. For instance, vertebrates have the telomeric repeat TTAGGG (Meyne et al., 1989), while in most plants, the sequence is TTTAGGG (Fajkus et al., 2005). The most noticeable difference in telomeres between organisms is differences in telomere length, which can be as short as 300 bps in yeast (*Saccharomyces cerevisiae*; Gatzbonton et al., 2006) and up to 150 kb in tobacco (*Nicotiana tabacum*; Fajkus et al., 1995). Within species, telomere sequences also display substantial heritable length variation. Several examples of telomere length polymorphisms and the underlying genes responsible for this variation have been identified in humans (*Homo sapiens*), yeast, and the roundworm *Caenorhabditis elegans* (Liti et al., 2009; Levy et al., 2010; Jones et al., 2012; Codd et al., 2013; Cook et al., 2016). In plants, variation in telomere length has also been observed between individuals (Burr et al., 1992; Shakirov and

Shippen, 2004; Maillet et al., 2006; Fulcher et al., 2015), between organs (Kilian et al., 1995), and between cell types (González-García et al., 2015). Quantitative trait locus (QTL) studies in *Arabidopsis thaliana* and maize (*Zea mays*) have indicated that natural variation in telomere length is a heritable complex trait (Burr et al., 1992; Brown et al., 2011; Fulcher et al., 2015). In *Arabidopsis*, a recent QTL analysis using MAGIC lines suggested that genes involved in ribosome biogenesis and cell proliferation, such as *NOP2A*, *RPL5A*, and *RPL5B*, may be involved in setting telomere length set points (Abdulkina et al., 2019).

A more puzzling question is this: What is the significance of natural variation in telomere length for organisms? Telomere length variation could be neutral and result from random genetic drift or random stochasticity in the activity of the telomerase. Alternatively, telomere length differences could have fitness effects that are subject to natural selection, possibly due to their association with cellular senescence, which has been implicated in controlling life span in yeast and animals (Aubert and Lansdorp, 2008; Kupiec, 2014). In mammals, for example, telomere shortening correlates with between-species differences in lifespan (Whittemore et al., 2019), suggesting that telomeres are involved in the aging process (Aubert and Lansdorp, 2008). Indeed, it has been suggested that the aging trajectory of telomere lengths could be a product of optimization of a life-history tradeoff (Young, 2018). This is by no means universal, as in *C. elegans*, no fitness differences or clear phenotypic consequences were associated with natural variation in telomere lengths (Cook et al., 2016). A recent hypothesis proposes that shorter telomeres are associated with a faster

“pace of life” in some organisms, in which energy is conserved for reproduction and away from somatic maintenance (Giraudeau et al., 2019).

While there is interest in the links between telomeres and life-history traits (e.g. aging) in animals, relatively little is known about how telomere length evolution affects plant life-history strategies. Aging in plants differs fundamentally from that in animals (Watson and Riha, 2011), and it is unclear whether the telomere-aging and “pace-of-life” models are also applicable to plants. Indeed, no specific hypotheses have been put forth to explain natural telomere length variation in plants; whether telomeres have an effect on plant life-history traits and are a target of natural selection remains an open question.

Here we describe an association between a life-history trait—flowering time—and natural telomere length variation in plants. Using whole-genome sequence data as well as experimental assays, we determine the extent of telomere length variation in three plant species: *A. thaliana*, rice (*Oryza sativa*), and maize (*Z. mays*). We use a genome-wide association (GWAS) mapping to identify genomic regions associated with natural telomere length variation in *A. thaliana*, including one that spans the telomerase gene *TERT*. We show that longer telomeres are found in plants that flower earlier in these three annual plant species.

Results

Genome-wide variation in *A. thaliana* tandem repeats

Satellite DNA sequences are repetitive sequences structured as arrays of DNA that are tandemly repeated in the genome, sometimes up to 10^6 copies. We examined genome-wide variation in satellite DNA repeat copy number in *A. thaliana* using the program k-Seek (Wei et al., 2014, 2018). k-Seek is an assembly-free method of identifying and quantifying k-mer repeats in unmapped short read sequence data, and k-mer counts are highly correlated with direct measurements of satellite repeat abundance (Wei et al., 2014).

We used whole-genome re-sequencing data from the 1001 *A. thaliana* Genome Consortium Project (Alonso-Blanco et al., 2016). We quantified genome-wide *A. thaliana* tandem repeat copy numbers by focusing on 483 individuals that were sequenced from leaves using identical protocols (designated as AraTha_{Kmer}; see Methods section for details). The quantity of each k-mer sequence is presented as copies per $1 \times$ read depth after GC normalization (Flynn et al., 2017). We set 50 bps as the minimum length of tandem repeat sequences that can be called.

Adding up k-mer copy numbers, the median total length of tandem repeats per individual is estimated at 341 kb. Across the population, individuals displayed over 25-fold differences in total tandem repeat lengths. The most abundant k-mer was the poly-A repeat, followed by the 7-mer AAACCCT (Figure 1A). Some k-mers, such as the AC repeat, had a wide range of variation between individuals, with a range of zero to thousands of copies. Our

computationally based estimates were qualitatively concordant with direct estimates of repeat copy number that used DNA gel blot analysis to characterize 1- to 4-mer variation in a single *A. thaliana* ecotype (Depeiges et al., 1995). For instance, DNA gel blot analysis showed that A and AG repeats were the most abundant 1-mer and 2-mer, respectively, while in the 3-mer class, AAG and ATC repeats were highly abundant. All of this was seen in our estimates as well. Furthermore, DNA gel blot analysis indicated that AAG was more abundant than AAC repeat, which we also observed as well, indicating that k-Seek is highly specific in quantifying the type of k-mer repeat.

Arabidopsis thaliana telomere length variation

The tandem repeat with the second highest abundance in the *A. thaliana* genome is the k-mer AAACCCT, which corresponds to the canonical telomere repeat sequence in plants [a reverse complement of the Arabidopsis TTAGGG telomere repeat, followed by tandem repetition] (Fajkus et al., 2005; Watson and Riha, 2010). There is a wide range in total copy numbers for the AAACCCT repeat, from 1,257 copies in *Arabidopsis* ecotype Ler-1 to 38,850 copies in ecotype IP-Fel-2 (Supplemental Data Set 1), with a median of 6,411 and a mean of $7,113.6 \pm 161.1$ (\pm standard error) copies (Fig. 1B).

We compared telomere repeat copy numbers inferred from k-Seek to directly measure telomere lengths in various *A. thaliana* accessions using the terminal restriction fragment (TRF) method (Fitzgerald et al., 1999). We experimentally measured the telomere length by TRF in 424 *A. thaliana* accessions (Supplemental Figure 1) and combined this dataset with data for 229 previously analyzed accessions (Fulcher et al., 2015). In our total dataset of 653 *A. thaliana* accessions, designated as AraTha_{TRF}, the mean telomere lengths ranged from 1,065.2 bp in Hov1-10 to 11,787.2 bp in IP-Pro-0 (Supplemental Data Set 2), with a median length of 3,533 bp and a mean length of $3,767.4 \pm 50.1$ bp (\pm standard error; Figure 1B).

A total of 112 accessions overlapped between the AraTha_{Kmer} and AraTha_{TRF} sample sets, and we found a significant positive correlation in log₁₀ telomere lengths from the two methods (Figure 1C; Pearson's $r = 0.58$ and $P = 2.3 \times 10^{-11}$, Spearman's $\rho = 0.55$ and $P = 1.9 \times 10^{-10}$, Kendall's $\tau = 0.39$ and $P = 8.8 \times 10^{-10}$). We also looked at genomic data from a second set of 201 accessions that were sequenced using a different protocol from the AraTha_{Kmer} set (Supplemental Data Set 1). This second genome dataset had 140 accessions in common with AraTha_{TRF}, and we found a significant positive correlation in estimated telomere lengths from the two methods as well (Pearson's $r = 0.33$ and $P = 8.5 \times 10^{-5}$; Spearman's $\rho = 0.34$ and $P = 4.1 \times 10^{-5}$; Kendall's $\tau = 0.24$ and $P = 2.7 \times 10^{-5}$). It should be noted that the estimates of total telomere lengths are generally higher using k-Seek compared to the direct TRF method, which may be attributed to interstitial telomere repeats that can be detected with the former method. Nevertheless, our analysis revealed significant correlations in estimated

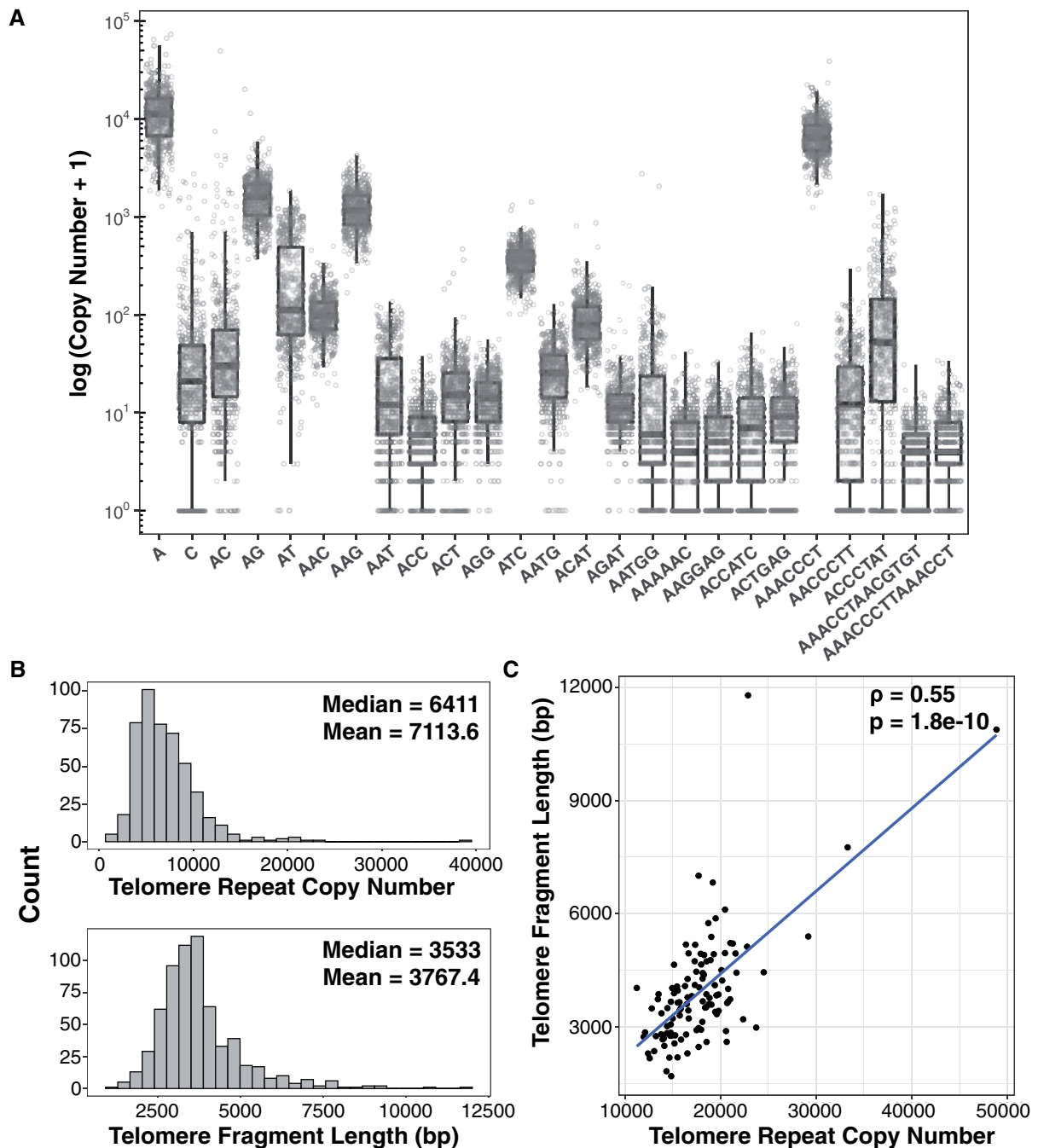


Figure 1 *Arabidopsis thaliana* tandem repeat profile. (A) Top 25 most abundant k-mers from the AraTha_{kmer} set. K-mers are ordered alphabetically, then by size. Whiskers in boxplot represent ± 1.5 interquartile range. (B) Distribution of estimates of telomere repeat copy numbers from the AraTha_{kmer} set (above) and terminal fragment lengths from the AraTha_{TRF} set (below). (C) Scatter plot of telomere lengths from the 112 accessions that were in common between the AraTha_{kmer} and AraTha_{TRF} sets. Spearman ρ and P -values are shown in the upper-right side.

telomere lengths from the k-Seek versus the TRF methods, and the use of either estimate gave similar results in our downstream analyses (see below).

GWAS mapping of telomere length variation implicates telomerase

We investigated whether the natural variation in *A. thaliana* telomere length has a genetic basis. We analyzed the

AraTha_{kmer} and AraTha_{TRF} sets separately by conducting genome-wide association mapping of telomere length variation. We used the FarmCPU method for GWAS analysis, which works well for identifying loci of complex traits that may be confounded by population structure (Liu et al., 2016). GWAS analysis revealed seven genomic regions with single-nucleotide polymorphisms (SNPs) significantly associated (after Bonferroni correction) with telomere length

variation in the AraTha_{Kmer} set, and seven significant GWAS hits in a separate analysis of the AraTha_{TRF} data (Figure 2 and see Supplemental Table 1 for genome coordinates).

Of these, one SNP was detected in both analyses ($P < 1 \times 10^{-10}$) and is located on chromosome 5 at position 5,538,242. This SNP is found at the 3'-UTR (untranslated region) of locus AT5G16850 (Figure 2), which corresponds to the *TERT* gene. The *TERT* gene is crucial in maintaining telomere length in *A. thaliana* (Fitzgerald et al., 1999) and other eukaryotes (Autexier and Lue, 2006). Besides the overlapping SNP at the *TERT* gene, the only other potentially overlapping SNPs were two GWAS-significant SNPs on chromosome 3 position 5,136,799 from the AraTha_{Kmer} set and position 5,295,758 from AraTha_{TRF} set (~159 kbp apart). However, none of the nonoverlapping 12 significant SNPs from both GWAS studies were in proximity (within 200 kbp) to known telomere-regulating genes. Nonetheless, two significant SNPs (chromosome 2 position 12,466,562 from AraTha_{TRF} dataset and chromosome 2 position 13,008,964 AraTha_{Kmer} dataset) from our GWAS analyses were also

located in a QTL region on chromosome 2 identified in a recombinant inbred line mapping study (Fulcher et al., 2015; Supplemental Table 1).

Arabidopsis thaliana telomere length is associated with flowering time variation

We examined if the telomere length distribution among ecotypes had a geographical basis. We compared the telomere lengths from the AraTha_{Kmer} and AraTha_{TRF} sets to each ecotype's natural geolocation. We detected a significant negative correlation between telomere length and latitude for both the AraTha_{Kmer} (Spearman's $\rho = -0.182$ and $P = 8.8 \times 10^{-5}$) and AraTha_{TRF} (Spearman's $\rho = -0.115$ and $P = 0.022$) datasets (Figure 3A), but for longitude, only the AraTha_{Kmer} set had a significant negative correlation [Spearman's $\rho = -0.165$ and $P = 3.8 \times 10^{-4}$] (Figure 3B).

In *A. thaliana*, life-history traits are often associated with geographic adaptation (Stinchcombe et al., 2004; Montesinos-Navarro et al., 2012). We hypothesized that telomere length polymorphisms occurred as a response to

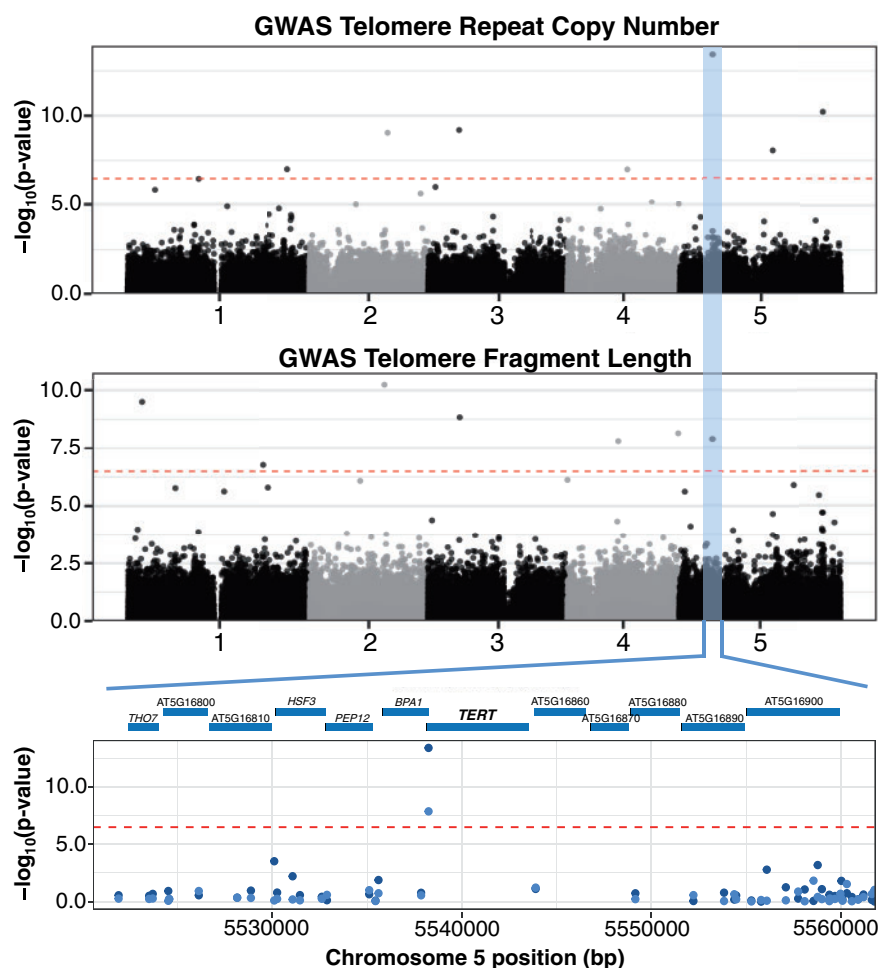


Figure 2 GWAS of *A. thaliana* telomere length variation. Manhattan plot of the genome-wide P -values testing the association of telomere copy number of the AraTha_{Kmer} set (top) and terminal fragment length of the AraTha_{TRF} set (middle) using the FarmCPU approach. Red dotted line indicates the Bonferroni-corrected significance threshold ($\alpha = 0.05$). The GWAS-significant region common in both the AraTha_{Kmer} and AraTha_{TRF} sets is highlighted in blue. Closeup of the region with the most significant GWAS SNP is shown at the bottom. Dark blue points are GWAS P -values from the AraTha_{Kmer} set, and light blue points are GWAS p -values from the AraTha_{TRF} set.

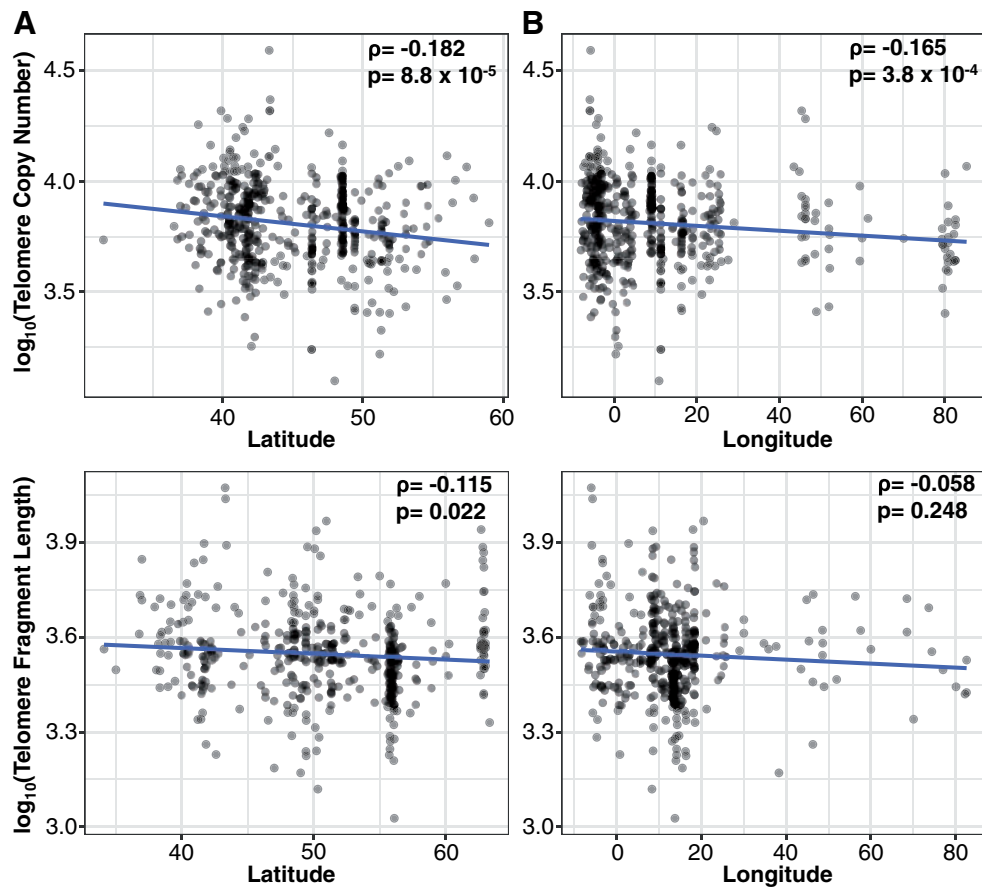


Figure 3 Geographic distribution of telomere lengths in *A. thaliana*. Correlation between telomere length and latitude/longitude in the AraTha_{Kmer} set (A) and AraTha_{TRF} set (B). Spearman's ρ and P-values are shown in the upper-right side in each plot.

adaptation to specific life-history strategies. We tested whether specific developmental traits associated with life history were correlated with a variation in telomere length by examining the developmental trait or telomere length phenotype for each accession. We compared telomere lengths from the AraTha_{Kmer} and AraTha_{TRF} sets to seven different developmental traits (see Figure 4A and Supplemental Figure 2). The AraTha_{Kmer} set had significant negative correlations with four traits: days to flowering at 10°C (Spearman's $\rho = -0.119$, $P < 0.009$), days to flowering at 16°C (Spearman's $\rho = -0.173$, $P < 1.6 \times 10^{-4}$), cauline leaf number (Spearman's $\rho = -0.125$, $P < 0.007$), and rosette leaf number (Spearman's $\rho = -0.152$, $P < 0.001$), and positive correlation with rosette branch number [Spearman's $\rho = 0.111$, $P < 0.03$] (Figure 4A). In the AraTha_{TRF} set, the same four traits had significant negative correlations with telomere length: days to flowering at 10°C (Spearman's $\rho = -0.223$, $P < 2 \times 10^{-6}$), days to flowering at 16°C (Spearman's $\rho = -0.210$, $P < 1 \times 10^{-5}$), cauline leaf number ($\rho = -0.178$, $P < 8 \times 10^{-4}$), and rosette leaf number [Spearman's $\rho = -0.211$, $P < 0.0001$] (Figure 4B; Supplemental Figure 2). Telomere length explains between 0.3% and 0.8% of variation

in flowering time at 10°C and 2.36 and 2.46% at 16°C. It should be noted that in *A. thaliana*, leaf number is developmentally correlated with flowering time.

To test whether these correlations were simply due to population structure, we fit a multiple linear regression model that included the first four axes of a principal component analysis of the SNP variation as additive variables. The results showed that in the AraTha_{Kmer} set, telomere copy number was a significant negative predictor for the traits days to flowering at 16°C ($P < 0.024$), cauline leaf number ($P < 0.034$), and rosette leaf number ($P < 0.003$), even when accounting for population structure. Also, in the AraTha_{TRF} set, telomere length was a significant negative predictor for days to flowering at 10°C ($P < 5 \times 10^{-4}$), days to flowering at 16°C ($P < 0.0015$), and rosette leaf number ($P < 0.0076$) after accounting for population structure. Telomere length in this set accounts for $\sim 0.5\%$ of flowering time variation at 10°C, but this increases to $\sim 2.4\%$ at 16°C. Together, these results suggest that telomere length is negatively associated with flowering time in this annual species, such that plants with longer telomeres flower earlier.

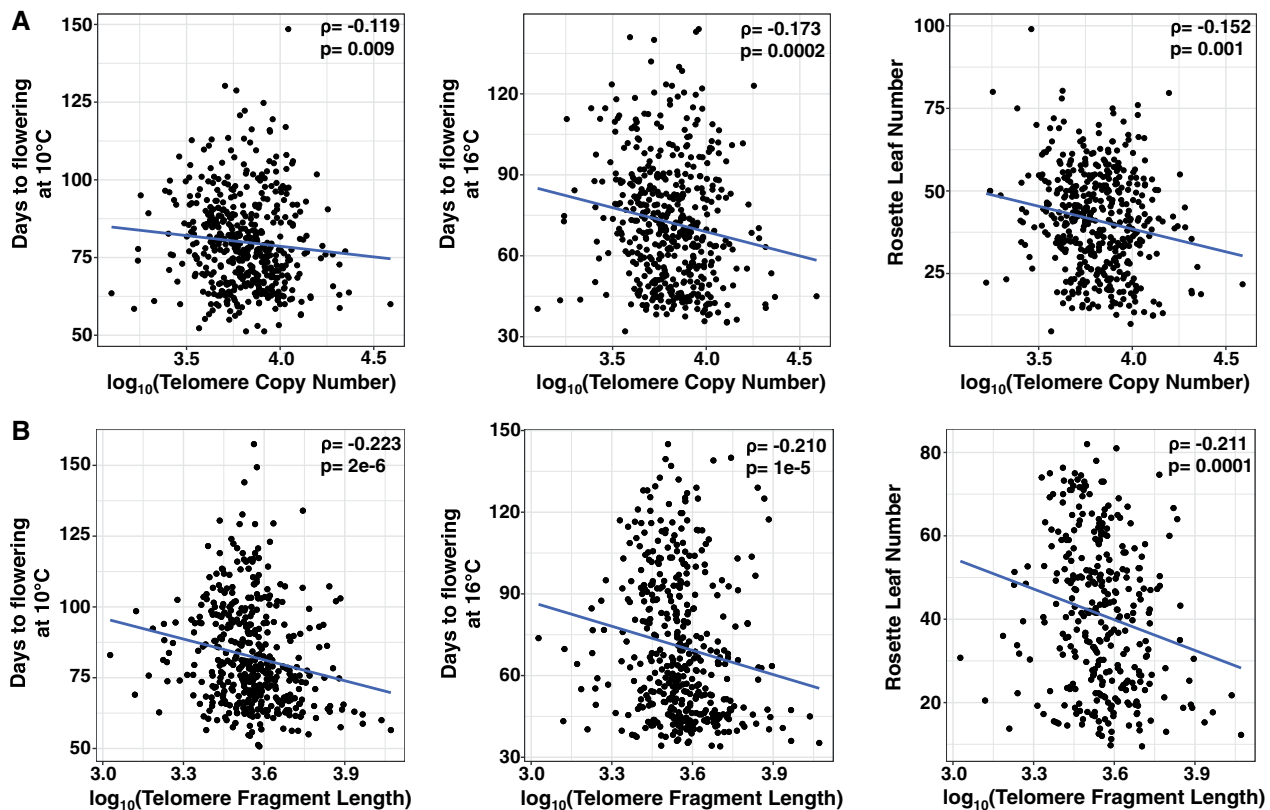


Figure 4 Association between telomere length and developmental traits in *A. thaliana*. Correlation between telomere copy number or length and various *A. thaliana* life-history traits for the AraTha_{Kmer} set (A) and AraTha_{TRF} set (B). Spearman's ρ and P-values are shown in the upper-right side in each plot.

Influence of *A. thaliana* genome size and telomere or k-mer repeats on flowering time

A recent study showed that chromosomal features other than telomeres, such as genome size, can affect flowering time in plants (Bilinski et al., 2018). We investigated the relationship between telomere length and genome size in *A. thaliana* using data obtained through direct TRF measurements. A previous flow cytometry study provided genome size estimates (Long et al., 2013) for 139 accessions in our AraTha_{TRF} set, and these samples showed no significant correlation between telomere length and genome size (Supplemental Figure 3). We also compared the copy number of ribosomal DNA, which is responsible for the majority of the *A. thaliana* genome size differences (Long et al., 2013), and also found no significant correlations with telomere length (Supplemental Figure 3). We did find a significant positive correlation between flowering time and genome size in this set (Spearman's $\rho = +0.228$, $P < 0.0042$), as has been observed in maize (Bilinski et al. 2018); interestingly, this relationship is opposite that of the correlation with telomere length.

We then examined other k-mer repeats and their relationship with the telomere repeat and flowering time variation in *A. thaliana*. Examining the computational predictions from the AraTha_{Kmer} set, the top 10 most abundant k-mers (Figure 1A) all had significant positive correlations with the

telomere repeat copy number (Supplemental Table 2). Examining each k-mer and its association with flowering time; however, there was no correlation with flowering time at 10°C, while there was a positive correlation with flowering time at 16°C for the k-mers A ($\rho = 0.155$, $P < 6.8 \times 10^{-4}$), AG ($\rho = 0.134$, $P < 3.42 \times 10^{-3}$), and AT ($\rho = 0.178$, $P < 9.77 \times 10^{-5}$). Note that the correlations for those three k-mers, like for genome size and ribosomal repeat lengths, were opposite that between telomere repeat length and flowering time. We then conducted a multiple linear regression of flowering time at 16°C with the k-mers A, AG, and AT (which were computationally estimated for abundance in this study) and the telomere repeat abundance in the model. The results showed that the telomere repeat was a significant negative predictor of flowering time at 16°C ($P < 5 \times 10^{-7}$), even after accounting for the abundance of the other three k-mers.

Evidence of selection within the genomic region associated with telomere length

The nonrandom geographical distribution of telomere lengths and the association with flowering time suggest that the length variation may be the result of natural selection. We conducted a selective sweep analysis focusing on the SNP region (chromosome 5 and position 5,538,242) that was associated with telomere length variation in GWAS

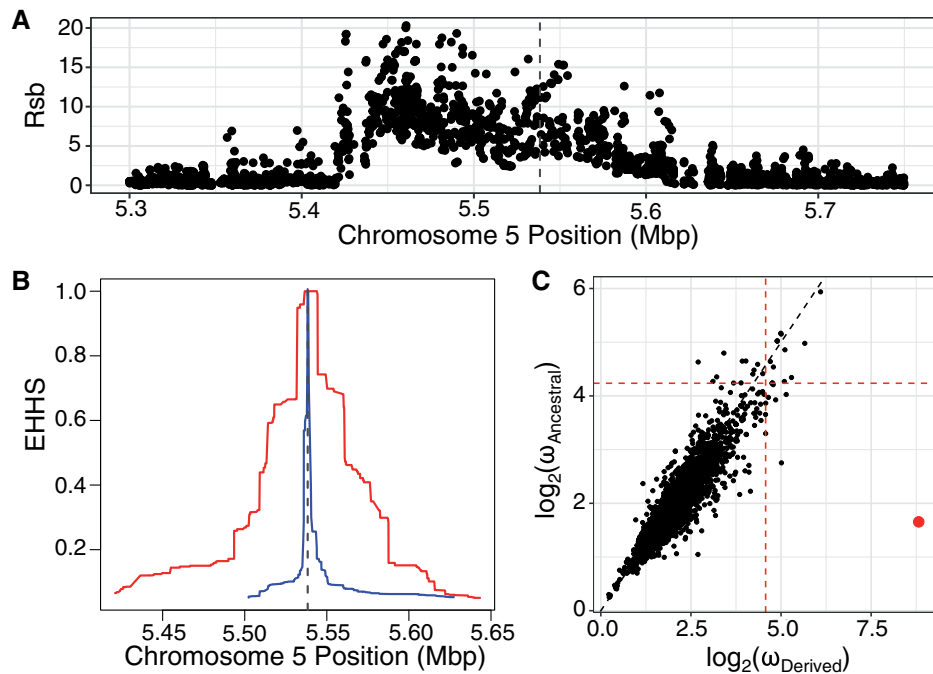


Figure 5 Selection on the GWAS-significant region found in both the AraTha_{Kmer} and AraTha_{TRF} sets. (A) Rsb statistics (\log_{10} P -value) comparing the EHHS of derived versus ancestral alleles on chromosome 5 near the GWAS-significant SNP (dotted line) region. (B) EHHS decay plot centered on site 5,538,242 of chromosome 5. Red line is for the derived allele (corresponding to the minor allele) and blue line is for the ancestral allele (corresponding to the major allele). (C) OmegaPlus-based selective sweep test statistics in 10-kbp windows for individuals carrying derived or ancestral allele. Red lines indicate the top 1% OmegaPlus statistic threshold. Red point indicates the OmegaPlus statistics for the window containing the *TERT* gene.

analyses of both the AraTha_{Kmer} and AraTha_{TRF} sets (Figure 2).

Using the entire 1,001 *A. thaliana* genome dataset, we calculated the integrated haplotype homozygosity score (iHS; Voight et al., 2006) and ω (Kim and Nielsen, 2004; Alachiotis et al., 2012) statistics, and found no evidence of a selective sweep across the entire population in this GWAS-significant SNP region. We then divided the sample population based on the allele status of the GWAS-significant SNP to examine group-specific evidence of selective sweeps. Our GWAS analysis indicated that individuals carrying the minor allele (frequency = 17% in the 1001 *A. thaliana* genome dataset) had longer telomeres than the others. We examined the site-specific extended haplotype homozygosity statistics (EHHS) between individuals carrying the major and minor alleles at the GWAS-significant SNP region. The ratio of EHHS between the populations (Rsb; Tang et al., 2007) statistic was elevated around the region encompassing the GWAS-significant SNP region (Figure 5A), and the increased EHHS occurred around the SNP for individuals carrying the minor allele (Figure 5B). We then calculated the ω statistic, which detects selective sweeps based on patterns of linkage disequilibrium (LD), and found that the GWAS-significant SNP region was a significant outlier within individuals carrying the minor SNP allele associated with longer telomeres, but not within the individuals carrying the major allele associated with shorter telomeres (Figure 5C). We also

conducted a bootstrap procedure to determine the significance of this sweep signature. None of the bootstrap replicates had a ω statistic as large as what we observed for the group of individuals carrying the minor SNP allele (Supplemental Figure 4).

Flowering time is also negatively correlated with telomere copy number in rice and maize

The association between telomere length and flowering time was unexpected, but it suggested that individuals with different telomere lengths had contrasting life-history strategies. We investigated if this correlation is found outside *A. thaliana* by examining the relationship between telomere length and flowering time in *O. sativa* and *Z. mays*. For each species, we analyzed whole-genome re-sequencing data from previous studies that also reported flowering time data (Flint-Garcia et al., 2005; Wang et al., 2018).

In rice (*O. sativa*) and maize (*Z. mays*), there was a wide variation in telomere copy number, and like *A. thaliana*, many of the differences appear to show population stratification. In maize, data are available that incorporate both whole-genome re-sequencing and flowering time (Flint-Garcia et al., 2005), and we were able to computationally estimate telomere copy numbers in this set of 277 maize genotypes using *k*-Seek. An earlier study directly measured telomere lengths in maize inbred lines (Burr et al. 1992), and 11 samples were common to both of these studies.

Although the sample overlap was low, we did find that in these 11 common genotypes, the correlation between the computational versus direct measurement of telomere length was very high (Spearman's ρ 0.95, $P < 1 \times 10^{-16}$), suggesting that our computational estimates also provide good estimates of telomere length in this species.

Most maize varieties are genetically classified as either from nonstiff-stalk (NSS) and stiff-stalk (SS) populations from temperate regions (Liu et al., 2003) or from the tropical/subtropical (TS) population. Our analysis of the 277 maize cultivars showed that NSS varieties had significantly higher telomere copy number than both SS and TS maize cultivars [Mann–Whitney U (MWU) test, $P = 0.0304$ and 0.0065 , respectively] (see Figure 6).

We also analyzed data for 2,952 rice varieties (Wang et al., 2018). This species displayed the most significant differences in telomere copy numbers between subpopulations, likely due to deep population structure in rice (Huang et al., 2012; Wang et al., 2018). Most rice varieties can be divided into *japonica* or *indica* subspecies (Wang et al., 2018), which

possess a significant genetic and physiological differentiation with each other (Zhao et al., 2011), so we analyzed each subpopulation separately (Figure 6). In *japonica*, the temperate *japonica* ($G_{J_{temp}}$) group had significantly higher telomere repeat copy numbers than both subtropical ($G_{J_{subtrp}}$) and tropical *japonica* ($G_{J_{trp}}$; MWU test, $P = 0.0051$ and 1.34×10^{-10} , respectively). In *indica* rice, the subpopulation XI-1A (from East Asia) had significantly higher telomere copy numbers compared to subpopulation XI-1B (modern varieties of diverse origin), XI-2 (from South Asia), and XI-3 (from Southeast Asia; MWU test, $P = 0.0046$, 2.38×10^{-20} , and 1.83×10^{-18} , respectively; see Figure 6).

Notably, in both rice and maize, the subpopulations with the highest telomere copy numbers (temperate *japonica* and NSS maize) were from temperate regions. It should be noted that in maize, the SS population is also temperate but has shorter telomeres, which may reflect the distinct recent breeding history of SS inbred maize.

Like in *A. thaliana*, we observed a significant negative correlation between telomere copy number and flowering

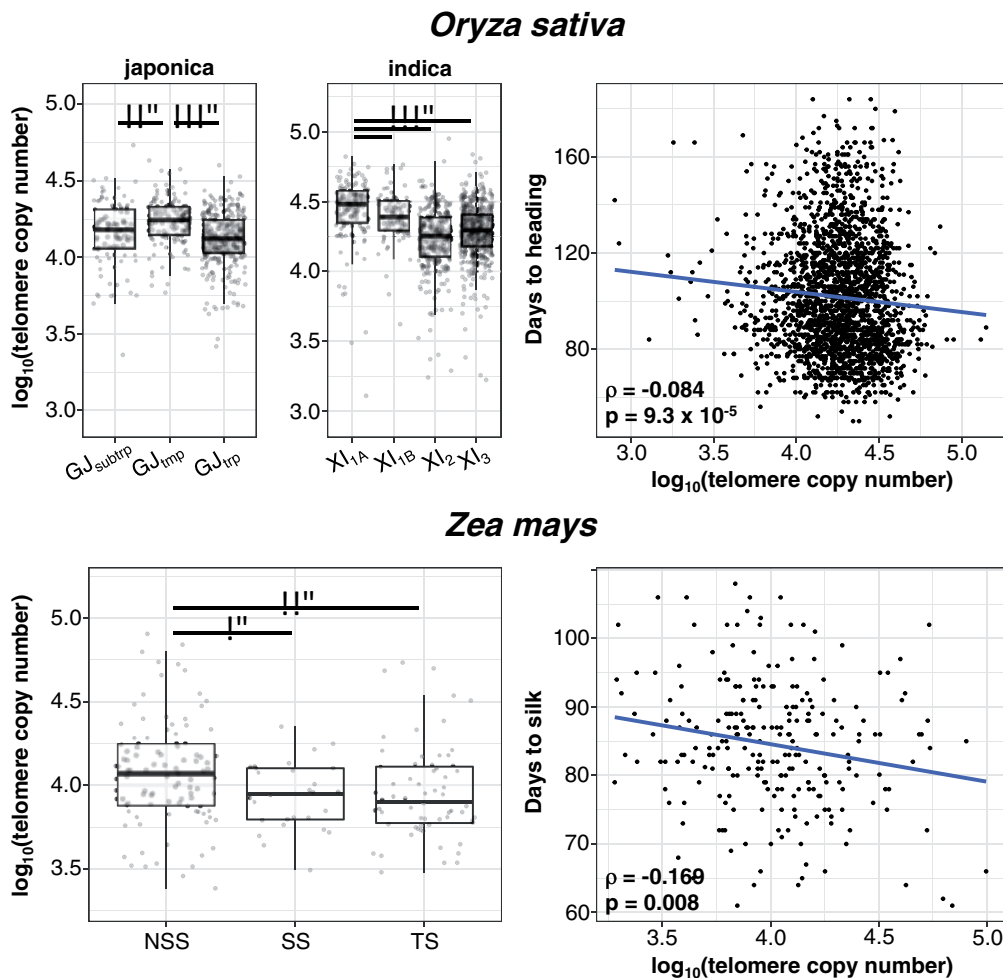


Figure 6 Association between telomere length and flowering time in rice (*O. sativa*) and maize (*Z. mays*). Telomere copy numbers per subpopulation are shown on the left. * $P < 0.05$, ** $P < 0.01$, and *** $P < 0.001$. Correlation between telomere copy number and flowering time is shown on the right. For maize, the days to silk measured in Aurora, NY in 2007 is shown. Spearman's ρ and P -values are shown in the lower-left side of each plot. Whiskers represent $\pm 1.5 \times$ interquartile range.

time in rice ($\rho = -0.084$, $P < 9.3 \times 10^{-5}$; see Figure 6). This correlation is even more pronounced within each rice subspecies (*ssp. japonica*, $\rho = -0.255$, $P < 5.3 \times 10^{-9}$ and *ssp. indica*, $\rho = -0.259$, $P < 2.9 \times 10^{-15}$). In maize, we obtained previously measured flowering time data that were collected in five field locations in the USA and over a 3-year period in some locations; there were data for a total of nine fields/seasons (Zhao et al., 2006). In seven cases, there were significant negative correlations between telomere repeat copy number and flowering time ($\rho = -0.123$ to -0.169 , $P < 0.008$ to 0.045 ; see Figure 6 for one example), one was marginally nonsignificant ($\rho = -0.130$, $P < 0.057$), and one still appeared negative but was nonsignificant ($\rho = -0.057$, $P < 0.43$; see Supplemental Table 3).

As in *A. thaliana*, telomere length explains a relatively small fraction of flowering time variation in these plants: a mean of 2.64% of the variation in maize (based on data from seven fields/seasons; see Supplemental Table 3), 5.39% of the variation for *japonica*, and 3.80% of the variation for *indica* rice. Despite the relatively low levels of flowering time variation explained, these correlations with telomere length are significant. To test whether these correlations were due simply to population structure, we fit a multiple linear regression model that included the first four axes of a principal component analysis of the SNP variation as additive variables. Similar to the case in *A. thaliana*, for both rice and maize, telomere length had a significantly negative effect on flowering time, even after accounting for population stratification ($P < 0.02$ for rice and $P < 0.033$ for maize).

GWAS of telomere repeat copy number variation in rice and maize using FarmCPU showed significant SNP markers in the *japonica* rice, *indica* rice, and maize populations (Supplemental Figure 5). There were 16, 11, and 9 SNPs in *indica* rice, *japonica* rice, and maize, respectively, that were significant after Bonferroni correction (Supplemental Table 4). We identified 19 rice and maize orthologs of known telomere-regulating genes (see Supplemental Table 5 for ortholog list) and compared their genomic positions to the GWAS-significant SNP markers; none of the significant SNPs were in close proximity to these telomere-regulating genes (in *japonica* rice, the closest GWAS-significant SNP was on chromosome 12 position 11,760,516 and near the gene *TERT* [< 395 kbp]; in maize, the closest GWAS-significant SNP was on chromosome 8 position 166,932,200 near the gene paralogs *RPL5A/RPL5B* [< 3.2 Mbp]).

No overlap of GWAS peaks for telomere length and flowering time

We examined whether telomere-regulating genes were in fact previously unrecognized flowering time QTLs, and vice versa. Using both AraTha_{Kmer} and AraTha_{TRF} individuals, we conducted GWAS of flowering time and compared the results to our GWAS results for telomere copy number (see Supplemental Tables 1 and 6 for significant SNPs from each GWAS analysis). We demarcated a 100-kb window centered on each significant telomere length SNP and examined

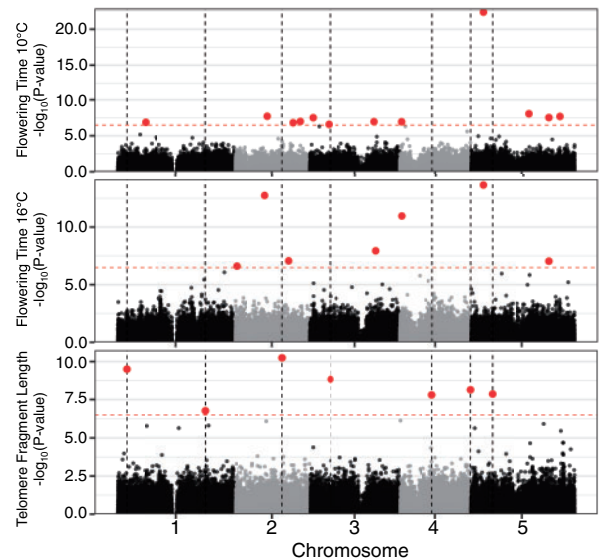


Figure 7 Genetic architecture underlying flowering time and telomere variation. GWAS results using the AraTha_{TRF} set for flowering time at 10°C (top), flowering time at 16°C (middle), and terminal fragment length (bottom). Red dotted line indicates the Bonferroni-corrected significance threshold ($\alpha = 0.05$). Dotted vertical lines are positions of the GWAS-significant SNPs for telomere fragment length. Red points indicate GWAS-significant SNPs.

whether any significant flowering time SNP was found within this window. No GWAS-significant SNPs were directly overlapping between the two traits (Figure 7 and see Supplemental Figure 6 for AraTha_{Kmer} set results). There was, however, a GWAS-significant SNP for telomere copy number on chromosome 5 position 15,389,625 in the AraTha_{Kmer} set and a GWAS-significant SNP for flowering time on chromosome 5 position 15,322,950 in the AraTha_{TRF} set, which are ~ 67 kb apart.

We also examined the genetic architecture underlying flowering time variation in rice and maize. Like in *A. thaliana*, we did not find any directly overlapping GWAS-significant SNP positions for telomere length and flowering time variation (see Supplemental Tables 4 and 7). The distances between the closest GWAS-significant SNP for telomere length and flowering time were ~ 434.2 kb for rice and ~ 26 Mb for maize.

Discussion

The links between telomere length and organismal life-history traits are tantalizing, especially since telomeres are linked to cellular senescence, aging, and disease in humans. Despite its central role in chromosomal stability, the drivers of telomere length variation and their phenotypic consequences remain unclear. This is particularly relevant for plants, where telomere length variation is not easily connected to aging and senescence, as observed in animals (Watson and Riha, 2011). In our analysis, we found that natural telomere length variation in three annual plant species is related to flowering time, one of the most crucial life-

history traits of plants. We found that in *Arabidopsis*, rice, and maize, individuals that had longer telomeres flowered earlier; finding this correlation in three distinct species from both dicots and monocots suggests that this relationship may be widespread. Indeed, we also observed a negative correlation between flowering time and telomere length in the polyploid plant rapeseed (*Brassica napus*), although this relationship was not significant in this species.

It should be noted that only a small amount of variation in flowering time is explained by telomere length in the three plant species. Telomere length accounts for approximately 0.5%–2.4% of flowering time variation in *Arabidopsis* and 3%–6% in maize and rice. The low fraction of variation in life-history traits explained by telomere length is perhaps not surprising: flowering time, for example, is a complex quantitative trait whose variation is largely explained by numerous major genes in the flowering time pathway (Salomé et al., 2011). What remains intriguing is that, despite the complex genetic architecture underlying flowering time variation, the effects of telomere length on this life-history trait were still observed, even after correcting for population structure accounting for local genetic differences. Moreover, this level of correlation seems to still be sufficiently large that it can be acted upon by natural selection, providing a key evolutionary link between telomeres and plant ecology and life history.

Moreover, while the levels of flowering time variation explained by telomere lengths may be low, similar levels are also observed for correlations of telomere length with life-span traits in animals. For example, the variation explained between telomere length and mortality traits in dogs ranges from 0.62% to 8.8% (Fick et al., 2012). In humans, three studies showed that the percentage of telomere length variation that can be explained by age ranges from 6.8% to 16% (Njajou et al., 2007; Fitzpatrick et al., 2007; Weischer et al., 2014), although one early study had it as high as 50% (Slagboom et al., 1994); the latter appears to be an outlier. These results indicate that in both plants and animals, the amounts of variation in life-history traits explained by telomere length (or vice versa) may be low but are nevertheless significant.

The precise molecular mechanisms behind the correlation between telomere length and flowering time remain unknown, but we can suggest several possibilities. First, genes that control telomere length may have pleiotropic effects and also affect flowering time (or vice versa). Our analysis did not find any overlap between significant GWAS peaks for these two traits, but this does not rule out the possibility of multiple pleiotropic genes with small effects. Second, early flowering and rapid reproduction may somehow affect chromosome stability (precisely how is unclear), and so longer telomeres are adaptive in early flowering lines. A third possibility is that longer telomeres may be adaptive due to some other selection pressure, but given the energy expenditure for telomere maintenance, this may result in a trade-off that leads to early flowering. Finally, perhaps greater telomerase

activity is adaptive in plants with rapid developmental rates and early flowering, and the longer telomeres in early flowering lines are merely an indication of higher levels of telomerase activity. In this latter case, it should be noted that in maize, for example, faster rates of cell differentiation in the shoot apical meristem are observed in plants with earlier flowering times (Bilinski et al., 2018; Leiboff et al., 2015), and telomerase is most active in differentiating tissues such as the meristem (Fitzgerald et al., 1996; Riha et al., 1998). All of these theories will need to be experimentally tested in the future.

In support of the hypothesis that the negative correlation between telomere length and flowering time may indeed be driven by adaptive evolution, we should note that there is evidence for a selective sweep at the *A. thaliana* *TERT* genomic region. Adaptation may also explain the significant latitudinal cline of telomere length variation in *Arabidopsis*, and we also found that longer telomeres are associated with other aspects of the spring cycling life-history strategy of this ruderal species, such as germination in response to cold. Moreover, longer telomeres are found in temperate-adapted varieties of rice (temperate *japonica*) and maize (non-stiff-stalk and stiff-stalk maize), which also flower significantly earlier in their growing seasons compared to TS varieties.

While we can advance these different hypotheses for the observed correlations, other explanations are possible. More work is required to understand why such correlations exist in plants. What is apparent is that the relationship of telomere length to life history is different in plants compared to other organisms. In mammals and other eukaryotic systems, shorter telomeres are generally associated with aging, in part because of the association of telomere shortening with cellular senescence (Urquidi et al., 2000). That clearly is not the case with the annual plant species we have studied. It has also been suggested that shorter telomeres are found in organisms with a faster “pace of life,” since the investment in somatic maintenance in such organisms is thought to be reduced to save energy for reproduction (Giraudeau et al., 2019). Again, our results suggest that this does not appear to be the case for these annual plants, as early flowering presumably due to faster developmental rates is associated with longer (and not shorter) telomeres.

Interestingly, the genetic architecture of telomere length variation is distinct in the three species we analyzed. In maize, an early QTL study examined telomere length variation and identified three loci associated with this trait (Burr et al., 1992), only one of which (a QTL on chromosome 4) may overlap with one of our nine GWAS hits. Natural telomere length variation has been more extensively studied in the model plant *A. thaliana*, and our GWAS found that three of our significant SNPs overlap with two previously identified telomere length QTL regions (Fulcher et al., 2015). Interestingly, genes previously associated with telomere set point regulation in *Arabidopsis*, including *NOP2A*, *RPL5A*, and *RPL5B* (Abdulkin et al., 2019), did not appear in our

GWAS, perhaps due to differences in the mapping populations employed or their allele frequency in the population.

Our GWAS mapping however, does span a key telomere-regulating gene in *A. thaliana*: the telomerase gene *TERT*. The *A. thaliana TERT* gene is involved in telomere elongation (Riha et al., 2001), and this locus also overlapped with a large QTL region for telomere length identified from a previous recombinant inbred line mapping study (Fulcher et al., 2015). *TERT* has also been identified in a human GWAS mapping study showing an association with leukocyte telomere length variation (Codd et al., 2013), pointing to cross-kingdom functional conservation. Determining whether it is indeed genetic variation at *TERT* that controls natural differences in telomere length variation in *A. thaliana* awaits future fine-mapping and molecular genetic investigations.

The lack of overlap between GWAS-significant SNPs across the three plant species suggests a divergence in the genetic architecture underlying telomere length regulation. However, there are caveats to our results. FarmCPU is computationally intensive. To reduce the computational burden in analyzing a large number of SNPs, we used LD pruned dataset for the GWAS. While this was suitable for *A. thaliana* and its small genome, for rice and maize, the LD pruning may have resulted in a coarse genome-wide landscape of the variation. However, it is worth noting that in maize, even with a reduced representation SNP set, the FarmCPU algorithm was able to detect SNPs in known flowering time-regulating genes, albeit by using a sample size that is 10-fold greater than our study (Liu et al., 2016). Hence, perhaps at least for our maize GWAS results, the analysis was underpowered due to the low sample size.

Our study complements recent work in identifying the effects of genome size and structure on life-history traits such as flowering time (Meagher and Vassiliadis, 2005). In maize, for example, genome size is positively correlated with flowering time (Jian et al., 2017), and changes in repetitive DNA sequences are associated with altitudinal adaptation (Bilinski et al., 2018). The negative correlation between genome size, repetitive DNA content, and cellular growth rate has been advanced as a plausible explanation for this phenomenon (Tenaillon et al., 2016; Bilinski et al., 2018). These studies, as well as our results on telomere variation, suggest that a variation in life-history strategy can indirectly influence chromosome and genome structure via selection. This opens up future areas of inquiry, including determining how widespread this phenomenon is, the relationship between telomere length and cell differentiation rate in plants, details of any selective advantage of telomere length in plant species with different life histories, and the precise molecular genetic mechanisms underlying telomere length polymorphisms in plant species.

Materials and methods

Plant materials and growth conditions

Seeds for the set of *A. thaliana* genotypes from the 1,001 Genome Project were purchased from ABRC (CS78942). Additional lines were obtained from the laboratory

collection of TEJ. Seeds were sown into a mixture of three parts Promix BX mycorrhizae soil, one part Profile Field and Fairway calcined clay, and one part Turface medium stabilizer. Plants were grown in a greenhouse at UT Austin under 16/8-h light/dark (150 μ mol light levels using Phantom Pro Double-ended 1,000-watt high-pressure sodium bulbs.), 21/18°C (day/night) conditions. Plant tissue for analysis was collected at the 5-week stage.

TRF analysis of telomere length analysis

Genomic DNA was extracted from individual whole plants ($n = 1-3$) and digested with the restriction enzyme Tru1I (Fermentas, Hanover, MD, USA) as previously described (Fitzgerald et al., 1999). [32 P] 5'-end-labeled or 5'-DIG-(T₃AG₃)₄ oligonucleotides were used as probes (Nigmatullina et al., 2016; Abdulkina et al., 2019). Radioactive signals were scanned with a Pharos FX Plus Molecular Imager (Bio-Rad), and nonradioactive signals were scanned with a GBox-F3 Imager (Syngene). Images were visualized with Quantity One v.4.6.5 software (Bio-Rad), and mean telomere length values (mean TRF used for GWAS) were calculated using the TeloTool program (Göhring et al., 2014), as described previously (Abdulkina et al., 2019). Overall, we measured telomere length in 424 *A. thaliana* accessions. Since our TRF method is identical to the one used in Fulcher et al. (2015), we added data for 229 accessions analyzed in their study to obtain our final dataset of 653 genotypes.

Analyzed genome sequences

We obtained the whole-genome re-sequencing data for *A. thaliana*, *O. sativa*, and *Z. mays* from previous published studies.

Arabidopsis thaliana: Genome sequences were obtained from the 1001 *A. thaliana* Genome Consortium (Alonso-Blanco et al., 2016) and are available at the NCBI SRA SRP056687. We grouped the 1,135 samples with the same genome sequencing protocols and analyzed the two most highly represented groups. This included the first group (designated as AraTha_{K_{mer}}) with 483 individuals prepared using leaf tissue and sequenced with 2 \times 100-bp read length on the Illumina HiSeq 2000 platform. The second group consists of 201 individuals prepared using leaf tissue and sequenced as 2 \times 101-bp read length on the Illumina HiSeq 2000 platform.

Oryza sativa: Genome sequencing data from Wang et al. (2018) were obtained at NCBI SRA PRJEB6180. All 3,000 samples were prepared from leaf tissue and sequenced as 2 \times 83 bps using Illumina HiSeq 2000. Only samples with greater than 5 \times genome coverage were used.

Zea mays: We analyzed the “Buckler-Goodman 282” panel of Flint-Garcia et al. (2005), which captures the genetic diversity of maize. We analyzed the most recent sequencing batch that re-sequenced the panel to a higher depth using 2 \times 150 bp on the Illumina HiSeq 10X platform (Bukowski et al., 2018). The data were obtained from the NCBI SRA PRJNA389800.

Identifying and quantifying tandem repeats

Sequencing reads were subjected to quality control using BBTools (<https://jgi.doe.gov/data-and-tools/bbtools/>). We used the `bbduk.sh` script version 37.66 using parameters `minlen = 25 qtrim=r trimq = 10 ktrim=r k = 25 mink = 11 hdist = 1 tpe tbo` to trim sequencing adapters and low-quality sequences.

We used k-Seek to quantify k-mers and to identify the total copy number for a tandemly repeating k-mer in a sample of an unmapped genome sequencing library (Wei et al., 2014, 2018). k-Seek requires the minimum repeating length to be 50 bp to avoid counting small microsatellites that are scattered across the genome. We note that because k-Seek is optimized to analyze “short” Illumina sequencing reads (< 150 bp), we will inevitably miss the characterization of large k-mer sequence-based satellite repeats (i.e. centromeric or ribosomal DNA repeats).

PCR-based library preparation is known to have a bias in underrepresenting high and low GC regions of the genome (Benjamini and Speed, 2012). To account for this bias, we implemented the method of Flynn et al., (2017) to correct for differences in GC content. Reads were mapped against the reference genome to first calculate the mean insert size using `bamPEFragmentSize` from `deeptools` version 3.3.0 package (Ramírez et al., 2016). The insert size was used to calculate the GC content of a given position in the genome, which was defined as the proportion of G or C bases of a given position plus the downstream fragment length of the library (Benjamini and Speed, 2012). The alignment was then used to calculate the average coverage of each GC content. We used `bwa-mem` version 0.7.16a-r1181 (Li, 2013) with default parameters to align paired end reads to the reference genome. The average coverage per GC content was then used to calculate the correction factor of Benjamini and Speed (2012) and applied to k-mer counts. We used scripts from Flynn et al., (2017; https://github.com/jmf422/Daphnia-MA-lines/tree/master/GC_correction) that implement the entire process.

For *A. thaliana* re-sequencing data, we used the reference genome TAIR10 from The Arabidopsis Information Resource. The genome sequences for *O. sativa* and *Z. mays*, however, were not ideal for implementing the GC content correction method. For *O. sativa*, samples were sequenced across multiple runs, suggesting that any differences in the sequencing run should also be implemented in the correction. For *Z. mays*, the genome coverage was relatively low (on average $\sim 5\times$), indicating that a coverage-based method of correction would not be ideal. Hence, for these two species, we only analyzed the telomere repeat, and for each sample, its telomere count was divided by average genome-wide coverage to account for differences in sequencing coverage between samples. The per-sample average coverage was obtained from Supplementary Data 2 of Wang et al., (2018) for *O. sativa* and was calculated using `bedtools` version 2.25.0 (Quinlan and Hall, 2010) `genomecov` program for *Z. mays*.

Genome-wide association study

For *A. thaliana*, the population VCF file was downloaded from the 1001 Genomes Project website (<https://1001genomes.org/>). For *O. sativa*, the population VCF was downloaded from the 3,000 Rice Genome Projects' `snp-seek` website (<https://snp-seek.irri.org/>; Mansueto et al., 2017). For *Z. mays*, the population VCF was downloaded from the Gigascience database (<http://dx.doi.org/10.5524/100339>; Bukowski et al., 2018).

The SNP files were initially filtered to exclude polymorphic sites that had > 10% of the individuals with a missing genotype and sites with < 5% minor allele frequency. The VCF files were converted to PLINK format using `vcftools` version 0.1.15 (Danecek et al., 2011), and the program `plink` ver. 1.9 (Chang et al., 2015) was used with the parameter `-indep-pairwise 100 5 0.5`, which scans the file in 100 variant count windows while shifting the window in 5 variants and LD pruning pairs of variants that have $r^2 > 0.5$. In the end, we used 173,688 SNPs for `AraThakmer` set, 153,845 SNPs `AraThaTRF` set, 585,026 SNPs for *indica* rice, 370,052 SNPs for *japonica* rice, and 3,048,120 SNPs in maize.

The LD-pruned PLINK file was converted to HAPMAP format for the GWAS using `GAPIT` version 2 (Tang et al., 2016). We took the \log_{10} of the telomere lengths (corresponding to telomere repeat copy number from k-Seek-based estimate or the telomere fragment length from the TRF-based estimate) to normalize the distribution. For GWAS mapping, we used `FarmCPU` (Liu et al., 2016), which is a mixed linear model (MLM) incorporating population structure and kinship but is robust to false-positive and false-negative associations compared to other MLM GWAS algorithms. We used four principal components to model the underlying population structure.

Orthologs of *A. thaliana* telomere-regulating genes were found in the rice and maize gene annotation using `Orthofinder` ver 2.3.12 (Emms and Kelly, 2019, 2015).

Selective sweep analysis of *A. thaliana*

Evidence of selective sweep was examined using the `iHS` (Voight et al., 2006), `Rsb` (Tang et al., 2007) and `OmegaPlus` methods (Alachiotis et al., 2012) with SNPs extracted from chromosome 5. The SNP files that were filtered to exclude polymorphic sites that had > 10% of the individuals with a missing genotype, and sites with < 5% minor allele frequency were used. Missing genotype imputation and SNP phasing were conducted with `Beagle` version 4.1 (Browning and Browning, 2016).

To calculate the `iHS` and `Rsb` statistics, we used the filtered and phased VCF file with the R program (`R Core Team, 2016`) `rehh` package (Gautier and Vitalis, 2012). `OmegaPlus` statistics were calculated with `OmegaPlus` version 3.0.3 with `-grid 2697` so that each grid would correspond to roughly 10,000 bp, and additional parameters `-minwin 5000 -maxwin 3,000,000 -no-singletons`. A bootstrap procedure was conducted to determine the significance of the `OmegaPlus` statistics. We randomly sampled individuals matching the sample size of the individuals carrying the

minor allele at the GWAS-significant region on chromosome 5. OmegaPlus statistics were then calculated for this randomized group, and this procedure was repeated 200 times to generate a bootstrap distribution.

Plant phenotype analysis

For *A. thaliana*, various developmental traits were obtained from Arapheno (<https://arapheno.1001genomes.org/>; Seren et al., 2017) with phenotype names FT10 (days to flowering at 10°C), FT16 (days to flowering at 16°C), CL (cauline axillary branch number), RL (leaf number), length (stem length), RBN (primary branch number), and diameter (flower diameter). Genome size and ribosomal DNA size estimates were taken from the github repository (<https://github.com/Gregor-Mendel-Institute/swedish-genomes/tree/master/files>) for the original study (Long et al., 2013).

For rice, we obtained flowering time data measured as part of the 3000 Rice Genome Project (Sanciangco et al., 2018; Wang et al., 2018), which was measured as number of days at which 80% of the plants were fully headed (code HDG_80HEAD). The data are available from <https://doi.org/10.7910/DVN/HGRSJG>. For maize, we obtained phenotype data from the Buckler-Goodman association panel. The phenotype file (traitMatrix_maize282NAM_v15-130212.txt) was downloaded from Panzea (<https://www.panzea.org/phenotypes>), and we only analyzed the days to silk trait (code GDDDaystoSilk).

For each plant genotype, the phenotypes we obtained were a single representative value that was a product of summarizing the phenotype values from multiple replicates. The replicate information, however, was not readily available; hence, we considered each phenotype value to be an overall representation of the genotype.

Association between the telomere length and plant phenotypes were conducted in R. Telomere lengths were \log_{10} -converted before the linear regression modeling. The multiple linear regression analysis was conducted using the `lm` function, and population structure information was obtained from the four principal components that were used in the GWAS analysis.

Accession numbers

Sequence data from this article can be found in the GenBank/EMBL libraries under the following accession numbers: PRJNA273563 for *A. thaliana*, PRJNA389800 for *Z. mays*, and PRJEB6180 for *O. sativa*. The genome-wide K-mer abundance estimation and SNP file used for GWAS for *A. thaliana*, rice, and maize are available from Zenodo data repository (<https://doi.org/10.5281/zenodo.4295944>).

Supplemental data

The following materials are available in the online version of this article.

Supplemental Figure 1. *Arabidopsis thaliana* accessions display different telomere length set points.

Supplemental Figure 2. Scatter plots of *A. thaliana* phenotypes including cauline leaf number, diameter of rosette,

rosette branch number, and length of flowering stem versus telomere copy number or terminal fragment length.

Supplemental Figure 3. Scatter plots of *A. thaliana* genome size and ribosomal DNA copy number versus terminal fragment length.

Supplemental Figure 4. Genome-wide ω statistic from 200 bootstrap groups.

Supplemental Figure 5. GWAS results for telomere repeat copy number in rice and maize.

Supplemental Figure 6. GWAS results for telomere copy number and days to flowering in *A. thaliana*.

Supplemental Table 1. GWAS-significant SNPs for telomere repeat copy number in the AraTha_{Kmer} set or telomere length in the AraTha_{TRF} set.

Supplemental Table 2. Top 10 most abundant k-mer repeats and association with the telomere repeat or flowering time in *A. thaliana*.

Supplemental Table 3. Correlation between maize telomere repeat copy number and days to silk measured in 9 different environments.

Supplemental Table 4. GWAS-significant SNPs for telomere repeat copy number in rice and maize.

Supplemental Table 5. Orthologs of *A. thaliana* telomere-regulating genes in rice and maize.

Supplemental Table 6. GWAS-significant SNPs for days to flowering in the AraTha_{Kmer} set and the AraTha_{TRF} set in *A. thaliana*.

Supplemental Table 7. GWAS of significant SNPs for days to flowering in rice and maize.

Supplemental Data Set 1. Telomere repeat copy numbers for the AraTha_{Kmer} set and the sequencing set with 101-bp read lengths.

Supplemental Data Set 2. Terminal fragment length for the AraTha_{TRF} set.

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