Depressive Symptoms Predict Change in Telomere Length and Mitochondrial DNA Copy Number Across Adolescence

Kathryn L. Humphreys, PhD, Lucinda M. Sisk, BA, Erika M. Manczak, PhD, Jue Lin, PhD, Ian H. Gotlib, PhD

Objective: Several studies have found associations between a diagnosis or symptoms of major depressive disorder and markers of cellular aging and dysfunction. These investigations, however, are predominantly cross-sectional and focus on adults. In the present study, we used a prospective longitudinal design to test the cross-sectional association between depressive symptoms in adolescents and telomere length (TL) as well as mitochondrial DNA copy number (mtDNA-cn).

Method: A total of 121 adolescents (mean age = 11.38 years, SD = 1.03; 39% male adolescents and 61% female adolescents) were followed for approximately 2 years. At baseline and follow-up, participants provided saliva for DNA extraction, from which measures of TL and mtDNA-cn were obtained. Depressive symptoms were obtained via the Children’s Depression Inventory.

Results: There was no association between depressive symptoms and markers of cellular aging at baseline; however, depressive symptoms at baseline predicted higher rates of telomere erosion ($\beta = -0.201, p = .016$) and greater increases in mtDNA-cn ($\beta = 0.190, p = .012$) over the follow-up period. Markers of cellular aging at baseline did not predict subsequent changes in depressive symptoms. Furthermore, including the number of stressful life events did not alter these patterns of findings.

Conclusion: These results indicate that depressive symptoms precede changes in cellular aging and dysfunction, rather than the reverse.

Key words: depression, adolescence, cellular aging, telomeres, mitochondrial DNA copy number

learn about the nature of the relationship between depression and TL.

Whereas TL has received considerable attention in relation to MDD, we know much less about another putative biological marker of cellular dysfunction: mitochondrial DNA copy number (mtDNA-cn). Mitochondria are organelles responsible for cellular energy production that contain multiple copies of their own, free-floating DNA. As previously noted, increased oxidative stress has been shown to be associated with aberrations in mtDNA-cn; the mechanism proposed to underlie this relation is an adaptive response to stress in which biogenesis of mitochondria increases. Further, TL and mtDNA-cn are considered markers of cellular aging; mice exposed to stress exhibited greater mtDNA-cn and decreased TL relative to nonstressed, age-matched control mice. Investigators have posited that processes involved in TL and mtDNA are co-regulated; unfortunately, few studies have assessed both mtDNA and TL within the same sample. Moreover, the directional nature of the association between depressive symptoms and cellular dysfunction is not clear; it has been posited that mitochondrial dysfunction may exacerbate or even play a causal role in depression. Although some studies with adults have found no relation between mtDNA-cn and depression, others have found that depression is characterized by reduced mtDNA counts, whereas still other investigations have found that depression is associated with increased mtDNA counts. Moreover, TL and mtDNA may not be linked in their changes related to depression. For example, one study found that high levels of depressive symptomatology were associated with decreased TL over 10 years, but not with changes in mtDNA.

The vast majority of studies in this area have focused on the relation between depression and cellular aging in adults. Given that the median age of MDD onset is in young adulthood, studying longitudinal patterns prior to most individuals’ first MDD episode reduces the likelihood of having associations influenced by recurrent MDD. In the present study, we examined the relation between depressive symptomatology and mtDNA-cn and TL in adolescents. We also examined fluctuations in levels of TL and mtDNA-cn over time, and tested whether changes in these markers of cellular aging predicted severity of depressive symptoms. Finally, given that TL has been found to be adversely affected by stress-responsive systems such as hypothalamic–pituitary–adrenal (HPA) axis activity, increased oxidative stress, increased inflammation, and a dysregulated autonomic nervous system, we covaried for levels of the number of stressful life events experienced by participants, testing whether stress exposure would explain levels of both depressive symptoms and cellular aging.

**METHOD**

**Participants**

We used flyers and local media to recruit individuals between 9 and 13 years of age on the basis of having experienced a range of early-life adversity (to avoid having a very low-stress community sample). The mean age of the sample in the present study at baseline (wave 1) was 11.38 years (SD = 1.03); 39% of participants were male, and 54% identified as white. We recruited only participants who were eligible to complete a neuroimaging scan (neuroimaging data are not included in this report). The study was approved by the Stanford University Institutional Review Board; participants and their parents gave assent and informed consent, respectively.

Participants were screened for initial inclusion/exclusion criteria through a telephone interview; potentially eligible individuals were then invited to the laboratory for in-person interviews and assessments. Inclusion criteria were that participants be between 9 and 13 years of age and proficient in English. Exclusion criteria were factors that would preclude a neuroimaging scan (eg, metal implants, dental braces), a history of major neurological or medical illness, severe learning disabilities that would make it difficult for participants to understand the study procedures, and, for female participants, the onset of menses. Girls and boys were selected to not differ in pubertal stage; consequently, boys were older than girls. Pubertal status was assessed via self-report Tanner staging.

Participants returned to the laboratory approximately 2 years after the initial assessment (wave 2; mean = 1.97 years, SD = 0.34). At wave 1, a total of 211 participants had complete data with both depressive symptom scores and saliva; at the time that we gave samples to the assaying laboratory at wave 2, a total of 122 participants had data with both depressive symptom scores and saliva data (of the original 211 participants, 43 did not return to the study after completing wave 1, and 48 contributed samples after the cutoff date for these analyses). In the present study, participants were included if they contributed saliva samples that yielded usable data across both time points, resulting in a final sample size of 121 participants (47 boys and 74 girls) with both depressive symptoms and saliva samples at both time points.

**Procedure**

At waves 1 and 2, participants attended laboratory sessions with a caregiver, during which they completed questionnaires,
provided saliva samples, and completed other assessments not included in the present study (eg, neuroimaging scans). Participants were compensated for their time.

**Measures**

*Children’s Depression Inventory–Short Form.* Participants reported on their symptoms of depression using the 10-item CDI Short Form (CDI-S), which was developed to assess depressive symptomatology in children and adolescents aged 8 to 17 years. The original CDI and the CDI-S yield comparable results. Responses to the CDI-S were scored on a 3-point scale, from 0 (occurs once in a while) to 2 (occurs frequently); we summed the 10 items to compute a total score. In this sample, the internal consistency of the CDI-S was acceptable ($\alpha = 0.74$).

*Stressful Life Events.* At the initial assessment, participants were interviewed about their lifetime exposure to 30 types of stressful events, using a modified version of the Traumatic Events Screening Inventory for Children (TESI-C). (For more information about our protocol, please see King et al. and Humphreys et al.) Only stressors rated by the panel as being above a “mild stressor” were included. For the purposes of the present analyses, to compute life stress we summed the number of stressful events reported in the participant’s life from birth to the time of the baseline assessment.

**Cellular Markers**

Genomic DNA was purified from 500 μl of saliva collected in the Oragene DNA Kit (DNA Genotek, Kanata, ON, Canada) with the DNA Agencourt DNA Advance Kit (catalog no. A48705; Beckman Coulter Genomics, Brea, CA) according to the manufacturer’s instructions. DNA was quantified by Quant-iT PicoGreen dsDNA Assay Kit (catalog no. P7589; Life Technologies, Grand Island, NY) and run on 0.8% agarose gels to check the integrity. DNA samples were stored at −80 °C. Parameters for the assaying of TL and mtDNA are presented in Supplement 1, available online.

**Data Analysis**

Cellular aging variables were examined for normality and outliers. One TL value was beyond 3 SD from the mean and was winsorized to the next closest value; no further transformations were conducted. We examined the association between depression scores and markers of cellular aging (ie, TL and mtDNA-cn, examined independently) both cross-sectionally and longitudinally, covarying baseline age, pubertal stage, and sex. Given the potential for number of stressful life events to be relevant to models, this variable was included as an additional covariate following the initial model. In addition, for the longitudinal models, we included the additional covariates of baseline cellular aging and the length of the interval between the baseline and follow-up assessments. We computed a slope to examine change in cellular aging markers as follows: ($\text{Wave 2 − Wave 1 values}$/$\text{Wave 2 − Wave 1 age}$). We used hierarchical linear ordinal least-squares regression to examine the association between depression scores, treated linearly, and markers of cellular aging, over and above the effects of the covariates. In addition, we used the general linear model to confirm effects using a group-based approach to conceptualizing depression, with a cutoff score of ≥3 on the CDI-S used as a screening cutoff for elevated depression likelihood. Effect sizes are reported below.

We additionally performed Benjamini–Hochberg multiple comparisons correction at a false discovery rate of 5% for all primary models. Details are provided in Table S1 and Supplement 1, available online.

**RESULTS**

**Participant Characteristics**

Demographic and clinical characteristics of the sample are presented in Table 1.

**Cross-Sectional Associations Between Depression and Cellular Aging**

We first examined whether depression scores predicted TL assessed at the same wave, controlling for age, pubertal stage, and sex. The full model was not significant ($F_{4,118} = 1.37, p = .250, R^2 = 0.044$): depression scores did not predict variance in TL over and above the effect of the covariates ($\hat{\beta} = -0.011, r_{118} = -0.12, p = .903, \Delta R^2 < 0.001$). Similar results were obtained for mtDNA-cn. Again, the full model was not significant ($F_{4,118} = 1.55, p = .191, R^2 = 0.050$): depression scores did not predict variance in mtDNA-cn over and above the effect of the covariates ($\hat{\beta} = -0.067, r_{118} = -0.73, p = .464, \Delta R^2 = 0.004$). The inclusion of stressful life events in the model did not change the patterns of results, and in neither model did stressful life events significantly predict TL or mtDNA-cn.

**Longitudinal Associations Between Depression and Cellular Aging**

Next, we examined whether depression scores at wave 1 predicted change in markers of cellular aging over a 2-year period. The full model was significant ($F_{6,113} = 6.54, p < .001, R^2 = 0.26$): depression scores predicted significant variance in change in TL over and above the effect of sex, pubertal stage, age at baseline, length of follow-up, and...
in Figure 1B, higher depression scores predicted greater increases in mtDNA-cn over adolescence. TL measures were correlated significantly at the 2 waves ($r_{118} = 0.60, p < .001$), but measures of mtDNA-cn were not ($r_{119} = 0.17, p = .064$).

We also used a group-based approach to examine the associations between depression and change in cellular aging, comparing participants who reported scores on the CDI-S of 3 or above (n = 40) with those who reported scores of 2 or below (n = 81). In an analysis of covariance with baseline age, pubertal stage, sex, number of stressful life events, follow-up duration, and baseline TL levels as covariates, depression group was significantly associated with TL slope ($F_{1,113} = 4.68, p = .033$, partial $\eta^2 = 0.04$). Children with scores above the depression cutoff had significantly greater telomere erosion than did children with scores below the cutoff (mean = −0.007, SE = 0.001, versus mean = −0.003, SE = 0.001; Cohen’s $d = −0.40$). Similarly, in the analysis of covariance conducted on wave 2 mtDNA-cn with baseline age, pubertal stage, sex, number of stressful life events, follow-up duration, and baseline mtDNA-cn as covariates, depression group was significantly associated with mtDNA-cn slope ($F_{1,114} = 5.84, p = .017$, partial $\eta^2 = 0.05$). Participants with scores above the depression cutoff had significantly greater change in mtDNA-cn than did those with scores below the cutoff (mean = 6.79, SE = 1.56, versus mean = 2.10, SE = 1.07; $d = 0.41$).

Next, we tested a reverse directionality model to examine whether baseline TL and mtDNA-cn predicted change in depressive symptoms over and above sex, pubertal stage, age at baseline, length of follow-up, and baseline depression scores. Neither TL ($\bar{\beta} = 0.008, t_{116} = 1.08$, $p = .281$, $\Delta R^2 = 0.01$) nor mtDNA-cn ($\bar{\beta} = −0.060, t_{116} = −0.73$, $p = .467$, $\Delta R^2 = 0.003$) at baseline significantly predicted change in depressive symptoms over the course of the follow-up period. Including stressful life events did not change the pattern of associations.

Finally, we examined whether changes in depressive symptoms over the period were associated with changes in TL and mtDNA-cn. After covarying for sex, pubertal stage, age at baseline, length of follow-up, baseline depression scores, and baseline cellular aging, change in depressive symptoms was not a significant predictor for either telomere erosion ($\bar{\beta} = −0.035, t_{112} = −0.36$, $p = .723$, $\Delta R^2 = 0.001$) or mtDNA-cn change ($\bar{\beta} = 0.152, t_{113} = 1.74$, $p = .084$, $\Delta R^2 = 0.002$).

**DISCUSSION**

In this study, we examined the prospective association between depression and changes in TL and mtDNA-cn in a
sample of 121 children and young adolescents. Perhaps not surprising given the mixed evidence regarding the associations between depression and cellular aging, we found that depressive symptoms were not associated cross-sectionally with either marker of cellular aging. There were, however, prospective associations between depressive symptomatology at baseline and both telomere erosion and increases in mtDNA-cn. In contrast, there was no association between baseline TL or mtDNA-cn and changes in depressive symptoms. These findings indicate that depressive symptoms may be causally related to changes in cellular aging. Although findings of prior research on cellular aging and depression have been mixed, meta-analyses have yielded a significant correlation between TL and depression, although these studies are primarily cross-sectional and thus have not examined the temporal nature of this association. Our findings examining the prospective association between depressive symptomatology and change in cellular aging are consistent with previous findings that the duration of participants’ longest episode of depression predicted mtDNA count, and that shorter TL was significantly associated with earlier age of onset of individuals’ most severe depressive episode.

Gotlib et al. found that healthy girls with a maternal history of depression had shorter TL than did their age-matched peers, suggesting that shorter TL portends depression. It is important to note, however, that this study was cross-sectional and operationalized “nondepressed” as not meeting diagnostic criteria for MDD. Given the present findings in an independent sample of young boys and girls, it appears that examining elevated levels of depressive symptoms is important in gaining a more comprehensive understanding of the association between depression and TL, as subclinical variations in depressive symptoms predict changes in cellular aging.

Previous studies have reported both higher and lower mtDNA-cn counts associated with depression. Far fewer studies have examined the relation between mtDNA-cn and psychopathology than is the case with TL. mtDNA-cn was positively correlated with anxiety, but not with depression, in a sample of individuals aged 13 to 17 years, suggesting specificity in the association with mtDNA-cn. Our findings that there was no cross-sectional association between depressive symptoms and mtDNA-cn adds to the growing number of studies examining this association, and adds a prospective lens that supports the formulation that depression symptoms precede changes in mtDNA-cn.

Differences between our findings and those of previous investigations may be due in part to study design, source of the DNA, methods for assessing depression, and age of our participants. For example, we used a prospective longitudinal design to examine the temporal nature of the relation between depressive symptoms and changes in cellular aging, rather than assessing simpler cross-sectional associations between these constructs. We also examined TL and mtDNA-cn from DNA extracted from saliva, and it is possible that cell type influences these markers. We assessed depressive symptomatology on a continuum, given formulations that depression is not best represented as a taxon. Nevertheless, given evidence that diagnosed depression is associated with shorter TL, it will be important in future research to examine the issue of the relation between cellular aging and depression assessed as a binary versus a continuous construct.
Both stress and depression have been found in previous research to be associated with shortened TL; our analyses indicate that depressive symptoms predicted TL over and above the effects of stress, which was not a significant predictor in cross-sectional or longitudinal analyses. The interaction of stress and MDD is likely to be complex; indeed, in a previous study, significant associations between stress and shortened TL were attenuated by controlling for depression status of the participants.

Although documenting the temporal association is important for establishing the potential for causality, we are unable to provide insight into the mechanism of such effects. However, depression has been implicated in chronic activation of the sympathetic nervous system and in increased blood glucose levels. Chronically elevated glucose levels have been shown to damage mitochondrial function, producing effects such as decreased quality control in mtDNA replication and increased reactive oxygen species production. In turn, evidence from in vitro studies indicates that increased circulation of reactive oxygen species has been shown to shorten TL, and to increase mtDNA-cn by damaging existing mitochondria and triggering replication. Cross-species research is most likely to be able to inform our understanding of causal and mechanistic processes regulating the relation between cellular aging and depressive symptoms.

We should note two limitations of this study, in addition to the possibility of cell type influencing the generalizability to other studies. First, we focused on depressive symptoms assessed with the CDI-S. Examining symptoms of other disorders that are associated with TL and/or mtDNA-cn will be important in determining the specificity and boundary conditions of the present findings. Second, although we found no evidence that stressful life events predicted change in cellular aging markers, we cannot rule out the possibility that intervening stressful events played a role in leading to patterns reported in this study. Given evidence of stress generation in depression, this remains a possible pathway to explain the link between depression and aging processes.

As a related point, although this sample has been previously characterized on the basis of type, severity, and timing of stressful experiences, its generalizability is not known.

Despite these limitations, the present study is important in demonstrating that depressive symptoms predict the rate of cellular aging in children and young adolescents. It will be important in future work to examine factors that may buffer individuals from experiencing accelerated cellular aging. For instance, one study found that lifestyle factors such as frequency of exercise and social support networks, termed “multisystem resiliency,” were positively associated with longer telomeres, and, conversely, that shortened telomeres were associated with depression only in individuals with lower levels of multisystem resiliency. Environmental factors that have been associated with the onset of depression have also been found to moderate risk related to cellular aging. For example, high-risk children whose parents were high in parental responsiveness had shorter TL than did their high-risk peers whose parents were less responsive. If replicated, these findings suggest that preventing depression will have long-term health benefits through reducing rates of telomere erosion and increases in mtDNA-cn.

REFERENCES
HUMPHREYS et al.


SUPPLEMENT 1

Telomere Length

Telomere length (TL) measurement assay was adapted from the method originally published by Cawthon and by Lin et al.1,2 The cycling profile for T (telomeric) polymerase chain reaction (PCR) consisted of the following: denature at 96 °C for 1 minute, denature at 96 °C for 1 second, anneal/extend at 54 °C for 60 seconds, with fluorescence data collection, 30 cycles. The cycling profile for S (single-copy gene) PCR consisted of: denature at 96 °C for 1 minute, denature at 95 °C for 15 seconds, anneal at 58 °C for 1 second, extend at 72 °C for 20 seconds, 8 cycles, followed by denature at 96 °C for 1 second, anneal at 58 °C for 1 second, extend at 72 °C for 20 seconds, hold at 83 °C for 5 seconds with data collection, 35 cycles.

The primers for the telomere PCR were tel1b (5'-CGGTTTG(TTTTGG)5GTT-3'), used at a final concentration of 100 nmol/L, and tel2b (5'-GGCTTG(CCTTAC)5CCT-3'), used at a final concentration of 900 nmol/L. The primers for the single-copy gene (human β-globin) PCR were hbg1 (5'-GCTTCTGACACAAGCTGTGTTCACTAGC-3'), used at a final concentration of 300 nmol/L, and hbg2 (5'-CACCAACTCATCAGTTACC-3'), used at a final concentration of 700 nmol/L. The final reaction mix contained 20 mmol/L Tris-HCl (pH 8.4), 50 mmol/L KCl, 200 μmol/L each dNTP, 1% dimethyl sulfoxide (DMSO), 0.4 × Syber Green I, 22 ng Escherichia coli DNA per reaction, 0.4 U of Platinum Taq DNA polymerase (Life Technologies, Carlsbad, CA) per 11-μL reaction and 7 ng of genomic DNA. Tubes containing 26, 8.75, 2.9, 0.97, 0.324, and 0.108 ng of a reference DNA (pooled human genomic DNA from buffy coat, Roche Diagnostics catalog no. 1691112001) were included in each PCR run so that the quantity of targeted templates in each research sample could be determined relative to the reference DNA sample by the standard curve method. The same reference DNA was used for all PCR runs.

TL was assayed in three batches. To control for interassay variability, eight control DNA samples were included in each run. In each batch, the T/S ratio of each control DNA was divided by the average T/S for the same DNA from 10 runs to obtain a normalizing factor. This was carried out for all eight samples, and the average normalizing factor for all eight samples was used to correct the participant DNA samples to get the final T/S ratio. The T/S ratio for each sample was measured two times. When the duplicate T/S value and the initial value varied by more than 7%, the sample was run a third time and the 2 closest values were reported. Using this method, the interassay coefficients of variation (CVs) were 2.3% for the first batch, 2.1% for the second batch, and 2.2% for the third batch. Intra-assay CVs were not collected for individual batches.

Mitochondrial DNA

Detection of a 69-bp fragment of the ND1 gene in mtDNA (nucleotides 3485–3553) and an 87-bp fragment of RNase P (TaqMan Copy Number Reference Assay, human, RNase P, catalog no. 4403328, Life Technologies) by TaqMan assays was used to determine the relative copy number of mtDNA per diploid nuclear genome. The primer and probe sequences for ND1 were:

- ND1-forward 5'-CCCTAAAAACCGCCACATCT-3'
- ND1-reverse 5'-GAGCGATGGGTAGAGCTAAGG-3'
- ND1-FAM probe 5' FAM-CCATACCCCTCTCATCACCGGCC-TAMRA-3'

All primers were ordered from IDT. RNase P probe was VIC labeled. The reaction contained 12.5 ng of genomic DNA, 100 nmol/L of ND1 probe, 300 nmol/L of ND1-forward primer and ND1-reverse primer each, 1X RNase P copy number Reference Assay, 1X LightCycler 480 Probe Master (Roche, catalog no. 04902343001) in a 10-μL reaction. All samples were run in triplicate wells in 384-well plates in a Roche Lightcyler 480. PCR condition was PCR cycle: 95 °C 10 minutes, 45 cycles of 95 °C 10 seconds, 60 °C 30 seconds, 72 °C 1 second with data acquisition at 72 °C. Crossing point (CP) for each well was derived by the LightCycler 480 program using the second derivation method. Relative copy number was calculated by the following formula:

Relative mtDNA copy number/diploid genome = (POWER[2, (CPND1-CPRNaseP)]+2

mtDN-cn was assayed in two batches. Inter- and intra-assay CVs were not collected for these batches, but QC samples run at the time that the assay was set up had an interassay CV of 3.4% and an intra-assay CV of 1.7%.

Benjamini–Hochberg False Discovery Rate Correction for Multiple Comparisons

We performed Benjamini–Hochberg false discovery rate correction for multiple comparisons, with a false discovery rate of 5% on primary models investigating the relationship between depressive symptoms and measures of cellular aging.3 We did not include models assessing group differences or probing the effect of including covariates such as number of stressful life events. All models that were significant in our original analysis survived correction. Table S1 summarizes this information. As demonstrated, the model ranked 2 was the last for which the p value was smaller than the Benjamini–Hochberg critical value, indicating that model 2 and all models ranked earlier than it remain statistically significant.
### TABLE S1 Benjamini–Hochberg (BH) False Discovery Rate Correction

<table>
<thead>
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<th>Model</th>
<th>Model p</th>
<th>Rank</th>
<th>BH critical value</th>
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<td>0.008</td>
</tr>
<tr>
<td>Depressive symptoms at T1 predicting change in TL from T1 to T2</td>
<td>.016</td>
<td>2</td>
<td>0.017</td>
</tr>
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<td>TL at T1 predicting change in depressive symptoms from T1 to T2</td>
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<tr>
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<td>0.033</td>
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<tr>
<td>mtDNAcn at T1 predicting change in depressive symptoms from T1 to T2</td>
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<td>0.042</td>
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<tr>
<td>Depressive symptoms at T1 predicting TL at baseline</td>
<td>.903</td>
<td>6</td>
<td>0.050</td>
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</tbody>
</table>

**Note:** mtDNA-cn = mitochondrial DNA copy number; TL = telomere length; T1 = time 1, first wave of data collection; T2 = time 2, second wave of data collection.

### SUPPLEMENTAL REFERENCES