Resilience factors may buffer cellular aging in individuals with and without chronic knee pain

Alisa J Johnson1,2, Ellen Terry1,2, Emily J Bartley1,2, Cynthia Garvan3, Yenisel Cruz-Almeida1,4, Burel Goodin5, Toni L Glover6, Roland Staud7, Laurence A Bradley8, Roger B Fillingim1,2, and Kimberly T Sibille1,4

Abstract
Telomere length, a measure of cellular aging, is inversely associated with chronic pain severity. While psychological resilience factors (e.g., optimism, acceptance, positive affect, and active coping) are associated with lower levels of clinical pain and greater physical functioning, it is unknown whether resilience may buffer against telomere shortening in individuals with chronic pain. Additionally, a broader conceptualization of resilience that includes social and biobehavioral factors may improve our understanding of the relationship between resilience, chronic pain, and health outcomes. In individuals with and without chronic knee pain, we investigated whether (1) psychological resilience would be positively associated with telomere length and if (2) a broader conceptualization of resilience including social and biobehavioral factors would strengthen the association. Seventy-nine adults, 45 to 85 years of age, with and without knee pain completed demographic, health, clinical pain, psychological, social, and biobehavioral questionnaires. Resilience levels were determined by summing the total number of measures indicating resilience based on published clinical ranges and norms. Blood samples were collected, and telomere length was determined. In regression analyses controlling for sex, race, age, and characteristic pain intensity, greater psychological resilience and psychosocial/biobehavioral resilience were associated with longer telomeres (p = .0295 and p = .0116, respectively). When compared, psychosocial/biobehavioral resilience was significantly more predictive of telomere length than psychological resilience (p < .0001). Findings are promising and encourage further investigations to enhance understanding of the biological interface of psychosocial and biobehavioral resilience factors in individuals with musculoskeletal chronic pain conditions.

Keywords
Resilience, telomeres, chronic pain, stress, osteoarthritis

Date Received: 12 February 2019; accepted: 5 March 2019
psychosocial, and behavioral. Importantly, a number of studies have reported an inverse relationship between TL and the biological and psychosocial stress experienced with persisting chronic pain. For example, in individuals reporting high stress and chronic pain, TL was shorter compared to those individuals with low stress and no chronic pain. Similarly, TL was shorter among individuals with fibromyalgia reporting higher pain and comorbid depression compared to those with low levels of pain and depression. More recently, in a larger cohort of individuals with and without chronic knee pain, TL was associated with chronic pain severity in a dose-response fashion. TL may also reflect the buffering effect of resilience factors in individuals with chronic pain.

An array of studies indicates the clinical benefit of several psychological factors in the experience of chronic pain. Qualities such as positive affect, dispositional optimism, active coping, acceptance, and purpose in life have been indicated as buffering against the negative sequelae of chronic pain. These psychological factors are frequently described as promoting “resilience” as a result of their being associated with lower clinical pain and greater physical function. In essence, resilience is “the process by which people bounce back from adversity and reintegrate and ideally grow from the experience.” Hence, these psychological factors appear to be protective, limiting the negative impact of living with chronic pain.

In addition to psychological factors, social factors may also promote resilience in individuals with chronic pain. Indeed, social support and social integration predict lower all-cause mortality rates in various populations. Additionally, positive social experiences and higher levels of self-reported social integration were associated with lower biological “wear and tear” recognized as allostatic load (AL). Further, there is evidence that emotional and social support may attenuate biological responses to environmental stress.

Resilience factors are typically regarded as psychological or psychosocial constructs. However, it may be more optimal to consider resilience from a more comprehensive biopsychosocial framework. For example, health behaviors such as regular exercise, moderate alcohol consumption, and being a non-smoker predict decreased morbidity and mortality in individuals with chronic widespread pain and have been linked with lower levels of inflammatory and metabolic biomarkers. Additionally, in individuals with or at risk for knee osteoarthritis (OA), lower omega-6:omega-3 polyunsaturated fatty acid (PUFA) ratios were associated with lower clinical pain, experimental pain sensitivity, and psychosocial distress, and greater physical functioning compared to those individuals with higher omega-6:omega-3 (O6:O3) PUFA ratio levels. Low O6:O3 PUFA ratios have also been associated with increased TL in overweight sedentary adults and may serve to enhance overall resilience. Thus, a broad conceptualization of resilience that includes psychosocial and biobehavioral resilience factors may be more informative in understanding the biological interface of resilience and chronic pain.

The aims of the current study were to determine (1) if greater levels of psychological resilience are associated with TL and (2) if a broader conceptualization of resilience that includes social and biobehavioral factors would improve the model. We hypothesized that (1) greater levels of psychological resilience and psychosocial/biobehavioral resilience would be associated with longer TL and that (2) greater levels of psychosocial/biobehavioral resilience would demonstrate stronger association with TL compared to psychological resilience.

Methods

Study design

The current investigation is a sub-study of a larger study, Understanding Pain and Limitations in Osteoarthritic Disease (UPLOAD). UPLOAD was a multi-site cross-sectional investigation conducted at the University of Florida (UF) and the University of Alabama at Birmingham (UAB). The study is limited to data collected at the University of Florida. All procedures were reviewed and approved by the UF Institutional Review Board (IRB201500906). Participants provided written informed consent. The Methods section is limited to the components applicable to the current study. A complete description of the overall study is available.

Participants

Participants in the current investigation were enrolled at the University of Florida during the period between 2010 and 2013, and had TL measures completed. The sample consisted of 79 community-dwelling adults between 45 and 85 years of age, with and without knee pain in the previous month, who self-identified as non-Hispanic Black (NHB) or non-Hispanic White (NHW). Individuals were excluded from the study for the following conditions: (1) prosthetic knee replacement or other clinically significant surgery to the painful knee; (2) uncontrolled hypertension (blood pressure > 150/95 mm Hg); (3) heart disease, congestive heart failure, or history of acute myocardial infarction; (4) severe peripheral neuropathy in which pain testing was contraindicated; (5) systemic rheumatic disorders including rheumatoid arthritis, systemic lupus erythematosus, gout, and fibromyalgia; (6) neurological diseases such as...
Parkinson’s disease, multiple sclerosis, stroke with loss of sensory or motor function, or uncontrolled seizures; (7) significantly greater pain in other body sites than in the knee; (8) chronic daily opioid use; (9) hospitalization within the preceding year for psychiatric illness; or (10) pregnant or nursing.

**Procedures**

Participants completed a standardized telephone screening to confirm initial eligibility. Eligible participants completed two sessions approximately two to three weeks apart. During Session 1, after written informed consent was obtained, participants completed a comprehensive health assessment, which involved the collection of demographic information, anthropometric measurements, health history information, and physical examination. During Session 2, blood samples were collected, and peripheral blood mononuclear cells were isolated to assess TL. Participants completed a number of self-report questionnaires to assess clinical pain, psychological, social, and behavioral factors.

**Measures**

**Chronic pain.** The Graded Chronic Pain Scale (GSPC) is a self-report questionnaire that assesses chronic pain intensity and pain-related disability. Characteristic pain intensity is determined by a response to three questions on a 0 to 10 numeric rating scale: current, worst, and average knee pain intensity during the past six months. Ratings from the three items are averaged and multiplied by 10 to calculate a score (0–100). The Graded Chronic Pain Scale (GCPS) has demonstrated test–retest reliability of .79 at 28 months and Cronbach’s α = 0.74. Response totals >2.87 indicated resilience.

**Psychological resilience.** Validated and recognized measures were used to operationalize the psychological resilience phenotype. For each measure, if the score fell within the positive/protective range based on clinical/normative values a score of 1 was added for resilience. A total psychological resilience value was determined based on the summative total from the combined measures. Dispositional optimism. The Life Orientation Test-Revised (LOT-R) is a 10-item questionnaire that assesses dispositional optimism and pessimism. The scale contains three items for optimism (e.g., “In uncertain times, I usually expect the best”) and pessimism (e.g., “If something can go wrong for me, it will”), with four non-contributing questions. Responses are based on a four-point Likert-type scale (0 = “strongly disagree” to 4 = “strongly agree”). The measure has demonstrated a test–retest reliability of .79 at 28 months and Cronbach’s α of .82. Response totals ≥18 indicated resilience.

**Positive and negative affect.** Positive and negative affect were assessed using the Positive and Negative Affect Scale (PANAS). The PANAS is a 20-item measure that consists of 10 positively valenced items (positive affect) and 10 negatively valenced items (negative affect). Items are self-rated on a five-point Likert-type scale (1 = “very slightly or not at all” to 5 = “extremely”) and summed to produce total scores for each affect component, with higher scores representing higher levels of the construct. The timeframe collected in this study was “to what extent you generally feel this way.” The PANAS has been demonstrated to be internally consistent (Cronbach’s α’s ≥ .84). Having high levels of trait positive affect and low levels of trait negative affect are considered as positive and resilience promoting. Response totals for positive affect ≥35 and negative affect ≤18.1 indicated resilience.

**Active coping.** Active coping was assessed using the Coping Strategies Questionnaire-Revised (CSQ-R), a 27-item measure of pain coping strategies. Participants rate how often they use each strategy on a 0 (“never do that”) to 6 (“always do that”) Likert scale. Items are summed for each domain separately with higher scores indicating greater use of that strategy. The CSQ-R has demonstrated acceptable reliability (Cronbach’s α’s = 0.72–0.86). Responses on the CSQ-R have been associated with arthritis pain and disability in both NHB and NHW individuals. Active coping was assessed using items from the distraction, ignoring pain sensations, distracting from pain, and coping self-statements domains. Based on prior findings, response totals >2.87 indicated resilience.

**Perceived stress.** Perceived stress was assessed through the 10-item Perceived Stress Scale (PSS), a reliable and valid scale designed to measure the role of non-specific appraised stress. Participants are asked to rate (0 = “never” to 4 = “very often”) statements asking about thoughts and feelings over the past month. A total perceived stress score is computed. Low levels of perceived stress are considered as a positive psychological factor and have been associated with TL. Based on recommended ranges, perceived stress scores ranging from 0 to 13 indicated resilience.

**Psychosocial/biobehavioral resilience.** In addition to the psychological measures described above, three additional validated and recognized measures were included to assess a psychosocial and biobehavioral conceptualization of resilience.
Social support. The Multidimensional Scale of Perceived Social Support (MSPSS) is a 12-item self-report measure used to assess the extent to which an individual perceives social support from family, friends, and significant others.56 Each item is rated on a seven-point Likert scale (1 = “very strongly disagree” to 7 = “very strongly agree”), for a total score ranging from 12 to 84, with higher scores indicating higher levels of perceived support. The MSPSS has been demonstrated to be a reliable measure across subscales (Cronbach’s α’s = 0.81–0.98).57,58 Based on published ranges, scores in the range of 49 to 84 indicated resilience.56

Tobacco use. Participants responded to a question regarding smoking status: never, former, or current. Responses endorsing never and former smoker status indicated resilience.

Waist-to-hip ratio. Participant’s waist circumference and hip circumference were measured using a measuring tape. The waist-to-hip ratio (WHR) is determined by dividing the wait circumference by the hip circumference. The World Health Organization defines abdominal obesity as a WHR of > .85 for women and > .90 for men.59 Risk and protective ranges were determined applying a generous approach. A WHR < .90 for both men and women was defined as indicating resilience.

TL analysis

TL analysis followed a standardized procedure as previously described.16

A blood sample was collected during the second study session and placed on ice. It was centrifuged at 4°C for 10 min at 3000 r/min. The blood was mixed with 1× phosphate-buffered saline, layered onto a volume of Lymphoprep solution that was contained in a centrifuge tube. After centrifugation, the lymphocyte band was separated, washed, and centrifuged to form a pellet. The pellet was re-suspended in 1× phosphate-buffered saline, and the sample is stored at -80°C. The DNA isolation was achieved using the Qiagen Flexigene kit. Lysis buffer was added to the sample before being mixed and centrifuged. The resulting pellet was re-suspended in denaturation buffer containing protease and incubated. DNA was then precipitated, washed, centrifuged, and re-suspended in hydration buffer. TL was analyzed by the Blackburn Lab, University of California San Francisco.50

The TL assay is adapted from the published original method by Cawthon.60,61 The telomere thermal cycling profile consists of Cycling for T(telomeric) polymerase chain reaction (PCR): 96°C for 1 min; denature at 96°C for 1 s, anneal/extend at 54°C for 60 s, with fluorescence data collection, 30 cycles. Cycling for S (single copy gene) PCR: PCR: 96°C for 1 min; denature at 95°C for 15 s, anneal at 58°C for 1 s, extend at 72°C for 8 cycles; followed by denature at 96°C for 1 s, anneal at 58°C for 1 s, extend at 72°C for 20 s, hold at 83°C for 5 cycles with data collection, 35 cycles. The primers for the telomere PCR are tel1b [5′-CGGGTTT(GTTTGG)2-GTT-3′], used at a final concentration of 100 nM, and tel2b [5′-GGCTTG(CCTTAC)2-CCT-3′], used at a final concentration of 900 nM. The primers for the single-copy gene (human beta-globin) PCR are hbg1 [5′-GCTTCTGACACA(ACTG)TTACTAQC-3′]; used at a final concentration of 300 nM, and hbg2 [5′-CACCACA(CTTAC)TCCACC-C-3′], used at a final concentration of 700 nM. The final reaction mix contains 20 mM Tris-HCl, pH 8.4; 50 mM KCl; 200 mM each dNTP; 1% DMSO; 0.4× Syber Green I; 22 ng E.coli DNA per reaction; 0.4 Units of Platinum Taq DNA polymerase (Invitrogen Inc.) per 11 microliter reaction; and 6 ng of genomic DNA. Tubes containing 26, 8.75, 2.9, 0.97, 0.324, and 0.108 ng of a reference DNA (a pooled samples of leukocyte genomic DNA from 100 female donors) are included in each PCR run so that the quantity of targeted templates in each research sample can be determined relative to the reference DNA sample by the standard curve method. The same reference DNA was used for all PCR runs.

To control for inter-assay variability, eight control DNA samples are included in each run. In each batch, the telomere to single copy gene (T/S) ratio of each control DNA is divided by the average T/S for the same DNA from 10 runs to get a normalizing factor. This is done for all eight samples, and the average normalizing factor for all eight samples is used to correct the participant DNA samples to get the final T/S ratio. The T/S ratio for each sample was measured twice. When the duplicate T/S value and the initial value vary by more than 7%, the sample was run the third time and the two closest values were reported. The average coefficient of variation for this study is 1.9%. The lab personnel who performed the assays received de-identified blood samples and were blind to demographic and clinical data.

To determine the conversion factor for the calculation of approximate base pair TL from T/S ratio, the above method was used to determine the T/S ratios, relative to the same reference DNA, for a set of genomic DNA samples from the human fibroblast primary cell line IMR90 at different population doublings, as well as with the telomerase protein subunit gene (hTERT) trans- fected into a lentiviral construct. The mean TRF length from these DNA samples was determined using
Southern blot analysis, and the slope of the plot of mean TRF length versus T/S for these samples served as the conversion factor for calculation of TL in base pairs from the T/S ratio. The equation for conversion from T/S ratio to base pairs for this study was base pairs $= 3274 + 2413 \times (T/S)$.

**Statistical analysis**

SAS version 9.4 (Cary, NC) was used for all analyses. Data were checked for distributional form and outliers. All testing was two sided using a .05 level of significance. Psychological resilience was operationalized based on the sum total of psychological measures indicating resilience (LOT-R, PANAS positive affect, PANAS negative affect, active coping (CSQ-R), and Perceived Stress Scale (PSS)). Psychosocial/biobehavioral resilience was operationalized based on the sum total of measures endorsed representing psychological resilience with the addition of three additional measures representing social and biobehavioral factors: MSPSS, tobacco use, and WHR.

Summary statistics were used to describe the sample. Consistent with telomere research recommendations, correlations were computed to assess known (age, sex, and race) and reported (optimism and WHR) associations with TL. Regression modeling was used to test Hypotheses 1 and 2. For Hypothesis 1, psychological resilience was entered as a predictor of TL, after controlling for relevant covariates: age, sex, and characteristic pain intensity. Similarly, for Hypothesis 2, the psychosocial/biobehavioral resilience composite was entered into a regression model as a predictor of TL, after controlling for relevant covariates: age, sex, race, and characteristic pain intensity. The strength of association between TL and the two resilience measures (i.e., psychological resilience and psychosocial/biobehavioral resilience) was compared using Meng et al.'s method.

**Results**

**Descriptive findings**

Of the 79 participants, 67% were females, 58% were NHW with mean age of 58.4 (±8.1) years. A description of the study sample is provided in Table 1. The mean (SD) TL was 1.04 (±0.20) with a range from 0.59 to 1.62. Anticipated relationships were demonstrated in that TL was significantly associated with age ($r_s = –31, p = .0045$) and sex ($r_s = –25, p = .0215$) such that females had longer TL, 1.07 (±0.21), than males, 0.97 (±0.18). As previously reported, TL was also significantly associated with optimism, LOT-R ($r_s = .24, p = .0341$), and WHR ($r_s = –.29, p = .0093$). A description of the reported ranges and sample-specific response patterns for the resilience measures are displayed in Table 2.

**Relationship between psychological resilience and TL**

In our regression analysis, we found that the psychological resilience was associated with longer telomeres after controlling for sex, age, race, and characteristic pain ($p = .030$, $b = .03$ (±.02), model $R^2 = .21$). Table 3 provides the full results for the tested models.

| Table 1. Demographic characteristics of the study sample. |
|----------------------------------------------------------|
| Demographics | % (N) or M (SD) |
| Age | 58.4 (8.1) |
| Sex |  |
| Female | 66% (53) |
| Male | 34% (26) |
| Ethnicity/race |  |
| Non-Hispanic Black | 43% (33) |
| Non-Hispanic White | 57% (46) |
| Characteristic pain intensity (GCPS) | 42.3 (25.9) |
| GCPS: Graded Chronic Pain Scale. |

**Relationship between psychosocial/biobehavioral resilience and TL**

The psychosocial/biobehavioral resilience composite was also positively associated with longer telomeres after controlling for sex, age, race, and characteristic pain ($p = .012$, $b = .03$ (±.01), model $R^2 = .23$). The full model is presented in Table 3.

**Comparison between psychological resilience and psychosocial/biobehavioral resilience**

In testing the strength of association between TL and each of the resilience composites, we found that the psychosocial/biobehavioral composite was significantly more associated with TL than the psychological resilience composite ($Z = –13.23, p < .0001$).

**Additional analysis**

Due to the strong association between WHR and TL, a post hoc analysis was conducted to evaluate the relationship between the psychological resilience composite and TL after adjusting for WHR. The psychological resilience composite remained significantly associated with TL with the addition of WHR as a covariate ($p = .03$, $b = .04$ (±.02), model $R^2 = .23$). Table 3 provides the full results for the tested model.

Additionally, to further assist with the interpretation of the clinical relevance, an analysis of covariance was completed comparing individuals with low levels of...
psychosocial/biobehavioral resilience (sum values in the 1–3 range) compared to those individuals with high levels of psychosocial/biobehavioral resilience (values in the 5–8 range). The T/S ratios for both groups were converted to base pairs as described in the Methods section. The groups differed by 256 base pairs. Applying findings from a prior publication, a difference of 256 base pairs suggests a general indication of an approximate 10 years difference in cellular aging.64 As base pairs were not directly measured in the current cohort, and the conversion formula is based on a series of DNA samples from a human primary cell line, not a direct comparison of the Southern blot method and quantitative PCR, interpretation is limited and can only provide a frame of reference for evaluating possible clinical relevance.

Discuss the implications of these findings in a clinical context.

**Discussion**

Our findings provide evidence that in individuals with and without knee pain, higher levels of both psychological and psychosocial/biobehavioral resilience were associated with longer telomeres. Importantly, the broader conceptualization of resilience including psychosocial and biobehavioral factors provided the strongest association with TL compared to psychological resilience alone. Thus, a resilience phenotype characterized by a greater array of psychosocial and biobehavioral protective factors is associated with longer telomeres.

**Psychological resilience and TL**

Psychological resilience has frequently been associated with enhanced emotional, cognitive, social, and physical...
functioning. Numerous publications report that positive psychological factors buffer the experience of chronic pain as indicated by lower report of clinical symptoms and functional limitations. Additionally, there is a growing body of evidence linking psychological resilience factors with biological benefits. A relationship between longer telomeres and positive psychological factors, particularly optimism, has been shown. In this study, we also found a correlation between optimism and TL. Our research contributes to the body of evidence indicating that psychological resilience factors have a biological interface.

**Psychosocial/biobehavioral resilience and TL**

A number of publications have addressed the role of social support in chronic pain, with most describing its benefits and a few noting that some forms of social support (e.g., the reinforcement of maladaptive behavior patterns) may be detrimental in chronic pain. However, the health benefits of social support have been consistently indicated, including associations with improved stress-related physiological functioning. Also, negative social experiences (e.g., social isolation and loneliness) are associated with multiple health risks, including inactivity, smoking, high blood pressure, and shorter telomeres in individuals who also demonstrated dysregulated parasympathetic functioning.

As expected, health behaviors, e.g., regular exercise, maintaining a healthy weight, healthy diet, moderate alcohol consumption, and not smoking, are also associated with lower levels of disease, biological burden, and death across a range of studies including those specific to individuals with chronic pain. Telomeres have been associated with an array of biopsychosocial positive factors, our findings further contribute to this body evidence. In our study, a linear relationship emerged such that an increase of psychosocial/biobehavioral measures representing resilience was significantly associated with longer telomeres.

**Additional considerations**

Telomeres can be conceptualized as a downstream measure of persisting overall stress system functioning. As such, a limited association between TL and specific psychological, social, and biobehavioral measures would be expected would not typically reach a “dose level” to contribute in a meaningful way to TL. However, combined, these factors characterize a resilience phenotype which could represent traits and behaviors of sufficient magnitude to buffer against telomere shortening. Our findings suggest that there is a positive linear relationship between TL and resilience characteristics in individuals with or at risk for knee OA. Additionally, we previously reported a dose-response relationship between chronic pain severity and TL, such that differences in TL between those with the highest chronic pain severity compared to individuals with no or low levels of chronic pain severity was 16 years of accelerated aging. Similarly, in the current study, based on conversion estimations, individuals with the lowest psychosocial/biobehavioral resilience scores compared to those with the highest psychosocial/biobehavioral resilience scores differed by approximately 10 years of cellular aging even after controlling for relevant covariates.

**Limitations and future directions**

There are a number of limitations warranting acknowledgment. As this investigation is a cross-sectional study with a relatively small sample, prospective investigations with a larger sample are necessary. Additionally, our sample is comprised of middle-aged and older adults, many who screened positive for or are at risk for knee OA, thus the generalizability is limited. Future investigations with individuals with differing chronic pain conditions will be necessary. Our resilience measures were selected from those available in a completed study. There are a number of other factors and measures that warrant evaluation in better understanding the biological interface of resilience in chronic pain conditions. Finally, telomere research is still a developing science with recognized limitations. Strategies to increase confidence were incorporated in the current study by working with a recognized and well-published lab, Blackburn Lab, University of California San Francisco, and evaluating known patterns between TL with age, sex, and ethnicity/race.

There are a number of additional strengths that are noteworthy in the current investigation. The UPLOAD study provides a well-characterized sample with a broad array of measures to answer the identified questions. The resilience measures were comprised of validated instruments and measures with recognized norms to define ranges for resilience. Characterizing a phenotype based on a combination of measures provides a stronger representation than any one measure alone. In regard to future directions, there are significant disparities in the clinical and functional limitations of knee OA in NHB and NHW. Investigations exploring potential race and ethnic group differences in resilience factors might highlight factors contributing to those disparities and illuminate possible targets for treatment. Lastly, if findings are replicated, the clinical implications are exciting. There is a strong body of evidence indicating resilience factors (psychological traits and health behaviors) are associated with lower rates of morbidity and mortality. Our findings suggest the biological interface of resilience factors is measurable. Hence, clinical strategies to increase
resilience and improve pain-related health outcomes may also not only buffer the biological burden of pain but findings suggest that we may be able to monitor and evaluate the biological benefits of various interventions.

Conclusions
In summary, our findings demonstrate a positive relationship between resilience factors and TL. Specifically, in middle-aged and older adults with or at risk for knee OA, greater psychosocial and biobehavioral resilience was significantly associated with longer telomeres. Further, we have previously shown a dose-response relationship between TL and increasing chronic pain severity. Current findings suggest that there is also a dose-response relationship between psychosocial/biobehavioral resilience factors and TL. Second, our findings support prior recommendations of the benefits of considering combining measures in better capturing the biological interface to particular phenotypes, in this study, resilience phenotypes. Third, our study has potential clinical relevance. If findings from the current study are replicated and extended into prospective analyses which indicate an influence on health outcomes, then psychosocial and biobehavioral resilience interventions and measures could be used to guide and evaluate treatment for chronic pain.

Authors’ Note
Toni L. Glover is now affiliated with School of Nursing, Oakland University, Rochester, MI, USA.

Acknowledgments
The authors thank UF UPLOAD participants, PRICE research team, collaborators at the University of Alabama at Birmingham, CTSI CRC RNs, the Pharmacogenomics Lab, and Cheryl Galloway for their help in this study. The authors also greatly appreciate the contributions of the Blackburn Lab and Jue Lin, PhD, for the completion of the telomere length analysis and description. The authors also extend their gratitude to Bruce McEwen, PhD, for supporting and informing their efforts in the investigations of the biological interface of chronic pain.

Declaration of Conflicting Interests
The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Funding
The authors disclosed receipt of the following financial support for the research, authorship, and/or publication of this article: This work was supported by the National Institutes of Health/ National Institute on Aging R37AG033906 and R01AG054370, the National Institute of Arthritis and Musculoskeletal and Skin Diseases, K23AR062099, the University of Florida Clinical and Translational Science Institute UL1TR000064, the American Pain Society Future Leaders in Pain Research Grant (KTS), NINDS K22NS102334 and UF McKnight Brain Institute Career Enhancement Award (ELT), 4R00AG052642-03 (EJB), and K01AG048259 (YCA). The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

ORCID iD
Alisa J Johnson http://orcid.org/0000-0003-4287-8703

References
1. Sibille KT, Witek-Janusek L, Mathews HL and Fillingim RB. Telomeres and epigenetics: potential relevance to chronic pain. Pain 2012; 153: 1789.
2. Epel ES, Blackburn EH, Lin J, Dhabhar FS, Adler NE, Morrow JD and Cawthon RM. Accelerated telomere shortening in response to life stress. Proc Natl Acad Sci U S A 2004; 101: 17312–17315.
3. Nettleton JA, Diez-Roux A, Jenny NS, Fitzpatrick AL and Jacobs DR, Jr. Dietary patterns, food groups, and telomere length in the Multi-Ethnic Study of Atherosclerosis (MESA). Am J Clin Nutr 2008; 88: 1405–1412.
4. Tzanetakou IP, Katsilambros NL, Benetos A, Mikhalidis DP and Perrea DN. “Is obesity linked to aging”? adipose tissue and the role of telomeres. Ageing Res Rev 2012; 11: 220–229.
5. Epel ES. Telomeres in a life-span perspective: a new “Psychobiomarker”? Curr Dir Psychol Sci 2009; 18: 6–10.
6. Tucker LA. Physical activity and telomere length in U.S. men and women: an NHANES investigation. Prev Med 2017; 100: 145–151.
7. Puterman E, Lin J, Krauss J, Blackburn EH and Epel ES. Determinants of telomere attrition over 1 year in healthy older women: stress and health behaviors matter. Mol Psychiatry 2015; 20: 529–535.
8. Schutte NS and Malouff JM. The relationship between perceived stress and telomere length: a meta-analysis. Stress Health 2016; 32: 313–319.
9. Schutte NS, Palanisamy SK and McFarlane JR. The relationship between positive psychological characteristics and longer telomeres. Psychol Health 2016; 31: 1466–1480.
10. Kiecolt-Glaser JK, Epel ES, Belury MA, Andridge R, Lin J, Glaser R, Malarkey WB, Hwang BS and Blackburn E. Omega-3 fatty acids, oxidative stress, and leukocyte telomere length: a randomized controlled trial. Brain, Behav, Immun 2013; 28: 16–24.
11. Ornish D, Lin J, Chan JM, Epel E, Kemp C, Weidner G, Marlin R, Frena SJ, Magbanua MJM, Daubenmier J, Estay I, Hills NK, Chainani-Wu N, Carroll PR and Blackburn EH. Effect of comprehensive lifestyle changes on telomerase activity and telomere length in men with biopsy-proven low-risk prostate cancer: 5-year follow-up of a descriptive pilot study. Lancet Oncol 2013; 14: 1112–1120.
12. Epel E. How “reversible” is telomeric aging? Cancer Prev Res 2012; 5: 1163.
13. Lin J, Epel E and Blackburn E. Telomeres and lifestyle factors: roles in cellular aging. Mutat Res 2012; 730: 85–89.
14. Sibille KT, Languea T, Burkley B, Gong Y, Glover TL, King C, Riley JL, 3rd, Leeuwenburgh C, Staud R, Bradley LA and Fillingim RB. Chronic pain, perceived stress, and cellular aging: an exploratory study. Mol Pain 2012; 8: 12–12.
15. Hassett AL, Epel E, Clauw DJ, Harris RE, Harte SE, Kairys A, Buyske S and Williams DA. Pain is associated with short leukocyte telomere length in women with fibromyalgia. J Pain 2012; 13: 959–969.
16. Sibille KT, Chen H, Bartley EJ, Riley J, Glover TL, King CD, Zhang H, Cruz-Almeida Y, Goodin BR, Sotolongo A, Petrov ME, Herbert M, Bulls HW, Edberg JC, Staud R, Redden D, Bradley LA and Fillingim RB. Accelerated aging in adults with knee osteoarthritis pain: consideration for frequency, intensity, time, and total pain sites. Pain Rep 2017; 2: e591.
17. Hassett AL and Finan PH. The role of resilience in the clinical management of chronic pain. Curr Pain Headache Rep 2016; 20: 39.
18. Karoly P and Ruehlman LS. Psychological “resilience” and its correlates in chronic pain: findings from a national community sample. Pain 2006; 123: 90–97.
19. Ong AD, Zautra AJ and Reid MC. Psychological resilience predicts decreases in pain catastrophizing through positive emotions. Psychol Aging 2010; 25: 516.
20. Sturgeon JA and Zautra AJ. Resilience: a new paradigm for adaptation to chronic pain. Curr Pain Headache Rep 2016; 20: 105–112.
21. Sturgeon JA and Zautra AJ. Psychological resilience, pain catastrophizing, and positive emotions: perspectives on comprehensive modeling of individual pain adaptation. Curr Pain Headache Rep 2013; 17: 317.
22. Resnick B. Resilience in older adults. Top Geriatric Rehabil 2014; 30: 155–163.
23. Lambert VA, Lambert CE, Klipple GL and Mewshaw EA. Relationships among hardness, social support, severity of illness, and psychological well-being in women with rheumatoid arthritis. Health Care Women Int 1990; 11: 159–173.
24. Kroenke CH, Kubzansky LD, Schernhammer ES, Holmes MD and Kawachi I. Social networks, social support, and survival after breast cancer diagnosis. J Clin Oncol 2006; 24: 1105–1111.
25. Berkman LF, Melchior M, Chastang J-F, Niedhammer I, Lecerf A and Goldberg M. Social integration and mortality: a prospective study of French employees of Electricity of France-Gas of France: the GAZEL Cohort. Am J Epidemiol 2004; 159: 167–174.
26. Holt-Lunstad J, Smith TB and Layton JB. Social relationships and mortality risk: a meta-analytic review. PLoS Med 2010; 7: e1000316.
27. Iribarren C, Jacobs DR, Kiefe CI, Lewis CE, Matthews KA, Roseman JM and Hulley SB. Causes and demographic, medical, lifestyle and psychosocial predictors of premature mortality: the CARDIA study. Soc Sci Med 2005; 60: 471–482.
28. Seeman TE, Singer BH, Ryff CD, Love GD and Levy-Storms L. Social relationships, gender, and allostatic load across two age cohorts. Psychosom Med 2002; 64: 395–406.
29. Coan JA, Schaefer HS and Davidson RJ. Lending a hand: social regulation of the neural response to threat. Psychol Sci 2006; 17: 1032–1039.
30. Che X, Cash R, Ng SK, Fitzgerald P and Fitzgibbon BM. A systematic review of the processes underlying the main and the buffering effect of social support on the experience of pain. Clin J Pain 2018; 34: 1061–1076.
31. Ruiz-Parraga GT and Lopez-Martinez AE. The role of experiential avoidance, resilience and pain acceptance in the adjustment of chronic back pain patients who have experienced a traumatic event: a path analysis. Ann Behav Med 2015; 49: 247–257.
32. Ruiz-Parraga GT, Lopez-Martinez AE, Estve R, Ramirez-Maestre C and Wagnild G. A confirmatory factor analysis of the Resilience Scale adapted to chronic pain (RS-18): new empirical evidence of the protective role of resilience on pain adjustment. Qual Life Res 2015; 24: 1245–1253.
33. Smith BW and Zautra AJ. Vulnerability and resilience in women with arthritis: test of a two-factor model. J Consult Clin Psychol 2008; 76: 799–810.
34. Sufieda A, Meissner W, Rosendahl J and Guntinas-Lichius O. Influence of depression, catastrophizing, anxiety, and resilience on postoperative pain at the first day after otolaryngological surgery: a prospective single center cohort observational study. Medicine 2016; 95: e4256.
35. Macfarlane GJ, Barnish MS and Jones GT. Persons with chronic widespread pain experience excess mortality: longitudinal results from UK Biobank and meta-analysis. Ann Rheum Dis 2017; 76: 1815–1822.
36. Sibille KT, Steingrimsdottir ÖA, Fillingim RB, Stubhaug A, Schirmer H, Chen H, McEwen BS and Nielsen CS. Investigating the burden of chronic pain: an inflammatory and metabolic composite. Pain Res Manag 2016; 2016: 7657329.
37. Sibille KT, King C, Garrett TJ, Glover TL, Zhang H, Chen H, Reddy D, Goodin BR, Sotolongo A and Petrov ME. Omega-6: omega-3 PUFA ratio, pain, functioning, and distress in adults with knee pain. Clin J Pain 2018; 34: 182–189.
38. King CD, Sibille KT, Goodin BR, Cruz-Almeida Y, Glover TL, Bartley E, Riley JL, Herbert MS, Sotolongo A, Schmidt J, Fessler BJ, Redden DT, Staud R, Bradley LA and Fillingim RB. Experimental pain sensitivity differs as a function of clinical pain severity in symptomatic knee osteoarthritis. Osteoarthritis Cartilage 2013; 21: 1243–1252.
39. Von Korff M, Ormel J, Keefe FJ and Dworkin SF. Grading the severity of chronic pain. Pain 1992; 50: 133–149.
40. Scheier MF, Carver CS and Bridges MW. Distinguishing optimism from neuroticism (and trait anxiety, self-mastery, and self-esteem): a reevaluation of the Life Orientation Test. J Pers Soc Psychol 1994; 67: 1063–1078.
41. Watson D, Clark LA and Tellegen A. Development and validation of brief measures of positive and negative affect: the PANAS scales. J Pers Soc Psychol 1988; 54: 1063–1070.

42. Crawford JR and Henry JD. The positive and negative affect schedule (PANAS): construct validity, measurement properties and normative data in a large non-clinical sample. Br J Clin Psychol 2004; 43: 245–265.

43. Strand EB, Zautra AJ, Thoresen M, Odegard S, Uhlig T and Finset A. Positive affect as a factor of resilience in the pain-negative affect relationship in patients with rheumatoid arthritis. J Psychosom Res 2006; 60: 477–484.

44. Zautra AJ, Johnson LM and Davis MC. Positive affect as a source of resilience for women in chronic pain. J Consult Clin Psychol 2005; 73: 212–220.

45. Sibille KT, Kindler LL, Glover TL, Staud R, Riley JL, 3rd and Fillingim RB. Affect balance style, experimental pain sensitivity, and pain-related responses. Clin J Pain 2012; 28: 410–417.

46. Hassett AL, Simonelli LE, Radvanski DC, Buyske S, Savage SV and Sigal LH. The relationship between affect balance style and clinical outcomes in fibromyalgia. Arthritis Rheum 2008; 59: 833–840.

47. Rosenstiel AK and Keefe FJ. The use of coping strategies in chronic low back pain patients: relationship to patient characteristics and current adjustment. Pain 1983; 17: 33–44.

48. Abbott A. The coping strategy questionnaire. J Physiother 2010; 56: 63.

49. Riley JL, 3rd and Robinson ME. CSQ: five factors or fiction? Clin J Pain 1997; 13: 156–162.

50. Robinson ME, Riley JL, 3rd, Myers CD, Sadler JJ, Kvaal SA, Geisser ME and Keefe FJ. The coping strategies questionnaire: a large sample, item level factor analysis. Clin J Pain 1997; 13: 43–49.

51. Monticone M, Ferrante S, Giorgi I, Galandra C, Rocca B and Foti C. The 27-item coping strategies questionnaire-revised: confirmatory factor analysis, reliability and validity in Italian-speaking subjects with chronic pain. Pain Res Manag 2014; 19: 153–158.

52. Jordan MS, Lumley MA and Leisen JC. The relationships of cognitive coping and pain control beliefs to pain and adjustment among African-American and Caucasian women with rheumatoid arthritis. Arthritis Care Res 1998; 11: 80–88.

53. Cohen S, Kamarck T and Mermelstein R. A global measure of perceived stress. J Health Soc Behav 1983; 24: 385–396.

54. Cohen S, Kamarck T and Mermelstein R. Perceived stress scale. Measuring stress: a guide for health and social scientists. Oxford: Oxford University Press 1994, pp. 235–283.

55. Cohen S and Janicki-Deverts D. Who’s stressed? Distributions of psychological stress in the United States in probability samples from 1983, 2006, and 2009. J Appl Soc Psychol 2012; 42: 1320–1334.

56. Zimet GD, Dahlem NW, Zimet SG and Farley GK. The multidimensional scale of perceived social support. J Pers Assess 1988; 52: 30–41.

57. Osman A, Lamis DA, Freedenthal S, Gutierrez PM and McNaughton-Cassill M. The multidimensional scale of perceived social support: analyses of internal reliability, measurement invariance, and correlates across gender. J Pers Assess 2014; 96: 103–112.

58. Zimet GD, Powell SS, Farley GK, Werkman S and Berkoff KA. Psychometric characteristics of the multidimensional scale of perceived social support. J Pers Assess 1990; 55: 610–617.

59. World Health Organization. Waist circumference and waist-hip ratio. Report of a WHO expert consultation, 8–11 December 2008. Geneva: World Health Organization.

60. Lin J, Epel E, Cheon J, Kroenke C, Sinclair E, Bigos M, Wolkowitz O, Mellon S and Blackburn E. Analyses and comparisons of telomerase activity and telomere length in human T and B cells: insights for epidemiology of telomere maintenance. J Immunol Methods 2010; 352: 71–80.

61. Cawthon R. Telomere measurement by quantitative PCR. Nucleic Acids Res 2002; 30: e47.

62. Lynch SM, Peek MK, Mitra N, Ravichandran K, Branas C, Spangler E, Zhou W, Paskett ED, Gehlert S, DeGraffinried C, Rebbeck TR and Rietham H. Race, ethnicity, psychosocial factors, and telomere length in a multicenter setting. PLoS One 2016; 11: e0146723.

63. Meng X-i, Rosenthal R and Rubin DB. Comparing correlated correlation coefficients. Psychol Bull 1992; 111: 172–175.

64. Muezzinler A, Zaineddin AK and Brenner H. A systematic review of leukocyte telomere length and age in adults. Ageing Res Rev 2013; 12: 509–519.

65. Thompson KA, bulls HW, Sibille KT, Bartley EJ, Glover TL, Terry EL, Vaughn IA, Cardoso JS, Sotolongo A, Staud R, Hughes LB, Edberg JC, Redden DT, Bradley LA, Goodin BR and Fillingim RB. Optimism and psychological resilience are beneficially associated with measures of clinical and experimental pain in adults with or at risk for knee osteoarthritis. Clin J Pain 2018; 34: 1164–1172.

66. Liu H, Zhang C, Ji Y and Yang L. Biological and psychological perspectives of resilience: is it possible to improve stress resistance? Front Hum Neurosci 2018; 12: 326.

67. Puterman E and Epel E. An intricate dance: life experience, multisystem resiliency, and rate of telomere decline throughout the lifespan. Soc Personal Psychol Compass 2012; 6: 807–825.

68. Holtzman S, Newth S and Delongis A. The role of social support in coping with daily pain among patients with rheumatoid arthritis. J Health Psycho 2004; 9: 677–695.

69. Gil KM, Keefe FJ, Crisson JE and Van Dalsen PJ. Social support and pain behavior. Pain 1987; 29: 209–217.

70. Wilson SJ, Woody A, Padin AC, Lin J, Malarkey WB and Kiecolt-Glaser JK. Loneliness and telomere length: immune and parasympathetic function in associations with accelerated aging. Ann Behav Med. Epub ahead of print 13 April 2018. DOI: 10.1093/abm/kay064.

71. Shankar A, McMunn A, Demakakos P, Hamer M and Steptoe A. Social isolation and loneliness: prospective associations with functional status in older adults. Health Psycho 2017; 36: 179–187.

72. Shankar A, McMunn A, Banks J and Steptoe A. Loneliness, social isolation, and behavioral and biological
health indicators in older adults. *Health Psychol* 2011; 30: 377–385.

73. Uchino BN. Social support and health: a review of physiological processes potentially underlying links to disease outcomes. *J Behav Med* 2006; 29: 377–387.

74. Dantzer R, Cohen S, Russo SJ and Dinan TG. Resilience and immunity. *Brain, Behav Immun* 2018; 74: 28–42.

75. Puterman E, Weiss J, Lin J, Schilf S, Slusher AL, Johansen KL and Epel ES. Aerobic exercise lengthens telomeres and reduces stress in family caregivers: a randomized controlled trial—Curt Richter Award Paper 2018. *Psychoneuroendocrinology* 2018; 98: 245–252.

76. Starkweather AR, Alhaeeri AA, Montpetit A, Brumelle J, Filler K, Montpetit M, Mohanraj L, Lyon DE and Jackson-Cook CK. An integrative review of factors associated with telomere length and implications for biobehavioral research. *Nurs Res* 2014; 63: 36–50.

77. National Institute of Environmental Health Sciences and National Institute on Aging. *Telomeres as sentinels for enviromental exposures, psychosocial stress, and disease susceptibility*. 2017. Research Triangle Park, NC: National Institute of Environmental Health Sciences and National Institute on Aging.