BRIEF REPORT

The Effect of Early Discrimination on Accelerated Aging Among African Americans

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Objective: This study examined the role of depressive symptoms in mediating the relationship between early life experiences of racial discrimination and accelerated aging in adulthood for African Americans (i.e., prediction over a 19-year period, from ages 10 to 29) after adjusting for gender and health behaviors.

Method: Longitudinal self-report data over 7 waves of data collection from the Family and Community Health Study were utilized. The sample included 368 African Americans with usable gene expression data to compute accelerated aging, as well as complete data on all self-report variables including racial discrimination (Schedule of Racist Events) and depression (Diagnostic Interview Schedule for Children—Version 4). Blood was collected by antecubital blood draws from participants at age 29. The proposed model was tested by path analysis.

Results: Findings revealed that high discrimination at ages 10–15 was associated with depression at ages 20–29 ($b = .19, p = .001$), controlling for depression at ages 10–15, which, in turn, was related to accelerated cellular-level aging ($b = .11, p = .048$) after controlling for gender, alcohol consumption, and cigarette use. The indirect effect of racial discrimination on aging through depression at ages 20–29 was significant ($b = .021, 95\%$ confidence interval [.001, .057]), accounting for 32.3\% of the total variance. Conclusion: These findings support research conceptualizations that early life stress due to racial discrimination lead to sustained negative affective states continuing into young adulthood that confer risk for accelerated aging and possibly premature disease and mortality in African Americans. These findings advance knowledge of potential underlying mechanisms that influence racial health disparities.

Keywords: discrimination, health, racism, African Americans, aging

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Research findings have suggested that early life stressful experiences have a lasting impact on biological risk for disease and premature mortality due to chronic diseases of aging (Berg, Simmons, Barr, Beach, & Philibert, 2017). Increasingly, researchers have suggested that early life exposure to psychosocial stressors may influence negative affective states (e.g., depression) and have effects on poor physical health through physiological dysregulation and increased wear and tear on body systems over time (Leger, Charles, & Almeida, 2018; Tomfohr, Pung, & Dimsdale, 2016).

An early and pervasive stressor related to health and mental health for African Americans is racial discrimination (Lewis, Cogburn, & Williams, 2015). Racial status is said to be one of the first social categories that young children learn, preceded only by...
learning to distinguish sex (Quintana & McKown, 2012). In addition, recent research with youth has suggested particularly strong effects of early exposure to discrimination (Gibbons et al., 2014), possibly due to stimulation of “fight or flight” responses and chronic vigilance for danger, disrupting the body’s regulatory systems and leading to increased vulnerability to chronic disease and depression (Williams, Neighbors, & Jackson, 2003). Physical dysregulation (e.g., autonomic nervous system dysfunction), particularly dysregulation of the hypothalamic-pituitary-adrenocortical axis, has also been associated with psychosocial stress and disease. Consistent with this process, Geronimus (2001) hypothesized that due to repeated exposure to race-related stressors, African Americans would be “weathered,” with greater exposure associated with increased acceleration of biological aging.

To measure accelerated aging, a recently developed gene expression index of aging developed by Peters et al. (2015) was utilized. This transcriptome index consists of 1,497 sites where the level of gene expression (amount of mRNA) is associated with increasing age. The sites included in the transcriptome index are located in gene pathways with known aging mechanisms, including metabolic functioning, immune senescence, and mitochondrial decline. This suggests that the transcriptome index assesses gene expression patterns of functional significance for aging and is associated with other biomarkers of metabolic—cardiovascular risk for chronic illness, such as systolic and diastolic blood pressure, total and HDL cholesterol, glucose levels, and body mass index (Peters et al., 2015). Accordingly, the transcriptome index appears well suited to capturing the process of accelerated aging and resulting risk for premature expression of chronic diseases of aging among African Americans and others.

Research has also established an association between depression and chronic conditions, especially cardiovascular disease (Penningx, 2017), with recent work indicating an association of major depression with accelerated aging (Han et al., 2018). In addition, sustained and chronic negative affective states in response to daily stressors are related to increases in chronic health conditions and functional limitations across 10 years (Leger et al., 2018). Smith-Bynum, Lambert, English, and Ialongo (2014) examined longitudinal trajectories of racial discrimination experiences among African Americans and found that African American adolescents who reported higher levels of discrimination were 4 times more likely to be in an increasing depression trajectory than were African American youth who reported consistently low levels of discrimination. Together these findings suggest the need to test the mediating role of sustained increases in depression on accelerated aging in an African American sample.

The current study builds on previous research demonstrating a link between racial discrimination and accelerated aging (Lee, Kim, & Neblett, 2017). This study also utilizes a sample that has been well characterized in prior reports. The present study hypothesized that racial discrimination would be related to accelerated aging such that a greater frequency of racial discrimination experiences during childhood would be associated with accelerated aging in adulthood for African Americans. This study also hypothesized that depressive symptoms would mediate the relationship between racial discrimination and accelerated aging. Gender was controlled for in this study due to previous research demonstrating gender differences in the report of stressful experiences that can influence health (Bale & Epperson, 2015).

Method

Participants

The hypotheses for the current study were tested using participants from the longitudinal Family and Community Health Study (FACHS), a multisite and nonclinical study of neighborhood and family effects on health and development in African American families (see Gibbons et al., 2014). Participants were recruited from rural, suburban, and metropolitan communities. Youth and their families were approached for participation in FACHS when youth were in the fifth grade (mean age = 10.56 years).

Procedure

The protocol and all study procedures were approved by the Institutional Review Board at the University of Georgia and Iowa State University. Written informed consent was obtained from all of the study respondents. To enhance rapport and cultural understanding, African American university students and community members served as field researchers to collect data from the families in their homes. Audio-enhanced, computer-assisted, self-administered interviews were used to assess all self-report material. At Wave 7 a certified phlebotomist performed antecubital blood draws at each participant’s home using a PAXgene tube. Samples were kept frozen at −80 °C until used for the analyses described later. All (N = 470) available samples were checked for quality and usability at the Rutgers repository (for further description of the FACHS sample and recruitment, see Appendix I in the online supplemental materials).

Measures

Racial discrimination. At Wave 1 of data collection, the target youths completed the 13-item revised version of the Schedule of Racist Events (Landrine & Klonoff, 1996) to assess frequency of specific discriminatory behaviors during the past year, including racially based slurs, insults, and physical threats. Cronbach’s alpha for the scale was .82.

Depressive symptoms ages 10–15. At each wave (1, 2, and 3) from ages 10 to 15, target youths completed the Diagnostic Interview Schedule for Children—Version 4 (DISC–IV; Shaffer et al., 1993) to assess frequency (all yes/no) of feeling sad, irritable, tired, restless, or worthless and other depressive symptoms (e.g., “In the last year, was there a time when you . . . often felt sad or depressed?”), Cronbach’s alpha for this measure exceeded .83 for each wave of data collected.

Depressive symptoms ages 20–29. In adulthood (Waves 5, 6, and 7), depression was assessed using the nine-item Composite International Diagnostic Interview (CIDI; Kessler et al., 1994) measure of depressive symptoms. Respondents were asked to report (0 = no; 1 = yes) whether they experienced symptoms of depression (e.g., “felt sad, empty, or depressed most of the day”) for at least a two-week period in the past year. Cronbach’s alpha for the scale was .83.

Control variables age 29. Health-related covariates were statistically controlled for in this study to exclude plausible rival explanations. In adulthood (Wave 7), alcohol consumption was measured by asking “During the past 12 months, how often have
you had a lot to drink, that is 3 or more drinks at one time?” Also in adulthood (Wave 7), cigarette use was measured by asking “How many cigarettes have you smoked in the last 3 months?” The response categories ranged from 1 (0 days) to 5 (all 7 days).

Transcriptional index of biological age. Biological age was measured using the transcriptomic clock developed by Peters et al. (2015). After excluding samples with poor quality (n = 81) or no amplification (n = 3), the Rutgers repository identified a total sample of N = 386 young adults with usable samples. Probe data yielded a microarray data set of 47,323 probes, which was filtered by removing probes with detection threshold of p ≤ .05, leaving 44,846 probes for analysis. After quantile normalization, data were log2-transformed and inspected visually for batch effects. Transcriptomic age was calculated using publicly available software (Transcription Age Prediction, 2016). Mean biological age was 29.49 (SD = .22). Accelerated aging was determined using the residual scores from the regression of biological age on chronological age. These residuals had a mean of 0, with positive scores indicating accelerated aging (for further information see Appendix II in the online supplemental materials).

Analytic Strategy

Initial data cleaning was conducted using SPSS; descriptive statistics and characterization of intercorrelations were conducted using Mplus; path modeling in Mplus was used to test the model. Steiger’s root-mean-square error of approximation (RMSEA < .05) and the comparative fit index (CFI > .90) were used to assess goodness of fit of the model. To evaluate the significance of the hypothesized indirect effect, we estimated the 95% confidence interval (CI) with bias-corrected and accelerated bootstrapping with 1,000 replications.

Results

All (N = 470) participants in the final sample with both self-report and a blood draw at age 29 (mean age = 29.49) self-identified as Black or African American. Of these, 368 provided usable data regarding gene expression as well as responses to all self-report data and therefore constitute the final analytic sample. The final sample (63% female) can be characterized as low to moderate in annual income (M = $20,991.39) and having moderate average educational attainment, with 91% graduating high school. In addition, most were employed (80.2%), had health insurance (79.3%), and had .72 children on average, with 64.4% of participants’ reporting a committed romantic partner relationship at age 29 (Supplemental Table 1 in the online supplemental materials displays means, standard deviations, and the zero-order correlation among the study variables). There was a significant correlation between racial discrimination and young adult depression (r = .236, p = .000). Young adult depression was significantly correlated with aging (r = .133, p = .011). The control variables showed no correlation with aging.

The results of the examination of indirect effects of racial discrimination on aging through young adult depression (ages 20–29) can be seen in Figure 1. The various fit indices, using root-mean-square error of approximation (RMSEA < .05) and the comparative fit index (CFI > .90) suggest that the model provides a good fit to the data, χ²(1, N = 368) = 1.716, p = .1902, RMSEA = .044, CFI = .986. Racial discrimination is related to young adult depression (β = .185, 95% CI [.084, .286], p = .000), which, in turn, is related to aging (β = .134, 95% CI [.028, .242], p = .011) after controlling for gender, alcohol consumption, and cigarette use. The test of the indirect effect of racial discrimination on aging through young adult depression was significant (β = .025, 95% CI [.005, .062]), accounting for 32.05% of the total variance.

Discussion

A main objective of this research was to longitudinally examine the role of depressive symptoms as a mechanism influencing the relationship between early life experiences of racial discrimination and accelerated aging among African Americans. Consistent with the study hypothesis, experiences of racial discrimination during childhood (ages 10–15) contributed to accelerated aging in adulthood (age 29), and effects were mediated through their impact on young adult depression (ages 20–29). Not shown, the effects of sex, alcohol consumption, and cigarette use are controlled in these analyses. The indirect effect is significant (β = .030, 95% confidence interval [.008, .070]), χ²(1, N = 368) = 1.716, p = .1902, comparative fit index = .986; root-mean-square error of approximation = .044. Values are standardized parameter estimates, and standard errors are in parentheses. Dashed arrow indicates the non-significant direct effect of discrimination on accelerated mRNA aging. * p ≤ .05. ** p ≤ .01.

Figure 1. Effects of early racial discrimination (ages 10–15) on accelerated aging (age 29) mediated through young adult depression (ages 20–29). Not shown, the effects of sex, alcohol consumption, and cigarette use are controlled in these analyses. The indirect effect is significant (β = .030, 95% confidence interval [.008, .070]), χ²(1, N = 368) = 1.716, p = .1902, comparative fit index = .986, root-mean-square error of approximation = .044. Values are standardized parameter estimates, and standard errors are in parentheses. Dashed arrow indicates the non-significant direct effect of discrimination on accelerated mRNA aging. * p ≤ .05. ** p ≤ .01.
increased depressive symptoms in young adulthood (ages 20–29), even when controlling for the effects of gender and health behaviors (i.e., alcohol consumption and cigarette use). The current findings are consistent with the hypothesis that poor health and health disparities during adulthood may result from stressful experiences earlier in life, such as early experiences of racial discrimination (Williams et al., 2003). These findings also suggest that when African American children experience racial discrimination their appraisal–coping responses could be depressive states, and in turn, these depressive states can influence biological risk in adulthood. Taking a life course perspective, this study highlights the need to consider childhood as a sensitive period during development whereby stressful adverse experiences, like racial discrimination, are biologically embedded in body systems in a lasting form that makes some African American children more susceptible to poor health later in life (Umberson, Williams, Thomas, Liu, & Thomeer, 2014). One marker of this biological embedding may be elevated levels of depressive symptoms in young adulthood, suggesting a potential area of therapeutic intervention.

Although this study makes several contributions to the available literature, some limitations should be noted. First, a limitation involves the sample utilized for the study. This study focused on African Americans living in towns and small cities and therefore was not a nationally representative sample. Second, the study did not provide diagnoses of depression, so effects attributed to depression may be due to negative affective processes more generally rather than being specific to depressive symptoms. Finally, the model tested was a simple mediational model and did not examine the range of potential buffers and resilience processes that may serve to protect against the effect of early racial discrimination exposures. Conversely, it also did not examine potential additive or multiplicative effects attributable to additional sources of early stress or developmental vulnerabilities. Future studies should address this complexity and better map the influence of different adverse experiences and protective processes on accelerated aging among marginalized and underserved populations.

References


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