Accelerated epigenetic aging mediates link between adverse childhood experiences and depressive symptoms in older adults: Results from the Health and Retirement Study

Eric T. Klopack a,⁎, Eileen M. Crimmins a, Steve W. Cole b, Teresa E. Seeman b, Judith E. Carroll b

a University of Southern California, United States
b University of California, Los Angeles, United States

ARTICLE INFO

Keywords:
- Aging
- Depression
- Adverse childhood events
- Epigenetic aging

ABSTRACT

Adverse childhood experiences (ACEs) increase risk for depression at subsequent ages and have been linked to accelerated biological aging. We hypothesize that accelerated epigenetic aging may partially mediate the link between ACEs and depression. This study examines three second-generation epigenetic aging measures (viz., GrimAge, PhenoAge, and DunedinPoAm38) as mediators of the link between ACEs and depressive symptoms in older adulthood. We utilize structural equation modeling to assess mediation in the Health and Retirement Study (N = 2672). Experiencing ACEs is significantly associated with an older GrimAge and a faster pace of aging via the DunedinPoAm38. Having an older GrimAge and faster DunedinPoAm38 pace of aging were also significantly associated with more depressive symptoms. PhenoAge was not significantly associated with depressive symptoms and was only associated with experiencing three ACEs. These associations were reduced by socioeconomic and lifestyle factors, including obesity and substance use. GrimAge explained between 9 and 14% of the association between ACEs and adult depressive symptoms, and DunedinPoAm38 explained between 2 and 7% of the association between ACEs and adult depressive symptoms. Findings indicate accelerated aging, as measured by GrimAge and DunedinPoAm38, is associated with ACEs and with depressive symptoms in older Americans. Findings also show these epigenetic aging measures mediate a portion of the association between ACEs and adult depressive symptoms. Epigenetic aging may represent a physiological mechanism underlying the link between early life adversity and adult depression. Weight maintenance and substance use are potentially important areas for intervention.

There is a wealth of evidence adverse childhood experiences (ACEs) have serious implications for mental health across the lifespan (Centers for Disease Contr, 2019). A large body of research investigating the “long arm of childhood” has pointed to a significant role of early life experiences raising lifetime risk for mood and anxiety disorders, as well as earlier onset of age related diseases (Kalmakis & Chandler, 2015). Experiencing more ACEs increases risk of major depressive disorder and depressive symptoms in adulthood (Merrick et al., 2017). This is important, as depression was recently shown to be the leading cause of disability worldwide (Friedrich, 2017), the estimated economic burden of adults with major depressive disorder in the US is $326.2 billion (Greenberg et al., 2021), and mental wellbeing is important for later life health (Moffitt & Caspi, 2019; Richmond-Rakerd et al., 2021). Emerging evidence suggests biological aging may be an important mediator linking ACEs and depression (Entringer & Epel, 2020; Mitchell et al., 2016; Sumner et al., 2019). Epigenetic aging—a methylation-based measure of aging—is thought to begin early in life and lead to health consequences later in life. Accumulating evidence suggests exposure to ACEs accelerates aging, setting one on a trajectory for early onset of aging related deterioration of function and multi-morbidities, thus contributing to health disparities across the life course (Belsky et al., 2017), and potentially risk for psychopathology (Wolkowitz et al., 2008), although data supporting this link is limited (Epel & Prather, 2018). In this model, early social stress alters cellular processes (e.g., metabolic activity, cellular stress, damage accumulation) that are associated with epigenetic age acceleration.

Abbreviations: ACEs, Adverse childhood events.
⁎ Corresponding author. 3715 McClintock Ave, Los Angeles, CA, 90089, United States.
E-mail address: klopack@usc.edu (E.T. Klopack).

https://doi.org/10.1016/j.ssmph.2022.101071
Received 17 December 2021; Received in revised form 11 March 2022; Accepted 12 March 2022
Available online 16 March 2022
2352-8273/© 2022 The Authors. Published by Elsevier Ltd. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).
Past research has shown epigenetic aging is accelerated in people with major depressive disorder (Ekaterina et al., 2021), depressive symptoms (Sumner et al., 2019), and PTSD (Yang et al., 2020); however, the role of early adversity in risk for psychiatric symptoms and the mechanisms that drive this are less clear. Epigenetic clocks summarize changes in DNAm-based markers associated with age and health outcomes common at older ages (Raffington et al., 2021). These clocks represent sets of DNAm sites that have been found to be differentially associated with chronological age and/or age-related biomarkers and diseases (Horvath & Raj, 2018). The first-generation clocks—HorvathAge and HannumAge—were trained to predict only chronological age and show relatively weak associations with clinical measures of physiological dysregulation and mortality (Horvath & Raj, 2018). In contrast, second generation clocks (e.g., GrimAge, PhenoAge), trained on age and age related morbidity/mortality-relevant biomarkers, and pace of aging measures (e.g., DunedinPoAm38), trained on change in age-relevant biomarkers, have been shown to be associated with chronic disease morbidity and mortality in adults (Fransquet et al., 2019).

Past research on the association between childhood adversity and epigenetic aging measures has been inconsistent. Past research has reported both significant (Colich et al., 2020; Jovanovic et al., 2017; Lawn et al., 2018; Marini et al., 2020) and null (Austin et al., 2018; Fiorito et al., 2017; Simons et al., 2016; Verhoeven et al., 2018) results using the first generation clocks. Research using the second generation clocks has been more consistently supportive (Belsky et al., 2020; George et al., 2021; Hamlat et al., 2021); however, prior research using the Health and Retirement Study suggests childhood adversity only accounts for a small proportion of PhenoAge (Liu et al., 2019). Past investigations have been limited because they typically investigate only one epigenetic aging measure in a given analysis. These measures were trained in different populations, using different aging criteria (e.g., biomarkers, chronological age), and/or in different tissues. Though they theoretically all measure aging (or pace of aging), there is very little overlap in the probes utilized. Though they tend to agree on direction of effect, they vary greatly in magnitude of association with psychiatric conditions; therefore, some scholars have suggested the epigenetic aging measures may be capturing different elements of the aging process (Oblak et al., 2021). Thus, research comparing multiple clocks is needed to identify which best characterize the biological mechanisms underlying this association. This research could help clarify the complex biological processes underlying psychiatric conditions (Oblak et al., 2021).

To our knowledge, only one study has directly tested epigenetic aging as a mediator of the link between childhood adversity and depressive symptoms (Sumner et al., 2019). This study was limited to measures of epigenetic aging using first generation clocks and was conducted in children. The current study builds on this past research by utilizing second generation clocks and pace of aging in a nationally representative sample of older Americans to test the hypothesis that early life adversity is related to accelerated epigenetic aging and test whether epigenetic aging mediates the effect of ACEs on depressive symptoms. This study advances this past research by 1) examining the association between ACEs and three validated epigenetic aging measures that have been shown to be predictive of a variety of age-related health outcomes in a nationally representative sample of older adults in the US and 2) assessing epigenetic aging as a mediator of the association between ACEs and depressive symptoms. To our knowledge, this is the first study examining either of these issues in a nationally representative sample of older US adults.

1. Methods and materials

1.1. Sample

We utilize data from the DNA methylation subsample from the Health and Retirement Study (HRS) 2016 Venous Blood Study (N = 4018) (Health & Retirement Study, 2021). Detailed methods for this sample are published elsewhere (Crimmins et al., 2017). This sample was designed to be representative of the U.S. population when weighted. Items used to construct ACEs scores were assessed as part of the psychosocial leave behind questionnaire at various waves and the life history surveys in 2015 and 2017. Depressive symptoms were assessed in the core survey in 2016 and 2018. About 1000 participants did not participate in the life history mail surveys and about 550 did not participate in the psychosocial leave behind questionnaire from 2006 to 2012. 2672 participants were included in the current study due to missingness in independent variables described below.

1.2. Measures

Epigenetic aging. The epigenetic aging measures were developed using different tissues, were trained on different outcomes, and vary in how predictive they are of certain outcomes (Crimmins et al., 2021). GrimAge is trained on DNAm surrogates of 7 plasma proteins and smoking pack years in multiple tissues (Lu et al., 2019). DunedinPoAm38 was trained on changes in a variety of health-related biomarkers and is meant to capture the current pace of epigenetic aging (Belsky et al., 2020), and PhenoAge was developed to estimate mortality risk using 9 markers of tissue and immune function and age in whole blood (Levine et al., 2018). GrimAge and PhenoAge are expressed in years of epigenetic age. Because DunedinPoAm38 is designed to assess pace of aging, it is expressed in years of epigenetic aging per chronological year. We use the epigenetic aging measures derived from the 2016 HRS Venous Blood Study which are available as restricted health data at https://hrdata.isr.umich.edu/data-products/epigenetic-clocks. More detailed information about the epigenetic aging measures is available elsewhere (Crimmins et al., 2020).

ACEs. An ACEs scale was constructed using 8 items that were included in the core survey and life history surveys. Items were selected that reflect the ACEs scale developed by the Centers for Disease Control and Prevention and childhood socioeconomic status (CDC, 2019). Specifically, participants reported on parental education (both parents < high school or one parent < high school if only one parent was reported), parental physical abuse (were you ever physically abused by either of your parents?), and parental alcohol and drug use (did either of your parents drink or use drugs so often that it caused problems in the family?) before age 18, as well as childhood separation from either parent (were you ever separated from your mother or father for 6 months or longer?), death of a parent (did one or both parents die?), living in a children’s home or orphanage (did you ever live in a children’s home or orphanage?), childhood poverty (would you say your family during that time was pretty well off financially, about average, or poor? poor = 1), parental separation or divorce (did your biological or adoptive parents separate or divorce?) before age 16. To address item nonresponse, participants were allowed to be missing on one item (coded as 0). Alternative coding schemes produced similar results. Because few participants reported more than 4 ACEs, this measure was top coded at 4 or more ACEs.

Consistent with past research on ACEs and health, this measure of ACEs was associated with greater physical limitations assessed using activities of daily life and instrumental activities of daily life and was associated with a greater chronic illness count after controlling for age,
race, and gender (see Table S6; outcome measures were taken from the 2016 RAND data).

**Depressive Symptoms** were assessed in the 2016 and 2018 core survey using the Center for Epidemiologic Studies Depression Scale (CES-D) (Radloff, 1977). The HRS uses an 8-item subset of the CES-D in which respondents report whether or not they experience 8 depressive symptoms over the past two weeks. This measure has been shown to have similar psychometric properties to the original CES-D (Turvey et al., 1999). 2016 scores were used for participants without 2018 scores.

**Socioeconomic Position.** Socioeconomic position was assessed using self-reports of education (categorized: 0–11 years, 12 years, 13–15 years, and 16+ years as the reference) and wealth (natural log transformed to account for skewness) using the wealth measure available in the HRS RAND data including all assets less debt (Bogliari et al., 2016).

**Lifestyle Factors.** We assessed the role of lifestyle factors using self-reported smoking (categorized: current smoker, past smoker, and non-smoker as the reference), BMI (categorized: ≥25 to <30 overweight, ≥30 to <35 obese I, ≥35 obese II or morbidly obese, and <25 normal and underweight as the reference), and alcohol use (categorized: 1–4 drinks per day drinking, 5+ drinks per day drinking, and non-drinker as the reference). BMI is an imperfect measure of lifestyle factors, so we estimated the SEMs with lifestyle controls with BMI replaced by physical activity (using the mean of the three physical activity items in HRS for 2016 or 2014 if 2016 data were missing). Results were similar to the original results with a nearly identical pattern of significant results. The sole difference in significance pattern was that GrimAge was not significantly associated with depressive symptoms (this association was significant in the model with BMI). Thus, BMI appears to capture similar variance as this physical activity measure.

**Controls.** In all models we control for chronological age, race/ethnicity (non-Hispanic white as reference), and gender (male as reference).

### 1.3. Plan of analysis

We began by estimating three structural equation models (SEMs) regressing each epigenetic clock first on ACEs and controls and regressing depressive symptoms on each epigenetic aging measure and controls. Mediation was tested using a bias corrected bootstrap procedure with 1000 draws. To assess the impact of lifestyle factors on these findings, we next estimated the same models controlling for lifestyle factors. Because all models were fully recursive, model fit statistics were all artificially perfect and therefore uninformative. Epigenetic aging measures and depressive symptoms were standardized in Mplus for the figures to ease comparison across measures using the STANDARDIZE command. Because the epigenetic measures all use meaningful scales, unstandardized results are discussed in the text below. All analyses were conducted in Mplus 8 (Muthén & Muthén, 1998).

We assessed gender differences by comparing models for men and women using the “GROUPING IS” command in Mplus. For each model in the study, we compared a model constraining all paths from ACEs to epigenetic measures and all paths from epigenetic measure to depressive symptoms to be equal for men and women with a model in which all these paths were freed. There was no significant $\chi^2$ change across all of these models suggesting that there is not a gender difference in these models.

2. Results

**Descriptive Statistics.** The weighted sample is 56.6% female and has a median age of 67 years. The weighted sample is 83.4% Non-Hispanic White, 8.4% Non-Hispanic Black, 5.7% Hispanic, and 2.5% Non-Hispanic Other Race. 10.4% of the weighted sample has less than 12 years of education, 31.5% has 12 years of education, 26.5% has 13–15 years of education, and 31.6% has 16+ years of education. The weighted sample had a median wealth of $232,119. 8.9% of the weighted sample are current smokers and 45.0% are former smokers. More than a third of the weighted sample is obese (35.4%), and 54.6% were non-drinkers. Descriptive statistics are shown in Table 1.

### ACEs and Depressive Symptoms.** The total effects of ACEs on depressive symptoms are shown in Table 2, Panel A. Results with epigenetic age and depressive symptoms standardized are shown in the tables and figure to ease comparison across measures, and unstandardized results are discussed in the text. The total effects of 2, 3, and 4 or more ACEs on depressive symptoms were statistically significant in the expected direction. Thus, in total, having 2 or more ACEs was associated with having more depressive symptoms compared to 0 ACEs. People who experienced 2 ACEs were expected to have 0.29 more depressive symptoms, and people who experienced 3 or 4+ ACEs were expected to experience 0.73 and 0.72 more depressive symptoms compared to similar peers who experienced 0 ACEs.

**GrimAge SEM Results.** SEM results for GrimAge are shown in Fig. 1, Panel A. Overall, having experienced childhood adversity was associated with an older GrimAge. Experiencing 1 or 2 ACEs was associated with having a GrimAge 0.87 and 0.88 years greater compared to those who experienced no ACEs. Having 3 ACEs was associated with a

| Variable | Mean/Proportion | SD | Range | Count |
|----------|-----------------|----|-------|-------|
| ACEs     |                 |    |       |       |
| 0 ACEs   | 0.34            |    |       | 775   |
| 1 ACE    | 0.28            |    |       | 711   |
| 2 ACEs   | 0.19            |    |       | 566   |
| 3 ACEs   | 0.11            |    |       | 357   |
| 4+ ACEs  | 0.09            |    |       | 263   |
| Depressive Symptoms | | | | |
| Race/Ethnicity | | | | |
| White, not | 0.83            |    |       | 1978  |
| Hispanic Black, not | 0.08            |    |       | 378   |
| Hispanic Hispanic | 0.06            |    |       | 257   |
| Other Race, not Hispanic | 0.03            |    |       | 59    |
| Gender | | | | |
| Male | 0.43            |    |       | 1064  |
| Female | 0.57            |    |       | 1608  |
| Education | | | | |
| Less than High School | 0.10            |    |       | 397   |
| High School | 0.32            |    |       | 876   |
| Some College | 0.27            |    |       | 680   |
| College or More | 0.32            |    |       | 719   |
| Wealth | 601,542         | 1,396,212 | -249,750–24,475,000 | |
| Smoker Status | | | | |
| Never Smoked | 0.46            |    |       | 1216  |
| Current | 0.09            |    |       | 252   |
| Smoker | | | | |
| Past Smoker | 0.45            |    |       | 1204  |
| BMI | | | | |
| Under/Nor | 0.27            |    |       | 725   |
| Weight | | | | |
| Overweight | 0.37            |    |       | 982   |
| Obese 1 | 0.22            |    |       | 581   |
| Obese 2 | 0.14            |    |       | 384   |
| Drinking | | | | |
| Non-drinker | 0.55            |    |       | 1588  |
| 1-4 Drinks per Day Drinking | 0.43            |    |       | 1031  |
| 5+ Drinks per Day Drinking | 0.02            |    |       | 53    |

Note: means and standard deviations are weighted using survey weights provided by HRS. Counts are not weighted.
GrimAge 1.47 years older than those who experienced no ACEs. Finally, those exposed to 4 or more ACEs were expected to have a GrimAge 2.18 years older than those with no ACEs.

GrimAge was significantly, positively associated with depressive symptoms. Increasing GrimAge was associated with increasing depressive symptoms ($b = 0.04$, $p < .001$). Thus, experiencing more ACEs was associated with greater epigenetic aging as measured by GrimAge, and more accelerated aging was associated with more depressive symptoms.

Indirect effects for the GrimAge model are shown in Table 2, Panel B. All four indirect effects that were tested were significant; however, because the total effect of 1 ACE was not significant, we focus on the other three indirect effects. In this model, GrimAge mediated about 14% of the effect of experiencing 2 ACEs on depressive symptoms, about 9% of the effect of 3 ACEs, and about 13% of the effect of 4+ ACEs on depressive symptoms.

**DunedinPoAm38 SEM Results.** SEM results for DunedinPoAm38 (Fig. 1, Panel B) were largely consistent with findings for GrimAge. People who experienced 1 ACE were expected to have a DunedinPoAm38 pace of aging about a tenth of year per chronological year faster than similar people who experienced no ACEs, and people who experienced 4 or more ACEs were expected to have a DunedinPoAm38 pace of aging about 4 tenths of year per chronological year faster than similar people who experienced no ACEs. The effects of experiencing 2 or 3 ACEs on DunedinPoAm38 were only marginally significant.

A faster pace of aging, as determined using the DunedinPoAm38, was significantly associated with more depressive symptoms. Having a DunedinPoAm38 pace of aging 1 year per chronological year greater than same-age peers was associated with a 1.18 higher score on this measure of depressive symptoms.

Indirect effects for DunedinPoAm38 are shown in Table 2, Panel C. Indirect effects for 1, 3, and 4 or more ACEs were significant; again, because the total effect of 1 ACE was not significant, we focus on the other indirect effects. DunedinPoAm38 appears to mediate about 2% of the effect of 3 ACEs and about 7% of the effect of 4+ ACEs on depressive symptoms.

**PhenoAge SEM Results.** Results for PhenoAge (Fig. 2, Panel C) were largely unsupportive. Having experienced 3 ACEs was associated with having a PhenoAge 1.46 years greater than similar peers who experienced 0 ACEs, but no other comparisons were significant. Additionally,

---

**Table 2 Direct and indirect effects from structural equation models.**

| Panel A. Total Effect of ACEs on Depressive Symptoms | Total Effect | SE | p   |
|-------------------------------------------------------|-------------|----|-----|
| 1 ACE                                                 | 0.048       | 0.055 | 0.385 |
| 2 ACEs                                                | 0.155       | 0.058 | 0.008 |
| 3 ACEs                                                | 0.393       | 0.112 | <0.001 |
| 4+ ACEs                                               | 0.387       | 0.108 | <0.001 |

| Panel B. Indirect Effects of ACEs on Depressive Symptoms via GrimAge | Indirect Effect | 95% CI    |
|---------------------------------------------------------------------|----------------|-----------|
| 1 ACE                                                               | 0.020          | 0.011–0.035|
| 2 ACEs                                                              | 0.021          | 0.007–0.039|
| 3 ACEs                                                              | 0.034          | 0.019–0.058|
| 4+ ACEs                                                             | 0.051          | 0.023–0.090|

| Panel C. Indirect Effects of ACEs on Depressive Symptoms via DunedinPoAm38 | Indirect Effect | 95% CI    |
|--------------------------------------------------------------------------|----------------|-----------|
| 1 ACE                                                                     | -0.001         | -0.009–0.003|
| 2 ACEs                                                                   | 0.003          | -0.001–0.017|
| 3 ACEs                                                                   | 0.009          | -0.001–0.024|
| 4+ ACEs                                                                  | 0.027          | 0.006–0.058|

| Panel D. Indirect Effects of ACEs on Depressive Symptoms via PhenoAge   | Indirect Effect | 95% CI    |
|-------------------------------------------------------------------------|----------------|-----------|
| 1 ACE                                                                    | 0.001          | 0.009–0.003|
| 2 ACEs                                                                  | 0.003          | -0.001–0.017|
| 3 ACEs                                                                  | 0.007          | -0.001–0.024|
| 4+ ACEs                                                                 | 0.002          | -0.003–0.011|

Note: CI = confidence interval. 95% CIs were estimated using a bias corrected bootstrap procedure with 1000 draws. Epigenetic aging and depressive symptoms, but not ACEs, are standardized using the “STANDARDIZE” command in Mplus.

**Fig. 1. SEM Results.** Note: $N = 2672$; all models are fully recursive, but not all paths are shown for ease of presentation. ACE = adverse childhood experience. Epigenetic aging and depressive symptoms, but not ACEs, are standardized using the “STANDARDIZE” command in Mplus. Unstandardized results are discussed in the text above. *$p < .05$.
PhenoAge was not significantly associated with depressive symptoms. Unsurprisingly given the non-significant main effects, there are no significant indirect effects of ACEs on depressive symptoms mediated by PhenoAge (shown in Table 2, Panel D).

**Socioeconomic and Lifestyle Factors.** Finally, we estimated the same SEMs including socioeconomic and lifestyle factors as controls to assess how much these factors affect the pathways linking ACEs with epigenetic aging and depressive symptoms. Full results are included in the supplementary material.

Controlling for socioeconomic and lifestyle factors reduced the effect of ACEs on GrimAge to non-significance compared to the model without socioeconomic and lifestyle factors because of the associations of GrimAge with years of education (0–11 years; $b = 0.66$, $p = .04$; 12 years; $b = 0.91$, $p < .001$; 13–15 years; $b = 0.94$, $p < .001$), wealth ($b = -0.96$, $p = .001$), high obesity (BMI $\geq 35$; $b = 1.43$, $p < .001$), smoker status (current smoker; $b = 6.96$, $p < .001$; past smoker; $b = 2.09$, $p < .001$), and heavy alcohol use (5+ drinks per day drinking; $b = 1.58$, $p = .02$). Including lifestyle factors reduced the association between GrimAge and depressive symptoms (Table S1; $b = 0.02$, $p = .04$). Wealth ($b = -0.48$, $p < .001$) was significantly associated with depressive symptoms.

The pace of aging, as measured by DunedinPoAm38, of people who experienced 1, 2, or 3 ACEs were not significantly different from the pace of aging of similar peers who experienced no ACEs when controlling for socioeconomic and lifestyle factors. However, having 4 or more ACEs remained significantly associated with a more accelerated pace of aging compared to no ACEs, even after controlling for socioeconomic and lifestyle factors ($b = 0.02$, $p = .003$). Obesity (BMI $\geq 30$ and $< 35$; $b = 0.01$, $p = .03$; BMI $\geq 35$; $b = 0.03$, $p < .001$), and smoker status (current smoker; $b = 0.12$, $p < .001$; past smoker; $b = 0.03$, $p < .001$) were all significantly associated with faster DunedinPoAm38 in this model. For the DunedinPoAm38 model, pace of epigenetic aging was no longer significantly associated with depressive symptoms ($b = 0.38$, $p = .45$) after controlling for socioeconomic and lifestyle factors. Wealth ($b = -0.50$, $p < .001$) was significantly associated with depressive symptoms in this model.

For the PhenoAge model, the only significant effect of ACEs on PhenoAge (3 ACEs) was reduced to non-significance after including lifestyle factors. Wealth ($b = -0.91$, $p = .03$), obesity (BMI $\geq 35$; $b = 1.58$, $p = .002$), and heavy alcohol use (5+ drinks per day drinking; $b = 2.78$, $p = .011$) were all associated with PhenoAge in this model.

### 2.1. First-generation clocks

As noted above, we focus on the second-generation clocks in the current study. As a supplemental analysis, we estimated the same models using the main first-generation clocks (viz., HorvathAge and HannumAge, see Fig. S3, Tables S4 and S5). Neither clock was significantly associated with ACEs or depressive symptoms before or after controlling for socioeconomic and lifestyle factors.

### 3. Discussion

This study builds on and addresses weakness of past research by investigating the association between early life adversity and 3 second-generation DNAm clocks in a representative sample of Americans over age 50 and formally testing whether these clocks mediate the association between ACEs and depressive symptoms. Results indicate exposure to early life adversity is associated with accelerated epigenetic aging, as measured by GrimAge, and a faster pace of aging as measured by the DunedinPoAm38, compared to similar, same-aged peers who experienced no early life adverse events. Moreover, accelerated GrimAge and a faster pace of aging were significantly associated with elevated depressive symptoms. In mediation models, these clocks each
mediated a portion of the total effect of ACEs on depressive symptoms. Controlling for socioeconomic and lifestyle factors such as smoking and obesity weakened all of these associations, suggesting these factors may play an important role in this biopsychological process.

This study builds on a large body of past research examining the association between ACEs and mental and physical health, and a few studies linking ACEs to epigenetic age acceleration. Our findings are consistent with past findings that suggest second generation clocks are more consistently associated with ACEs in adults (George et al., 2021), potentially because these clocks were trained to predict phenotypic indicators of accelerated aging and are more consistently associated with age-related health problems (Horvath & Raj, 2018). This study also builds on this past body of research examining the association between ACEs and epigenetic clocks by utilizing a representative sample of US adults over the age of 50. These findings suggest that the association between ACEs and epigenetic aging may be non-linear. That is, experiencing 1 or 4 or more ACEs is associated with DunedinPoAm38 before controlling for lifestyle factors and only with 4 or more ACEs after controlling for lifestyle factors. It is possible that experiencing a very high number of ACEs is particularly damaging and may directly affect aging independent of health behaviors. Further research is needed to assess these findings.

These results suggest epigenetic effects of ACEs may persist into later life. One proposed pathway may be inflammation, as epigenetic aging—and senescent cells in particular—is a source of inflammatory signaling. Steeper inflammatory trajectories have been observed in adults who were exposed to child abuse (Renna et al., 2021). Inflammatory signaling from accelerated epigenetic aging may be a key pathway by which exposure to ACEs affects psychiatric symptoms in adulthood. These paths may be bidirectional, as past research and theory suggests stress, epigenetic aging, psychiatric disorders may mutually reinforce one another (Epe & Prather, 2018). Further research investigating this pathway is needed.

Another potential mechanism linking ACEs to accelerated epigenetic aging is socioeconomic and lifestyle factors. We found the effect of fewer than 4 ACEs was no longer significant and the effect of 4+ ACEs on GrimAge and DunedinPoAm38 was reduced after controlling for socioeconomic and lifestyle effects. Specifically, GrimAge was affected by education, wealth, high obesity, smoking, and alcohol use; whereas, DunedinPoAm38 was consistently affected by BMI and smoking. Thus, a portion of the effects of ACEs on epigenetic aging may be mediated by socioeconomic and lifestyle consequences of childhood adversity. This is consistent with a large body of research linking childhood adversity with BMI (Chu et al., 2018; Pudrovska & Anikputa, 2014; Wickrama et al., 2014), health risk behaviors (Campbell et al., 2016; Duffy et al., 2018; Merrick et al., 2017), and socioeconomic attainment (Font & Maguire-Jack, 2016; Liu et al., 2013; Yang et al., 2017) and research suggesting the effect of ACEs on depressive symptoms is mediated by these socioeconomic and lifestyle factors (Hughes et al., 2017; Jones et al., 2018; Merrick et al., 2017; Wickrama et al., 2014). Additionally, BMI (Visser et al., 1999), health risk behaviors (Shiel et al., 2014), and socioeconomic attainment (Castagne et al., 2020; Chen & Miller, 2013) have all been linked to inflammation, a potential pathway linking ACEs to depressive symptoms (Howren et al., 2009). Thus, these lifestyle factors, especially health behaviors, may be a useful area for intervention.

We also found GrimAge mediates about 8-13% of the association between 2 or more ACEs and depressive symptoms in later life. This finding is consistent with past research that found DNAm age significantly mediated the association between childhood threat exposure and depressive symptoms in children (Summer et al., 2019). Further research is needed to examine the mechanisms linking ACEs, DNAm age, and psychiatric symptoms across the life course. The association between GrimAge and depressive symptoms was substantially reduced and the association between DunedinPoAm38 was reduced to non-significance after controlling for socioeconomic and lifestyle factors. Obesity and smoking were relevant for both clocks, suggesting interventions targeting these factors may be useful for breaking the link between ACEs and depressive symptoms in later life.

Similar to a past study using the HRS, PhenoAge was not significantly associated with childhood adversity. This may be a result of the ways in which the different clocks were trained. PhenoAge was trained to predict mortality risk using markers of tissue and immune function (albumin, creatinine, serum glucose, CRP, lymphocyte percent, mean (red) cell volume, red cell distribution width, alkaline phosphatase, white blood cell count) (Levine et al., 2018). These indicators may be less sensitive to childhood conditions or may be more affected by recent stress and adversity compared to the indicators used to train GrimAge (DNAm surrogate measures of adrenomedullin, cystatin c, leptin, TIMP-1, pack years, plasminogen activation inhibitor 1 (PAI-1), growth differentiation factor 15 (GDF15), and beta 2 microglobulin (B2M) regressed on time to death)(Levine et al., 2018) and DunedinPoAm38 (rates of change in 18 biomarkers associated with aging) (Belinsky et al., 2020). Similarly, past research using the HRS has shown that GrimAge and DunedinPoAm38 are more consistently associated with smoking and educational attainment (Crimmins et al., 2021), two important mechanisms identified in past literature linking ACEs to depressive symptoms (discussed above). The use of different individual clocks—each of which has different strengths and weaknesses—in individual studies may be a cause of the mixed findings in past research. More research like the current study comparing multiple clocks is needed to clarify what clocks are most responsive to certain exposures and are most predictive of psychiatric outcomes.

The current study has some key limitations. This is a study of US older adults. Further research using international samples is needed. ACEs were assessed using a retrospective self-report and may be biased by recall and social desirability; however, past research has established the validity and reliability of retrospective recall of childhood adversity (Brewin et al., 1993). Not all major ACEs, as defined by the CDC, were available. Particularly, we do not have a measure of neglect, which has been shown to be associated with adult mental health (Horwitz et al., 2001). Future research using the complete CDC ACEs scale is needed to validate these findings. We are not able to control for early life epigenetic age or psychiatric symptoms before ACEs exposure. We focused on BMI as a mediator of the effect of ACEs on epigenetic aging. Future studies should investigate whether modifiable health behaviors that affect BMI (e.g., diet, physical activity) mediate this association. Our measure of depressive symptoms does not capture clinically diagnosed depressive disorder. Because this was a population study, obtaining clinical diagnoses for all participants would be impractical and would substantially increase subject burden. Future work should validate our results in a well characterized clinical psychiatric population.

The current findings have important implications for biological psychiatric research, providing novel evidence accelerated DNAm aging may be a plausible biological mechanism explaining how childhood adversity may lead to elevated risk for depression. Moreover, these results suggest interventions aimed at healthy weight maintenance, smoking cessation, and alcohol abuse may be an important target for reducing the mental health risk and epigenetic aging risk associated with ACEs.

**Author statement**

Eric T. Klopack: Conceptualization, Formal analysis, Writing - Original Draft, Eileen M. Crimmins; Writing - Review & Editing, Supervision, Steve W. Cole: Writing - Review & Editing, Teresa E. Seeman: Writing - Review & Editing, Judith E. Carroll: Conceptualization, Writing - Review & Editing, Supervision.

**Declaration of competing interest**

The authors report no conflicts of interest.
E.T. Klopack et al.

Merrick, M. T., Ports, K. A., Ford, D. C., Affif, T. O., Gershoff, E. T., & Grogan-Kaylor, A. (2017). Unpacking the impact of adverse childhood experiences on adult mental health. Child Abuse & Neglect, 69(1), 10–19. https://doi.org/10.1016/j.chiabu.2017.03.016

csiuab.2017.03.016

epsel.

Miller, G. E., Chen, E., & Parker, K. J. (2011). Psychological stress in childhood and susceptibility to the chronic diseases of aging: Moving toward a model of behavioral and biological mechanisms. Psychological Bulletin, 137(6), 959–997. https://doi.org/10.1037/a0024768

Miller, A. H., Maletic, V., & Raison, C. L. (2009). Inflammation and its discontents: The role of cytokines in the pathophysiology of major depression. Biological Psychiatry, 65(9), 732–741. https://doi.org/10.1016/j.biopsych.2008.11.029

Mitchell, C., Schnepper, L. M., & Notterman, D. A. (2016). DNA methylation, early life environment, and health outcomes. Pediatric Research, 79(1), 212–219. https://doi.org/10.1016/0887-8795(92)90339-9

Moffitt, T. E., & Caspi, A.’s opportunity to prevent the rising burden of age-related disease. JAMA Psychiatry, 76(5), 461–462. https://doi.org/10.1001/jamapsychiatry.2019.0037

Muthen, L. K., & Muthén, B. O. (1998). Mplus user’s guide (7th ed.). Muthén & Muthén.

Obal, L., van der Zaag, J., Higgins-Chen, A. T., Levine, M. E., & Boks, M. P. (2021). Childhood abuse histories predict steeper inflammatory trajectories across time. Brain, Behavior, and Immunity, 91, 541–545. https://doi.org/10.1016/j.bbi.2020.11.012

Pudrovská, T., & Anikputa, B. (2014). Early-life socioeconomic status and mortality in later life: An integration of four life-course mechanisms. Journal of Gerontology: Series B, 69(3), 451–460. https://doi.org/10.1093/geronb/gbt122

Raffington, L., Belsky, D. W., Rothari, M., Malanchini, M., Tucker-Drob, E. M., & Harden, K. P. (2021). Socioeconomic disadvantage and the pace of biological aging in children. Pediatrics. https://doi.org/10.1542/peds.2020-024406

Raison, C. L., Capuron, L., & Miller, A. H. (2006). Cytokines sing the blues: Inflammation and the pathogenesis of depression. Trends in Immunology, 27(1), 24–31. https://doi.org/10.1016/j.it.2005.11.006

Renna, M. E., Peng, J., Shrouf, M. R., Madison, A. A., Andridge, R., Alfan, C. M., Povoski, S. P., Lipari, A. M., Malarkey, W. B., & Kiecolt-Glaser, J. K. (2021). Childhood abuse histories predict steeper inflammatory trajectories across time. Brain, Behavior, and Immunity, 91, 541–545. https://doi.org/10.1016/j.bbi.2020.11.012

Richmond-Rakerd, L. S., D’Souza, S., Milne, B. J., Caspi, A., & Moffitt, T. E. (2021). Longitudinal associations of mental disorders with physical diseases and mortality among 2.3 million New Zealand citizens. JAMA Network Open, 4(1), Article e2033484. https://doi.org/10.1001/jamanetworkopen.2020.33484

Shiel, M. S., Katki, H. A., Freedman, N. D., Purdue, M. P., Wentzensen, N., Trabert, B., Kitahara, C. M., Furr, M., Li, Y., Kemp, T. J., Goedert, J. J., Chang, C. M., Engels, E. A., Caporaso, N. E., Pinto, L. A., Hidesheim, A., & Chaturvedi, A. K. (2014). Cigarette smoking and variations in systemic immune and inflammation markers. Journal of the National Cancer Institute, 106(1). https://doi.org/10.1093/jnci/djz094

Simons, R. L., Lei, M. K., Beach, S. R. H., Philibert, R. A., Cutrana, C. E., Gibbons, F. X., & Barr, A. B. (2016). Economic hardship and biological weathering: The epigenetics of aging in a US sample of black women. Social Science & Medicine, 150(1), 192–200. https://doi.org/10.1016/j.socscimed.2015.12.001

Slavich, G. M., & Irwin, M. R. (2014). From stress to inflammation and major depressive disorder: A social signal transduction theory of depression. & Link to external site, this link will open in a new window Psychological Bulletin, 140(3), 774–815. https://doi.org/10.1037/a0035302

Sumner, J. A., Colich, N. L., Uddin, M., Armstrong, D., & McLaughlin, K. A. (2019). Early experiences of threat, but not deprivation, are associated with accelerated biological aging in children and adolescents. Biological Psychiatry, 85(3), 268–278. https://doi.org/10.1016/j.biopsych.2018.09.008

Turvey, C., Wallace, R. B., & Herzog, R. (1999). A revised CES-D measure of depressive symptoms and a DSM-based measure of major depressive episodes in the elderly. International Psychogeriatrics, 11(2), 139–148. https://doi.org/10.1017/S1041610299005694

Verhoeff, J. E., Yang, R., Wolkowitz, O. M., Bersani, F. S., Lindqvist, D., Mellen, S. H., Yehuda, R., Florry, J. D., Lin, J., Abu-Amara, D., Makotkine, I., Marmar, C., Jett, M., & Hamann, R. (2018). Epigenetic age in male combat-exposed war veterans: Associations with posttraumatic stress disorder status. Complex Psychiatry, 4(2), 90–99. https://doi.org/10.1015/s41380-020-001431

Visser, M., Bouter, L. M., McQuillan, G. M., Wener, M. H., & Harris, T. B. (1999). Elevated C-reactive protein levels in overweight and obese adults. JAMA, 282(22), 2131–2135. https://doi.org/10.1001/jama.282.22.2131

Wickramp, K. A. S., Kwon, J. A., Lee, T. K. (2014). Early socioeconomic adversity and young adult physical illness: The role of body mass index and depressive symptoms. Journal of Adolescent Health, 55(4), 556–563. https://doi.org/10.1016/j.jadohealth.2014.04.006

Wolkowitz, O. M., Epp, E. S., & Mellon, S. (2008). When blue turns to grey: Do stress and depression accelerate cell aging? World Journal of Biological Psychiatry, 9(1), 2–5. https://doi.org/10.1080/15622970701875601

Yang, Y., Gerken, K., Schopp, K., Boen, C., & Harris, K. M. (2017). Early-life socioeconomic status and adult physiological functioning: A life course examination of biosocial mechanisms. Biodemography and Social Biology, 65(2), 87–103. https://doi.org/10.1080/19485565.2017.1279536

Yang, R., Wu, G. W. Y., Verhoeff, J. E., Gautam, A., Reus, V. I., Kang, J. I., Flory, J. J., Abu-Amara, D., Hood, L., Doyle, F. J., Yehuda, R., Marmar, C. R., Jett, M., Marmar, R., & Wolkowitz, O. M. (2020). A DNA methylation clock associated with age-related illnesses and mortality is accelerated in men with combat PTSD. Molecular Psychiatry, 1–11. https://doi.org/10.1038/s41380-020-0753-z