Pos
traumatic stress disorder (PTSD) is prevalent in US military veterans and has been linked to premature mortality, which can be predicted by accelerated epigenetic aging, a marker of biological aging.\(^1\)

Recent studies have reported associations between PTSD and accelerated epigenetic aging.\(^1\) GrimAge, a composite of DNA methylation (DNAm)-based markers linked to health- and lifespan, is the strongest predictor of various age-related metrics (eg, time to death, to coronary heart disease, and to cancer) relative to other measures of epigenetic aging.\(^3\) We examined the relationship between PTSD and its specific symptoms, other medical and psychiatric conditions, and GrimAge in a large, nationally representative sample of male US veterans.

**METHODS**

**Participants**

The sample included 1,132 male European-American US veterans who participated in the National Health and Resilience in Veterans Study, which surveyed a nationally representative sample of US veterans (see Supplementary Material).

**Assessments**

**GrimAge.** GrimAge is a composite measure of epigenetic aging based on 8 DNAm surrogates of plasma proteins that are associated with mortality or morbidity, and a DNAm-based estimator of smoking pack-years. Accelerated GrimAge (ie, greater epigenetic than actual age) was operationalized as a residual GrimAge acceleration ≥ 5 years relative to chronologic age (mean = 8.3 years, standard deviation = 2.2, range = 5–16).

**PTSD symptoms.** PTSD symptoms were assessed using the PTSD Checklist-Specific Stressor Version (PCL-S); scores ≥ 35 were indicative of probable PTSD.

**RESULTS**

A total of 18.3% (95% confidence interval [CI] = 16.0–20.8%) of the sample had accelerated GrimAge. Relative to veterans without accelerated GrimAge, those with accelerated GrimAge were less likely to be married/partnered and to have an annual household income > $60,000/year and more likely to be combat veterans, obese, and current smokers and to screen positive for current alcohol use disorder and PTSD; they also reported more cumulative traumas and medical conditions (Table 1).

When these variables were entered into a relative regression analysis, PTSD was associated with 2-fold greater odds of accelerated GrimAge while college graduate or higher education was associated with lower odds of this outcome; none of the other variables were significant (all P values > 0.20). PTSD remained significantly associated with accelerated GrimAge even after additionally adjusting for smoking status (risk ratio = 2.00, 95% CI = 1.01–3.99).

PTSD was specifically associated with an acceleration in DNAm surrogates of tissue inhibitor metalloproteinase-1 (TIMP-1), \(\beta\)-2 microglobulin (B2M), and growth differentiating factor-15 (GDF-15) (Supplementary Table 1).

Greater severity of trauma-related detachment (ie, feeling distant or cut off from others, OR = 1.73, 95% CI = 1.22–2.45, \(P = .002\); Supplementary Figure 1) and sleep disturbance (OR = 1.51, 95% CI = 1.16–1.95, \(P = .002\)) were independently associated with accelerated GrimAge.
DISCUSSION

Results of this nationally representative study of male European-American US veterans revealed that PTSD is associated with 2-fold greater odds of accelerated GrimAge, which averaged nearly a full decade. This association was independent of a broad range of other factors, including smoking.

PTSD was associated with an acceleration in DNA methylation age surrogates of TIMP-1, B2M, and GDF-15. TIMP-1 has been linked to abnormalities in long-term potentiation in the prefrontal cortex; B2M, with increased stress-induced gene expression in individuals with psychiatric disorders; and GDF-15, a stress-induced cytokine, with inflammation and physical dysfunction.4

Greater severity of 2 specific PTSD symptoms—detachment and sleep disturbance—were independently associated with accelerated GrimAge. Potential mechanisms linking these particular symptoms to accelerated GrimAge include dysregulation of hypothalamic-pituitary-adrenal axis, neuroendocrine, and inflammatory responses.5,6 While larger and prospective studies are needed to disentangle temporal and mechanistic associations between PTSD and accelerated GrimAge, these findings underscore the utility of transdiagnostic, symptom-level approaches to identifying dimensions of psychopathology linked to epigenetic aging.8

Collectively, the results of this study extend prior work1 to suggest that PTSD is associated with accelerated epigenetic aging in veterans. Limitations of our study include the cross-sectional design; focus on male, European-American veterans, which may limit generalizability of the findings; and use of self-report assessment instruments. Further research is needed to replicate these results in larger, more diverse samples and evaluate the efficacy of interventions for PTSD in forestalling premature biological aging in veterans and other high-risk populations.

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**Supplementary material:** Available at Psychiatrist.com.

**REFERENCES**


See supplementary material for this brief report at Psychiatrist.com.
Supplementary Material

Brief Report Title: Association of Symptoms of Posttraumatic Stress Disorder and GrimAge, an Epigenetic Marker of Mortality Risk, in US Military Veterans

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List of Supplementary Material for the brief report

1. **Methods**

2. **Table 1** DNAm Surrogate Markers by PTSD Screening Status

3. **Figure 1** Probability of accelerated GrimAge as a function of severity of detachment symptoms of PTSD

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This Supplementary Material has been provided by the author(s) as an enhancement to the published report. It has been approved by peer review; however, it has undergone neither editing nor formatting by in-house editorial staff. The material is presented in the manner supplied by the author.
Supplementary Methods

Participants

The National Health and Resilience in Veterans Study (NHRVS) is a nationally representative survey of U.S. military veterans. The NHRVS sample was drawn from KnowledgePanel, a research panel of more than 50,000 households that is maintained by Ipsos, a survey research firm. KnowledgePanel® is a probability-based, online non-volunteer access survey panel of a nationally representative sample of U.S. adults that covers approximately 98% of U.S. households. Panel members are recruited through national random samples, originally by telephone and now almost entirely by postal mail. Households are provided with access to the Internet and computer hardware if needed. KnowledgePanel® recruitment uses dual sampling frames that include both listed and unlisted telephone numbers, telephone and non-telephone households, and cell-phone-only households, as well as households with and without Internet access.

Of the 4,750 veterans who were in KnowledgePanel® when the NHRVS was fielded (veteran status was assessed using a general demographic questionnaire), 3,408 (71.7%) responded to an invitation to participate and completed a screening question to confirm their study eligibility (current or past active military service). Of these respondents, 3,199 (93.5%) confirmed their current or past active military service, 3,157 (92.6%) complete a confidential, 60-minute online survey, and 2,397 (70.3%) consented to and provided a saliva sample for genotyping. Given that the vast majority of U.S. veterans are male and European-American and to increase statistical power and avoid confounding analyses with low numbers of female and ethnically diverse veterans, epigenetic analyses were limited to male European-American veterans in the genotyped cohort (n=1,132).

Demographic data of survey panel members are assessed regularly by Ipsos using the same set of questions used by the U.S. Census Bureau. To permit generalizability of study results to the entire population of U.S. veterans, the Ipsos statistical team computed post-stratification weights using the following benchmark distributions of U.S. military veterans from the 2011 Current Veteran Population Supplemental Survey of the U.S. Census Bureau’s American Community Survey: age, gender, race/ethnicity, Census Region, metropolitan status, education, household income, branch of service, and years in service. An iterative proportional fitting (raking) procedure was used to produce the final post-stratification weights. All participants provided informed consent and the study was approved by the Human Subjects Committee of the VA Connecticut Healthcare System.

Assessments

Cumulative trauma burden. The Trauma History Screen (THS)\(^1\) was used to assess exposure to the lifetime occurrence of 14 potentially traumatic events; the NHRVS additionally assessed exposure to life-threatening illness or injury. The sum of potentially traumatic events endorsed, ranging from 0–15, was used as an index of lifetime trauma burden.
**PTSD symptoms.** PTSD symptoms were assessed using the PTSD Checklist-Specific Stressor Version (PCL-S); score ≥ 35, which have been recommended for general population samples,² were indicative of a positive screen (Cronbach’s α in the current sample=0.94). Participants were asked to respond to PCL-S in relation to their ‘worst’ trauma endorsed on the THS.

**Combat veteran.** Combat veteran status was assessed with the following question: “Did you ever serve in a combat or war zone?” and the Combat Exposure Scale,³ a 7-item self-report measure that assesses wartime stressors experienced by combatants.

**Childhood physical or sexual abuse.** Childhood physical or sexual abuse was assessed using two items from the THS: “Hit or kicked hard enough to injure – as a child” and/or “Forced or made to have sexual contact – as a child.” A three-level variable was created based on responses to these items: No physical or sexual childhood abuse; physical abuse; and physical and sexual abuse.

**Years since index trauma.** Years since index trauma was assessed by subtracting current age from age of index trauma.

**Number of medical conditions.** Sum of number of medical conditions endorsed in response to question: “Has a doctor or healthcare professional ever told you that you have any of the following medical conditions?” (e.g., arthritis, cancer, diabetes, heart disease, asthma, kidney disease). Range: 0-24 conditions.

**Body mass index (BMI).** BMI was calculated based on self-reported height and weight using the standard formula weight (kg)/height (m²). Obesity was defined as BMI ≥30.00, consistent with CDC guidelines.⁴

**Current major depressive disorder.** Major depressive disorder symptoms were assessed using the two depressive symptoms items of the PHQ-4,⁵ which assessed symptoms occurring in the past two weeks; score ≥ 3 was indicative of a positive screen for major depressive disorder (Cronbach’s α in the current sample=0.90).⁵

**Current alcohol use disorder.** Alcohol use disorder was assessed using the Alcohol Use Disorders Identification Test-Consumption (AUDIT-C), a validated measure used to screen for alcohol use disorder.⁶ The AUDIT-C consists of 3 questions that assess severity of alcohol consumption and yield a total score ranging from 0 to 12. A score of 5 or higher was considered as indicative of probable alcohol use disorder.⁷,⁸

**Current smoker.** Current smoking status was assessed using a question that asked whether veterans had ever smoked cigarettes; response options were “Yes, in the past;” “Yes, currently;” and “Never.”

**GrimAge calculation**

GrimAge is a composite epigenetic biomarker based on the DNAm surrogates of plasma proteins that are known to be associated with mortality or morbidity, and a DNAm-based estimator of
smoking pack-years. A two-stage procedure was performed to develop GrimAge. First, DNAm-based surrogate biomarkers of smoking pack-years and 88 plasma proteins previously identified to be linked to mortality were identified. Second, time-to-death was regressed on chronological age, sex, and DNAm-based biomarkers of smoking pack-years and the 12 plasma proteins that exhibited a correlation $r>0.35$ with their respective DNAm-based surrogate marker in step 1. The plasma protein surrogates that were selected by an elastic net regression model were leptin, cystatin C, tissue inhibitor metalloproteinases 1 (TIMP1), adrenomedullin (ADM), beta-2-microglobulin (B2M), growth differentiation factor-15 (GDF-15), and plasminogen activation inhibitor 1 (PAI-1). The resultant mortality risk estimate of the regression model was then transformed linearly into units of years. The rationale for selecting these proteins and details of analytical procedures that generate GrimAge have been described in detail previously.9

In the current study, we operationalized accelerated GrimAge as a residual GrimAge acceleration of 5 or more years relative to chronologic age (mean in full sample=8.3 years, SD=2.2, range=5-16). This magnitude difference is clinically meaningful (i.e., approximates an average 5-10 year greater acceleration of epigenetic relative to chronological aging) and permits comparability to prior studies.10-12 We calculated DNAm age for each individual based on salivary DNA samples profiled with the Illumina Infinium EPIC array and Horvath age estimation algorithm.13 In the present sample, chronological age correlated strongly with GrimAge ($r=0.91; p<0.001$).

Cell proportion estimation analysis was conducted using a modified version of the Houseman method,14 which yielded estimates of each cell type proportion (e.g., CD14, CD34, and buccal cells) in the peripheral saliva samples. Principal component analysis was conducted to adjust for population stratification using the Barfield method;15 the first 10 principal components were included in analyses.

**Data Analysis**

First, we compared baseline characteristics of veterans with or without PTSD using chi-square and independent-samples t-tests. Second, we conducted a multivariable relative risk regression analysis to identify variables associated with accelerated GrimAge variables associated with accelerated GrimAge at the $p<0.05$ level in bivariate analyses were entered into this analysis; PTSD screening status and variables shown in Table 1, as well as cell type proportions (CD34, CD14, and buccal) and 10 ancestry principal components, were entered into this analysis. Given that GrimAge includes a measure of smoking pack-years, analyses were first conducted without adjusting for smoking status; we then conducted a sensitivity analysis to determine whether any significant associations were robust to smoking status. Third, we conducted a second regression analysis with individual PTSD symptoms entered as independent variables to identify symptom(s) that were independently associated with accelerated GrimAge; alpha for this analysis was Bonferroni-corrected to 0.0029 (0.05/17 symptoms). Fourth, we conducted a multivariate analysis of covariance to evaluate how PTSD related to component aspects of GrimAge; this analysis adjusted for variables identified as significant correlates of accelerated GrimAge in the regression model.
Supplement References


### Supplementary Table 1. DNA methylation surrogate markers by PTSD screening status

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<th>Positive Screen for PTSD</th>
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<td></td>
<td>N=1,019 (weighted 88.5%)</td>
<td>N=113 (weighted 11.5%)</td>
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<td>DNAm PACKYRS</td>
<td>Weighted mean (SE)</td>
<td>Weighted mean (SE)</td>
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<td>6.45 (0.46)</td>
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<td>114.67 (509.18)</td>
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<td>219.29 (52.89)</td>
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<td>DNAm B2M</td>
<td>1886.92 (3892.29)</td>
<td>21033.80 (7887.71)</td>
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<td>DNAm GDF-15</td>
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<td>DNAm PAI-1</td>
<td>284.29 (131.89)</td>
<td>160.71 (267.28)</td>
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*Note. SE=standard error of the mean; PTSD=posttraumatic stress disorder; DNAm=DNA methylation; PACKYRS=smoking pack-years; TIMP-1=tissue inhibitor metalloproteinase 1; ADM=adrenomedullin; B2M(beta-2 microglobulin; GDF-15=growth differentiation factor-15; PAI-1=plasminogen activation inhibitor 1.

Analysis is adjusted for education, current smoking status, cell type proportions (CD34, CD14, and buccal), and top 10 ancestry principal components.

Significant difference: *=p<0.05; **p<0.01.
Supplementary Figure 1. Probability of accelerated GrimAge as a function of severity of detachment symptoms of PTSD

Note. Circles represent mean probabilities; error bars represent 95% confidence intervals. Mean probability of accelerated GrimAge in the sample was 0.18.