UCSF UC San Francisco Previously Published Works

Title

Internal capsule microstructure mediates the relationship between childhood maltreatment and PTSD following adulthood trauma exposure.

Permalink

https://escholarship.org/uc/item/3mq414z9

Journal Molecular Psychiatry, 28(12)

Authors

Wong, Samantha Lebois, Lauren Ely, Timothy <u>et al.</u>

Publication Date

2023-12-01

DOI

10.1038/s41380-023-02012-3

Peer reviewed

ARTICLE OPEN

Check for updates

Internal capsule microstructure mediates the relationship between childhood maltreatment and PTSD following adulthood trauma exposure

Samantha A. Wong ^[1], Lauren A. M. Lebois ^{[1,2}, Timothy D. Ely ^[1], Sanne J. H. van Rooij ^[0], Steven E. Bruce ^[0], Vishnu P. Murty⁵, Tanja Jovanovic⁶, Stacey L. House⁷, Francesca L. Beaudoin^{8,9}, Xinming An¹⁰, Donglin Zeng¹¹, Thomas C. Neylan ^[0], Gari D. Clifford ^{13,14}, Sarah D. Linnstaedt ^[0], Laura T. Germine ^[0], ^{2,15,16}, Kenneth A. Bollen¹⁷, Scott L. Rauch^{2,15,18}, John P. Haran ^[0], Alan B. Storrow ^[0], Christopher Lewandowski ^[0], Paul I. Musey Jr. ^[0], Paul I. Hendry²³, Sophia Sheikh ^[0], Christopher Lewandowski ^[0], Paul I. Musey Jr. ^[0], Robert A. Swor³⁰, Lauren A. Hudak³¹, Jose L. Pascual^{32,33}, Christopher W. Jones²⁴, Brittany E. Punches ^[0], Sola C. Merchant³⁷, Robert M. Domeier³⁸, Niels K. Rathlev³⁹, Brian J. O'Neil⁴⁰, Paulina Sergot⁴¹, Leon D. Sanchez^{37,42}, Mark W. Miller ^[0], ^{43,44}, Robert H. Pietrzak^{45,46}, Jutta Joormann⁴⁷, Deanna M. Barch ^[0], Diego A. Pizzagalli ^[0], Steven E. Harte^{49,50}, James M. Elliott^{51,52,53}, Ronald C. Kessler ^[0], Karestan C. Koenen ^[0], Samuel A. McLean ^[0], Steven J. Ressler ^[0], James F. Stevens ^[0], Steven ^[0], Steven ^[1], Steven ^[1], Jennifer S. Stevens ^[0], ^{3,58^[2]} and Nathaniel G. Harnett ^[0], ^{1,2,58^[2]}

© The Author(s) 2023

Childhood trauma is a known risk factor for trauma and stress-related disorders in adulthood. However, limited research has investigated the impact of childhood trauma on brain structure linked to later posttraumatic dysfunction. We investigated the effect of childhood trauma on white matter microstructure after recent trauma and its relationship with future posttraumatic dysfunction among trauma-exposed adult participants (n = 202) recruited from emergency departments as part of the AURORA Study. Participants completed self-report scales assessing prior childhood maltreatment within 2-weeks in addition to assessments of PTSD, depression, anxiety, and dissociation symptoms within 6-months of their traumatic event. Fractional anisotropy (FA) obtained from diffusion tensor imaging (DTI) collected at 2-weeks and 6-months was used to index white matter microstructure. Childhood maltreatment load predicted 6-month PTSD symptoms (b = 1.75, SE = 0.78, 95% CI = [0.20, 3.29]) and inversely varied with FA in the bilateral internal capsule (IC) at 2-weeks (p = 0.0294, FDR corrected) and 6-months (p = 0.0238, FDR corrected). We observed a significant indirect effect of childhood maltreatment load on 6-month PTSD symptoms through 2-week IC microstructure (b = 0.37, Boot SE = 0.18, 95% CI = [0.05, 0.76]) that fully mediated the effect of childhood maltreatment load on PCL-5 scores (b = 1.37, SE = 0.79, 95% CI = [-0.18, 2.93]). IC microstructure did not mediate relationships between childhood maltreatment and depressive, anxiety, or dissociative symptomatology. Our findings suggest a unique role for IC microstructure as a stable neural pathway between childhood trauma and future PTSD symptoms following recent trauma. Notably, our work did not support roles of white matter tracts previously found to vary with PTSD symptoms and childhood trauma exposure, including the cingulum bundle, uncinate fasciculus, and corpus callosum. Given the IC contains sensory fibers linked to perception and motor control, childhood maltreatment might impact the neural circuits that relay and process threat-related inputs and responses to trauma.

Molecular Psychiatry (2023) 28:5140-5149; https://doi.org/10.1038/s41380-023-02012-3

INTRODUCTION

Childhood trauma is a well-established risk factor for development of trauma and stress-related disorders in adulthood. Early life stress may interact with stressors in adulthood to increase an individual's risk for posttraumatic stress disorder (PTSD), major depression, substance use, or behavioral disorders [1]. Furthermore, childhood trauma is associated with variability in brain circuits known to play a role in PTSD, which could represent potential neural signatures of PTSD susceptibility. However, limited work to date has investigated neural correlates of how earlier childhood trauma augments posttraumatic reactions after a

A full list of author affiliations appears at the end of the paper.

Received: 14 October 2022 Revised: 17 February 2023 Accepted: 22 February 2023 Published online: 17 March 2023

trauma sustained as an adult. Identifying the neurobiological correlates of childhood trauma related risk for acute stress reactions in adulthood may advance neuroscience-based approaches for prediction and prevention of PTSD development.

PTSD is thought to be partially driven by dysfunction of threat learning neurocircuitry – particularly the prefrontal cortex, hippocampus, and amygdala – as a result of a traumatic experience [2–4]. White matter tracts such as the cingulum bundle, uncinate fasciculus, and fornix/stria terminalis interconnect threat neurocircuitry regions and are thought to be involved in PTSD-related dysfunction (See [5] for review), potentially due to

experience-dependent changes in tract microstructure [6]. In line with this reasoning, previous PTSD research has investigated Fractional Anisotropy (FA) as one of several measures to index white matter microstructure derived from Diffusion Tensor Imaging (DTI). Greater FA indicates greater linearity in the flow of water molecules due to constraint by myelinated tracts. Individuals with PTSD show reduced FA of the cingulum bundle and uncinate fasciculus [7–11], which interconnects the prefrontal cortex, amygdala, and hippocampus, although there is some heterogeneity in findings [12, 13]. Successful psychotherapy for PTSD appears to lead to increased FA in tracts such as the cingulum and fornix [14]. Further, studies of recent trauma exposure suggest variability in these same tracts are related to future development of PTSD such that lower FA is generally related to greater PTSD symptom severity [15-18]. Taken together, the previous work suggests white matter tracts of core threat neurocircuitry are related to the development and expression of PTSD symptoms.

Despite the importance of threat neurocircuitry white matter tracts, emergent research in childhood and adult trauma suggests that PTSD-related white matter alterations may additionally occur within other tracts [19-21]. Previous retrospective and metaanalytic DTI studies demonstrate that childhood trauma exposure is associated with alterations in FA both within threat neurocircuitry tracts and sensory integration tracts such as the anterior thalamic radiation, superior longitudinal fasciculus, inferior frontooccipital fasciculus, optic radiations, and arcuate fasciculus [20-27]. Further, recent meta-analyses from the PGC-PTSD and ENIGMA groups found that the largest reduction in FA for individuals with PTSD was not within threat neurocircuitry tracts, but instead within the tapetum of the corpus callosum [28]. Perception and integration of sensory stimuli is necessary for appropriate threat learning [29]. The prior findings thus suggest trauma and PTSD-related FA reductions may extend outside threat neurocircuitry and encompass regions necessary for stimulus perception.

Limited research exists on the interrelationship between childhood trauma, white matter microstructure, and posttraumatic outcomes following a more recent trauma, though it may improve our understanding of the biological basis of PTSD. However, previous studies have found relationships between childhood trauma, brain structure, and stressors in adulthood [30-33]. In one study, total childhood trauma exposure moderated the effect of later combat exposure on FA within the hippocampal component of the cingulum, with greater childhood trauma and combat exposure related to decreased FA [30]. In a longitudinal study of young adults, uncinate fasciculus FA values at baseline moderated the relationship between recent stressors (e.g., break up with romantic partner, failing a course, or financial problems) and mood and anxiety symptoms at follow up among those with higher reported childhood maltreatment [31]. Limited work, however, has considered potential associations with white matter tracts outside threat neurocircuitry, which may be important in light of recent findings of PTSD-related FA reductions.

The present study investigated whether, among recent trauma survivors, brain white matter microstructure mediated the effect of childhood maltreatment exposure on posttraumatic dysfunction. Given prior findings in both studies on threat neurocircuitry of PTSD and emergent work implicating sensory and other white matter tracts, we assessed FA across white matter tracts using a whole-brain approach following previous work by the PGC-ENIGMA consortium [28]. We hypothesized that white matter FA at 2 weeks post-trauma, in general, would be negatively associated with childhood maltreatment load. We further hypothesized that white matter FA associated with childhood maltreatment would mediate associations between childhood maltreatment and posttraumatic outcomes after a recent trauma. Our findings highlight a neural pathway through which childhood trauma may confer risk for acute stress reactions in adulthood and shed light on white matter markers of susceptibility for PTSD.

MATERIALS AND METHODS

Participants

Participants were recruited as part of the AURORA study, a longitudinal multisite investigation of adverse neuropsychiatric segualae [34]. Participants included in this investigation have been reported on in previous work [35-38]. However, the investigation described here is the first to consider the relationships of childhood maltreatment exposure, white matter microstructure, and later posttraumatic outcomes. As detailed in our prior reports [34], enrollment occurred at emergency departments (ED) and focused on those presenting within the 72 h following exposure to a qualifying trauma (physical or sexual assault, motor vehicle accident, fall >10 feet, mass casualty incident, or other life-threatening traumatic event reported on a screener question and agreed upon as a plausible qualifying event by the study staff). Participants were included if they were Englishspeaking, between 18 and 75 years-old, and able to consent and follow study procedures. Participants were recruited regardless of prior PTSD symptoms or diagnosis and were asked to report retrospectively on prior PTSD (and other disorders) symptoms in the emergency department. General exclusion criteria for the AURORA study have been described previously [34]. MRI collection exclusion criteria were having metal or ferromagnetic implants, history of seizure or epilepsy, history of Parkinson's disease, dementia, or Alzheimer's disease, current pregnancy, and/or declining to complete the MRI. From the beginning of study enrollment in September 2017 to July 2020, MRI data were collected within ~2 weeks of trauma exposure for 439 participants and DTI data were available from 353 participants. Participants were excluded for MRI quality issues (n = 37) (e.g., motion artefact, anatomical barriers, or low-quality data). The present analyses focused on participants who completed both DTI and the abbreviated Childhood Trauma Questionnaire (described below) at 2-weeks and posttraumatic outcome measures at 6-months post qualifying trauma and excluded participants missing a required guestionnaire (n = 153). A total of 202 participants were retained for final analyses. Further analyses of DTI data from a subset of 85 participants (n = 111 collected, n = 26 excluded) collected at a 6-month follow-up imaging session also were completed. All participants provided informed consent as approved by the Biomedical IRB at UNC Chapel Hill through the office of Human Research Ethics, the central IRB for all study sites.

Baseline surveys and socio-demographics

Participants completed a baseline assessment in the ED that included selfreported trauma characteristics and demographic characteristics [34]. Age, sex assigned at birth, race/ethnicity, highest education level, marital status, employment status, and total household income were obtained in the ED baseline surveys. Patients were also asked if they hit their head or experienced a head injury during the event that brought them to the ED (n = 86 endorsed).

Childhood maltreatment load

An abbreviated 11-item version of the Childhood Trauma Questionnaire-Short Form (CTQ-SF; [39, 40]) was used to index childhood maltreatment. Items were selected from the CTQ-SF to capture maltreatment subscales while minimizing participant burden (individual questions selected provided in the supplementary information). Items selected to capture childhood maltreatment showed high internal reliability (Cronbach's a = 0.92). The questionnaire was administered two weeks after the qualifying trauma. Items were self-reported on a 5-point Likert scale (0: never, 1: rarely, 2: sometimes, 3: often, 4: very often). The maltreatment subtypes evaluated include emotional abuse (sub-score range: 0 to 8), physical abuse (sub-score range: 0 to 8), sexual abuse (sub-score range: 0 to 12), emotional neglect (sub-score range: 0 to 8), and physical neglect (sub-score range: 0 to 8). Total possible summed scores ranged from 0 to 44. We indexed childhood maltreatment load as the endorsements of moderate to extreme levels of each maltreatment subtype [41-43]. Moderate to extreme abuse or neglect for other maltreatment subtypes were defined as a subtype score of 4 or above. Moderate to extreme sexual abuse was defined as a sub-score of 3 or above on sexual abuse items. These cutoffs were modified for the abbreviated assessment from clinical cutoffs previously suggested [39]. The sum of moderate to extreme maltreatment types was used to index total childhood maltreatment load.

Lifetime trauma

Lifetime trauma exposure was assessed with the Life Events Checklist (LEC-5), an established 17-item instrument assessing exposure to 17 traumatic life events [44, 45]. Participants completed the LEC-5 at 8-weeks after their qualifying exposure. Participants indicated if a selection of traumatic experiences happened to them personally, if they witnessed it happen to someone else, learned about it happening to someone close to them, or was exposed to details about it as part of their job. A modified total LEC score (mLEC-5, range: 0 to 17) was calculated by summing the types of traumatic events endorsed, regardless of exposure modality. Although participants could endorse experiencing a life event in multiple ways (e.g., "happened to me," "vitnessed it"), any exposure to a given traumatic life event resulted in the maximal score of one for the modified total LEC score.

Posttraumatic outcomes

Posttraumatic dysfunction was assessed in terms of PTSD, depression, anxiety, and dissociation symptoms at 6-months following the index trauma. PTSD symptoms were assessed using the Posttraumatic Stress Disorder (PTSD) checklist for DSM-5 (PCL-5), a psychometrically rigorous 20-item questionnaire on symptom presence and severity [46, 47]. Participants rated symptom severity on a scale of 0 (not at all) to 4 (extremely). Depression symptoms were assessed with the 8-item Patient-Reported Outcomes Measurement Information System (PROMIS) Depression instrument, short form 8b [48, 49]. A total raw score was computed from summing the individual items and then converted to a T-score [50]. Anxiety symptoms were assessed with 4-items from the PROMIS Anxiety bank [49, 51]. Participants rated how often they felt tense, worried about things, had trouble relaxing, or felt anxious on a scale of 1 (none of the time) to 5 (all or almost all of the time), and item scores were summed to create a total anxiety score. Dissociation was assessed using a modified 2-item Brief Dissociative Experiences Scale (DES-B-Modified, [52]). Participants rated how often they felt people, objects, or the world around them seemed unreal, and how often they felt they were looking through a fog so that people and things seemed unclear on a scale from 1 (none of the time) to 5 (all or almost all of the time). A sum of the two questions was used as an index of dissociation severity.

Diffusion tensor imaging

Diffusion weighted imaging (DWI) data were collected across five sites (Table S1). Data processing was similar to prior reports [36, 53], following recommendations of the ENIGMA consortium (http:// the enigma.ini.usc.edu/protocols/dti-protocols/). To ensure quality data, raw data were visually inspected, and we calculated metrics of temporal signalto-noise ratio and outlier maximum voxel intensity as in a prior report [54]. Participants who demonstrated both: (a) TSNR values lower than 4.88 and (b) maximum voxel intensities greater than 5000 were removed from analyses to retain the maximum number of participants while removing low-quality data. Briefly, motion and eddy current effects in the DWI data were reduced using the 'eddy' subroutine in FSL and susceptibility effects were corrected for using nonlinear warping of the DWI data to the participant's T1-weighted anatomical scan [55-57]. Tract-Based Spatial Statistics (TBSS) processing was used as implemented in the ENIGMA-DTI working group processing standards to extract FA values across white matter regions [58, 59]. First, FA maps were non-linearly registered to the standard ENIGMA FA map in Montreal Neurological Institute (MNI) standard space [59]. The ENIGMA FA skeleton map was then projected onto each subjects FA maps in standard space. Finally, regional FA values were extracted from the John's Hopkins University (JHU) White matter atlas [60] and used in group level analyses. We also extracted axial diffusivity (AD), radial diffusivity (RD) and mean diffusivity (MD) for exploratory follow-up analyses (see Supplementary Information).

Statistical analysis

Statistical analyses were performed with IBM SPSS Statistics for Macintosh, Version 28 [61]. Participant demographics, trauma histories, and symptoms were evaluated with chi-square tests, Pearson's correlations, and independent sample t-tests for differences across imaging sites. Linear regressions covarying for MRI scanner site, age, and sex at birth assessed effects of childhood maltreatment load and posttraumatic outcomes on FA in bilateral white matter tracts. These tests were conducted for the 18 individual white matter tracts included in the JHU atlas. FA was examined due to its predominance in the literature. Relationships with AD, RD, and MD were examined in exploratory follow-up analyses for significant tracts

in the FA analysis (see Supplementary information). Identical follow-up tests evaluated the contribution of the subcomponents of tracts significantly associated with childhood maltreatment load. A nominal significance threshold was set at p < 0.05, 2-tailed. False discovery rate (FDR) correction using the Benjamini-Hochberg method was used to control for multiple comparisons and maintain $\alpha = 0.05$. For statistically significant models where subcomponent data was available (e.g., the anterior limb of the internal capsule), identical follow-up models were completed with separate FDR correction using the Benjamini-Hochberg method. Linear models covarying for MRI scanner site, age, and sex at birth evaluated effects of summed exposure to moderate to extreme threat (physical, emotional, and sexual abuse) and deprivation (emotional and physical neglect) components of childhood maltreatment load, as well as their interaction, on major bilateral white matter tracts significantly associated with childhood maltreatment load. Tracts that showed a significant association with childhood trauma were also included in subsequent mediation analyses, conducted using the PROCESS macro version 4 [62], including childhood trauma load, posttraumatic outcomes at 6-months, and a mediator of white matter microstructure. For mediation analyses, we completed bootstrapping with 5000 permutations to obtain 95% bias-corrected confidence intervals as an inferential test of direct and indirect effects. Lastly, univariate effects of childhood maltreatment load on 6-month bilateral white matter tracts significantly associated with childhood maltreatment at 2-weeks were evaluated with ANOVA, in models covarying for scanner site, age, and sex assigned at birth.

RESULTS

Participant characteristics

Participant demographics and trauma characteristics are detailed in Table 1. Samples from the imaging sites were well matched across sex assigned at birth, age, educational attainment, employment, total family income, and marital status (Table S2). Further, each MRI scanning site sample had similar distributions of participants' qualifying traumas and proportions of individuals who hit their head as part of the trauma (Supplementary Information). Participant racial/ethnic identity significantly differed by site (p < 0.001).

Childhood and lifetime trauma load among participants are detailed in Table 2 and distribution of childhood maltreatment load scores in Table S3. On average, participants endorsed greater than one moderate to extreme childhood maltreatment type, with emotional abuse (32.7%) being the most frequently endorsed followed by sexual abuse (24.8%) and emotional neglect (24.8%). There were no significant site differences in the prevalence of any maltreatment subtype or the average number of moderate to extreme maltreatment subtypes endorsed (Table S4). However, there were significant site differences in the modified total LEC score (p = 0.002). Associations between childhood maltreatment load, modified total LEC score, and 6-month symptom scores at each site are shown in Table S5. Participants did not significantly differ in 6-month PTSD, depression, anxiety, or dissociation symptoms across sites (Table S6). Of note, participants included in the present analyses reported significantly lower 6-month PTSD symptoms and total childhood trauma scores than those excluded due to MRI issues but did not differ in childhood maltreatment load (Supplementary Information).

Childhood maltreatment and white matter

Childhood maltreatment load was associated with FA of several white matter tracts (Table 3). Following FDR correction, childhood maltreatment load negatively varied with FA in bilateral internal capsule (IC) at 2-weeks post-trauma, after covarying for sex assigned at birth, scanner site, and age (Table 3). Given the significant relationship between the IC and childhood maltreatment load, we considered the contribution of the IC subcomponents including the Posterior Limb of the IC (PL-IC), the Retrolenticular Part of the IC (RL-IC), and the Anterior Limb of the IC (AL-IC) in identical models and found childhood maltreatment load significantly negatively varied with all 3 IC subcomponents after FDR correction, though the PL-IC was the strongest

5142

	5 1		
			Overall (<i>n</i> = 202)
	Sex Assigned at Birth, <i>N</i> (%)	Male	72 (35.6)
		Female	130 (64.4)
	Age, M (SD)	-	35.4 (12.99)
	Highest Grade, N (%)	Some HS/Below	9 (4.5)
		High School/GED	52 (25.7)
		Some College	70 (34.7)
		Associate Degree	21 (10.4)
		Bachelor's Degree	36 (17.8)
		Graduate Degree	14 (6.9)
	Employment, N (%)	Employed	140 (69.3)
		Retired	6 (3)
		Homemaker	9 (4.5)
		Student	9 (4.5)
		Unemployed, Disabled/ Other	38 (18.8)
	Total Family	TFI ≤\$19000	54 (26.7)
	Income (TFI), <i>N</i> (%)	\$19001≤TFI≤\$35000	69 (34.2)
		\$35001≤TFI≤\$50000	28 (13.9)
		\$50001≤TFI≤\$75000	19 (9.4)
		\$75001≤TFI≤\$100000	12 (5.9)
		TFI≥\$100001	19 (9.4)
	Race-Ethnicity,	Hispanic	26 (12.9)
	N (%)	Non-Hispanic White	72 (35.6)
		Non-Hispanic Black	92 (45.5)
		Non-Hispanic Other	11 (5.4)
	Marital, N (%)	Married	36 (17.8)
		Separated	3 (1.5)
		Divorced	27 (13.4)
		Widowed	2 (1.0)
		Never Married	134 (65.8)
	ED Event, N (%)	Motor Vehicle Collision	151 (74.8)
		Physical Assault	18 (8.9)
		Sexual Assault	1 (.5)
		Fall > = 10 feet	3 (1.5)
		Non-motor Collision	8 (4.0)
		Fall <10 / unknown	9 (4.5)
		Burns	1 (.5)
		Animal Related	7 (3.5)
		Other	4 (2)
	Hit Head, N (%)	No	98 (48.5)
		Yes	86 (42.6)
		Missing	18 (8.9)

Table 1. Demographics and trauma characteristics.

ED Emergency Department.

contributor to the effect (Table 3; Fig. 1). Exploratory follow-up analyses with other diffusivity metrics revealed childhood maltreatment load was also associated with RD in the PL-IC (Table S7). Additional statistical analyses found only the threat (physical, emotional, and sexual abuse; $\beta = -0.19$, p = 0.01), not deprivation (emotional and physical neglect; $\beta = -0.05$, p = 0.53), component of childhood maltreatment load significantly contributed to the observed effect on the IC when both dimensions were included in

Table 2. Childhood and lifetime trauma load.

		Overall (<i>n</i> = 202)
# Mlx Types (0–5), M (SD)	-	1.21 (1.53)
Emotional Abuse, M (SD)		2.50 (2.63)
Mod-Extreme Emotional Abuse, N (%)	No	136 (67.3)
/ M (SD)	Yes	66 (32.7)
Sexual Abuse, M (SD)		1.66 (2.91)
Mod-Extreme Sexual Abuse, N (%) / M	No	152 (75.2)
(SD)	Yes	50 (24.8)
Physical Abuse, M (SD)		1.56 (2.46)
Mod-Extreme Physical Abuse, N (%)	No	158 (78.2)
/ M (SD)	Yes	44 (21.8)
Emotional Neglect, M (SD)		1.86 (2.36)
Mod-Extreme Emotional Neglect, N (%)	No	152 (75.2)
/ M (SD)	Yes	50 (24.8)
Physical Neglect, M (SD)		1.49 (2.15)
Mod-Extreme Physical Neglect, N (%)	No	168 (83.2)
/ M (SD)	Yes	34 (16.8)
CTQ Total (0-44), M (SD)	-	9.91 (9.91)
mLEC-5 Score (0–17), M (SD)		6.77 (4.65)

Note: *Mlx* Maltreatment; *CTQ* Childhood Trauma Questionnaire (11 item); *mLEC-5 Score* modified total Life Events Checklist Score.

 Table 3.
 Significant univariate effects in childhood maltreatment load model (2-Week).

	β	t-statistic	FDR adjusted <i>p</i> -values
IC	-0.21	-3.195	0.029**
—PL-IC	-0.21	-3.114	0.006**
-RL-IC	-0.18	-2.655	0.013**
—AL-IC	-0.13	-2.237	0.026**
CR	-0.14	-2.276	0.215
G-CC	-0.12	-2.155	0.194
SS	-0.14	-2.001	0.211

Reported tracts were significant at a nominal p < 0.05. Benjamini–Hochberg adjusted p-values were calculated by multiplying raw p-values by m/i (i = rank, m = total number of tests). Internal Capsule subcomponents were included in a follow-up, identical model, and FDR corrected with Benjamini–Hochberg. The PL-IC, RL-IC, and AL-IC were tested in follow-up models given the significant IC effect and separately FDR corrected. **Significant After Benjamini–Hochberg (p < 0.05) Adjustment.

IC Internal Capsule, *CR* Corona Radiata, *G-CC* Genu of the Corpus Callosum, *SS* Sagittal Stratum, *PL-IC* Posterior Limb of the Internal Capsule, *RL-IC* Retrolenticular Part of Internal Capsule, *AL-IC* Anterior Limb of Internal Capsule.

an identical model as described above. The interaction between threat and deprivation was not significant. We further conducted sensitivity analyses to determine if associations between IC FA and childhood maltreatment remained while controlling for prior (i.e., endorsed pre-trauma) PCL-5 scores or mLEC-5 scores. Inclusion of either covariate did not impact the relationship between IC FA and childhood maltreatment load (see Supplementary information).

We performed follow-up analyses to test whether associations between childhood maltreatment and FA of the IC were also observed at 6-months post-trauma. Childhood maltreatment load negatively predicted bilateral IC FA indexed 6-months after trauma

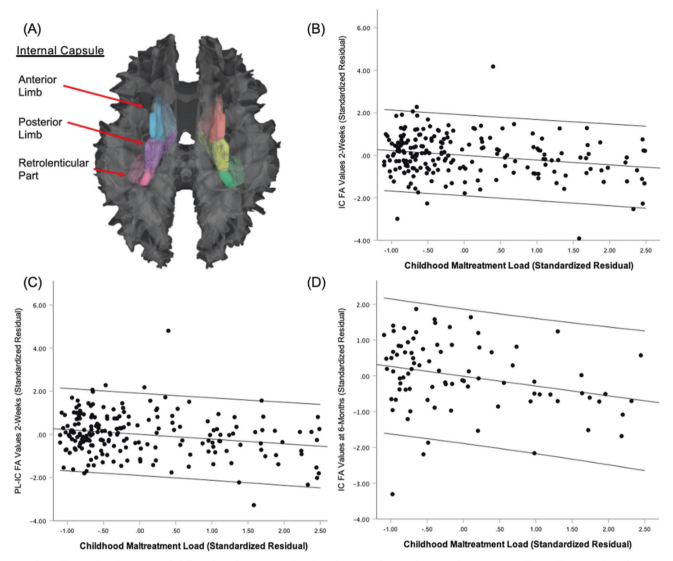


Fig. 1 The relationship between childhood maltreatment load and internal capsule FA values at 2-weeks and 6-months. The internal capsule and its subcomponents are displayed on 3D rendering of human white matter tracts (**A**) Standardized residual plot of the regression of Childhood Maltreatment Load on 2-Week Internal Capsule FA Values depicts the significant negative effect (**B**). Standardized residual plot of Childhood Maltreatment Load on 2-Week Posterior-Limb of the Internal Capsule FA Values depicts the significant negative effect (**C**). Standardized residual plot of the regression of Childhood Maltreatment Load on Internal Capsule FA Values indexed 6-months post-trauma depicts the significant negative effect (**D**).

(Table 4; Fig. 1). In further analyses of IC subparts, negative predictive relationships of childhood maltreatment load with PL-IC and AL-IC microstructure were significant following Benjamini–Hochberg FDR correction (Table 4).

Mediation analyses: childhood maltreatment, IC microstructure, and 6-month PTSD symptoms

Mediation analyses revealed a total effect of childhood maltreatment load on PCL-5 scores at 6-months (b = 1.75, SE = 0.78. 95% CI = [0.20, 3.29]). We found a significant indirect effect of childhood maltreatment load on 6-month PCL-5 scores through IC microstructure (b = 0.37, Boot SE = 0.18, 95% CI = [0.05, 0.76]) that completely mediated the effect of childhood maltreatment load on PCL-5 scores (b = 1.37, SE = 0.79, 95% CI = [-0.18, 2.93]) (Fig. 2). Similar analyses were performed with the total childhood maltreatment score, and we observed similar results (see Supplementary information).
 Table 4.
 Univariate effects in childhood maltreatment load model (6-Month).

	β	t-statistic	FDR adjusted <i>p</i> -value
IC	-0.28	-2.552	0.026**
—PL-IC	-0.31	-2.946	0.016**
—AL-IC	-0.24	-2.318	0.031**
-RL-IC	-0.21	-1.889	0.063

Benjamini–Hochberg adjusted *p*-values were calculated by multiplying raw *p*-values by *m/i* (i = rank, *m* = total number of tests). Internal Capsule subcomponents were included in a follow-up, identical model. The Benjamini–Hochberg FDR-correction included all 6-month tests. **Significant After Benjamini–Hochberg (*p* < 0.05) Adjustment.

PL-IC Posterior Limb of the Internal Capsule, *IC* Internal Capsule, *AL-IC* Anterior Limb of Internal Capsule, *RL-IC* Retrolenticular Part of Internal Capsule.

5144

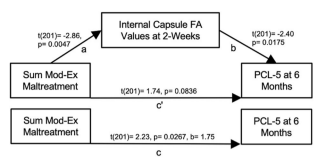


Fig. 2 Mediation model of the effect of childhood maltreatment load on 6-month PCL-5 through Internal Capsule FA values indexed 2-weeks post-trauma. The indirect effect is significant based on a 5000 permutation, bootstrapped 95% confidence interval (i.e., path ab; b = 0.37, Boot SE = 0.18, 95% CI = [0.05, 0.76]), completely mediating the effect of childhood maltreatment load on PCL-5 scores (i.e., path c'; b = 1.37, SE = 0.79, 95% CI = [-0.18, 2.93]).

Exploratory mediation analyses assessed if findings were specific to future PTSD symptoms or if similar relationships were observed with other posttraumatic outcomes including depression, anxiety, and dissociation (Supplementary Information; Figure S1). Although there was a total effect of childhood maltreatment load on 6-month PROMIS-Depression (b = 1.29, SE = 0.48, 95% CI = [0.34, 2.24]), the indirect effect of childhood maltreatment load on 6-month PROMIS-Depression through IC microstructure (b = 0.15, Boot SE = 0.12, 95% CI = [-0.07, 0.39]) was not significant and did not mediate the effect of childhood maltreatment load on 6-month PROMIS-Depression scores (b = 1.14, SE = 0.49, 95% CI = [0.17, 2.11]). There was a total effect of childhood maltreatment load on 6-month PROMIS-Anxiety (b = 0.43, SE = 0.20, 95% CI = [0.05, 0.82]); however, neither the indirect effect of childhood maltreatment load on 6-month PROMIS-Anxiety through IC microstructure (b = 0.05, Boot SE = 0.05, 95% CI = [-0.04, 0.14]) nor the direct effect of childhood maltreatment load on 6-month PROMIS-Anxiety (b = 0.39, SE = 0.20, 95% CI = [-0.01, 0.78]) met statistical significance. No total effect of childhood maltreatment load on dissociation emerged (b = 0.14, SE = 0.08, 95% CI = [-0.01, 0.29]).

DISCUSSION

To our knowledge, the present study is the first to investigate the relationship between childhood maltreatment load and white matter microstructure with posttraumatic symptoms in the early aftermath of trauma. We observed robust relationships between childhood maltreatment load and fractional anisotropy (FA) in the internal capsule (IC) in the early aftermath of an acute trauma event (2-weeks) and 6-months later. Furthermore, variations in IC FA values at 2-weeks fully mediated the relationship between childhood maltreatment load and later posttraumatic symptoms at 6-months. The mediation was specific to posttraumatic symptoms and not observed for depressive, anxiety, or dissociative symptomatology. Given that childhood maltreatment was related to IC microstructure at 2-weeks and 6-months following the adulthood traumatic event, our findings may point to IC FA values as a stable biomarker of later posttraumatic dysfunction and suggest a potential neurobiological pathway through which childhood trauma could confer risk for acute stress reactions in adulthood. Additionally, this study did not reproduce effects of white matter tracts previously found to vary with PTSD symptoms and childhood trauma exposure, including the cingulum bundle, uncinate fasciculus, and corpus callosum.

Our findings implicate the IC as a critical substrate for the effects of childhood trauma on PTSD development. The IC is a dense fiber bundle that contains several projections including the

corticospinal tract, frontopontine and corticofugal fibers, the anterior and superior thalamic radiation, the optic radiation, and the auditory radiation [63, 64]. Anatomically, the IC is limited laterally by the pallidum and medially by the thalamus, the head of the caudate nucleus, and the corticospinal tract [65]. The IC further appears to contain fibers for both medial (hippocampal formation, mammillary bodies, anterior thalamic nuclei, and cingulate gyrus) and basolateral (orbitofrontal cortex, dorsomedial thalamic nucleus, amygdala, and anterior temporal cortex) limbic circuits [66].

The present findings may be related to dysfunctional stimulus processing during PTSD. Although threat processing and its neural substrates are commonly dysregulated in PTSD, these components are dependent on the ability to perceive and integrate sensory stimuli. Recent work suggests variability in structure of visual processing regions, such as the ventral visual stream, is associated with susceptibility to PTSD symptom development [36, 67]. This pathway supports important processes, such as object recognition, integral to threat learning and includes core threat-related regions such as the amygdala and medial PFC [68]. In the current study, we found that higher childhood maltreatment load was associated with lower FA of the IC and its subcomponents. The IC encompasses occipital connections between the higher order visual cortex and temporal lobe [64], as well as components of major motor tracts and somatosensory relays from the thalamus to the cortex [63]. Prior work found trauma-exposed children and adults with childhood maltreatment histories had reduced FA of the IC and its component tracts, including the optic radiations and left anterior thalamic radiation [25, 69]. Speculatively, reduced FA of the IC may reflect disrupted white matter myelination and membrane integrity in fibers that transmit visual sensory information and contribute to altered perception and processing of threat-related information, which, in turn, may contribute to PTSD-related disruptions. Disruption of the IC could further be related to altered ability to consolidate. encode, or retrieve sensory components of trauma memories leading emotion dysregulation. In line with such reasoning, ischemic damage to the IC can lead to cognitive and behavioral alterations such as agitation and impaired attention [63], and deep brain stimulation of the ventral IC/ventral striatum enhances prefrontal cortex driven cognitive control [70]. However, corresponding data on visual processing was not collected in the present study, and specific interpretations of IC function should thus be tempered. Taken together with prior literature, the present results suggest childhood maltreatment has a pronounced effect on IC microstructure which may confer risk for PTSD-related dysregulation following subsequent trauma.

Of note, we did not observe effects in canonical threat circuitry often associated with PTSD. Past studies have not typically considered childhood maltreatment when evaluating white matter markers of PTSD susceptibility [15-18], and it is possible that reduced IC FA may be a sequela of childhood maltreatment exposure. Moreover, although we analyzed imaging data from over 200 participants, we could have been underpowered to detect all effects with our unbiased whole-brain analytic approach. There are likely different biological subtypes of PTSD that are not accounted for here and such heterogeneity may have decreased our ability to detect associations in other tracts. For example, subtypes that show stronger intrusive symptoms or disruptions in emotional memory may be more associated with canonical threat neurocircuitry. Notably, varied white matter microstructure in the IC and its component tracts has been previously implicated in PTSD [11], with recent works suggesting a role in predicting PTSD in the acute aftermath of trauma exposure [16] and in treatment response [71]. Further investigation of the role of childhood maltreatment load in the relationship between white matter microstructure and PTSD development might assist in developing robust predictive models.

5146

The findings of the current work should be interpreted with several considerations. First, we assessed childhood maltreatment load with a retrospective self-assessment. Since we did not query the age participants experienced childhood trauma, we could not assess the role of the developmental timing of trauma on the observed effects. Future longitudinal work is needed to investigate white matter microstructural variability in children and recently traumatized adults with more granular information on childhood trauma exposure and timing. Further, although we used items from the childhood trauma questionnaire, which is itself a validated and broadly used tool, and prospective research suggests the reliability of such retrospective reporting [40, 72-74], we were unable to administer the full questionnaire as to minimize participant burden within the parent study. It would also be beneficial to investigate these associations in longitudinal studies of childhood trauma as opposed to using purely retrospective reports. Secondly, data that could be related to hypothesized contributions of the IC to sensory processes were not available, and thus the specific functional role of variability within IC microstructure in relation to PTSD is unclear. Future research considering the specific targets and functional outcomes of variable IC microstructure among those with childhood trauma would further clarify the present findings. Lastly, our analyses do not consider potential protective or socioeconomic factors that may contribute to early life stress load or resiliency. We did, however, consider the site at which MRI scanning occurred, which largely accounted for participant race and ethnicity. Given the relationships between racial discrimination, neighborhood disadvantage, and socioeconomic status with white matter microstructure [75-77], a critical next step will be to understand how these factors and protective agents impact white matter markers of PTSD susceptibility. As participants in this work reported substantial childhood maltreatment and low levels of PTSD symptoms prior to the presenting trauma, resiliency factors may be especially critical to decipher. Relatedly, it will also be important to consider potentially salient factors such as prenatal exposures and genetics.

The present study of recent trauma survivors examined the relationship of childhood maltreatment load with white matter microstructure and posttraumatic symptoms in the early aftermath of trauma. Childhood maltreatment load stably, inversely varied with the FA of the IC following the acute trauma event. Further, the FA of the IC in the early aftermath of an acute trauma event mediated the relationship between childhood maltreatment load and PTSD symptoms 6-months following the adulthood trauma exposure. These findings suggest a unique role for IC microstructure as a neural pathway between childhood trauma and future PTSD symptoms following a recent trauma. Furthermore, these data suggest DTI imaging may assist in revealing neural signatures of risk for later stress-related dysfunction in those with earlier childhood trauma.

REFERENCES

- Kessler RC, McLaughlin KA, Green JG, Gruber MJ, Sampson NA, Zaslavsky AM, et al. Childhood adversities and adult psychopathology in the WHO World Mental Health Surveys. Br J Psychiatry J Ment Sci. 2010;197:378–85.
- Fitzgerald JM, DiGangi JA, Phan KL. Functional neuroanatomy of emotion and its regulation in PTSD. Harv Rev Psychiatry. 2018;26:116–28.
- Lanius RA, Frewen PA, Vermetten E, Yehuda R. Fear conditioning and early life vulnerabilities: two distinct pathways of emotional dysregulation and brain dysfunction in PTSD. Eur J Psychotraumatology. 2010;1:5467.
- Rauch SL, Shin LM, Phelps EA. Neurocircuitry models of posttraumatic stress disorder and extinction: human neuroimaging research—past, present, and future. Biol Psychiatry. 2006;60:376–82.
- 5. Harnett NG, Goodman AM, Knight DC. PTSD-related neuroimaging abnormalities in brain function, structure, and biochemistry. Exp Neurol. 2020;330:113331.
- Sampaio-Baptista C, Johansen-Berg H. White matter plasticity in the adult brain. Neuron. 2017;96:1239–51.

- Fani N, King TZ, Jovanovic T, Glover EM, Bradley B, Choi K, et al. White matter integrity in highly traumatized adults with and without post-traumatic stress disorder. Neuropsychopharmacology. 2012;37:2740–6.
- Kim SJ, Jeong DU, Sim ME, Bae SC, Chung A, Kim MJ, et al. Asymmetrically altered integrity of cingulum bundle in posttraumatic stress disorder. Neuropsychobiology. 2006;54:120–5.
- O'Doherty DCM, Ryder W, Paquola C, Tickell A, Chan C, Hermens DF, et al. White matter integrity alterations in post-traumatic stress disorder. Hum Brain Mapp. 2018;39:1327–38.
- Sanjuan PM, Thoma R, Claus ED, Mays N, Caprihan A. Reduced white matter integrity in the cingulum and anterior corona radiata in posttraumatic stress disorder in male combat veterans: a diffusion tensor imaging study. Psychiatry Res. 2013;214:260–8.
- Schuff N, Zhang Y, Zhan W, Lenoci M, Ching C, Boreta L, et al. Patterns of altered cortical perfusion and diminished subcortical integrity in posttraumatic stress disorder: an MRI study. NeuroImage. 2011;54:S62–8.
- Abe O, Yamasue H, Kasai K, Yamada H, Aoki S, Iwanami A, et al. Voxel-based diffusion tensor analysis reveals aberrant anterior cingulum integrity in posttraumatic stress disorder due to terrorism. Psychiatry Res Neuroimaging. 2006;146:231–42.
- Zhang L, Li W, Shu N, Zheng H, Zhang Z, Zhang Y, et al. Increased white matter integrity of posterior cingulate gyrus in the evolution of post-traumatic stress disorder. Acta Neuropsychiatr. 2012;24:34–42.
- Kennis M, van Rooij SJH, Tromp DPM, Fox AS, Rademaker AR, Kahn RS, et al. Treatment outcome-related white matter differences in veterans with posttraumatic stress disorder. Neuropsychopharmacology. 2015;40:2434–42.
- Harnett NG, Ference EW, Knight AJ, Knight DC. White matter microstructure varies with post-traumatic stress severity following medical trauma. Brain Imaging Behav. 2020;14:1012–24.
- Hu H, Zhou Y, Wang Q, Su S, Qiu Y, Ge J, et al. Association of abnormal white matter integrity in the acute phase of motor vehicle accidents with posttraumatic stress disorder. J Affect Disord. 2016;190:714–22.
- Weis C, Fitzgerald J, Huggins A, Miskovich T, Isely K, Hanson J, et al. T30. White matter integrity in individuals at-risk for PTSD development: a longitudinal investigation. Biol Psychiatry. 2019;85:S140–1.
- Weis CN, Huggins AA, Miskovich TA, Fitzgerald JM, Bennett KP, Krukowski JL, et al. Acute white matter integrity post-trauma and prospective posttraumatic stress disorder symptoms. Front Hum Neurosci. 2021;15:742198.
- Daniels JK, Lamke JP, Gaebler M, Walter H, Scheel M. White matter integrity and its relationship to PTSD and childhood trauma-a systematic review and metaanalysis: white matter integrity. Depress Anxiety. 2013;30:207–16.
- Siehl S, King JA, Burgess N, Flor H, Nees F. Structural white matter changes in adults and children with posttraumatic stress disorder: A systematic review and meta-analysis. NeuroImage Clin. 2018;19:581–98.
- 21. Teicher MH, Samson JA. Annual research review: enduring neurobiological effects of childhood abuse and neglect. J Child Psychol Psychiatry. 2016;57:241–66.
- Benedetti F, Bollettini I, Radaelli D, Poletti S, Locatelli C, Falini A, et al. Adverse childhood experiences influence white matter microstructure in patients with bipolar disorder. Psychol Med. 2014;44:3069–82.
- Choi J, Jeong B, Rohan ML, Polcari AM, Teicher MH. Preliminary evidence for white matter tract abnormalities in young adults exposed to parental verbal abuse. Biol Psychiatry. 2009;65:227–34.
- Huang H, Gundapuneedi T, Rao U. White matter disruptions in adolescents exposed to childhood maltreatment and vulnerability to psychopathology. Neuropsychopharmacology. 2012;37:2693–701.
- Lim L, Howells H, Radua J, Rubia K. Aberrant structural connectivity in childhood maltreatment: A meta-analysis. Neurosci Biobehav Rev. 2020;116:406–14.
- Lu S, Wei Z, Gao W, Wu W, Liao M, Zhang Y, et al. White matter integrity alterations in young healthy adults reporting childhood trauma: A diffusion tensor imaging study. Aust N. Z J Psychiatry. 2013;47:1183–90.
- Teicher MH, Samson JA, Sheu YS, Polcari A, McGreenery CE. Hurtful words: association of exposure to peer verbal abuse with elevated psychiatric symptom scores and corpus callosum abnormalities. Am J Psychiatry. 2010;167:1464–71.
- Dennis EL, Disner SG, Fani N, Salminen LE, Logue M, Clarke EK, et al. Altered white matter microstructural organization in posttraumatic stress disorder across 3047 adults: results from the PGC-ENIGMA PTSD consortium. Mol Psychiatry [Internet]. 2021 Aug [cited 2021 Aug 31]; Available from: http://www.nature.com/articles/ s41380-019-0631-x
- Gilmartin MR, Balderston NL, Helmstetter FJ. Prefrontal cortical regulation of fear learning. Trends Neurosci. 2014;37:455–64.
- Banihashemi L, Wallace ML, Sheu LK, Lee MC, Gianaros PJ, Mackenzie RP, et al. Childhood maltreatment moderates the effect of combat exposure on cingulum structural integrity. Dev Psychopathol. 2017;29:1735–47.

- Hanson JL, Knodt AR, Brigidi BD, Hariri AR. Lower structural integrity of the uncinate fasciculus is associated with a history of child maltreatment and future psychological vulnerability to stress. Dev Psychopathol. 2015;27:1611–9. 4pt2
- Kuo JR, Kaloupek DG, Woodward SH. Amygdala volume in combat-exposed veterans with and without posttraumatic stress disorder: a cross-sectional study. Arch Gen Psychiatry. 2012;69:1080.
- Woodward SH, Kuo JR, Schaer M, Kaloupek DG, Eliez S. Early adversity and combat exposure interact to influence anterior cingulate cortex volume in combat veterans. NeuroImage Clin. 2013;2:670–4.
- McLean SA, Ressler K, Koenen KC, Neylan T, Germine L, Jovanovic T, et al. The AURORA Study: a longitudinal, multimodal library of brain biology and function after traumatic stress exposure. Mol Psychiatry. 2020;25:283–96.
- 35. Harnett NG, van Rooij SJH, Ely TD, Lebois LAM, Murty VP, Jovanovic T, et al. Prognostic neuroimaging biomarkers of trauma-related psychopathology: resting-state fMRI shortly after trauma predicts future PTSD and depression symptoms in the AURORA study. Neuropsychopharmacology. 2021;46:1263–71.
- Harnett NG, Finegold KE, Lebois LAM, van Rooij SJH, Ely TD, Murty VP, et al. Structural covariance of the ventral visual stream predicts posttraumatic intrusion and nightmare symptoms: a multivariate data fusion analysis. Transl Psychiatry. 2022;12:321.
- Steuber ER, Seligowski AV, Roeckner AR, Reda M, Lebois LAM, van Rooij SJH, et al. Thalamic volume and fear extinction interact to predict acute posttraumatic stress severity. J Psychiatr Res. 2021;141:325–32.
- Stevens JS, Harnett NG, Lebois LAM, van Rooij SJH, Ely TD, Roeckner A, et al. Brain-based biotypes of psychiatric vulnerability in the acute aftermath of trauma. Am J Psychiatry. 2021;178:1037–49.
- Bernstein DP, Fink L, Handelsman L, Foote J, Lovejoy M, Wenzel K, et al. Initial reliability and validity of a new retrospective measure of child abuse and neglect. Am J Psychiatry. 1994;151:1132–6.
- Bernstein DP, Stein JA, Newcomb MD, Walker E, Pogge D, Ahluvalia T, et al. Development and validation of a brief screening version of the Childhood Trauma Questionnaire. Child Abus Negl. 2003;27:169–90.
- Kim YJ, Rooij SJH, Ely TD, Fani N, Ressler KJ, Jovanovic T, et al. Association between posttraumatic stress disorder severity and amygdala habituation to fearful stimuli. Depress Anxiety. 2019;36:647–58.
- Powers A, Ressler KJ, Bradley RG. The protective role of friendship on the effects of childhood abuse and depression. Depress Anxiety. 2009;26:46–53.
- Stanhope KK, Suglia SF, Boulet SL, Powers A, Michopoulos V. Childhood trauma and postpartum care use, estimating mediation by posttraumatic stress disorder and depressive symptoms. Ann Epidemiol. 2022;76:1–6.
- Gray MJ, Litz BT, Hsu JL, Lombardo TW. Psychometric properties of the life events checklist. Assessment. 2004;11:330–41.
- Weathers FW, Blake DD, Schnurr PP, Kaloupek DG, Marx BP, Keane TM. The life events checklist for DSM-5 (LEC-5). Instrument available from the National Center for PTSD at www.ptsd.va.gov. 2013.
- Blevins CA, Weathers FW, Davis MT, Witte TK, Domino JL. The posttraumatic stress disorder checklist for *DSM-5* (PCL-5): development and initial psychometric evaluation: posttraumatic stress disorder checklist for *DSM-5*. J Trauma Stress. 2015;28:489–98.
- Weathers FW, Litz BT, Keane TM, Palmieri PA, Marx BP, Schnurr PP. The PTSD checklist for DSM-5 (PCL-5). Scale available from the National Center for PTSD at www.ptsd.va.gov. 2013.
- Amtmann D, Kim J, Chung H, Bamer AM, Askew RL, Wu S, et al. Comparing CESD-10, PHQ-9, and PROMIS depression instruments in individuals with multiple sclerosis. Rehabil Psychol. 2014;59:220–9.
- Pilkonis PA, Choi SW, Reise SP, Stover AM, Riley WT, Cella D, et al. Item banks for measuring emotional distress from the Patient-Reported Outcomes Measurement Information System (PROMIS[®]): depression, anxiety, and anger. Assessment. 2011;18:263–83.
- PROMIS Health Organization, PROMIS Cooperative Group. PROMIS Depression Scoring Manual. In: HealthMeasures: Patient-Reported Outcomes Measurement Information System (PROMIS) Scoring Manuals [Internet]. 2015. Available from: http://www.healthmeasures.net/index.php?option=com_content&view=article& id=180&/temid=994
- 51. PROMIS Cooperative Group, PROMIS Health Organization. PROMIS Anxiety Scoring Manual. In: HealthMeasures: Patient-Reported Outcomes Measurement Information System (PROMIS) Scoring Manuals [Internet]. 2015. Available from: http://www.healthmeasures.net/index.php?option=com_content&view=article& id=180<emid=994
- 52. Dalenberg CJ, Carlson EB. New versions of the dissociative experiences scale: the DES-R (revised) and the DES-B (brief). Annual Meeting of the International Society for Traumatic Stress Studies. Montreal, Quebec. 2010.
- Harnett NG, Stevens JS, Rooij SJH, Ely TD, Michopoulos V, Hudak L, et al. Multimodal structural neuroimaging markers of risk and recovery from posttrauma anhedonia: A prospective investigation. Depress Anxiety. 2021;38:79–88.

- Roalf DR, Quarmley M, Elliott MA, Satterthwaite TD, Vandekar SN, Ruparel K, et al. The impact of quality assurance assessment on diffusion tensor imaging outcomes in a large-scale population-based cohort. NeuroImage. 2016;125:903–19.
- Andersson JLR, Sotiropoulos SN. An integrated approach to correction for offresonance effects and subject movement in diffusion MR imaging. NeuroImage. 2016;125:1063–78.
- Smith SM, Jenkinson M, Woolrich MW, Beckmann CF, Behrens TEJ, Johansen-Berg H, et al. Advances in functional and structural MR image analysis and implementation as FSL. NeuroImage. 2004;23:S208–19.
- 57. Wang S, Peterson DJ, Gatenby JC, Li W, Grabowski TJ, Madhyastha TM. Evaluation of field map and nonlinear registration methods for correction of susceptibility artifacts in diffusion MRI. Front Neuroinformatics [Internet]. 2017;11. Available from: http://journal.frontiersin.org/article/10.3389/fninf.2017.00017/full
- Smith SM, Jenkinson M, Johansen-Berg H, Rueckert D, Nichols TE, Mackay CE, et al. Tract-based spatial statistics: Voxelwise analysis of multi-subject diffusion data. NeuroImage. 2006;31:1487–505.
- Jahanshad N, Kochunov PV, Sprooten E, Mandl RC, Nichols TE, Almasy L, et al. Multi-site genetic analysis of diffusion images and voxelwise heritability analysis: A pilot project of the ENIGMA-DTI working group. NeuroImage. 2013;81:455–69.
- Hua K, Zhang J, Wakana S, Jiang H, Li X, Reich DS, et al. Tract probability maps in stereotaxic spaces: analyses of white matter anatomy and tract-specific quantification. NeuroImage. 2008;39:336–47.
- IBM Corp. IBM SPSS Statistics for Macintosh, Version 28.0. Armonk, NY: IBM Corp; 2021.
- 62. Hayes AF. Introduction to mediation, moderation, and conditional process analysis: A regression-based approach. New York, NY: Guilford publications; 2017.
- Emos MC, Agarwal S. Neuroanatomy, internal capsule. In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing; 2021 [cited 2021 Nov 4]. Available from: http://www.ncbi.nlm.nih.gov/books/NBK542181/
- Zarei M, Johansen-Berg H, Jenkinson M, Ciccarelli O, Thompson AJ, Matthews PM. Two-dimensional population map of cortical connections in the human internal capsule. J Magn Reson Imaging. 2007;25:48–54.
- Sullivan EV, Zahr NM, Rohlfing T, Pfefferbaum A. Fiber tracking functionally distinct components of the internal capsule. Neuropsychologia. 2010;48:4155–63.
- Livingston KE, Escobar A. Anatomical bias of the limbic system concept: a proposed reorientation. Arch Neurol. 1971;24:17.
- 67. Harnett NG, Stevens JS, Fani N, van Rooij SJH, Ely TD, Michopoulos V, et al. Acute posttraumatic symptoms are associated with multimodal neuroimaging structural covariance patterns: a possible role for the neural substrates of visual processing in posttraumatic stress disorder. Biol Psychiatry Cogn Neurosci Neuroimaging. 2022;7:129–38.
- Kravitz DJ, Saleem KS, Baker CI, Ungerleider LG, Mishkin M. The ventral visual pathway: an expanded neural framework for the processing of object quality. Trends Cogn Sci. 2013;17:26–49.
- Monteleone AM, Monteleone P, Esposito F, Prinster A, Ruzzi V, Canna A, et al. The effects of childhood maltreatment on brain structure in adults with eating disorders. World J Biol Psychiatry. 2019;20:301–9.
- Widge AS, Zorowitz S, Basu I, Paulk AC, Cash SS, Eskandar EN, et al. Deep brain stimulation of the internal capsule enhances human cognitive control and prefrontal cortex function. Nat Commun. 2019;10:1536.
- Graziano RC, Vuper TC, Yetter MA, Bruce SE. Treatment outcome of posttraumatic stress disorder: A white matter tract analysis. J Anxiety Disord. 2021;81:102412.
- Bifulco A, Brown GW, Harris TO. Childhood Experience of Care and Abuse (CECA): a retrospective interview measure. J Child Psychol Psychiatry. 1994;35:1419–35.
- Bifulco A, Brown GW, Lillie A, Jarvis J. Memories of childhood neglect and abuse: corroboration in a series of sisters. J Child Psychol Psychiatry. 1997;38:365–74.
- Hardt J, Rutter M. Validity of adult retrospective reports of adverse childhood experiences: review of the evidence. J Child Psychol Psychiatry. 2004;45:260–73.
- Bell KL, Purcell JB, Harnett NG, Goodman AM, Mrug S, Schuster MA, et al. White matter microstructure in the young adult brain varies with neighborhood disadvantage in adolescence. Neuroscience. 2021;466:162–72.
- Fani N, Harnett NG, Bradley B, Mekawi Y, Powers A, Stevens JS, et al. Racial discrimination and white matter microstructure in trauma-exposed black women. Biol Psychiatry. 2022;91:254–61.
- Shaked D, Leibel DK, Katzel LI, Davatzikos C, Gullapalli RP, Seliger SL, et al. Disparities in diffuse cortical white matter integrity between socioeconomic groups. Front Hum Neurosci. 2019;13:198.

ACKNOWLEDGEMENTS

The investigators wish to thank the trauma survivors participating in the AURORA Study. Their time and effort during a challenging period of their lives make our efforts to improve recovery for future trauma survivors possible. This project was supported by NIMH under K00MH119603 and U01MH110925, the US Army MRMC, One Mind, and The Mayday Fund. The content is solely responsibility of the authors and does

5148

not necessarily represent the official views of any of the funders. Support for title page creation and format was provided by AuthorArranger, a tool developed at the National Cancer Institute. Data and/or research tools used in the preparation of this manuscript were obtained from the National Institute of Mental Health (NIMH) Data Archive (NDA). NDA is a collaborative informatics system created by the National Institutes of Health to provide a national resource to support and accelerate research in mental health. Dataset identifier(s): NIMH Data Archive Digital Object Identifier (DOI) 10.15154/1528044. This manuscript reflects the views of the authors and may not reflect the opinions or views of the NIH or of the Submitters submitting original data to NDA.

AUTHOR CONTRIBUTIONS

Design and conceptualization of study: RCK, KCK, SM, and KJR. Design and conceptualization of analysis: SAW and NGH. Data acquisition, recruitment, and logistics: LDS, TJ, PIM, MJS, SLH, FLB, XA, DZ, TCN, GDC, SDL, LTG, KAB, SLR, JPH, ABS, CL, PLH, SS, CWJ, BEP, RAS, LAH, JLP, EH, AMC, CP, DAP, RCM, RMD, BJO, PS, NKR, SEB, MWM, RHP, JJ, DMB, DAP, SHE, JME, RCK, KCK, SAM, KJR. Data processing and statistical analyses: SAW, NGH, VPM, SEB, TJ, LAML, TDE, JSS, and SVR. Data interpretation: SAW, NGH, JSS, and KJR. Drafting of the manuscript: SW, NGH, JSS, and KJR. All authors revised the manuscript critically for important intellectual context and agreed to be accountable for all aspects of the work in ensuring that questions investigated and resolved.

COMPETING INTERESTS

LAML reports unpaid membership on the Scientific Committee for the International Society for the Study of Trauma and Dissociation (ISSTD), grant support from the National Institute of Mental Health (NIMH), K01 MH118467, and the Julia Kasparian Fund for Neuroscience Research. LAML also reports spousal IP payments from Vanderbilt University for technology licensed to Acadia Pharmaceuticals unrelated to the present work. TCN has received research support from NIH, VA, and Rainwater Charitable Foundation, and consulting income from Jazz Pharmaceuticals. In the last 3 years GDC has received research funding from the NSF, NIH and LifeBell AI, and unrestricted donations from AliveCor Inc. Amazon Research, the Center for Discovery. the Gates Foundation, Google, the Gordon and Betty Moore Foundation, MathWorks, Microsoft Research, Nextsense Inc, One Mind Foundation, the Rett Research Foundation, and Samsung Research. GDC has financial interest in AliveCor Inc and Nextsense Inc. He also is the CTO of MindChild Medical and CSO of LifeBell AI and has ownership in both companies. These relationships are unconnected to the current work. SLR reports grants from NIH during the conduct of the study; personal fees from SOBP (Society of Biological Psychiatry) paid role as secretary, other from Oxford University Press royalties, other from APP (American Psychiatric Publishing Inc.) royalties, other from VA (Veterans Administration) per diem for oversight committee, and other from Community Psychiatry/Mindpath Health paid board service, including equity outside the submitted work; other from National Association of Behavioral Healthcare for paid Board service; and Leadership roles on Board or Council for SOBP, ADAA (Anxiety and Depression Association of America), and NNDC (National Network of Depression Centers). SS has received funding from the Florida Medical Malpractice Joint Underwriter's Association Dr. Alvin E. Smith Safety of Healthcare Services Grant; Allergan Foundation; the NIH/NIA-funded Jacksonville Aging Studies Center (JAX-ASCENT; R33AG05654); and the Substance Abuse and Mental Health Services Administration (1H79Tl083101-01); and the Florida Blue Foundation. CWJ has no competing interests related to this work, though he has been an investigator on studies funded by AstraZeneca, Vapotherm, Abbott, and Ophirex. JJ receives consulting payments from Janssen Pharmaceuticals. DMB has received function from the NIMH, NIDA, and the American Foundation for Suicide Prevention, and consults for Boehringer Ingelheim. Over the past 3 years, DAP has received consulting fees from Albright Stonebridge Group, Boehringer Ingelheim, Compass Pathways, Concert Pharmaceuticals, Engrail Therapeutics, Neumora Therapeutics (former BlackThorn Therapeutics), Neurocrine Biosciences, Neuroscience Software, Otsuka Pharmaceuticals, and Takeda Pharmaceuticals; honoraria from the Psychonomic Society (for editorial work) and Alkermes, and research funding from NIMH, Dana Foundation,

Brain and Behavior Research Foundation, and Millennium Pharmaceuticals. In addition, he has received stock options from Neumora Therapeutics (former BlackThorn Therapeutics), Compass Pathways, Engrail Therapeutics, and Neuroscience Software. SEH has no competing interests related to this work, though in the last three years he has received research funding from NIH, Aptinyx, and Arbor Medical Innovations, and consulting payments from Aptinyx, Heron Therapeutics, and Memorial Sloan Kettering Cancer Center, JME reports support from the National Institutes of Health (NIH) through Grant Numbers R01HD079076 & R03HD094577: Eunice Kennedy Shriver National Institute of Child Health & Human Development; National Center for Medical Rehabilitation Research. He also reports funding from New South Wales Health, Spinal Cord Injury Award (2020-2025) and consulting fees (<\$15,000 per annum) from Orofacial Therapeutics, LLC. In the past 3 years, RCK was a consultant for Cambridge Health Alliance, Canandaigua VA Medical Center, Holmusk, Partners Healthcare, Inc., RallyPoint Networks, Inc., and Sage Therapeutics. He has stock options in Cerebral Inc., Mirah, PYM, and Roga Sciences. KCK research has been supported by the Robert Wood Johnson Foundation, the Kaiser Family Foundation, the Harvard Center on the Developing Child, Stanley Center for Psychiatric Research at the Broad Institute of MIT and Harvard, the National Institutes of Health, One Mind, the Anonymous Foundation, and Cohen Veterans Bioscience. She has been a paid consultant for Baker Hostetler, Discovery Vitality, and the Department of Justice. She has been a paid external reviewer for the Chan Zuckerberg Foundation, the University of Cape Town, and Capita Ireland. She has had paid speaking engagements in the last 3 years with the American Psychological Association, European Central Bank. Sigmund Freud University – Milan, Cambridge Health Alliance, and Coverys. She receives royalties from Guilford Press and Oxford University Press. SAM serves as a consultant for Walter Reed and for Arbor Medical Innovations, KJR has performed scientific consultation for Bioxcel, Bionomics, Acer. Takeda, and Jazz Pharma; serves on Scientific Advisory Boards for Sage and the Brain Research Foundation, and he has received sponsored research support from Takeda, Brainsway and Alto Neuroscience. NGH reports grant support from the National Institute of Mental Health, K00 MH119603.

ADDITIONAL INFORMATION

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41380-023-02012-3.

Correspondence and requests for materials should be addressed to Jennifer S. Stevens or Nathaniel G. Harnett.

Reprints and permission information is available at http://www.nature.com/ reprints

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http:// creativecommons.org/licenses/by/4.0/.

© The Author(s) 2023

¹Division of Depression and Anxiety, McLean Hospital, Belmont, MA, USA. ²Department of Psychiatry, Harvard Medical School, Boston, MA, USA. ³Department of Psychiatry and Behavioral Sciences, Emory University School of Medicine, Atlanta, GA, USA. ⁴Department of Psychological Sciences, University of Missouri—St. Louis, St. Louis, MO, USA. ⁵Department of Psychology, Temple University, Philadelphia, PA, USA. ⁶Department of Psychiatry and Behavioral Neurosciences, Wayne State University, Detroit, MI, USA. ⁷Department of Emergency Medicine, Washington University School of Medicine, St. Louis, MO, USA. ⁸Department of Epidemiology, Brown University, Providence, RI, USA. ⁹Department of Emergency Medicine, Brown University, Providence, RI, USA. ¹⁰Institute for Trauma Recovery, Department of Anesthesiology, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA. ¹¹Department of Biostatistics, Gillings School of Global Public Health, University of North Carolina, Chapel Hill, NC, USA. ¹²Department of Psychiatry Atlanta, GA, USA. ¹⁴Department of Biomedical Engineering, Georgia Institute of Technology and Emory University, Atlanta, GA, USA. ¹⁵Institute for Technology in Psychiatry McLean Hospital, Belmont, MA, USA. ¹⁶The Many Brains Project, Belmont, MA, USA. ¹⁷Department of Psychology and Neuroscience & Department of Sociology, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA. ¹⁸Department of Psychiatry, McLean Hospital, Belmont, MA, USA. ¹⁹Department of Emergency Medicine, University of Massachusetts Chan Medical School, Worcester, MA, USA. ²⁰Department of Emergency Medicine, Vanderbilt University Medical Center, Nashville, TN, USA. ²¹Department of Emergency Medicine, Henry Ford Health System, Detroit, MI, USA. 22 Department of Emergency Medicine, Indiana University School of Medicine, Indianapolis, IN, USA. ²³Department of Emergency Medicine, University of Florida College of Medicine—Jacksonville, Jacksonville, FL, USA.²⁴Department of Emergency Medicine, Cooper Medical School of Rowan University, Camden, NJ, USA.²⁵Department of Emergency Medicine, Ohio State University College of Medicine, Columbus, OH, USA.²⁶Ohio State University College of Nursing, Columbus, OH, USA. ²⁷Department of Emergency Medicine, University of Alabama School of Medicine, Birmingham, AL, USA. ²⁸Department of Surgery, Division of Acute Care Surgery, University of Alabama School of Medicine, Birmingham, AL, USA. ²⁹Center for Injury Science, University of Alabama at Birmingham, Birmingham, AL, USA. ³⁰Department of Emergency Medicine, Oakland University William Beaumont School of Medicine, Rochester, MI, USA. ³¹Department of Emergency Medicine, Emory University School of Medicine, Atlanta, GA, USA. ³²Department of Surgery, Department of Neurosurgery, University of Pennsylvania, Philadelphia, PA, USA. ³³Perelman School of Medicine, University of Pennsylvania, Philadelphia, PA, USA. ³⁴Department of Surgery, Division of Traumatology, Surgical Critical Care and Emergency Surgery, University of Pennsylvania, Philadelphia, PA, USA, 35 Department of Emergency Medicine, Wayne State University, Ascension St. John Hospital, Detroit, MJ, USA, ³⁶Department of Emergency Medicine, Massachusetts General Hospital, Boston, MA, USA. ³⁷Department of Emergency Medicine, Brigham and Women's Hospital, Boston, MA, USA. ³⁸Department of Emergency Medicine, Saint Joseph Mercy Hospital, Ypsilanti, MI, USA. ³⁹Department of Emergency Medicine, University of Massachusetts Medical School-Baystate, Springfield, MA, USA. ⁴⁰Department of Emergency Medicine, Wayne State University, Detroit Receiving Hospital, Detroit, MI, USA. ⁴¹Department of Emergency Medicine, McGovern Medical School at UTHealth, Houston, TX, USA. ⁴²Department of Emergency Medicine, Harvard Medical School, Boston, MA, USA. ⁴³National Center for PTSD, Behavioral Science Division, VA Boston Healthcare System, Boston, MA, USA. 44 Department of Psychiatry, Boston University School of Medicine, Boston, MA, USA. ⁴⁵National Center for PTSD, Clinical Neurosciences Division, VA Connecticut Healthcare System, West Haven, CT, USA. ⁴⁶Department of Psychiatry, Yale School of Medicine, New Haven, CT, USA. 47 Department of Psychology, Yale University, New Haven, CT, USA. 48 Department of Psychological & Brain Sciences, Washington University in St. Louis, St. Louis, MO, USA. 49 Department of Anesthesiology, University of Michigan Medical School, Ann Arbor, MI, USA. 50 Department of Internal Medicine-Rheumatology, University of Michigan Medical School, Ann Arbor, MI, USA. ⁵¹Kolling Institute, University of Sydney, St Leonards, NSW, Australia. ⁵²Faculty of Medicine and Health, University of Sydney, Northern Sydney Local Health District, Camperdown, NSW, Australia. ⁵³Physical Therapy & Human Movement Sciences, Feinberg School of Medicine, Northwestern University, Chicago, IL, USA. 54 Department of Health Care Policy, Harvard Medical School, Boston, MA, USA. 55 Department of Epidemiology, Harvard T.H. Chan School of Public Health, Harvard University, Boston, MA, USA. 56 Department of Emergency Medicine, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA. ⁵⁷Institute for Trauma Recovery, Department of Psychiatry, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA. ⁵⁸These authors contributed equally: Jennifer S. Stevens, Nathaniel G. Harnett. Memail: jennifer.stevens@emory.edu; nharnett@mclean.harvard.edu