ARTICLES

Brain-Based Biotypes of Psychiatric Vulnerability in the Acute Aftermath of Trauma

Jennifer S. Stevens, Ph.D., Nathaniel G. Harnett, Ph.D., Lauren A.M. Lebois, Ph.D., Sanne J.H. van Rooij, Ph.D., Timothy D. Ely, B.A., Alyssa Roeckner, B.S., Nico Vincent, B.S., Francesca L. Beaudoin, M.D., Ph.D., Xinning An, Ph.D., Donglin Zeng, Ph.D., Thomas C. Neylan, M.D., Gari D. Clifford, D.Phil., Sarah D. Linnstaedt, Ph.D., Laura T. Germine, Ph.D., Scott L. Rauch, M.D., Christopher Lewandowski, M.D., Alan B. Storrow, M.D., Phyllis L. Hendry, M.D., Sophia Sheikh, M.D., Paul I. Musey, Jr., M.D., John P. Haran, M.D., Ph.D., Christopher W. Jones, M.D., Brittany E. Punches, Ph.D., Michael S. Lyons, M.D., M.P.H., Michael C. Kurz, Meghan E. McGrath, M.D., Jose L. Pascual, M.D., Ph.D., Elizabeth M. Datner, M.D., Anna M. Chang, M.D., Claire Pearson, M.D., David A. Peak, M.D., Robert M. Domeier, M.D., Brian J. O’Neil, M.D., Niels K. Rathlev, M.D., Leon D. Sanchez, M.D., M.P.H., Robert H. Pietrzak, Ph.D., M.P.H., Jutta Joormann, Ph.D., Deanna M. Barch, Ph.D., Diego A. Pizzagalli, Ph.D., John F. Sheridan, Ph.D., Beatriz Luna, Ph.D., Steven E. Harte, Ph.D., James M. Elliott, Ph.D., Vishnu P. Murty, Ph.D., Tanja Jovanovic, Ph.D., Steven E. Bruce, Ph.D., Stacey L. House, M.D., Ph.D., Ronald C. Kessler, Ph.D., Karestan C. Koenen, Ph.D., Samuel A. McLean, M.D., M.P.H., Kerry J. Ressler, M.D., Ph.D.

Objective: Major negative life events, such as trauma exposure, can play a key role in igniting or exacerbating psychopathology. However, few disorders are diagnosed with respect to precipitating events, and the role of these events in the unfolding of new psychopathology is not well understood. The authors conducted a multisite transdiagnostic longitudinal study of trauma exposure and related mental health outcomes to identify neurobiological predictors of risk, resilience, and different symptom presentations.

Methods: A total of 146 participants (discovery cohort: N = 69; internal replication cohort: N = 77) were recruited from emergency departments within 72 hours of a trauma and followed for the next 6 months with a survey, MRI, and physiological assessments.

Results: Task-based functional MRI 2 weeks after a motor vehicle collision identified four clusters of individuals based on profiles of neural activity reflecting threat reactivity, reward reactivity, and inhibitory engagement. Three clusters were replicated in an independent sample with a variety of trauma types. The clusters showed different longitudinal patterns of post-trauma symptoms.

Conclusions: These findings provide a novel characterization of heterogeneous stress responses shortly after trauma exposure, identifying potential neuroimaging-based biotypes of trauma resilience and psychopathology.

AJP in Advance; doi: 10.1176/appi.ajp.2021.20101526

The diathesis-stress model of psychopathology has remained one of the most well-supported theories addressing the causes of mental disorders. In combination with predisposing factors, antecedent stressors increase risk for the onset and recurrence of depression (1), schizophrenia (2), insomnia (3), and posttraumatic stress disorder (PTSD) (4). However, stress and its severity or chronicity alone cannot account for the wide variety of different types of mental health outcomes that can follow major stressful life events (5), theoretically driven by existing individual differences (6). These variations and their biological bases are not well captured by existing definitions of psychiatric disorders. In the present study, our objective was to discover brain-based profiles to map heterogeneity following a stressor in a nationwide longitudinal study of trauma exposure and subsequent mental health outcomes, the Advancing Understanding of Recovery After Trauma (AURORA) study (7).

Neuroimaging is an attractive tool for mapping symptoms to biology. Previous efforts to account for heterogeneity have often explored brain-wide patterns of activation or connectivity to identify “biotypes,” subtypes of a particular form of psychopathology that differ in their neurophysiological features (8–10). The identification of such subgroups may in turn improve our understanding of variance in outcomes and response to treatment. However, previous work has defined the neuroimaging features of interest on the basis of their association with either a specific symptom type or the response to treatment. These backward
inferences constrain the solution to features that already have high relevance to a diagnostic category, potentially excluding features that contribute to atypical symptom profiles and raising concerns related to overfitting (11). Taking a complementary approach, we constructed a forward inference model, examining neuroimaging profiles in the acute posttrauma period and then investigating their association with the emergence of later symptoms. The goal was to identify posttrauma biotypes with relevance to overall stress vulnerability and resilience but not specific to a particular diagnosis or symptom.

Neural models of stress vulnerability involve hyperreactivity of regions involved in threat detection and the fear response, such as the amygdala, insula, and dorsal anterior cingulate cortex (dACC) (12–15). In addition, both chronic depression and PTSD emerging following a stressful event appear to be preceded by low reward reactivity in affective-evaluative regions, including the nucleus accumbens (NAcc), amygdala, and orbitofrontal cortex (OFC) (16–20). Finally, reduced pretrauma or early posttrauma engagement of regions involved in inhibition, including the ventromedial prefrontal cortex (vmPFC) and hippocampus, is also predictive of greater subsequent PTSD and lower resilience (12, 21–24). Therefore, an early posttrauma profile of co-occurring high threat reactivity, low reward reactivity, and low inhibition would likely be predictive of later chronic symptoms of PTSD and depression. However, it is not yet clear whether these features co-occur within particular individuals or groups. Building a brain-based model of individual differences in the response to major stressors is critical for efforts to construct effective intervention and prevention strategies for stress-related psychiatric disorders.

Here, we collected functional MRI (fMRI) scans in a regionally diverse cohort of civilian trauma survivors 2 weeks posttrauma (7). Participants reported on symptoms of psychopathology through mobile surveys over the first 6 months posttrauma. fMRI-based phenotypes used in the biotyping analysis were motivated by previous longitudinal studies of stress exposure and included all brain regions previously linked with vulnerability to poststress psychopathology within the domains of threat responsivity, reward responsivity, and inhibition/impulsivity as described above. Participants engaged in fMRI tasks that were simple in their design and interpretation and that have been widely used to probe threat (12), reward (25), and response inhibition (26). Multivariate profiles of regional activation were entered into a hierarchical clustering analysis to identify brain-based groupings of individuals in the early posttrauma period, indicative of distinct biotypes. We predicted that fMRI-based clusters would be associated with different patterns of subsequent posttrauma symptoms across PTSD, dissociation, anxiety, depression, and impulsivity. Finally, to better understand whether the clusters overlap with widely known biomarkers of chronic posttraumatic pathology, such as deficits in fear inhibition (27) and extinction (28), we tested whether the clusters differed in these features in a fear-potentiated startle paradigm conducted on the same day as the fMRI scan.

**METHODS**

**Participants**

Participants were recruited from emergency departments as part of a multisite longitudinal study of adverse neuropsychiatric sequelae of trauma (7). Twenty-two emergency departments within the Northeast, Southern, Mid-Atlantic, and Midwest regions of the United States enrolled patients within 72 hours of trauma exposure. All participants were ages 18–75, able to speak and read English, oriented to time and place, and physically able to use a smartphone, and they had possessed a smartphone for more than 1 year. Potential participants were excluded if they had a solid organ injury greater than grade 1 or a significant hemorrhage, required a chest tube or general anesthesia, or were likely to be admitted for >72 hours. MRI scans and psychophysiology data were collected a mean of 18 days (SD=6) later at a laboratory visit at McLean Hospital (Belmont, Mass.), Emory University (Atlanta), Temple University (Philadelphia), or Wayne State University (Detroit), which were each located in proximity to multiple enrolling sites. Written informed consent was obtained as approved by each site’s institutional review board.

Data collection for the AURORA study is ongoing. The discovery cohort included an initial sample that was restricted to motor vehicle-related traumas for participants with at least 8 weeks of follow-up data by March 2019 (94 patients completed the MRI visit). Data for the replication cohort were separate from the discovery cohort because these data had not yet been released by the time of the initial analysis. A second freeze and release of the survey data was broadened to include all trauma types with at least 8 weeks of follow-up data by mid-October 2019. Unique participants in this second freeze made up the replication cohort (additional participants, N=108). After quality control, 69 participants in the discovery cohort and 77 in the replication cohort were retained for analyses. The study participants’ demographic characteristics are presented in Table 1.

**Demographic Variables and Psychiatric Assessment**

Trauma severity was measured using an injury severity score, as well as participants’ subjective ratings of their chances of dying. Assessments of pretrauma risk factors included a general physical health status assessment, childhood maltreatment assessment, and demographic variables. Assessments of posttrauma outcomes, including PTSD symptoms, depression symptoms, dissociative symptoms, anxiety symptoms, and impulsivity, were assessed for the pretrauma period (queried in the emergency department) and at 2 weeks (days 7–21), 8 weeks (days 46–67), 3 months (days 77–104), and 6 months (days 168–195) posttrauma. Measures and scoring details are summarized in the online supplement.
MRI

Acquisition. Brain imaging data were acquired on four separate Siemens 3-T MRI scanners using the two-dimensional echo-planar blood-oxygen-level-dependent sequence for functional scans and a magnetization-prepared rapid acquisition gradient echo T1-weighted image for structural scans. Site-specific sequence parameters are presented in Table S5 in the online supplement.

fMRI tasks. The three fMRI tasks (Figure 1) included a threat task designed to probe reactivity to social threat cues (12), an inhibition task, which was a modified version of Liebenluft’s stop-signal task (26), and a reward task, which was a short version of Delgado’s monetary reward task (25).

Preprocessing and analysis. Full preprocessing information is reported in the online supplement. Functional images were preprocessed with fMRIPrep, version 1.2.2 (29). Echo-planar imaging scans were coregistered to the T1-weighted images, then spatially realigned, slice-time corrected, and normalized to the 2009 ICBM-152 template. Volume-wise motion and other sources of artifact were corrected using ICA-AROMA (30). To handle cases in which motion was likely too high for effective independent component analysis correction, we also implemented an overall motion threshold for any run with >15% of volumes showing $1-mm framewise displacement. Images were then smoothed with a 6-mm kernel. Site-by-site quality metrics are plotted in Figure S3 in the online supplement.

The final sample was restricted to participants with good-quality data across all three fMRI tasks (threat, inhibition, and reward). We did not interpolate any data point because the goal of the clustering analysis was to identify existing patterns of activation across the three tasks. Participants were excluded for falx calcification (discovery cohort, N=0; replication cohort, N=5); discontinuing the scan before completing all three tasks (discovery cohort, N=0; replication cohort, N=7); discontinuing the scan before completing all three tasks (discovery cohort, N=0; replication cohort, N=7); discontinuing the scan before completing all three tasks (discovery cohort, N=0; replication cohort, N=7); discontinuing the scan before completing all three tasks (discovery cohort, N=0; replication cohort, N=7).
N=4; replication cohort, N=7); superthreshold head motion on one or more tasks (discovery cohort, N=11; replication cohort, N=5); technical reasons, such as problems with stimulus display on one or more tasks (discovery cohort, N=2; replication cohort, N=4); or low behavioral performance on either the inhibition or reward task (<75% of trials receiving a button press, indicating sleepiness or low effort; discovery cohort, N=8; replication cohort, N=8). The analysis therefore included 69 participants in the discovery cohort and 77 in the replication cohort.

Statistical modeling of the fMRI data and region of interest definitions are detailed in the online supplement. Regions of interest were defined anatomically and included the left and right amygdala, insula, subgenual anterior cingulate cortex (sgACC), dACC (threat: fearful > neutral faces), nucleus accumbens (NAcc), OFC, amygdala (reward: monetary gain > loss), hippocampus, and vmPFC (inhibition: no-go > go).

Fear-potentiated startle. Participants completed fear-acquisition and extinction tasks on the same day as the MRI scan. Details on the data acquisition and paradigm are presented in the online supplement.

Clustering Analysis
Analyses were conducted in R, version 3.6.3, with RStudio, version 1.2.1335. All tests were two-tailed and used a significance threshold of 0.05, with family-wise error correction as
noted in the Results section. Clustering was conducted on data from the regions of interest extracted from the three fMRI tasks, using hierarchical agglomerated clustering, with the cluster package, version 2.1.0, following Ward’s criterion (agnes function). This is a bottom-up method designed to preserve the existing structure of the data without imposing assumptions of linearity, appropriate for exploratory analysis. The optimal number of clusters was determined using silhouette (31) and distance (32) methods. Nonparametric bootstrapping using the fpc package, version 2.2.5, was applied to the cluster solutions, with 1,000 iterations. After the initial hierarchical clustering, the data were randomly resampled with replacement. In each bootstrap, clustering was performed on the resampled data, and the new cluster most similar to each original cluster was identified by saving the maximum Jaccard coefficient (indexing similarity) for each old-new comparison (33). This was repeated, and a mean permuted Jaccard coefficient was computed across all the bootstraps by cluster. Permuted Jaccard coefficient therefore represents the proportion of individuals from each original cluster solution that were again clustered together in the permuted data. A permuted Jaccard coefficient of 0.6–0.75 indicates stable clusters, >0.75 represents high stability, and <0.50 is thought to indicate cluster instability (33); clusters were considered reconstituted on any bootstrap with a permuted Jaccard coefficient >0.60.

The replication was assessed quantitatively using a train and test approach. We trained a simple k-nearest-neighbors (knn function) model (34) with the class, version 7.3, package using the discovery cohort data, labeled using the cluster labels from the hierarchical clustering solution. We applied this knn model (“test”) in the replication cohort to obtain a new set of labels. We then compared these new labels to the de novo hierarchical clustering of the replication cohort in caret, version 6.0 (35).

Analysis of Posttrauma Outcomes by Cluster
Because clustering produced some small cells within-cohort (Ns as low as 11), we combined the clusters that replicated between the discovery and replication cohorts for further characterization. Cluster assignments from the initial cluster solutions were retained, rather than reclustering in a combined data set.

Chi-square tests (categorical variables) or one-way analyses of variance (ANOVAs) (continuous variables) were used to assess whether demographic factors or trauma-related factors differed between the cluster groups.

Given the multiple overlapping adverse mental health outcomes of trauma, we used multivariate analysis of variance (MANOVA) to test whether the pattern of subsequent mental health outcomes varied across the cluster groups. The outcome was a vector of standardized scores for PTSD symptoms, depression symptoms, dissociation, anxiety, and impulsivity. Predictors included cluster, assessment time point (time-invariant term for the 2-week, 8-week, 3-month, and 6-month posttrauma assessments; linear and quadratic terms), cluster-by-time point interaction, cohort, and a random effect for participant. Post hoc tests separating each outcome type were conducted using linear mixed models in the lme4 package, with the same set of predictors used in the MANOVA. To test whether the fMRI-based clustering provided incremental information above and beyond pretrauma symptom levels, we conducted secondary models including pretrauma symptoms that participants reported in the emergency department. Initial AURORA study findings indicated that among sociodemographic risk factors, pretrauma symptom levels were the strongest predictor of later PTSD and depression symptom severity (36, 37).

Finally, we tested whether a dimensional model of the 2-week fMRI data outperformed cluster assignment in predicting posttrauma outcomes using the first three principal components from the principal component analysis of the nine regions of interest in the combined discovery and replication data set as dimensional predictors of later outcomes. These models used the same structure as the cluster-based analyses. Model fit for dimensional and cluster-based models were directly compared.

Fear-potentiated startle during fear conditioning and extinction was also collected on the same day as the MRI scan (acquisition details are presented in the online supplement). ANOVAs tested whether the cluster groups varied in fear-potentiated startle responses during either acquisition or extinction as a function of cluster, cohort, block, and conditioned stimulus (CS) type and interactions between cluster, block, and CS type.

RESULTS
Covariance Among the fMRI Tasks and Regions of Interest
To assess for feature redundancy, we examined the covariance structure between the tasks and regions of interest. Regions of interest showed positive within-task covariance but not between tasks (Figure 2A,B). The small to moderate correlations suggested that each task and region would contribute unique variance to a clustering analysis. Interestingly, similar regions were uncorrelated from one task to another; for example, participants’ amygdala reactivity to threat was not correlated with amygdala reactivity to reward (r=0.00, p=0.97). This suggests that the “crude” factor (everything correlates with everything) (38) was very low across this set of tasks.

Clustering of Individuals Using Task-Based fMRI
2 Weeks Posttrauma
Hierarchical clustering was first applied in the discovery sample, with 69 survivors of motor vehicle accidents. A four-cluster solution was identified (Figure 2C; see also Figure S1A,B in the online supplement). Silhouette results suggested an optimal clustering with two groups (k=2) but with only a small decrement in width for k=4, whereas Hartigan’s distance metric showed an optimal gain in cluster
FIGURE 2. Functional MRI (fMRI) profiles of four clusters among trauma survivors in the Advancing Understanding of Recovery After Trauma study in the discovery (N=69) and replication (N=77) cohorts.

(A) Brain imaging: 
- Cluster 1: Reactive/Disinhibited 
- Cluster 2: Low Reward/High Threat 
- Cluster 3: High Reward 
- Cluster 4: Inhibited

(B)idebar: 
- Cluster 1: Reactive/Disinhibited 
- Cluster 2: Low Reward/High Threat 
- Cluster 3: High Reward 
- Cluster 4: Inhibited

(E) Heatmap: 
- Contrast Estimate, Z

(F) Scatter plot: 
- PC1 - Threat 
- PC2 - Reward

(G) Cluster distribution: 
- Cluster 1, 2, 3, 4

(H) Cluster distribution: 
- Cluster 1, 2, 3, 4

BRAIN-BASED BIOTYPES OF PSYCHIATRIC VULNERABILITY AFTER TRAUMA
COHESIVENESS AT K = 4. EXAMINATION OF fMRI ACTIVATION PATTERNS (FIGURE 2E,G) INDICATED THAT INDIVIDUALS IN CLUSTER 1 (PERMUTED JACCARD COEFFICIENT = 0.52) SHOWED HIGH ACTIVITY TO BOTH THREAT AND REWARD, WITH LITTLE ENGAGEMENT OF REGULATORY REGIONS IN THREAT OR INHIBITION. THEREFORE, WE CLASSIFIED THE CLUSTER 1 GROUP AS “REACTIVE/DISINHIBITED.” INDIVIDUALS IN CLUSTER 2 (PERMUTED JACCARD COEFFICIENT = 0.54) SHOWED THREAT RESPONSIVENESS DOMINATED BY THE sgACC BUT LOW REWARD REACTIVITY, AND WE CLASSIFIED THIS GROUP AS “LOW REWARD/HIGH THREAT.” INDIVIDUALS IN CLUSTER 3 (PERMUTED JACCARD COEFFICIENT = 0.52) SHOWED NO REACTIVITY TO THREAT, NOR ENGAGEMENT OF THE vmPFC OR HIPPOCAMPUS DURING INHIBITION, BUT VERY HIGH RESPONSIVITY TO REWARD, AND WE CLASSIFIED THIS GROUP AS “HIGH REWARD.” FINALLY, INDIVIDUALS IN CLUSTER 4 (PERMUTED JACCARD COEFFICIENT = 0.56) SHOWED MARKED DEACTIVATION TO THREAT IN THE AMYGDALA, dACC, AND INSULA, SOME ACTIVATION OF THE HIPPOCAMPUS IN THE INHIBITION TASK, AND LITTLE REACTIVITY TO REWARD, AND WE CLASSIFIED THIS GROUP AS “INHIBITED.”

THE REPETITION COHORT INCLUDED 77 PARTICIPANTS WITH A VARIETY OF DIFFERENT TRAUMA TYPES, INCLUDING INTERPERSONAL TRAUMAS. HERE, THE MOST FAVORABLE CLUSTERING SOLUTION INCLUDED THREE GROUPS, WITH AGREEMENT BETWEEN THE SILLUETTE AND DISTANCE METRICS AT K = 3 (FIGURE 2D; SEE ALSO FIGURE SIC AND D IN THE ONLINE SUPPLEMENT). THE GROUPS APPEARED TO BE CONSISTENT WITH CLUSTER 1 (REACTIVE/DISINHIBITED; PERMUTED JACCARD COEFFICIENT = 0.53), CLUSTER 2 (LOW REWARD/HIGH THREAT; PERMUTED JACCARD COEFFICIENT = 0.73), AND CLUSTER 4 (INHIBITED; PERMUTED JACCARD COEFFICIENT = 0.55) FROM THE DISCOVERY SAMPLE (FIGURE 2F,H). THERE WAS A STRIKING ABSENCE OF A HIGH REWARD-LIKE PHENOTYPE; INDIVIDUALS WHO SHOWED HIGH REWARD REACTIVITY ALSO SHOWED HIGH THREAT REACTIVITY.

THE INCLUSION OF HIGHER-IMPACT TRAUMAS MAY HAVE PUSHED REWARD-RESPONSIVE INDIVIDUALS TOWARD HIGHER THREAT REACTIVITY. TO TEST THIS, WE COMBINED BOTH COHORTS AND EXAMINED EFFECTS OF EITHER INJURY SEVERITY OR INTERPERSONAL VIOLENCE ON THREAT ACTIVITY IN THE AMYGDALA. INJURY SEVERITY POSITIVELY PREDICTED AMYGDALA ACTIVITY (F = 4.58, DF = 1, 144, P = 0.03; SEE ALSO FIGURE S2 IN THE ONLINE SUPPLEMENT), WHEREAS INTERPERSONAL COMPARED WITH NONINTERPERSONAL TRAUMA DID NOT (P = 0.61). HIGH REWARD WAS THEREFORE LIKELY SUBSUMED UNDER THE REACTIVE/DISINHIBITED PHENOTYPE, RELATED TO HIGHER-ACUITY TRAUMAS.

In the quantitative assessment of replication for clusters 1, 2, and 4, we assessed the extent to which a model trained on the clustering solution from the discovery cohort could predict the clustering solution within the replication cohort. The model trained on the discovery cohort data had 65.0% (95% CI = 53.2, 75.5) accuracy in predicting the original hierarchical clustering-based labels in the replication cohort, compared with a 45.4% no-information rate (P = 0.0005, Kappa = 0.45). This indicated that the clustering solution in the replication cohort could be recapitulated above and beyond chance levels using only the features of the discovery cohort solution.

THE CLUSTERS WERE UNRELATED TO DEMOGRAPHIC, HEALTH-RELATED, TRAUMA-RELATED, OR SITE-SPECIFIC FACTORS IN FOLLOW-UP TESTING (SEE THE ONLINE SUPPLEMENT) AND THEREFORE APPEARED TO REFLECT COVERT NEUROCOGNITIVE FEATURES.

PROSPECTIVE TRAJECTORIES OF MENTAL HEALTH AMONG THE FOUR CLUSTERS

We next assessed trauma-related outcomes across the clusters in a combined sample of 125 individuals from the clusters that replicated across both cohorts (see Figure S3 in the online supplement). The clusters showed different multivariate symptom profiles posttrauma (F = 2.25, DF = 2, 948, P = 0.013) (Figure 3A). Although assessment time point was included as a factor in the model, there was no interaction of cluster by time point on the symptom profile (P = 0.82). Follow-up tests were then performed for each symptom type separately. FIRST, THERE WAS AN EFFECT OF CLUSTER ON THE LONGITUDINAL MODEL OF PTSD SYMPTOMS (Wald χ² = 6.47, P = 0.039), WITH THE HIGHEST SYMPTOMS IN THE REACTIVE/DISINHIBITED CLUSTER (FIGURE 3B). THE EFFECT OF CLUSTER WAS REDUCED WHEN PRETRAUMA PTSD SYMPTOMS WERE ADDED AS A PREDICTOR IN THE MODEL (CLUSTER EFFECT: χ² = 5.10, P = 0.078; PRETRAUMA PTSD SEVERITY EFFECT: χ² = 20.19, P < 0.001).

SECOND, THERE WAS AN EFFECT OF CLUSTER ON THE LONGITUDINAL MODEL OF ANXIETY SYMPTOMS (Wald χ² = 6.23, P = 0.044) THAT WAS HIGHER IN THE REACTIVE/DISINHIBITED CLUSTER (FIGURE 3E). THIS EFFECT HELD AFTER INCLUDING PRETRAUMA ANXIETY SYMPTOMS AS A PREDICTOR IN THE MODEL (CLUSTER EFFECT: χ² = 6.07, P = 0.048; PRETRAUMA ANXIETY SEVERITY EFFECT: χ² = 72.37, P < 0.001), SUGGESTING THAT CLUSTER INFORMATION PROVIDED UNIQUE PREDICTIVE VALUE ABOVE BASELINE SYMPTOMS.

* Panels A and B show the region-of-interest covariance matrices revealing linear associations between z-scored contrast estimates extracted from the nine regions of interest across three tasks: threat, inhibition (inhib), and reward. For threat reactiveness, participants passively viewed fearful and neutral face stimuli. For threat reactiveness, fMRI activation was extracted from the amygdala (amyg), dorsal anterior cingulate cortex (dACC), insula, and subgenual anterior cingulate cortex (sgACC) for the contrast of fearful > neutral faces. For reward reactiveness, activation was extracted from the amygdala, nucleus accumbens (NAcc), and orbitofrontal cortex (OFC) for the contrast of gain > loss trials. For response inhibition, activation was extracted from the hippocampus (hipp) and ventromedial prefrontal cortex (vmPFC) for the contrast of no-go > go trials. Matrices are ordered hierarchically, such that regions that are more strongly associated with one another are adjacent. Significant associations are indicated on a red and blue color scale, thresholded at a p value < 0.05, uncorrected. Panels C and D show the dendrograms illustrating the final cluster solution with four clusters in the discovery cohort and three clusters in the replication cohort. Panels E and F show cluster differences (mean and standard deviation) for standardized contrast estimates extracted from the regions of interest across the threat (fearful > neutral faces), inhibition (no-go > go), and reward (gain > loss) contrasts. Panels G and H show individual subjects plotted along summary dimensions that reflect variance associated with primarily threat (principal component [PC] 1) and primarily reward (PC2); color reflects cluster assignment. Principal components were not used in the clustering analysis but are used in the graphs to illustrate graphically the cluster features and are described in further detail in Table S1 in the online supplement. Three-dimensional animated plots showing the inhibition dimension (PC3) are presented in Figure S5 in the online supplement.
FIGURE 3. Future patterns of mental health and fear learning in the four cluster groups among trauma survivors in the Advancing Understanding of Recovery After Trauma study.

A) PTSD

B) PTSD Symptoms

C) Impulsivity Symptoms

D) Depression Symptoms

E) Anxiety Symptoms

F) Dissociation Symptoms

G) Fear-Potentiated Startle (CS-NA) - CS+

H) Fear-Potentiated Startle (CS-NA) - CS–
Cluster did not predict depressive (p=0.19) or dissociative symptoms (p=0.86) or impulsivity (p=0.96) (Figure 3C,D,F).

Cluster-Based Compared With Dimensional Models for Predicting Longitudinal Posttrauma Outcomes

We tested the utility of the discrete clusters against a dimensional model of the fMRI data for predicting longitudinal trajectories of stress-related symptoms. Dimensional fMRI predictors were continuous covariates reflecting threat reactivity, reward reactivity, and inhibition. Models with these covariates as predictors of later posttrauma symptoms showed negligible improvement in the model fit over the cluster-based models (see Table S3 in the online supplement). The individual fMRI dimensions were not linearly associated with any posttrauma outcome, with the exception of a negative association between the inhibition-related fMRI dimension and later dissociative symptoms (p=0.044). In models that included both clusters and dimensions competing for the variance in posttrauma outcomes, cluster assignment still predicted subsequent PTSD symptoms (p=0.012), whereas the three fMRI dimensions did not (all p values >0.05). For anxiety, neither cluster (p=0.080) nor dimensions (all p values >0.05) were significant in the head-to-head model. In summary, the dimensional model did not provide better predictive value than the cluster-based models.

Convergent Validity With Fear-Learning Phenotypes

On the day of the MRI scan, participants also completed a fear-potentiated startle paradigm that included fear conditioning and extinction. During fear conditioning, effects of CS (F=12.73, df=1, 468, p=0.0004) and the CS-by-block interaction (F=6.06, df=1, 468, p=0.003) suggested that fear conditioning occurred and that discrimination between the CS+ and CS− developed across acquisition. There was a significant cluster-by-block interaction (F=4.13, df=4, 468, p=0.003), such that the low reward/high threat cluster showed the highest fear-potentiated startle responses to both the CS+ and CS− at the beginning of fear conditioning, but this cluster was comparable to the other cluster groups by the end of the task (Figure 3G). There was no cluster-by-CS interaction (p=0.82). During fear extinction, startle responses to the CS+ and CS− showed the expected decline over time (block effect: F=28.79, df=3, 623, p=0.02 × 10−14), indicating the presence of extinction learning, but there was no interaction of CS by block (p=0.64), indicating no difference in the extinction pattern for CS+ compared with CS−. This was consistent with findings from previous studies of chronic PTSD using this startle paradigm (39). There was again an interaction of cluster by block, indicating different rates of extinction in the different cluster groups (F=2.35, df=6, 623, p=0.03). The low reward/high threat cluster showed the highest fear-potentiated startle responses to both the CS+ and CS− at the beginning of extinction, decreasing to become comparable to the other cluster groups by the end of the task (Figure 3H).

Voxel-Wise Whole-Brain Comparison of Cluster Groups

Finally, to identify brain regions outside the primary regions of interest included in the clustering, we conducted whole-brain analysis in the combined sample of 125 participants (Figure 4; see also Table S4 in the online supplement), comparing the three replicated cluster groups within the threat, reward, and inhibition fMRI tasks. The reactive/disinhibited cluster showed greater activation than the other two cluster groups in a mesopontine cluster overlapping with the median raphe nucleus and ventral tegmental area, as well as the hypothalamus, dACC, and insula in response to threat cues. In contrast, the low reward/high threat cluster showed greater activation in the left and right amygdala, hippocampus, and insula in response to threat cues. The reactive/disinhibited cluster also showed greater reactivity than the other two cluster groups in the amygdala, hippocampus, and rostral anterior cingulate cortex in response to reward. The inhibited cluster showed no region of greater activation compared with the other two cluster groups.

DISCUSSION

In a well-characterized cohort followed longitudinally in the aftermath of trauma, we identified participant clusters in a manner that was agnostic to standard diagnostic categories for posttrauma outcomes using fMRI across several neurocognitive dimensions of interest, including threat, reward, and inhibition. In the discovery cohort of motor vehicle accident survivors, four clusters were

---

4 The clusters showed differences in a multivariate profile of outcomes from 2 weeks to 6 months posttrauma (F=2.26, df=3, 1206, p=0.008). Panel A shows mental health profiles for each cluster, revealing standardized values for each outcome rescaled to a 0–1 scale. Because there was no interaction with time point, cluster profiles are collapsed across the 2-week, 8-week, 3-month, and 6-month study visits. Panel B shows how the clusters differed in the longitudinal model of posttraumatic stress disorder (PTSD) symptom severity, with the highest symptoms in the reactive/disinhibited cluster. The mean PTSD Checklist total score for each cluster over the assessment time points is shown, and gray shading shows 95% confidence intervals. The clusters showed no differences for univariate longitudinal models of depression symptoms (panel C), dissociative symptoms (panel D), or impulsivity (panel F). Panel E shows how the clusters differed in the longitudinal model of anxiety symptom severity, with highest symptoms among individuals in the inhibited cluster. Panel G shows the fear-potentiated startle response during the fear-conditioning paradigm conducted 2 weeks posttrauma. Fear conditioning to the conditioned stimulus (CS)+ danger cue and the CS− safety cue are shown over the course of three experimental blocks, with an overlay showing the main effect of CS type. The low reward/high threat cluster group showed significantly elevated fear-potentiated startle in response to both CS+ and CS− at the beginning of the task compared with the other clusters. Panel H shows the fear extinction task results, revealing fear-potentiated startle to the CS+ over the early and late trials of the task, with the overlay illustrating the main effect of block. Responses to the CS− showed a significant decrease over time, consistent with extinction, but there were no differences between the four clusters. mo.=months; NA=noise alone.
observed, and three were replicated in a cohort with a wider variety of index traumas. Given the timing at 2 weeks posttrauma, the clustering likely reflects a combination of traits that predate the trauma, as well as acute stress responses in the wake of the traumatic event. The clusters were not related to the demographic characteristics and background variables of the participants (e.g., gender, childhood trauma) but could still plausibly reflect pretrauma factors, such as genetics, family history, or temperament (40). Our findings confirmed the hypothesis that clusters may be associated with different posttrauma outcomes: different longitudinal patterns emerged over the first 6 months posttrauma, with the reactive/disinhibited cluster associated with subsequent heightened symptoms of PTSD/hypervigilance and anxiety. In addition, these findings represent an important step toward defining a neuroimaging-based longitudinal prediction model for stress-related resilience and vulnerability.

Our results suggest that an unsupervised forward-inference model is tractable for modeling heterogeneity in stress-related psychiatric outcomes, despite the lack of constraints on the model. We used very simple and transparent tools for clustering, uninformed by psychiatric symptoms or diagnoses, and found strong evidence for overlap (65%) in the cluster solutions arising from two fully independent hierarchical clustering solutions in different subcohorts. This level of overlap is consistent with cluster replication levels seen in larger brain-based clustering efforts, such as in the Adolescent Brain Cognitive Development study (41), and suggests that multivariate task-based fMRI data contain consistent information about individual differences. In fact, a major value of this study is its demonstration of task-based fMRI as a useful tool for mapping psychiatric heterogeneity. Task-based fMRI may not be needed to identify unitary biomarkers, such as for a diagnosis (e.g., PTSD) or a symptom (e.g., hyperarousal), where resting-state MRI is
likely preferable for its rich information on neural circuit function and low barriers to translation. However, for the purposes of resolving heterogeneity, task fMRI shows clear strengths. For example, the signal from the amygdala was clearly uncorrelated in threat tasks compared with reward tasks (Figure 2A,B), and this variability was important in resolving the clusters. Such information would not have been apparent from analyses of intrinsic network activity.

Interpretation of the Biotypes
The reactive/disinhibited cluster was the most interesting candidate as a risk-related biotype. Individuals in this cluster showed threat hyperreactivity, particularly in the insula and dACC, accompanied by high NAcc reward reactivity, as well as higher subsequent PTSD symptoms. This was partly contrary to previous findings showing that lower NAcc reward response predicted subsequent PTSD (16). Threat and reward reactivity have rarely been assessed concurrently in previous studies of trauma and related outcomes. However, preclinical findings indicate that interacting pathways regulate both threat and reward reactivity. For example, stress-related hyperactivity of the basolateral amygdala can directly influence NAcc function via direct efferent projections (42) and change reward-seeking behaviors (43). Participants in the reactive/disinhibited cluster also showed greater activation in the reticular nuclei (median raphe, ventral tegmental area) during the threat task. The reticular formation stimulates wakefulness and arousal (44,45). The role of these nuclei in dopamine and serotonin synthesis may point toward tailored intervention opportunities for the future; early studies are exploring dopaminergic modulation to treat PTSD (46).

The fMRI features observed in the low reward/high threat cluster included moderate responsivity to threat dominated by sgACC activation, along with markedly low reactivity to reward. Reduced reward responsivity in the NAcc, amygdala, and OFC is characteristic of major depressive disorder (47), as is sgACC hyperreactivity to sadness-inducing stimuli. The heightened fear-potentiated startle shown by this group during early fear acquisition and extinction is consistent with patterns previously observed in comorbid PTSD and major depressive disorder (27), which is more common posttrauma than each disorder alone (48). This cluster group also showed greater threat reactivity in the amygdala, insula, and hippocampus in whole-brain analyses compared with the other cluster groups. Together, the findings suggest the possibility that PTSD-related symptom groups may be divided into a low reward/high threat group driven more by cortical function, and a reactive/disinhibited group, driven more by brainstem nuclei.

Participants in the inhibited cluster appeared to be most consistent with active coping, with low threat reactivity accompanied by relatively high vmPFC and hippocampus engagement during inhibition. Individual features of this pattern have previously been associated with resilience. For example, lower amygdala threat reactivity predicts lower future PTSD symptoms (12,13). Similarly, greater vmPFC and hippocampal activation during inhibition has been associated with resilience (22,26). Our findings indicate that some individuals show the combined profile in the acute posttrauma period. However, this was the smallest cluster, and it is possible that inhibition as an adaptation to the stress of the index trauma had not yet fully emerged by 2 weeks posttrauma. As data collection continues, the AURORA study will include additional neuroimaging 6 months posttrauma, allowing a window into the further development of these profiles.

Participants in the high reward cluster showed high reward reactivity, low threat reactivity, and low inhibition, a pattern suggesting preserved positive affect in the context of low top-down regulation. However, this pattern may only be observed when the emotional impact of trauma is relatively low, and this cluster was not observed in the replication cohort.

Limitations
To reduce participant burden, symptom assessments were abbreviated and based on self-report. This limited our ability to directly compare the outcomes to gold-standard assessments of psychiatric disorders. However, future work could apply these biotypes to archival data with interview-based assessments to extend our findings. Additionally, with sample sizes of 69 participants in the discovery cohort and 77 in the replication cohort, the study may be underpowered. However, we are encouraged that three clusters were replicated, indicating generalizability even across different types of trauma.

CONCLUSIONS
Neuroimaging phenotypes emerging in the early aftermath of trauma are associated with risk of or resilience to trauma-related psychopathology. Contrary to our initial predictions that heightened threat and blunted reward reactivity may reflect stress vulnerability, a cluster showing heightened reactivity to both threat and reward was associated with the subsequent maintenance of the highest levels of PTSD symptoms. Heightened reward reactivity in the early aftermath of a major stressor may be an undetected risk mechanism for the development of stress-related disorders. The biotypes identified here, with further development to assess normative values and precision, may provide important information about targeted interventions to address different forms of future stress-related psychopathology.

AUTHOR AND ARTICLE INFORMATION
Department of Psychiatry and Behavioral Sciences, Emory University, Atlanta (Stevens, van Rooij, Ely, Roecner, Vincent); Division of Depression and Anxiety, McLean Hospital, Belmont, Mass. (Harnett, Lebois, Ressler); Department of Psychiatry, Harvard Medical School, Boston (Harnett, Lebois, Pizzagalli, Ressler); Departments of Emergency Medicine and Health Services, Policy, and Practice, Alpert Medical School of Brown University, Rhode Island Hospital, and the Miriam Hospital, Providence, R.I. (Beaudoin); Department of Anesthesiology, Institute of Trauma Recovery, University of North Carolina, Chapel Hill (An,
Linnstaedt, McLean); Department of Biostatistics, Gillings School of Global Public Health, University of North Carolina, Chapel Hill (Zeng); Departments of Psychiatry and Neurology, University of California, San Francisco (Neylan); Department of Biomedical Informatics, Emory University School of Medicine, Atlanta (Clifford); Institute for Technology in Psychiatry (Germaine) and Department of Psychiatry (Rauch), McLean Hospital, Belmont, Mass.; Department of Emergency Medicine, Henry Ford Health System, Detroit (Lewandowski); Department of Emergency Medicine, Vanderbilt University Medical Center, Nashville, Tenn. (Storrow); Department of Emergency Medicine, University of Florida College of Medicine, Jacksonville (Hendry, Sheikh); Department of Emergency Medicine, Indiana University School of Medicine, Indianapolis (Musey); Department of Emergency Medicine, University of Massachusetts Medical School, Worcester (Haran); Department of Emergency Medicine, Cooper Medical School of Rowan University, Camden, N.J. (Jones); Department of Emergency Medicine, College of Medicine and College of Nursing, University of Cincinnati, Cincinnati (Punches); Department of Emergency Medicine and Center for Addiction Research, University of Cincinnati College of Medicine, Cincinnati (Lyons); Departments of Emergency Medicine and Surgery, Division of Acute Care Surgery, University of Alabama School of Medicine, Birmingham (Kurz); Center for Injury Science, University of Alabama, Birmingham (Kurz); Department of Emergency Medicine, Boston Medical Center, Boston (McGrath); Departments of Surgery (Pascual) and Neurosurgery (Pascual), Perelman School of Medicine, University of Pennsylvania, Philadelphia; Department of Emergency Medicine, Einstein Health Care Network, Philadelphia (Datner); Department of Emergency Medicine, Jefferson University Hospitals, Philadelphia (Chang); Department of Emergency Medicine, Wayne State University, Detroit (Pearson); Department of Emergency Medicine, Massachusetts General Hospital, Boston (Peak); Department of Emergency Medicine, Saint Joseph Mercy Hospital, Ann Arbor, Mich. (Domeier); Department of Emergency Medicine, Wayne State University School of Medicine, Detroit (O’Neill); Department of Emergency Medicine, University of Massachusetts Medical School–Baystate, Springfield (Rathley); Department of Emergency Medicine, Beth Israel Deaconess Medical Center, Boston (Sanchez); Department of Emergency Medicine, Harvard Medical School, Boston (Sanchez); Department of Psychiatry, Yale School of Medicine, and U.S. Department of Veterans Affairs National Center for Posttraumatic Stress Disorder, Veterans Affairs Connecticut Healthcare System, New Haven, Conn. (Pietrzak); Department of Psychology, Yale University, New Haven, Conn. (Joorman); Department of Psychological and Brain Sciences, Washington University, St. Louis (Barch); Department of Biosciences and Neuroscience and Institute for Behavioral Medicine Research, Ohio State University Wexner Medical Center, Columbus (Sheridan); Department of Psychiatry, University of Pittsburgh, Pittsburgh (Luna); Departments of Anesthesiology and Internal Medicine–Rheumatology, University of Michigan Medical School, Ann Arbor (Harte); the Kolling Institute of Medical Research, Northern Clinical School, University of Sydney, and Faculty of Medicine and Health, University of Sydney, Sydney, Australia (Elliott); Physical Therapy and Human Movement Sciences, Feinberg School of Medicine, Northwestern University, Chicago (Elliott); Department of Psychology, Temple University, Philadelphia (Murty); Department of Psychiatry and Behavioral Neurosciences, Wayne State University, Detroit (Jovanovich); Department of Psychological Sciences, University of Missouri, St. Louis (Bruce); Department of Emergency Medicine, Washington University School of Medicine, St. Louis (House); Department of Health Care Policy, Harvard Medical School, Boston (Kessler); Department of Epidemiology, Harvard T.H. Chan School of Public Health, Boston (Koene); Department of Emergency Medicine, University of North Carolina, Chapel Hill (McLean).

Send correspondence to Dr. Stevens (jennifer.stevens@emory.edu).

Dr. Jones has received funding from AstraZeneca, Hologic, Janssen, and Roche Diagnostics. Dr. Peak has received consulting fees from Akili Interactive Laboratories, BlackThorn Therapeutics, Boehringer Ingelheim, Posit Science, and Takeda Pharmaceuticals, and he has received an honorarium from Alkermes. Dr. Kessler has received funding for epidemiological studies from Sanofi-Aventis, and he has served as a consultant for DataStat, Sage Pharmaceuticals, and Takeda. Dr. Ressler has received consulting fees or sponsored research support from Alkermes, BrainsWay, and Genomind, and he serves on scientific advisory boards for Janssen, Takeda, and Verily. The other authors report no financial relationships with commercial interests.

Presented in part at the annual meeting of the American College of Neuropsychopharmacology, December 8–11, 2019, Orlando, Fl.

Dr. Sheikh is supported by the Florida Medical Malpractice Joint Underwriter’s Association’s Dr. Alvin E. Smith Safety of Healthcare Services, the NIH/National Institute on Aging Jacksonville Aging Studies Center (grant R33AG05654), and the Florida Blue Foundation. Dr. Elliott is supported by NIH (grants R01HD079076 and R03HD094577), the Eunice Kennedy Shriver National Institute of Child Health and Human Development, and the National Center for Medical Rehabilitation Research.

Supported by the Mayday Fund, NIMH (grants U01 MH110925, K00 MH119603, and K01 MH118467), the One Mind Foundation, and the U.S. Army Medical Research and Material Command. Data and/or research tools used in the preparation of this study were obtained from the NIMH Data Archive. The NIMH Data Archive is a collaborative informatics system created by NIH to provide a national resource to support and accelerate research in mental health (data set identifier, 10.15154/1521266).

The authors thank the trauma survivors who participated in the AURORA study.

This article reflects the views of the authors and may not reflect the opinions or views of NIH or of the submitters of original data to the NIMH Data Archive.

Received October 23, 2020; revisions received March 9 and April 21, 2021; accepted May 27, 2021; published online October 14, 2021.

REFERENCES


27. Jovanovic T, Norholm SD, Blanding NQ, et al: Impaired fear inhibition is a biomarker of PTSD but not depression. Depress Anxiety 2010; 27:244–251
42. McDonald AJ: Topographical organization of amygdaloid projections to the caudatoputamen, nucleus accumbens, and related striatal-like areas of the rat brain. Neuroscience 1991; 44:15–33