Cortisol response to an experimental stress paradigm prospectively predicts long-term distress and resilience trajectories in response to active police service

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\textbf{A B S T R A C T}

Heterogeneity in glucocorticoid response to experimental stress conditions has shown to differentiate individuals with healthy from maladaptive real-life stress responses in a number of distinct domains. However, it is not known if this heterogeneity influences the risk for developing stress related disorders or if it is a biological consequence of the stress response itself. Determining if glucocorticoid response to stress induction prospectively predicts psychological vulnerability to significant real life stressors can adjudicate this issue. To test this relationship, salivary cortisol as well as catecholamine responses to a laboratory stressor during academy training were examined as predictors of empirically identified distress trajectories through the subsequent 4 years of active duty among urban police officers routinely exposed to potentially traumatic events and routine life stressors (N = 234). During training, officers were exposed to a video vignette of police officers exposed to real-life trauma. Changes in salivary 3-methoxy-4-hydroxyphenylglycol (MHPG) and cortisol in response to this video challenge were examined as predictors of trajectory membership while controlling for age, gender, and baseline neuroendocrine levels. Of officers who followed trajectories of resilience and recovery over 4 years mounted significant increases in cortisol in response to the experimental stressor, while those following a trajectory of chronic increasing distress had no significant cortisol change in response to the challenge. MHPG responses were not associated with distress trajectories. Cortisol response prospectively differentiated trajectories of distress response suggesting that a blunted cortisol response to a laboratory stressor is a risk factor for later vulnerability to distress following significant life stressors.

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1. Objectives of the study and background

Biological response to a stressor is multifaceted, including processes that rapidly increase preparedness for harm as well as slower processes for regaining homeostasis to prevent adverse biological consequences of a prolonged physiological stress response (Conrad, 2011). Rapid responses to a stressor occur in the sympathetic adrenal medullary pathway (SAM), which activate physiological responses such as increased heart rate and blood pressure in service of defensive behavior such as fight or flight (Schedlowski et al., 1993). However, this level of arousal can have lasting physiological consequences if unchecked. Stressors are also associated with increases in cortisol 15–20 min post-stressor. Stressor-related cortisol responses have been shown to aid in braking sympathetic stress responses once the perceived threat has
been removed (Munck et al., 1984), and may be important in regaining physiological homeostasis following a significant stressor (Yehuda and LeDoux, 2007). These neurobiological pathways are thought to be highly conserved across species as part of the threat response survival circuitry (LeDoux, 2012). As such, identifying their relationship to heterogeneous stress-response phenotypes may be informative as to how threat circuitry functioning leads to clinically relevant long-term outcomes such as resilience, recovery, and chronic stress.

Heterogeneity in cortisol responses to stressful experimental conditions has been identified in a diverse set of clinical stress reactions. It has been proposed that cortisol plays a regulatory role that may be of broad relevance for understanding the relationship between stressor exposure and pathological stress responses. Specifically, a blunted cortisol response, in which individuals do not increase their secretion of cortisol in response to an experimental stressor, has been observed in a number of clinical contexts. Very low income women who demonstrated a blunted cortisol response to a stressor were more likely to experience elevated depression symptomatology (Burke et al., 2005). Children who received an initial diagnosis of attention deficit hyperactive disorder (ADHD) who demonstrated this effect were more likely to maintain the diagnosis one year later (King et al., 1998). Alcohol and poly-drug substance abusing men were more likely to demonstrate a blunted cortisol response to an experimental stressor compared to healthy controls (Lovatti et al., 2000), and abstaining alcohol and poly-substance abusers who demonstrated this abnormality have been shown to be more likely to relapse (Junghanns et al., 2003). Individuals with schizophrenia, a neurobiological disorder in which symptoms are exacerbated by stress, have also demonstrated a blunted cortisol response compared to healthy controls (Brenner et al., 2009; Jansen et al., 1998; Jansen et al., 2000). This effect has also been observed in other contexts in which stress can exacerbate medical symptomatology including tinnitus (Hébert and Lupien, 2007), fatigue among breast cancer survivors (Bower et al., 2005), and flare-ups of inflammatory disorders such as allergic asthma and atopic dermatitis (Buske-Kirschbaum et al., 1997, 2003). In preclinical settings, rodents that vary genetically in the corticosterone (the rodent analogue to cortisol) response varied significantly in prolonged behavior stress responses following exposure to a predator stress condition, with blunted corticosterone responses associated with prolonged stress responses (Cohen et al., 2006). Together, we are presented with a transdiagnostic picture in which the inability to mount an effective cortisol response to an experimental stressor is associated with poorer real-life stress management. In most of these studies, heart rate variability was also examined but not found to be predictive of the outcome of interest. This suggests that the ability to biologically regain homeostasis may be more impactful on longer-term stress adaptation than the initial SAM stress response itself.

While these findings support the hypothesis that the management of stress may have strong biological underpinnings rooted in HPA-axis activity, many of the studies to date have focused on populations who have already expressed pathological stress responses. A number of studies have examined cortisol response immediately following trauma exposure as a prospective predictor of subsequent PTSD development but have not presented with consistent evidence of a relationship between cortisol soon after the trauma and later PTSD (Bonne et al., 2003; Heinrichs et al., 2005; McFarlane et al., 2011; Shalev et al., 2008).

The cortisol awakening response has been examined prospectively in military populations but failed to predict later PTSD development (van Zuiden et al., 2011). In an earlier report from this cohort, the cortisol awakening response was found to prospectively predict both peri-traumatic reactions as well as acute stress disorder symptom severity 12-months into active police service (Inslicht et al., 2011). As such, controversy remains about the assertion that cortisol response is a dimension underlying diverse clinical outcomes despite theory to support the assertion that the cortisol response to significant life stressors will ultimately impact long term patterns of adaptation (Yehuda and LeDoux, 2007). Determining if cortisol responses to an experimental stressor prospectively predict responses to real-life stressors may be informative as to the relationship between underlying threat circuitry functioning and long term psychological responses. In the current study we examine this relationship in urban police officers, as they represent a generally healthy population at the time of academy training who will encounter repeated duty related stressors in the course of their careers.

Police officers have been shown to be susceptible to adverse consequences of both routine work stressors and exposure to potentially traumatic events (PTEs) including the development of posttraumatic stress disorder (PTSD) (Marmar et al., 2006), depression (Wang et al., 2010), pathological sleep disturbances (Neylan et al., 2002), anxiety, somatization, alcohol abuse, and aggressive behavior (Gershon et al., 2002). Despite the significant psychological risks associated with police work, evidence indicates that there is significant and meaningful heterogeneity in distress responses among officers exposed to similar conditions. Empirical studies have demonstrated that police officers, along with others who are similarly exposed such as military personnel and first responders, follow a limited set of trajectories of symptom or distress responses including Resilience (little or no long term emotional distress), Recovery (significant distress followed by remission), Chronic Distress (high levels of distress or pathology that do not abate and may increase over time) and, when the stressor is anticipated Anticipatory Distress characterized by elevated pre-event stress that declines following the event is also commonly observed (Bonanno, 2004; Bonanno et al., 2012a; Bonanno et al., 2012b; Galatzer-Levy and Bonanno, 2012; Galatzer-Levy et al., 2013, 2012). These patterns have been found in response to events as varied as terrorist attacks (Bonanno et al., 2005), disease epidemics (Bonanno et al., 2008), traumatic injury (deRoon-Cassini et al., 2010), deployment to warzones (Dickstein et al., 2010), and traumatic loss (Bonanno, 2004), and have been observed in response to life stressors including breast cancer diagnosis (Lam et al., 2010), job loss (Galatzer-Levy et al., 2010), and even child birth (Galatzer-Levy et al., 2011b). When the population is repeatedly exposed to significant stressors, the chronic stress trajectory shows progressive growth in symptomatology (Reactive-Worsening) rather than a chronic stable elevated trajectory (Galatzer-Levy et al., 2011a).

Our previous work has demonstrated that heterogeneous patterns of stress response among police officers do not demonstrate a linear relationship with either routine work stress or PTE exposure (Galatzer-Levy et al., 2013). Specifically, the previous work found that continuous counts of routine work stress and PTEs were non-significant in association with the trajectories of general distress under study in the current work. This does not indicate that such events are not impactful but rather that stress responses among officers, a routinely exposed population, are not characterized by a positive linear increase in association with the number of routine work stressors or PTEs. As such, stressor exposure may be a necessary condition for the emergence of adaptive or maladaptive responses rather than a driving force behind the pattern of response. Neurobiological responses are a strong candidate as a factor impacting individual differences as there is consistent evidence for heterogeneity in the neurobiological (Cohen et al., 2006) and behavioral response (Galatzer-Levy et al., 2012) to identical stress conditions.
Using a prospective longitudinal design in which police officers were followed from academy training through 4 years of active duty service, we tested the hypothesis that a blunted cortisol response to an ecologically valid stress induction during academy training would prospectively predict membership in the Reactive-Worsening distress trajectory. Secondly, we examined if catecholamine responses to the same experimental condition predicted membership in the Reactive-Worsening trajectory to test the hypothesis that the initial adrenergic response would predict outcomes.

2. Materials and methods

2.1. Participants and procedures

Officer recruits from 4 urban police departments (New York City, San Francisco, Oakland, San Jose) were recruited into a large prospective study of bio-psycho-social predictors of stress responses to critical incident exposure. Trainees who had previously served in the military, law enforcement, or emergency services were excluded. Procedures were approved by the University of California, San Francisco Institutional Review Board and a Federal Certificate of Confidentiality was obtained. Participants were evaluated on general distress and cortisol and catecholamine reactivity at baseline (academy training), and general distress, PTSD symptoms, and depression symptoms at 12, 24, 36, and 48 months after commencement of active duty police service. Prior to the initial assessment, study procedures were described in detail and written informed consent was obtained (for a full description of recruitment procedures see (McCaslin, 2008)).

This study utilized a subsample of officers from the parent study who had data on general emotional distress available on at least 3 of 5 time points, including during training and across 48 months of active duty service (N = 234) drawn from a larger cohort of police officers followed longitudinally from academy training (N = 400). Of the 234 participants, n = 106 reported exposure to a PTE at least once by 6 months, n = 208 by 12 months, n = 221 by 24 months, n = 224 by 36 months, and n = 217 by 48 months. We previously identified trajectories of general distress using Latent Class Growth Analysis (LCGA). The previously identified trajectories included Resilience, characterized by healthy adaptation from baseline to 4 years into active duty (76.7%), Reactive-Worsening (15.8%), characterized by consistent growth in distress across all 4 years, Recovering (4.6%), demonstrating sharp growth in distress from baseline though 2 years of service that remitted at roughly the same rate from 2 years to 4 years of service, and Anticipatory Distress (2.9%), characterized by high distress at baseline prior to initiation of active duty that diminishes in distress over time (Fig. 1). We previously demonstrated that individuals in these trajectories were significantly different with regard to PTSD and depression symptomatology (Galatzer-Levy et al., 2013). For a full description of modeling and sampling procedures see (Galatzer-Levy et al., 2013).

2.2. Measures

2.2.1. Critical Incident History Questionnaire (CIHQ)

The Critical Incident History Questionnaire (CIHQ) is a 39-item self-report measure designed to assess exposure to PTSD criterion A events typically encountered by police officers in the line of duty (for a full description see Weiss et al., 2002). The CIHQ was administered at 12, 24, 36, and 48 months either in-person or through the mail. Mean life threat through 48 months was 11.58 incidents (SD = 13.98). In the current analysis we utilized a dichotomous variable indicating exposure to a life-threatening event in the first 12 months as a covariate, as this may also predict stress response patterns.

2.2.2. Hopkins Symptom Checklist 90-R Global Severity Index (SCL-90-R GSI)

Self-reported distress from psychological symptoms was measured using a combination of 29 items from multiple scales of the Symptom Checklist-90-R that make up the Global Severity Index (GSI) (Derogatis and Melisaratos, 1983). The SCL-90-R was administered at baseline, 12, 24, 36, and 48 months either in-person or through the mail.

2.2.3. Salivary cortisol and salivary 3-methoxy-4-hydroxyphenylglycol (MHPG)

The details of the critical incident video have been described in an earlier report from this study (Apfel et al., 2011). Briefly, participants observed an innocuous travelogue video for 10 min, followed by a 20-min critical incident depiction video, then a 20-min travelogue video again during the response period. The critical incident video contained real-life footage of 14 incidents involving police officers that were edited into one continuous 20-min segment depicting police-related scenes including an officer being hit by a car, an officer being mauled by a dog, and an officer being killed while attempting to defuse a bomb. Saliva was collected at three time points; T1 (Baseline) was a baseline measure obtained immediately following the first travelogue video and just prior to the viewing the video stressor, T2 (Initial Cortisol Response) was a measure obtained immediately after the conclusion of the 20-min video stressor, and T3 (Final Cortisol Response) which was obtained at the conclusion of the 20-min response period (see Fig. 2). There was no food intake, exercise, or smoking for a minimum of 2 h before the stress challenge, and use of prescription and over-the-counter medication was systematically assessed (for a complete description of sample collection, processing, and analysis of samples see Otte et al. (2005)). Because of significant evidence that the cortisol response to a laboratory stressor peaks between 20 and 40 min after stress onset (Dickerson and Kemeny, 2004), as well as evidence for greatest MHPG change in the same timeframe (Otte et al., 2005), we computed change scores as T2 cortisol and MHPG measurements minus T3 measurements. MHPG and cortisol levels at T1 were used as covariates to capture variance associated with individual differences in MHPG and cortisol that are not associated with the video challenge stress response. We measured MHPG in saliva because it is less invasive than plasma measures and correlates with plasma MHPG, which increases in response to acute stressors and is unaffected by beta-blockade, suggesting that it is a measure of central noradrenergic activity (Hamer et al., 2007). Salivary MHPG also correlates strongly with MHPG in cerebrospinal fluid, providing further support for its use as a proxy for central
noradrenergic metabolism (Reuster et al., 2002). All values were log-transformed to address significant skewness and kurtosis in both. Transformations normalized the distributions of all measures.

2.3. Data analytic plan

Our data analysis plan consisted of two steps. Step 1 aimed to determine if changes in cortisol and catecholamines in response to the video stressor predicted trajectories of general distress through 48-months of active duty. To test this both catecholamine and cortisol change scores across the response period (T3–T2) were used as predictors of previously identified trajectories of general distress (Galatzer-Levy et al., 2013) using a multinomial logistic regression while controlling for baseline levels of both hormones as well as other possible confounding variables. Change scores only provide a gross estimate of the magnitude of change. However, they do not provide evidence of significant levels of change. To determine if significant levels of change occurred during the response period, those change scores that demonstrate significance as predictors of trajectories were examined using a repeated measures ANOVA with T2 and T3 scores entered as a within subjects factor and class membership entered as a between subjects factor. This analysis log transformed values of T2 and T3 cortisol were tested as a within subjects factor. Corrections were made for multiple comparisons using Least Squared Differences. Exposure at 12-months was examined as a dichotomous variable to determine if individuals exposed earlier in police service would fall into distress trajectories. Total critical incident exposure and work stress exposure across the 48-months has already been examined and found to be non-significant in association with the trajectories (Galatzer-Levy et al., 2013).

3. Results

In the previous trajectory analysis, we compared solutions with linear parameters only and linear and quadratic parameters to assess which parameters best fit the data. Ultimately a four-class solution with linear and quadratic parameters best fit the data based on a significant reduction the information criteria, entropy, the BLRT, and conformity with theory and parsimony (see Fig. 2 for the classes and their proportions). Though reductions in the information criteria continued to be observed through five classes with linear and quadratic weights (reduction in AIC = 43.34, BIC = 29.52, SSBIC = 42.15), the BLRT was significant at $p < .05$ for a four vs. three class solution but not a five vs. four class solution. This was true both when applying linear weights only ($p = .95$) and linear + quadratic weights ($p = .36$). Further, the addition of a fifth class served to split a small class into two parallel classes, and as such was a less parsimonious and less interpretable solution (Galatzer-Levy et al., 2013).

Next, we explored the data descriptively by generating means and standard deviations or frequencies for our variables of interest, including gender and exposure, age, baseline cortisol and MHPG levels, cortisol and MHPG levels at T2 and T3. Both log-transformed values of cortisol and MHPG that are used in analyses are presented as well as raw scores on these variables for comparison to other studies (See Table 1). Descriptive statistics indicated that there was an increase in cortisol among those in the Resilient, Recovery, and Anticipatory Distress classes and a decrease among those in the Reactive-Worsening class. Similar trends were observed for MHPG with the exception of the Anticipatory Distress class which showed a decrease.

To test if the observed change in cortisol and MHPG significantly differed by class, we regressed class membership on our covariates in a multinomial logistic regression using the Auxiliary option in MPlus 6.12 on the modeled classes. This analysis revealed no significant differences by age, gender, or trajectory class in the probability of exposure at 12-months. Further, no significant difference was observed between the classes on T1 cortisol or T1 MHPG levels or MHPG change from T2 to T3. However, individuals in the Reactive-Worsening class demonstrated significantly less change in cortisol response from T2 to T3 when the Resilient class was used as

\[ \text{Note: } *p < .05; **p < .001; \text{ Results reveal that those in the Resilient and the Recovering class demonstrated cortisol change across the response period while those in the Reactive-Worsening class did not.} \]
the references class \((Est/SE = -1.95; p < .05)\) and significantly less change compared to the Recovery class when the Reactive-Worsening class was set as the reference class \((Est/SE = 2.14; p < .05; \text{see Table 2 for complete results})\).

3.1. Post-hoc analyses

To determine if there was significant change in cortisol levels separately by class in response to the video stressor, a Repeated Measures Analysis of Variance (ANOVAs) was conducted to test for mean level change in cortisol from T2 to T3 with probable class membership as a fixed effect. In this analysis, log transformed cortisol scores at T2 and T3 were used as a within subjects factor. This analysis revealed that those in the Resilient and Recovery classes demonstrated significant increases in cortisol across these time points \([\text{Resilient: } F (1,106) = 11.59; p < .001; \text{Recovery } F(1, 8) = 6.03; p < .05]\) while those in the Reactive-Worsening distress class did not show a significant increase in cortisol response from T2 to T3 \([F(1,30) = 0.11; p = .74; \text{Fig. 2}]\).

4. Discussion

In the current investigation we tested the hypothesis that a blunted cortisol response would prospectively predict long-term stress response patterns. Secondly, we examined whether catecholamine responses during the same time frame would predict these patterns. We tested this hypothesis by examining if trajectories were predicted by change scores in cortisol and catecholamine levels from immediately following a video stressor to 20 min later \((\text{capturing the cortisol response as well as a sensitive period of catecholamine change})\). Because cortisol and MHPG levels can be affected by a number of individual characteristics, we included gender, age, and baseline cortisol and MHPG levels as covariates. Further, because patterns of stress response can potentially be explained by initial exposure rather than cortisol and MHPG, exposure by 12-months was also included as a covariate.

Compared to those who demonstrated Resilient and Recovering trajectories, those who demonstrated consistent growth in stress \((\text{Reactive-Worsening class})\) demonstrated significantly lower levels of cortisol change in response to the video challenge stressor prior to exposure to real life stressors. MHPG responses to the video challenge and levels of MHPG and cortisol just prior to the video challenge did not differentiate classes. Further, post-hoc analyses revealed that individuals in both the Recovering and Resilient classes demonstrated significant increases in cortisol in response to the video stress challenge while those in the Reactive-Worsening class did not demonstrate a significant change, indicating that they did not mount a cortisol response to the video challenge while the others did.

This study has several limitations. Stress hormone levels are variable for a number of reasons including individual differences in age, gender, medication, substance use, psychiatric comorbidity, and time of day that levels are recorded. We attempted to address

<table>
<thead>
<tr>
<th>Variables in model</th>
<th>Trajectory membership</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Resilient ((n = 178))</td>
</tr>
<tr>
<td></td>
<td>Mean (SD)/%</td>
</tr>
<tr>
<td>Gender (male)</td>
<td>91.5%</td>
</tr>
<tr>
<td>Age</td>
<td>27.34 (4.92)</td>
</tr>
<tr>
<td>Exposure</td>
<td>80.9%</td>
</tr>
<tr>
<td>Baseline Cortisol</td>
<td>6.17 (0.51)</td>
</tr>
<tr>
<td>Time 2 Cortisol</td>
<td>6.01 (0.52)</td>
</tr>
<tr>
<td>Time 3 Cortisol</td>
<td>6.11 (0.60)</td>
</tr>
<tr>
<td>Baseline MHPG</td>
<td>1.63 (0.43)</td>
</tr>
<tr>
<td>Time 2 MHPG</td>
<td>1.63 (0.43)</td>
</tr>
<tr>
<td>Time 3 MHPG</td>
<td>1.66 (0.42)</td>
</tr>
<tr>
<td>Baseline Cortisol Raw Score</td>
<td>5.60 (2.90)</td>
</tr>
<tr>
<td>Time 2 Cortisol Raw Score</td>
<td>5.78 (3.31)</td>
</tr>
<tr>
<td>Time 3 Cortisol Raw Score</td>
<td>5.77 (3.05)</td>
</tr>
<tr>
<td>Baseline MHPG Raw Score</td>
<td>540.13 (277.60)</td>
</tr>
<tr>
<td>Time 2 MHPG Raw Score</td>
<td>478.63 (414.32)</td>
</tr>
<tr>
<td>Time 3 MHPG Raw Score</td>
<td>578.70 (805.53)</td>
</tr>
</tbody>
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Note: Exposure is a dichotomous item indicating presence/absence of exposure to a life threatening event in the first year of police duty; All cortisol and MHPG values are log-transformed unless labeled as Raw Scores.

Table 2
Multinomial logistic regression for predictors of class membership \((N = 234)\).
this limitation by controlling age, gender, and baseline hormone levels which will account for random individual variability, and by examining change scores in response to a standardized task. Further, it remains unclear what the true impact of critical incidents and routine life stressors are on patterns of response. In the current study we found that exposure to a life-threatening event in the first year was not predictive of trajectory, and in the previous study we found that whether individuals were exposed to trauma throughout the 48 months was similarly not predictive of the trajectories (Galatzer-Levy et al., 2013). However, this may be because there is not much variability in dichotomous measures of exposure, as this is a highly exposed population overall. Conversely, continuous measures of exposure may not be predictive in this context as the population overall is highly and consistently exposed leading to truncated variability clustered in the high exposure range. A further limitation is that the relationship between cortisol response and early childhood trauma exposure was not explored. There is evidence that childhood trauma has a significant impact on HPA axis reactivity (Heim et al., 2000) and glucocorticoid receptor sensitivity (Klengel et al., 2013). As such, childhood trauma may explain observed differences by class on cortisol response to the video stressor. Unfortunately, though the Early Trauma Inventory Self Report—Short Form [ETISR-SF; (Bremner et al., 2007)] was collected in this sample, significant missingness in >50% of cases precluded these analyses. This represents a significant limitation that should be addressed in future studies. In addition, an alternative explanation of the results is that those in the Reactive-Worsening class simply mounted a more rapid cortisol response and recovery and as such, their response was not captured in the sampling window. This is plausible given that glucocorticoid hypersensitivity as well as glucocorticoid resistance have been identified in PTSD (Bachmann et al., 2005; Yehuda et al., 1991).

Finally, in previous work, we demonstrated that individuals in the Recovery class demonstrated both the highest levels of distress and PTSD symptomatology in years 1 through 3 and did not differ from those in the Reactive-Worsening class on levels of PTSD symptomatology at 48-months (Galatzer-Levy et al., 2013). However, PTSD symptom levels were primarily below the clinical cut-off in this sample limiting interpretations as to the role of cortisol or MHPG response in relation to psychopathology. As such, the present results do not clarify how the cortisol response influences patterns PTSD response.

Despite these limitations, the current study provides evidence that a blunted cortisol response prospectively predicts the subsequent development of chronic non-remitting stress responses. This finding has both theoretical and practical relevance. First, such tasks are easy to implement and may provide useful information regarding likely outcomes among populations that are likely to be exposed to PTEs and other significant stressors such as police, firefighters, first responders, and service members. Second, these findings provide evidence that a blunted cortisol response may be a pre-existing risk factor for poor adaptation to stressors. This informs the understanding of the role of individual differences in HPA-axis responses to stress as they relate to the development and long-term course of stress pathology and resilience. It is important to note that post-hoc analyses demonstrated that those on the Recovery and Resilient trajectories both demonstrated significant increases in cortisol response to the video challenge while those in the Reactive-Worsening class demonstrated a non-significant cortisol response. This may indicate that cortisol response influences more than the short term stress response; it also influences the long-term course of adaptation to stress. As such, identification of individuals with a blunted cortisol response may potentially aid in the early identification of those who are likely to follow a chronic course and informs our understanding of the neurobiological role in psychological vulnerability to pathological stress responses.

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**Contributors**

Isaac R. Galatzer-Levy conceptualized the manuscript, conducted the analysis, interpreted the result, wrote the manuscript text, and revised the manuscript following review.

Maria M. Steenkamp aided in writing and reanalysis for review.

Meng Qian aided in post-hoc data analysis.

Sabra Inslicht designed the experimental paradigm.

Clare Henn-Haase aided in interpretation of behavioral data.

Christian Otte aided in the interpretation of neuroendocrine results.

Rachel Yehuda aided in the interpretation of neuroendocrine results and the design of the experiment.

Thomas C. Neylan aided in the interpretation of behavioral and neuroendocrine results.

Charles R. Marmar aided in interpretation of results and writing of the manuscript.

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