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Behavioral/Cognitive

Distinct Trajectories of Cortisol Response to Prolonged Acute Stress Are Linked to Affective Responses and Hippocampal Gray Matter Volume in Healthy Females

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The development of robust laboratory procedures for acute stress induction over the last decades has greatly advanced our understanding of stress responses in humans and their underlying neurobiological mechanisms. Nevertheless, attempts to uncover linear relationships among endocrine, neural, and affective responses to stress have generally yielded inconsistent results. Here, 79 healthy females completed a well established laboratory procedure of acute stress induction that was modified to prolong its effect. Endocrinological and subjective affect assessments revealed stress-induced increases in cortisol release and negative affect that persisted 65 and 100 min after stress onset, respectively, confirming a relatively prolonged acute stress induction. Applying latent class linear mixed modeling on individuals’ patterns of cortisol responses identified three distinct trajectories of cortisol response: the hyper-response (n = 10), moderate-response (n = 21), and mild-response (n = 48) groups. Notably, whereas all three groups exhibited a significant stress-induced increase in cortisol release and negative affect, the hyper-response and mild-response groups both reported more negative affect relative to the moderate-response group. Structural MRI revealed no group differences in hippocampal and amygdala volumes, yet a continuous measure of cortisol response (area under the curve) showed that high and low levels of stress-induced cortisol release were associated with less hippocampal gray matter volume compared with moderate cortisol release. Together, these results suggest that distinct trajectories of cortisol response to prolonged acute stress among healthy females may not be captured by conventional linear analyses; instead, quadratic relations may better describe links between cortisol response to stress and affective responses, as well as hippocampal structural variability.

Key words: cortisol; hippocampus; HPA; mood; stress

Introduction
Stress sensitivity is a key element in the etiology and pathophysiology of psychopathology (Harkness et al., 2015). Accordingly, extensive scientific effort has been devoted to the characterization of the neural and physiological responses to acute stress that accompany the typical stress-induced temporary shift toward more

Significance Statement
Despite substantial research, it is unclear whether and how individual neuroendocrine stress response patterns are linked to affective responses to stress and structural variability in neuroendocrine regulatory brain regions. By applying latent class linear mixed modeling on individuals’ patterns of cortisol responses to a prolonged acute stressor, we identified three distinct trajectories of cortisol response. Relative to the group showing a moderate cortisol response, groups characterized by hyper and mild cortisol response were both associated with more negative affect. Moreover, a continuous measure of cortisol response showed that high and low levels of stress-induced cortisol release correlated with reduced hippocampal gray matter volume. Given that neuroendocrine stress responses are conceptualized as biomarkers of stress susceptibility, these insights may have clinical implications.
negative affective state. This work established that the physiological response to acute stress involve the activation of endocrine stress-response systems, most prominently the hypothalamic-pituitary-adrenal (HPA) axis (Herman et al., 2016). As the end point of the HPA system, salivary cortisol is the most frequently used variable to evaluate endocrine stress response in laboratory settings, with a cortisol increase ≥2.5 mmol/L after stress induction typically taken as a threshold for "stress response" (Foley and Kirschbaum, 2010). A wealth of laboratory acute stress procedures established that ≈50–80% of individuals can be classified as "stress responders" based on this cutoff (Dickerson and Kemeny, 2004). Interestingly, however, in most cases, links between stress-induced cortisol and affective responses are not clear, with the majority of studies not reporting significant differences in affective responses to stress among stress responders compared with nonresponders (Campbell and Ehler, 2012). Further, only ~25% of studies report a linear relation between cortisol and affective responses to stress (Campbell and Ehler, 2012).

The endocrine stress-response systems have also been the focus of neuroimaging research, particularly targeting the hippocampus and amygdala structures because of their pivotal roles in HPA regulation. Results have been mostly inconsistent, with hippocampal volume being positively associated (Pruessner et al., 2007), negatively associated (Cho, 2001), or not associated (Liu et al., 2012) with the magnitude of cortisol response to stress. Similarly mixed results also emerged with regard to amygdala volume (Klimes-Dougan et al., 2014; Cacciaglia et al., 2017). Together, attempts to link patterns of stress-induced cortisol response to both affective responses and structural variability in HPA-regulating brain regions have yielded mixed results. One potential explanation for these inconsistencies may relate to the fact that studies typically report on mean cortisol response or use predetermined values to classify individuals as stress responders versus nonresponders, thus disregarding the important role of individual differences in determining stress sensitivity (Monroe and Simons, 1991; Liu, 2015). In addition, given that current laboratory protocols induce stress for a relatively short time period (typically <20 min), it is possible that induced effects were too brief to allow sufficient endocrine and emotional variability to evolve.

To address these limitations, the first aim of the current study was to induce acute stress effectively for a relatively prolonged time period among healthy females while capturing individual endocrine and affective response patterns. To this end, healthy females completed the Maastricht Acute Stress Test (MAST), a robust laboratory acute stress procedure (Smeets et al., 2012) that was modified to prolong its effect by informing participants upon task completion that, due to their poor performance, they would need to repeat the task later in the session. Our second aim was to identify distinct trajectories of cortisol response to such prolonged acute stress without a priori assumptions regarding the number, size, or pattern of change of these trajectories. This was accomplished by applying latent class linear mixed modeling (LCMM) on individuals' patterns of cortisol responses. Our final aim pertained to investigating in a subsample (n = 69) with MRI data potential links between cortisol response to stress and structural variability with a priori hypotheses relating to key regions implicated in HPA regulation such as the amygdala and hippocampus. Overall, we hypothesized that applying a data-driven approach on cortisol patterns of response to stress may provide a more accurate account for individual variability and that these insights may enable linking stress-induced cortisol responses with affective responses and with structural variability in regions implicated in HPA regulation.

Materials and Methods

Participants. A total of 88 right-handed psychiatrically, medically, and neurologically healthy female participants were included. Only females were investigated to avoid potential sex-dependent variability in HPA axis stress response (Kudielka and Kirschbaum, 2005). All participants were recruited using community advertisements. Exclusion criteria included any current or past psychiatric disorder as assessed by a Structured Clinical Interview for the DSM-IV (SCID; First et al., 2005). In addition, individuals were excluded for five or more lifetime exposures to any illegal substance, as well as due to recent use of illegal drugs, psychotropic medications, or nicotine. For a complete summary of relevant demographic characteristics, please see Table 1.

Study procedure. Participants were tested in individual sessions between 11:00 A.M. and 4:00 P.M. to minimize the effects of diurnal variation on endogenous cortisol levels (Blascovich et al., 2011). The stress procedure itself occurred between 1:00 P.M. and 2:00 P.M. Further, to allow for controlled saliva collection, participants were asked not to brush their teeth and to refrain from food, drinks, and intense physical exercise at least 1 h before the test phase. None of the participants reported to have violated these directives. Upon arrival, participants received information about the study and the measurements that would be taken and provided written informed consent to a protocol approved by...
the Partners Institutional Review Board. Next, participants completed two tasks, one probabilistic reward task (PRT) and one reaction time (RT) task (these data will be reported separately) and a clinical interview to determine eligibility (SCID). After the interview, participants completed the modified MAST, which was followed again by administration of the two tasks (Fig. 1). After this first laboratory session, participants were asked to return to the laboratory within ~1 month to complete an MRI scanning session (mean days = 25, SD = 21). The scanning session included an fMRI task (described in Treadway et al., 2017), as well as a high-resolution anatomical scan.

MAST. Stress was induced via a modified version of the MAST (Smeets et al., 2012), a laboratory acute stress procedure that was shown previously to yield robust endocrine and affective stress responses among healthy individuals (Smeets et al., 2012). The MAST consists of a 5 min preparation phase and a 10 min acute stress phase that combines the physical aspects of immersing one hand in ice-cold water from the cold pressor test with the unpredictability, uncontrollability, negative social preparation phase and a 10 min acute stress phase that combines the visual analog mood scale (VAMS) at the same six time points as the saliva sample collection devices for Biological Health Psychology at Brandeis University (Direc-

T1-weighted MPRAGE images (TR = 2200 ms; TE = 1.54 ms; FOV = 230 mm; matrix = 192 × 192; resolution = 1.22 mm²; 144 slices). MRI data were analyzed using the voxel-based morphometry (VBM) module of the Computational Anatomy Toolbox (CAT12) (http://www.neuro.uni-jena.de/cat/) for SPM12 (Wellcome Department of Cognitive Neurology). VBM analysis incorporated the following preprocessing steps: (1) spatial registration to a reference brain, (2) tissue classification (segmentation) into gray and white matter and CSF, (3) bias correction of intensity nonuniformities, and (4) smoothing (8 mm).

Statistical analysis. Of the 88 participants who were recruited, saliva samples of nine participants were excluded from analyses due to missing data, leaving a total sample size of 79 participants for cortisol and α-amylase analyses. Ten additional participants did not undergo structural MRAI, leaving a total sample size of 69 for MRI analyses. Cortisol responses were log transformed before statistical analysis to reduce skewness. Main effects of time were tested using a repeated-measures ANOVA with the six sampling time points as a within-subject factor. A similar approach was implemented for α-amylase. Regarding the VAMS, ratings were transformed so that higher scores indicate greater negative affect. To probe parallel endocrine and affective patterns, only the six VAMS ratings that were assessed alongside saliva samples were included in the analyses. VAMS ratings were analyzed using a repeated-measures ANOVA with the six sampling time points and the five VAMS scales as within-subject factors. Repeated-measures ANOVAs were also implemented separately for self-report measures of state anxiety (STAI-S), positive affect (PANAS-PA), and negative affect (PANAS-NA) with the three time points as a within-subject factor. For all post hoc tests, α was set at 0.05 and adjusted (Bonferroni) for multiple comparisons.

LCMM, also called the growth mixture model (Bauер and Curran, 2003), was used to identify distinct classes (i.e., groups) of participants featuring similar trajectories of cortisol response to stress. Specifically, we tested whether the model that best fits our data included two, three, or four classes of distinct trajectories of cortisol response. Within LCMM, time was modeled as polynomials while allowing for linear, quadratic, and cubic trajectories to be derived empirically based on trajectory subgroups. For continuous outcomes such as cortisol values, the LCMM is an extension of the standard linear mixed model for handling various subpopulations of longitudinal trajectories (O’Brien and Fitzmaurice, 2005). Importantly, this approach captures all the heterogeneity in individual trajectories and identifies subgroups of participants with similar profiles of trajectories independently of observed participant’s characteristics. In other words, LCMM estimates the number of distinct trajectories of cortisol response that best capture variability in the data without requiring a priori assumptions regarding the number, size, or pattern of change of these trajectories. Bayesian information criterion (BIC) was used to compare different models, allowing for two versus three versus four classes and determine the optimal number based on variability in the data (Nyland et al., 2007). Low BIC values indicate a better fit of the model to the data. Analyses were performed using RStudio version 2005).
Overall effect of stress. Change in cortisol (A), α-amylase (B), VAMS rating (C), state anxiety (D), positive affect (E), and negative affect (F) throughout the session. Across all measures, there was a significant effect of time driven by stress-induced increase in cortisol release and negative affect. Notably, cortisol levels and negative affect were still elevated 65 and 100 min after stress onset, respectively, reflecting a relatively prolonged acute stress induction. Participants were fully debriefed and their mood had returned to baseline before they left the laboratory. *p < 0.05.

Results

Overall effect of stress

Repeated-measures ANOVA with cortisol responses to stress revealed a main effect of time ($F_{(5,390)} = 47.25$, $p < 0.001$), with a strong quadratic effect ($F_{(1,78)} = 40.70$, $p < 0.001$). Post hoc analyses revealed significant stress-induced increase in mean cortisol levels from before the MAST ($T_{0 \text{ min}}$) to 25 ($T_{25 \text{ min}}$), 50 ($T_{50 \text{ min}}$), and 65 min ($T_{65 \text{ min}}$) (all $p < 0.001$), but not 100 min after its completion, although it did not reach the significance level ($T_{100 \text{ min}}$; $p = 0.2$) (Fig. 2A). For α-amylase, the main effect of time was also significant ($F_{(5,390)} = 9.29$, $p < 0.001$). Post hoc tests revealed some increase from before the MAST to slightly after its completion, although it did not reach the significance level ($T_{25 \text{ min}}$; $p = 0.151$). There was, however, a significant decrease in α-amylase levels from 25 min after MAST onset to 40 min later ($T_{25 \text{ min}} > T_{65 \text{ min}}; p < 0.001$; Fig. 2B).

When considering VAMS ratings, repeated-measures ANOVA also resulted in a highly significant main effect of time, indicating an overall increase in negative affect across all VAMS scales ($F_{(3,234)} = 96.93$, $p < 0.001$), with the expected quadratic effect ($F_{(1,46)} = 124.40$, $p < 0.001$). Mirroring the cortisol results, negative mood across all scales was elevated 25 ($T_{25 \text{ min}}$), 50 ($T_{50 \text{ min}}$), and 65 min ($T_{65 \text{ min}}$) after stress onset relative to before stress ($T_{0 \text{ min}}$) (all $p < 0.001$). Furthermore, unlike cortisol, negative mood was still elevated at the final sampling time point 100 min after stress onset relative to before stress ($T_{100 \text{ min}} > T_{0 \text{ min}}; p < 0.001$), yet it was
significantly less negative at that point relative to 65 min after stress onset (after an additional 35 min that included the relief component: $T_{+65\text{ min}} > T_{+100\text{ min}}$, $p = 0.001$) (Fig. 2C).

Repeated-measures ANOVA on self-reported anxiety state (STAI-S) revealed a main effect of time ($F_{(2,162)} = 80.62$, $p < 0.001$) due to significantly higher anxiety levels after the MAST ($T_{+25\text{ min}}$) compared with both arrival ($T_{-80\text{ min}}$) and session completion ($T_{+100\text{ min}}$, all $p < 0.001$). Interestingly, anxiety levels were still reported anxiety state (STAI-S) revealed a significantly higher at session completion ($T_{+100\text{ min}}$). The model allowed for three latent classes also had a good discrimination ability, with $<15\%$ of participants a posteriori classified in other classes than the one initially assigned (see Table 2 for full statistic description).

Mix-effect ANOVA with the three groups from LCMM as a between-subjects factor and the five cortisol sampling time points as a within-subjects factor revealed, as expected, a significant

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**LCMM**

Approximately 72% of participants from our sample (57/79) could be classified as cortisol responders based on a cortisol increase $\geq 2.5\text{ nmol/L}$, suggesting that the laboratory procedure succeeded in activating HPA response. Despite these robust overall effects, patterns of cortisol response throughout the session greatly differed across participants, showing various patterns of change over time (Fig. 3A). Accordingly, individual cortisol response patterns were analyzed via LCMM aiming to estimate whether two, three, or four distinct trajectories of cortisol response best captured variability in the data without requiring any priori assumptions. BIC criteria comparing the different LCMM models yielded BIC$_1 = 424$, BIC$_3 = 413$, and BIC$_4 = 416$, indicating that a model allowing for three latent classes was optimal. These three classes were labeled based on their distinct trajectories of cortisol response to stress as the hyper-response ($n = 10$), moderate-response ($n = 21$), and mild-response ($n = 48$) groups. Figure 3B depicts the estimated mean trajectories of the three groups. Note that in the second-best model (4-class model), the fourth class was small (7.6% of the sample), thus contributing very little value beyond the 3-class model. The model allowing for three latent classes also had a good discrimination ability, with $<15\%$ of participants a posteriori classified in other classes than the one initially assigned (see Table 2 for full statistic description).

Mixed-effect ANOVA with the three groups from LCMM as a between-subjects factor and the five cortisol sampling time points as a within-subjects factor revealed, as expected, a significant

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**Figure 3.** LCMM. A, Individual patterns of cortisol response throughout the session ($n = 79$). B, Applying LCMM on these data revealed that the model that best fit our data included three latent classes, labeled based on their distinct trajectories of cortisol response to stress as the hyper-response ($n = 10$), moderate-response ($n = 21$), and mild-response ($n = 48$) groups. Note that all three groups exhibited a significant stress-induced increase in cortisol release 25 and 50 min after stress onset. *$p < 0.05$.

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**Table 2. LCMM model statistics**

<table>
<thead>
<tr>
<th>Between-model comparison</th>
<th>Log likelihood</th>
<th>NPM$^a$</th>
<th>BIC</th>
<th>Class 1$^b$</th>
<th>Class 2$^b$</th>
<th>Class 3$^b$</th>
<th>Class 4$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class 1 21 (26)</td>
<td>-185.89</td>
<td>12</td>
<td>424.22</td>
<td>40.5%</td>
<td>59.5%</td>
<td>11.69</td>
<td></td>
</tr>
<tr>
<td>Class 2 10 (13)</td>
<td>-169.81</td>
<td>17</td>
<td>413.89</td>
<td>26.6%</td>
<td>12.7%</td>
<td>60.7%</td>
<td>11.4%</td>
</tr>
<tr>
<td>Class 3 48 (61)</td>
<td>-159.67</td>
<td>22</td>
<td>415.47</td>
<td>7.6%</td>
<td>26.6%</td>
<td>11.4%</td>
<td>54.4%</td>
</tr>
</tbody>
</table>

Within the three-class model: fixed effects allowing for cubic time ($t$) trends

<table>
<thead>
<tr>
<th>n (%)</th>
<th>Coefficient</th>
<th>SE</th>
<th>Wald</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class 1 21 (26)</td>
<td>t $^1$</td>
<td>8.47</td>
<td>3.06</td>
<td>2.78</td>
</tr>
<tr>
<td></td>
<td>t$^2$</td>
<td>32.88</td>
<td>5.51</td>
<td>5.97</td>
</tr>
<tr>
<td></td>
<td>t$^3$</td>
<td>0.13</td>
<td>1.84</td>
<td>0.07</td>
</tr>
<tr>
<td>Class 2 10 (13)</td>
<td>t $^1$</td>
<td>-40.52</td>
<td>3.46</td>
<td>-11.69</td>
</tr>
<tr>
<td></td>
<td>t$^2$</td>
<td>-34.27</td>
<td>3.95</td>
<td>-8.67</td>
</tr>
<tr>
<td></td>
<td>t$^3$</td>
<td>-12.14</td>
<td>1.62</td>
<td>-7.51</td>
</tr>
<tr>
<td>Class 3 48 (61)</td>
<td>t $^1$</td>
<td>4.29</td>
<td>2.52</td>
<td>1.71</td>
</tr>
<tr>
<td></td>
<td>t$^2$</td>
<td>-17.07</td>
<td>4.02</td>
<td>-4.25</td>
</tr>
<tr>
<td></td>
<td>t$^3$</td>
<td>-3.97</td>
<td>1.36</td>
<td>-2.92</td>
</tr>
</tbody>
</table>

Within the three-class model: mean of posterior probabilities (n) in each class

<table>
<thead>
<tr>
<th>Class 1</th>
<th>Class 2</th>
<th>Class 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class 1 (n = 21)</td>
<td>91</td>
<td>4</td>
</tr>
<tr>
<td>Class 2 (n = 10)</td>
<td>85</td>
<td>8</td>
</tr>
<tr>
<td>Class 3 (n = 48)</td>
<td>4</td>
<td>1</td>
</tr>
</tbody>
</table>

$^a$Number of model parameters.

$^b$Posterior proportion for each class.
main effect of time, which was not pursued by post hoc tests. The ANOVA further revealed a main effect of groups ($F_{(2,76)} = 6.17, p = 0.003$) due to significantly higher overall cortisol release in the hyper-response class relative to the mild-response class ($p = 0.009$). Critically, a significant group $\times$ time interaction ($F_{(8,304)} = 36.92, p < 0.001$) also emerged, which was followed up by within- and between-group post hoc comparisons. Within-group analyses revealed a significant stress-induced increase in mean cortisol release from before the MAST ($T_{(0 \ min)}$) to both 25 ($T_{(+25 \ min)}$) and 50 ($T_{(+50 \ min)}$) minutes after MAST onset (all $p < 0.05$) in all three classes (Fig. 3B). Between-group comparison, however, revealed that the moderate-response class exhibited significantly increased cortisol release 25 min after MAST onset ($T_{(+25 \ min)}$) relative to both the mild-response and hyper-response classes ($p < 0.001$ and $p = 0.008$, respectively), whereas the hyper-response class exhibited significantly increased cortisol release 50 ($T_{(+50 \ min)}$) and 65 ($T_{(+63 \ min)}$) minutes after MAST onset relative to both the mild-response and moderate-response classes ($p < 0.001$) (Fig. 3B).

Effects of stress by cortisol class

When considering the VAMS ratings of the three LCMM groups, mixed-effect ANOVA revealed a main effect of time ($F_{(4,304)} = 23.96, p < 0.001$), a group $\times$ scale interaction ($F_{(8,1216)} = 2.74, p = 0.006$), and a group $\times$ time $\times$ scale interaction ($F_{(32,1216)} = 2.46, p < 0.001$), which was pursued for each scale separately. This resulted in three of the five scales showing a significant group $\times$ time interaction: happy–sad, $F_{(8,304)} = 2.06, p = 0.039$; relaxed–tense, $F_{(8,304)} = 3.10, p = 0.002$; friendly–hostile, $F_{(8,304)} = 2.15, p = 0.031$. For the happy–sad scale, post hoc tests indicated that these results were driven by significantly higher sadness scores 50 min ($T_{(+50 \ min)}$) after stress onset in the hyper-response class relative to both the mild-response and moderate-response classes ($p = 0.006, p = 0.003$, respectively) (Fig. 4A). A similar pattern emerged with regard to the relaxed–tense scale except that increased tension among hyper-response relative to the mild-response and moderate-response classes occurred 25 min after stress onset ($T_{(+25 \ min)}$) ($p = 0.05$ and $p = 0.023$, respectively; Fig. 4B). For the friendly–hostile scale, the analysis revealed that the mild-response class was driving the effect, exhibiting a trend toward more hostility 25 min after stress onset ($T_{(+25 \ min)}$) compared with the moderate-response class ($p = 0.080$; Fig. 4C). For state anxiety (STAI-S) and positive affect (PANAS-PA), the mixed-effect ANOVAs with LCMM groups revealed no effect of groups (Fig. 4D, E). For negative affect (PANAS-NA), however, there was a significant group $\times$ time interaction ($F_{(4,148)} = 3.17, p = 0.016$) driven by significantly more negative affect 25 min after stress onset in the mild-response class compared with the moderate-response class ($p = 0.037$) (Fig. 4F).

Structural variability and cortisol class

Mixed-effect ANOVA with the three groups from LCMM as a between-subjects factor and left and right hippocampal gray matter volume as a within-subjects factor revealed only a significant main effect of side due to overall more gray matter volume on the right compared with the left side ($F_{(1,66)} = 33.52, p < 0.001$), with no group or group $\times$ side effects ($p = 0.99$ and $p = 0.36$, respectively, data not shown). Analyses of amygdala gray matter volume yielded similar results, including a main effect of side (right > left; $F_{(1,66)} = 138.04, p < 0.001$) and no group or group $\times$ side effects ($p = 0.99, p = 0.16$, respectively, data not shown).

Structural variability and cortisol AUC

Hierarchical regressions with the continuous cortisol response measure AUCg separately for participants’ gray matter volumes in the left and right amygdala and hippocampus revealed a significant relation only for the right hippocampus. Further, a significant $F$ change in the hierarchical regression model ($F_{(1,66)} = 12.56, p = 0.001$; significant after Bonferroni correction; $p < 0.00625$) indicated that high and low AUCg values were both associated with less right hippocampal gray matter volume compared with moderate AUCg levels ($r = 0.421$; Fig. 5). Similar results appeared when using AUCi as the dependent measure.
the regression, showing only a significant, quadratic relation between AUCi magnitude and right hippocampal gray matter volume ($F_{1,66} = 4.86, p = 0.031; r = 0.262$). AUCi results, however, did not survive multiple-comparisons correction and should therefore be considered with caution.

Discussion

The overarching goal of the current study was threefold. First, we aimed to evaluate an acute stress laboratory procedure specifically designed to yield a prolonged effect. Toward this end, participants completed the MAST, a robust laboratory procedure for acute stress induction (Smeets et al., 2012), and were told immediately afterward that they would soon need to repeat the task due to their poor performance. Results indicated that this revised version of the MAST yielded stress-induced increase in cortisol release 25, 50, and 65 min after stress onset, as well as a shift toward more negative affect in all of these time points and up to 100 min after stress onset. This represents a relatively prolonged period of acute stress induction compared with previous reports, including studies using the original MAST procedure. Given the inherent delay between stress onset and increase in salivary cortisol and uncertainties whether individuals differ in such latency, efficient manipulations that induce a sustained acute stress response could be useful in future research.

Trajectories of cortisol response to prolonged acute stress

By probing individual patterns of stress-induced cortisol release over this relatively prolonged time period and analyzing these patterns using LCMM, our second aim was to investigate the number of distinct trajectories of cortisol response that best captured variability in our data without a priori assumptions. This analysis revealed three distinct trajectories of cortisol response, labeled as the hyper-response, moderate-response, and mild-response groups, all exhibiting a significant, quadratic increase in cortisol release. Generalization of this finding should proceed with caution. First, because only healthy females were included, no predictions can be made regarding cortisol patterns that may emerge in response to stress in males or among psychiatric samples. More critically, it is possible that specific sample characteristics influenced the final number of classes identified. In fact, we expect that a higher number of cortisol trajectories may emerge among larger and more environmentally and/or genetically diverse samples. Although the exact number of classes may vary based on sample characteristics, our results point to the importance of adequately modeling individual differences in cortisol response to stress and including multiple measures of the response trajectory when probing the endocrine stress response. The endocrine stress-response system, most prominently the HPA axis, acts via tightly regulated negative feedback loops that control the onset, magnitude, and duration of stress response activation (Joëls and Baram, 2009). Accordingly, these different parameters of the stress response system may all contribute to determine its physiological and affective outcomes.

Linking cortisol and affective responses to prolonged acute stress

In parallel with variability in cortisol stress response patterns, individuals are also known to differ in their affective responses to stress, including in measures of response threshold, amplitude, and rise time to peak (Davidson, 2000). In the current study, we found that, although all three cortisol groups experienced a shift toward a more negative affective state, individuals in the moderate-response class exhibited less stress-induced sadness and tension relative to individuals in the hyper-response class, as well as less hostility (trend) and negative affect relative to individuals in the mild-response class. Such quadratic association between the magnitude of cortisol response to stress and affective responses may account for the paucity of studies reporting a linear link between the two (Campbell and Ehlert, 2012). This quadratic pattern is also consistent with the well-established inverted U-shape relationship between basal levels of cortisol and cognitive performance, with beneficial effects of moderately elevated cortisol levels on cognition (de Kloet et al., 1999). Why both “too little” and “too much” cortisol are associated with reduced cognitive performance and/or heightened affective response to stress is not clear. One potential explanation may relate to the notion that, at least for some susceptible individuals, repeated hypersecretion of cortisol may have caused desensitization of the HPA axis, eventually resulting in reduced HPA sensitivity (Heim et al., 2000). This interpretation remains speculative in regard to our sample, however, because we did not assess participants’ lifetime exposure to stress. It has been suggested that lifetime exposure to stress itself may affect stress vulnerability via a quadratic pattern, challenging the assumption that stress and negative outcomes show a simple, linear relationship (Liu, 2015). Our findings suggest that cortisol and affective response to stress may also exhibit a nonlinear relation.

Linking cortisol responses to prolonged acute stress and hippocampal structural variability

The third aim of the current study was to investigate potential relations between cortisol response to stress and structural variability in the amygdala and hippocampus, key HPA regulatory regions. Resembling the link between cortisol and affective responses, a quadratic link was also present between a continuous measure of cortisol response to stress (AUCg) and hippocampal gray matter volume. Specifically, both high and low levels of stress-induced cortisol release were found to be associated with...
less right hippocampal gray matter volume compared with moderate cortisol release. This finding may be discussed in light of extensive preclinical evidence showing that exposure to cumulative stress may translate to morphological damage, most pronouncedly in the hippocampus (Sapolsky et al., 1990; Joëls et al., 2004). Increased susceptibility of the hippocampus to the effects of stress was attributed to its important role in stress regulation, as reflected by its substantial number of cortisol receptors (McEwen, 1999). In humans, reduced hippocampal volume has been the most commonly described neural structural abnormality in people who were exposed to traumatic stress and consequently developed an anxiety disorder such as posttraumatic stress disorder (Smith, 2005). Accordingly, reduced hippocampal volume among individuals exhibiting mild cortisol response may further link such a response pattern to repeated hyposecretion of cortisol. This interpretation, again, could be tested in studies assessing participants’ lifetime exposure to stress.

Limitations and conclusions

A few limitations of our study should be emphasized. First, it is not clear why α-amylase levels did not increase in response to stress in the current study nor why only some affective scales showed a significant relation with cortisol trajectories whereas others did not. Second, our LCMM-based classes did not differ in their hippocampal and amygdala gray matter volumes and the link between cortisol response and hippocampal volume only emerged when treating cortisol response as a continuous measure. This could relate to our modest sample size for these analyses, with only seven participants with MRI data in the hyper-response class (20 in the moderate-response class and 42 in the mild-response class). Whether the link between cortisol response to prolonged acute stress, affective responses, and hippocampal gray matter volume is categorical or continuous by nature is a topic for future research. A third limitation relates to the fact that factors that were shown previously to affect emotional and cortisol responses to acute psychosocial stress in healthy volunteers, including menstrual cycle phase (Duchesne and Pruessner, 2013), contraceptive usage (Roche et al., 2013), and personality traits (Childs et al., 2014), did not interact with cortisol classes in the current study ($\chi^2$ tests revealed no effects of menstrual cycle phase or contraceptives use on cortisol class: $\chi^2 = 1.82$ and 2.83, respectively; mixed-effect ANOVA revealed no interaction of NEO five-factor personality scores with cortisol class: $p = 0.82$). These inconsistencies could stem from the fact that previous work did not apply LCMM to classify cortisol response, but rather relied on mean group measures. Alternately, it may also be the case that additional factors we did not control for may have influenced current results, including age (Hostinar et al., 2014), fatigue (Bower et al., 2005) and body shape (Epeel et al., 2000), to name a few. Whereas controlling for all potential contributors to the stress response is practically impossible, future studies may choose to assess multiple factors and try to incorporate them into multidimensional models to account for additional variability in the data.

Our results suggest that investigating cortisol responses to stress without a priori assumptions regarding pattern of change can uncover distinct trajectories of cortisol response to prolonged acute stress among healthy females that would have been overlooked in conventional analyses parsing participants into cortisol responders and nonresponders. Most critically, identifying such distinct cortisol trajectories enabled us to discover the often-hidden link between stress-induced cortisol release and patterns of affective responses to stress as well as hippocampal structural variability. Given that neuroendocrine stress responses are conceptualized as biomarkers reflecting individual differences in stress resilience and susceptibility to psychopathology and disease (Feder et al., 2009), these insights regarding individual differences in trajectories of cortisol response to stress may have clinical implications.

References


References