Anxiety and Stress Alter Decision-Making Dynamics and Causal Amygdala-Dorsolateral Prefrontal Cortex Circuits During Emotion Regulation in Children

Stacie L. Warren, Yuan Zhang, Katherine Duberg, Percy Mistry, Weidong Cai, Shaozheng Qin, Sarah-Nicole Bostan, Aarthi Padmanabhan, Victor G. Carrion, and Vinod Menon

ABSTRACT

BACKGROUND: Anxiety and stress reactivity are risk factors for the development of affective disorders. However, the behavioral and neurocircuit mechanisms that potentiate maladaptive emotion regulation are poorly understood. Neuroimaging studies have implicated the amygdala and dorsolateral prefrontal cortex (DLPFC) in emotion regulation, but how anxiety and stress alter their context-specific causal circuit interactions is not known. Here, we use computational modeling to inform affective pathophysiology, etiology, and neurocircuit targets for early intervention.

METHODS: Forty-five children (10–11 years of age; 25 boys) reappraised aversive stimuli during functional magnetic resonance imaging scanning. Clinical measures of anxiety and stress were acquired for each child. Drift-diffusion modeling of behavioral data and causal circuit analysis of functional magnetic resonance imaging data, with a National Institute of Mental Health Research Domain Criteria approach, were used to characterize latent behavioral and neurocircuit decision-making dynamics driving emotion regulation.

RESULTS: Children successfully reappraised negative responses to aversive stimuli. Drift-diffusion modeling revealed that emotion regulation was characterized by increased initial bias toward positive reactivity during viewing of aversive stimuli and increased drift rate, which captured evidence accumulation during emotion evaluation. Crucially, anxiety and stress reactivity impaired latent behavioral dynamics associated with reappraisal and decision making. Anxiety and stress increased dynamic casual influences from the right amygdala to DLPFC. In contrast, DLPFC, but not amygdala, reactivity was correlated with evidence accumulation and decision making during emotion reappraisal.

CONCLUSIONS: Our findings provide new insights into how anxiety and stress in children impact decision making and amygdala-DLPFC signaling during emotion regulation, and uncover latent behavioral and neurocircuit mechanisms of early risk for psychopathology.

Keywords: Anxiety, Causal amygdala-DLPFC circuits, Children, Decision-making dynamics, Emotion regulation, Stress

https://doi.org/10.1016/j.biopsych.2020.02.011

Childhood is a vulnerable period for the development of symptoms and syndromes of anxiety, ranging from typical developmental experiences to pathological experiences (1). Nearly all affective disorders have an onset in childhood, and affective disorders are the most frequent mental disorders in children and adolescents (1,2). Cognitive and neural models of anxiety have implicated impaired emotion regulation in the etiology and maintenance of anxiety disorders (3–5). Few studies have investigated the cognitive and neural dynamics of emotion regulation in children and how these processes are altered by anxiety and stress reactivity, which is important for the development of clinically useful biomarkers for early diagnosis and treatment implementation. Here, we investigate how anxiety and stress reactivity impact latent behavioral emotion regulation processes and dynamic causal neural circuits in a developmentally homogeneous group of children.

Reappraisal is a widely used strategy for altering emotional reactivity to aversive stimuli and events by reinterpreting the meaning or significance of the experience (6). Reappraisal relies on a frontoparietal network linking the amygdala with the prefrontal cortex (PFC) regions involved in cognitive control, including the dorsolateral PFC (DLPFC), ventrolateral PFC (VLPFC), and medial PFC (7–11). The amygdala detects and encodes emotionally salient stimuli and mediates threat learning and vigilance (12). The DLPFC, along with coordinated interactions with other PFC regions, regulates adaptive...
responses to emotionally negative stimuli, thereby attenuating or heightening affective response (13–17). The DLPFC is of particular interest because it plays a critical role in supporting mental representations of affective states and their manipulation in working memory, processes that are essential for emotion regulation (6,18). Brain imaging studies of emotion regulation have also identified the DLPFC as a locus of deficits in several psychiatric disorders (19). Lastly, the DLPFC has an outsized role as an intervention target in brain stimulation studies that are designed to alleviate treatment-resistant anxiety and mood disorders in adults (20–23). Thus, convergent lines of evidence indicate that DLPFC dysfunction plays a prominent role in anxiety- and stress-related emotion dysregulation. Understanding the effects of anxiety and stress reactivity on causal amygdala-DLPFC circuits during emotion regulation in children may provide new insights into pathophysiological mechanisms of vulnerability to affective disorders.

Attentional control theory posits that excessive anxiety biases bottom-up signals from the amygdala, disrupting cognitive functions associated with the DLPFC, limiting attentional resources necessary for emotion regulation (24). This theory has not been empirically tested within a causal circuit analysis framework. As anxiety is associated with a bias toward negative interpretations, anxious individuals may not have the prerequisite abilities to alter distorted perceptions (25). However, extant behavioral studies suggest that anxious individuals perform similarly to nonanxious individuals on emotion reappraisal tasks (26,27). Thus, observed behavioral measures (reaction time, accuracy) may not be adequate to uncover dynamic processes driving reappraisal and its modulation by anxiety and stress (28). Computational modeling approaches are needed to uncover latent behavioral and neuronal processes and their links to emotional reappraisal and reactivity in children (6,29–31).

Here, we use computational modeling to determine how individual differences in anxiety and stress reactivity influence latent behavioral dynamics and causal bottom-up and top-down neural signaling between the amygdala and DLPFC during emotion regulation. A developmentally homogeneous (10–11 years of age) group of children performed an emotion reappraisal task during which they downregulated their emotional experience of aversive visual images (6,31). Trait-like measures of anxiety (worry) and stress reactivity (temperament), which are associated with appraising situations as more stressful and threatening and are risk factors for developing anxiety disorders (32,33), were assessed in each child. Drift-diffusion modeling (DDM) (34) was used to dissociate observed behavior into latent dynamic processes that represent distinct cognitive-affective components, including initial bias during viewing of aversive stimuli and a subsequent decision-making process during evaluation of emotional reaction. We tested the hypothesis that anxiety and stress reactivity negatively impact both of these latent emotion regulation processes. As worry is cognitively demanding, we hypothesized that anxiety would have broad effects on latent emotion regulation decision-making processes.

To investigate the role of causal amygdala-DLPFC circuits in emotion regulation, we used a state-space multivariate dynamical systems (MDS) model (35) to compute directional (bottom-up and top-down) causal interactions between amygdala and DLPFC in (latent) quasi-neuronal space, unconfounded by regional variations in hemodynamic response. We hypothesized that anxiety and stress in children would be associated with greater bottom-up causal interactions from the amygdala, reflecting a maladaptive influence on DLPFC cognitive control mechanisms. Alternatively, greater top-down causal interactions from the DLPFC might reflect an adaptive role in ameliorating the effects of anxiety and stress. Finally, we hypothesized that the DLPFC would be sensitive to decision making and evaluation of reactivity to aversive stimuli during emotion regulation.

METHODS AND MATERIALS

Participants

A total of 76 children were recruited from a suburban public school district in northern California as part of a larger study examining health and wellness in a historically low socioeconomic status and high-adversity community. Participants were excluded from the present study if they demonstrated excessive head motion during functional magnetic resonance imaging (fMRI) acquisition (n = 12), if they failed to engage in the task or if their behavioral data could not be acquired owing to equipment failure (n = 8), or if they did not complete clinical measures (n = 11). Our final sample size included 45 participants (Figure 1A; Supplement). Participant demographics are summarized in Table S1.

Clinical Measures of Anxiety and Stress Reactivity

The anxiety subscale from the Behavior Assessment System for Children, Second Edition (36) was used to evaluate predominantly worry-related anxiety symptoms. The involuntary response to stress subscale from the Response to Stress Questionnaire (37) was used to assess physiological and/or temperamental reactions to stressors (stress reactivity).

fMRI Experimental Design and Emotion Regulation Task

Consistent with well-validated procedures (31), participants were trained on the experimental paradigm prior to scanning. Participants were told that they would see an instructional cue followed by an image. For LOOK cues, participants were asked to notice their feelings toward the picture. For LESS cues, participants were asked to reappraise aversive images by telling themselves a story to make the pictures seem less negative, or more positive (Supplement).

During fMRI acquisition, participants completed 2 scanning runs, each consisting of 30 experimental trials. Each trial began with a 2-second instructional cue word (LOOK or LESS), followed by an aversive or neutral image appearing for 7.5 seconds, followed by a rating scale appearing for 2 seconds (Figure 2A). Participants rated their emotional state for the following conditions: looking at neutral images and responding naturally (LOOK; neutral condition), looking at aversive images and responding naturally (LOOK; aversive condition), and reappraising aversive images (LESS; reappraisal condition). There were 20 trials in each of the 3 task conditions: neutral, aversive, and reappraisal. The rating scale consisted of
numbers 1 (okay) to 4 (very bad). Participant ratings served as a behavioral index of reappraisal effectiveness. Reappraisal success was computed using the following equation: \((\frac{m_{\text{aversive}} - m_{\text{reappraisal}}}{m_{\text{aversive}}}) \times 100\), with higher scores indicating better reappraisal ability.

### fMRI Data Acquisition and Preprocessing

Images were preprocessed using a standard SPM12 pipeline. For each participant, contrast images corresponding to aversive versus neutral, reappraisal versus neutral, and reappraisal versus aversive task conditions were generated using a general linear model (GLM) and an omnibus F-test to identify amygdala and prefrontal cortex (PFC) regions of interest (ROIs). Time series were extracted from each ROI and were used to estimate causal interactions between amygdala and PFC ROIs using a multivariate dynamical state-space model. Finally, the relations among latent behavioral dynamics, anxiety/stress, and casual brain circuit measures were examined. DDM, drift-diffusion modeling, DLPFC, dorsolateral PFC.

---

**Figure 1.** Schematic view of participant selection procedure and data analysis pipeline. (A) Children (10–11 years of age) were excluded if they failed to engage in the task or if their behavioral data could not be acquired owing to equipment failure \((n = 8)\), demonstrated excessive motion during functional magnetic resonance imaging (fMRI) acquisition \((n = 12)\), or did not complete clinical measures \((n = 11)\), yielding a final sample size of 45 participants. (B) Reappraisal success and latent behavioral dynamics were computed using behavioral data. The Behavior Assessment System for Children, Second Edition (BASC) anxiety and Response to Stress Questionnaire stress reactivity subscales were the clinical measures of interest. Brain responses to task conditions were estimated using a general linear model (GLM) and an omnibus F test to identify amygdala and prefrontal cortex (PFC) regions of interest (ROIs). Time series were extracted from each ROI and were used to estimate causal interactions between amygdala and PFC ROIs using a multivariate dynamical state-space model. Finally, the relations among latent behavioral dynamics, anxiety/stress, and casual brain circuit measures were examined. DDM, drift-diffusion modeling, DLPFC, dorsolateral PFC.

**Figure 2.** Functional magnetic resonance imaging (fMRI) experimental design, behavioral performance, and emotion regulation abilities. (A) Children 10–11 years of age were presented with cues (LOOK or LESS) followed by neutral or aversive images. They were asked to notice their feelings toward the picture when LOOK was presented, and to reappraise aversive images by telling themselves a story to make the pictures seem less negative or more positive when LESS was presented. The neutral condition consisted of viewing a neutral picture, the aversive condition consisted of viewing an aversive picture, and the reappraisal condition consisted of reframing an aversive picture as less negative. Following the presentation of a neutral or aversive image, a rating scale consisting of numbers from 1 (okay) to 4 (very bad) was shown for children to indicate their emotional evaluation. (B) Children reported neutral stimuli to be significantly less unpleasant than the aversive images, regardless of the instructional cue, and reported the aversive stimuli to be significantly less unpleasant during the reappraisal condition. ***\(p < .001\). IAPS, International Affective Picture System.
linear model. An omnibus $F$ test was used to identify brain regions showing significant group-level responses to reappraisal versus neutral or aversive versus neutral task conditions, with a height threshold $p < .005$ and familywise error corrections for multiple comparisons at $p < .01$ (minimum cluster size = 87 voxels or 696 mm$^3$). Activation peaks in bilateral amygdala, DLPFC, and other PFC regions were identified and used to construct 6-mm-sphere regions of interest (ROIs) for subsequent dynamic causal and latent brain-behavior analyses (Figure 1B, Figure S2, Supplement).

**Computational Modeling of Latent Behavioral Dynamics During Emotion Regulation**

The emotion evaluation process was modeled as a drift diffusion process, in which evidence accumulates over time, resulting in a decision when a decision threshold is reached. The evaluations were coded as positive (ratings of 1 or 2) or negative (ratings of 3 or 4) (Figure 3A). The initial bias represented the starting point for the drift-diffusion process, and it captures the initial reaction during image viewing, prior to the decision window. The drift rate parameter ($d$) characterizes evidence accumulation, with higher values indicating a greater proportion of positive responses, and higher absolute values of the drift rate characterizing faster responses. For this task, drift rate indexes not only evidence accumulation, but also the decision to make an evaluative response (“okay” to “very bad”) when presented with an image. The decision threshold parameter ($a$) captures response caution, or the degree of confidence required to conclusively evaluate the emotion, with higher values characterizing slower and more consistent responses. The decision threshold for an individual was allowed to vary by instruction—viewing (LOOK) versus reappraisal (LESS). The drift rate and initial bias could vary by instruction (LOOK, LESS) and stimulus type (neutral, aversive, and reappraisal). The nondecision time, reflecting perceptual processes prior to evidence accumulation, for each individual was fixed across instructions and stimulus types. Initial bias for the aversive reappraisal condition was constrained to lie between viewing neutral and aversive conditions. The DDM was implemented within a Bayesian inference framework.

![Drift-diffusion model of latent behavioral dynamics.](image)

(A) Illustration of a single trial of the drift-diffusion process, in which the random walk represents noisy evidence accumulation over time for a positive versus negative evaluation of the stimulus. When the evidence accumulation process hits either decision boundary (separated by the decision threshold), a response is made. The initial bias captures the bias toward positivity or negativity that is built up over the 7500-ms stimulus window and acts as a starting point for the random walk. The drift rate captures the rate of evidence accumulation during the 2000-ms response window. (B) Children showed significantly greater initial bias under the reappraisal than the aversive condition. Initial bias was highest in the neutral condition. (C) Children showed significantly higher drift rates under the reappraisal than the aversive condition. The drift rate was highest in the neutral condition. *$p < .05$; **$p < .001$.
framework using JAGS (38). Model fit was validated by comparing the posterior predictive model emotion evaluations and response times under the 3 different conditions to the actual values (Supplement).

**Computational Modeling of Dynamic Causal Interactions Between the Amygdala and DLPFC**

We used MDS, a state-space model for estimating context-dependent causal interactions between multiple brain regions while accounting for regional variation in hemodynamic responses (35). MDS has been validated using extensive simulations (35,39,40). See the Supplement for details of the computational model and variational Bayes solution used to infer model parameters.

**RESULTS**

**Behavioral Performance and Emotion Regulation Abilities**

Children rated their emotional reaction to negative stimuli during reappraisal and aversive task conditions, and to stimuli in a neutral task condition. Stimuli were rated as less unpleasant during the reappraisal condition than during the aversive condition ($t_{44} = -3.57, p = .001$) (Figure 2B). Stimuli were rated as more unpleasant during the aversive ($t_{44} = 13.66, p < .001$) and reappraisal ($t_{44} = 8.55, p < .001$) conditions than the neutral condition. Results demonstrate that children are able to modulate their negative affective ratings of aversive stimuli, with more positive evaluations reflecting a higher degree of reappraisal success.

**Latent Behavioral Dynamics During Emotion Regulation**

A novel implementation of DDM was used to determine 1) initial bias, which captures the initial reaction during viewing of aversive images, prior to the decision window; 2) the drift rate, which captures the ability to regulate emotion evaluation during the response (decision) window; and 3) a decision threshold, which measures response caution (Figure 3A). Children showed a greater initial bias during the reappraisal condition ($0.51 \pm 0.11$) than during the aversive condition ($0.48 \pm 0.14$) ($t_{44} = 3.62, p < .001$) (Figure 3B). Children also showed higher drift rates during the reappraisal condition ($0.22 \pm 0.88$) than during the aversive condition ($-0.12 \pm 0.79$) ($t_{44} = 2.67, p = .011$) (Figure 3C). Children did not show a significant difference in the decision threshold parameter during the reappraisal condition ($2.83 \pm 1.1$) than during the aversive condition ($2.83 \pm 1.08$) or the neutral condition ($2.83 \pm 1.08$). Results show that emotion regulation was characterized by increased positivity bias while viewing images during the reappraisal condition and by higher drift rate during the decision period when evaluating their emotional reaction.

**Latent Behavioral Dynamics During Decision Making Are Correlated With Reappraisal Scores**

We determined whether DDM-derived latent cognitive parameters are related to reappraisal success. Individual reappraisal scores were correlated with a change in drift rate between the reappraisal and aversive conditions ($t_{42} = 12.96, r = .89, p < .001$) (Figure S1). Hierarchical linear regressions included reappraisal success as the dependent variable and changes in initial bias, drift rate, and decision threshold under reappraisal versus aversive conditions as the independent variables, and revealed an excellent model fit (adjusted $R^2 = .78, F_{4,41} = 53.64, p < .001$). Change in drift rate from the aversive to reappraisal conditions was the only independent variable that contributed unique variance, and thus emerged as the dominant predictor ($t_{41} = 11.36, \beta = .99, p < .001$) (see Supplemental Results). Results suggest that success in emotion regulation is characterized by decision making during the postviewing, response period and not by initial bias during reappraisal.

**Anxiety and Stress Impair Latent Behavioral Dynamics During Viewing and Evaluation**

Anxiety scores were negatively correlated with initial bias ($t_{43} = -2.18, r = -.32, p = .035$), drift rate ($t_{43} = -2.36, r = -.34, p = .023$), and decision threshold ($t_{43} = -2.28, r = -.33, p = .028$) during reappraisal (Figure 4A–C). Stress reactivity was negatively correlated with the decision threshold during reappraisal ($t_{43} = -2.34, r = -.34, p = .024$) (Figure 4D). Anxiety and stress reactivity were not correlated with reappraisal success ($p > .4$). Results demonstrate that anxiety and stress impair latent behavioral dynamics of emotion regulation and that DDM captures their influence on behavior in ways that traditional response selection and reaction time measures by themselves do not.

**Brain Areas Activated During Reappraisal and Aversive Emotion Processing**

An omnibus $F$ test contrasting reappraisal versus neutral or aversive versus neutral conditions revealed significant activation in bilateral amygdala, DLPFC, dorsomedial PFC (DMPFC), VLPFC, posterior parietal cortex, posterior cingulate cortex, and occipital cortex (Figure 5, Figure S2, Tables S2–S5), consistent with previous reports (7–10,41) (Supplemental Results).

**Anxiety Increases Causal Interactions Between the Amygdala and DLPFC During Emotion Regulation**

To identify amygdala and DLPFC ROIs for causal circuit and latent brain-behavior analyses, we used task-related activation identified by the $F$ test, as described above, thereby avoiding biases associated with selection of regions specific to either task condition. We also conducted additional control analyses using multiple PFC regions (DLPFC, VLPFC, DMPFC, and anterior insula) identified by the $F$ test (Table S5), with false discovery rate (FDR) corrections for multiple comparisons. We then used MDS to compute task condition–specific causal circuit interactions between amygdala and DLPFC ROIs within each hemisphere. A contrast of the strength of causal interactions between the reappraisal and aversive conditions was used to probe how anxiety influences causal circuit interactions during emotion regulation. Both forward (amygdala → DLPFC) and backward (DLPFC → amygdala)
links in both hemispheres were tested, with FDR corrections ($p < .05$) for multiple comparisons.

The strength of causal influence from right amygdala to right DLPFC (Figure 6A) during emotion regulation (reappraisal vs. aversive conditions) was positively correlated with the Behavior Assessment System for Children, Second Edition anxiety subscale ($t_{42} = 3.28, r = .45$, FDR-corrected $p = .008$) (Figure 6B). No such effects were observed in the reverse connectivity pattern (DLPFC / amygdala). Post hoc analysis of left amygdala / DLPFC link with anxiety showed a marginally significant effect ($t_{42} = 1.84, r = .27$, uncorrected $p = .074$).

We conducted an additional analysis using bilateral VLPFC, DMPFC, and anterior insula regions that also showed significant activation associated with emotion processing (Table S6). Again, only the right amygdala → DLPFC link was significantly correlated with anxiety (FDR-corrected $p = .033$).

**Stress Reactivity Increases Causal Interactions Between the Amygdala and DLPFC During Emotion Regulation**

Next, we found that the strength of causal influence from right amygdala → DLPFC was positively correlated with stress reactivity ($t_{42} = 3.04, r = .42$, FDR-corrected $p = .016$) (Figure 6C). No such effects were observed for the reverse direction (DLPFC → amygdala).

**The Amygdala-DLPFC Causal Circuit Is a Common Pathway for Anxiety and Stress During Emotion Regulation**

To disentangle the roles of anxiety and stress in their relation to amygdala → DLPFC causal interaction during emotion regulation, we conducted additional analyses using residualized anxiety, derived by regressing stress out from anxiety, and residualized stress, derived by regressing anxiety out from anxiety.
stress. The strength of causal influence from the right amygdala to the right DLPFC was correlated with neither residualized anxiety ($t_{42} = 1.54, r = .23, p = .13$) nor residualized stress ($t_{42} = 1.06, r = .16, p = .30$). Structural equation modeling revealed a significant relationship between the strength of right amygdala → DLPFC causal interactions and a latent factor underlying anxiety and stress (Figure 6D). These results suggest that shared variance between anxiety and stress reactivity drives bottom-up amygdala → DLPFC signaling during emotion regulation (Supplemental Results).

### Right DLPFC Reactivity Is Correlated With Latent Behavioral Dynamics

Finally, we investigated the role of the DLPFC in decision making during emotion regulation. Activation in the right DLPFC was correlated with the difference in drift rate between the reappraisal and aversive conditions ($t_{42} = 2.48, r = .36, p = .017$) (Figure 7). No such relation was observed with amygdala response or causal interactions between amygdala and right DLPFC ($ps > .05$). Results demonstrate that the right DLPFC, rather than amygdala, reactivity underlies evidence accumulation and decision making during emotion regulation.

### DISCUSSION

Although anxiety and stress are known risk factors for the development of affective disorders, the behavioral and neurocircuit mechanisms that potentiate maladaptive emotion regulation behaviors are poorly understood. We used computational tools to investigate how anxiety and stress impact latent decision-making processes and dynamic causal amygdala-PFC interactions during emotion regulation in children. A National Institute of Mental Health Research Domain Criteria approach (42,43) allowed us to capture dimensional and shared representations of childhood anxiety and stress reactivity. We found that emotion regulation in children is characterized by increased initial bias during viewing of aversive stimuli and a more positive evaluation of their emotional reaction to negative stimuli during reappraisal. Anxiety impaired multiple latent behavioral dynamic measures, including initial bias and decision making during evaluation, and stress reactivity resulted in less confident, more impulsive decision making. State-space circuit modeling revealed that directed causal influences from amygdala to DLPFC, but not the reverse, were exacerbated by anxiety and stress reactivity during reappraisal. Furthermore, DLPFC, but not amygdala,
reactivity was correlated with weak evidence accumulation during emotion reappraisal. Control analyses confirmed the specificity of our findings with respect to the amygdala and DLPFC, and their functional circuit interactions. Our findings reveal latent dynamic behavioral and neurocircuit mechanisms of early pathophysiology during childhood.

**Anxiety and Stress Impair Latent Behavioral Dynamics During Emotion Regulation**

We devised a novel DDM to disentangle 3 latent components supporting emotion regulation: 1) initial reaction while viewing aversive stimuli, captured by changes in initial bias; 2) decision making during emotion evaluation, captured by changes in drift rate; and 3) response caution, captured by changes in decision threshold (44) (Figure 3A).

DDM revealed that children with higher anxiety demonstrated lower positivity bias (initial reaction), lower drift rate (lower ability to regulate), and lower decision threshold (less consistent and controlled evaluation) during reappraisal. These results point to lower and less consistent positivity ratings under reappraisal for higher anxiety scores and highlight latent mechanisms by which anxiety impacts the reappraisal process (Figure 4). Stress reactivity effects were only observed in relation to decision threshold, indicating that reappraisal is associated with less cautious and more impulsive decision making in children with higher reactive stress responses. These effects were specific to latent behavioral dynamic measures, as overt ratings of reappraisal were not correlated with clinical measures of anxiety or stress reactivity. Thus, DDM provided a more sensitive measure of anxiety and stress reactivity effects on behavior than traditional response times.

![Figure 6](image6.png)

**Figure 6.** Anxiety and stress increase causal interactions between the amygdala (AMY) and dorsolateral prefrontal cortex (DLPFC) during emotion regulation. (A) The strength of causal influence from the right amygdala to the right DLPFC during emotion regulation was positively correlated with Behavior Assessment System for Children, Second Edition anxiety (B) and Response to Stress Questionnaire stress reactivity (C) scores. (D) Structural equation modeling revealed that shared variance between anxiety and stress drives right amygdala → DLPFC interactions during emotion regulation.
Anxiety and Stress Alter Emotion Dynamics and Circuits

Figure 7. Right dorsolateral prefrontal cortex (DLPFC) reactivity increases with evidence accumulation during emotion regulation. Increased activation in right DLPFC (reappraisal vs. aversive conditions) was correlated with evidence accumulation during evaluation of emotional reaction, as assessed by change in drift rate between the conditions.

Anxiety and Stress Are Correlated With Causal Right Hemisphere Amygdala → DLPFC Interactions During Emotion Regulation

We focused on dissociations between bottom-up and top-down causal interactions between the amygdala and DLPFC, as this pathway is theorized to be critical for regulating reactivity to negative emotions (8,18,46). Our causal circuit analysis addressed an important gap in the literature, as the effects of anxiety and stress on this core pathway have been poorly understood. Our analysis revealed that the strength of dynamic causal interaction from right amygdala to right DLPFC was enhanced by both anxiety and stress reactivity during emotion regulation. Crucially, top-down influences from the DLPFC to amygdala were not correlated with anxiety or stress, highlighting the specificity of bottom-up signaling from amygdala to DLPFC. These effects were also specific to the DLPFC, as amygdala interactions with the insula, ventromedial PFC, and DMPFC were not correlated with anxiety and specific to right hemisphere amygdala-DLPFC interactions. We also found that variance shared between anxiety and stress reactivity drives right amygdala-to-DLPFC signaling, suggesting that these interactions reflect a transdiagnostic circuit in childhood. Furthermore, asymmetric involvement of right amygdala-DLPFC circuits is consistent with right hemispheric dominance for anxiety and anxiety-related processes observed in adults (47–51).

Our findings that both anxiety and stress are associated with bottom-up, context-dependent functional signaling from the amygdala are consistent with attentional control theory, which posits that excessive anxiety biases bottom-up signals from the amygdala (24). Our findings add important developmental dimensions to emerging neurobiological models of anxiety and stress close to the age at which these symptoms manifest, suggesting that early adverse experiences are associated with modulation of causal dynamics in amygdala-DLPFC circuitry, rather than with amygdala reactivity itself.

Right DLPFC Reactivity Drives Evidence Accumulation and Decision Making During Emotion Regulation

We next investigated DLPFC involvement in decision making during emotion regulation, given its central role in cognitive and affective control (8). Right DLPFC activity was modulated by drift rate, reflecting the efficiency of evidence accumulation and the decision-making process during emotion regulation. These effects were specific to DLPFC reactivity, as drift rate did not modulate amygdala activation or causal interactions between the amygdala and DLPFC. Furthermore, reaction time and response selection were not associated with DLPFC activity, demonstrating that latent behavioral dynamic measures provide new insights into the role of the DLPFC that cannot be obtained from overt behavioral measures. While DDM has been widely used to investigate perceptual decision making, to our knowledge, no previous studies have examined decision-making processes associated with emotion regulation. It is noteworthy that aversive stimuli remained perceptually unchanged while the child reappraised their negative content, revealing a novel aspect of DLPFC function based on internally generated cognitive control processes during reappraisal. These findings further highlight the role of the DLPFC in children’s decision making during emotion regulation.

Integrative View of Findings

Increased right amygdala → DLPFC connectivity with anxiety and stress raises the question of whether such signaling reflects adaptive or maladaptive function. Our findings suggest that enhanced signaling in this pathway represents “hijacking” a key cortical circuit involved in cognitive control. If it were primarily an adaptive function that signaled the need for more top-down control, we would expect that causal DLPFC → amygdala connectivity would be negatively correlated with anxiety and stress, which we did not find evidence for. More likely, this signaling is not effectual in increasing top-down control. Moreover, based on latent behavioral findings that anxiety and stress reactivity impair emotion regulation decision-making dynamics, enhanced anxiety- and stress-related causal amygdala-DLPFC signaling points to ineffective engagement of cognitive control.

Clinical Implications

Anxiety disorders are generally chronic and persist into adulthood, and childhood is a critical period for their onset. Our findings may represent a promising target for understanding early pathophysiology and are in line with the recent focus on identifying early psychological and biological dimensional factors that cut across diagnoses to explain mental illness (52). Bottom-up causal amygdala-DLPFC signaling may represent a critical transdiagnostic circuit as trait-like aspects of negative emotional reactivity, including cognitive (worry) and...
temperamental (stress reactivity), are present in some form across all anxiety disorders and related psychopathologies (53,54). Our characterization of trait-like anxiety and stress reactivity effects on a circumscribed functional circuit in the developing brain may provide a fruitful approach for understanding the emergence and course of early pathological anxiety and related disorders.

Conclusions

Our study provides new insights into how anxiety and stress reactivity in children impact latent decision-making processes, dynamic causal interactions between the amygdala and DL-PFC, and DL-PFC reactivity during emotion regulation. Over time, it is likely that these dynamics would impoverish cognitive control processes anchored in the right DL-PFC, rendering it a node of vulnerability and a target for intervention. Our identification of a common circuit that impacts cognitive-emotional function in preadolescence may contribute to improved early treatment of anxiety disorders and related psychopathology.

ACKNOWLEDGMENTS AND DISCLOSURES

This work was completed in partnership with the Ravenswood City, Alum Rock, and Orchard school districts and Pure Edge, Inc., and was supported by the Lucile Packard Foundation for Children’s Health (to VC); National Institutes of Health Grants Nos. EB022907, NS086085, and MH121069 (to VM); the Stanford Maternal Child Health Research Institute through the Transdisciplinary Initiatives Program (to VM); and a Stanford Institute for Computational & Mathematical Engineering GPU computing seed grant (to VM).

We thank Dr. Lang Chen for valuable input on data analysis.

The authors report no biomedical financial interests or potential conflicts of interest.

ARTICLE INFORMATION

From the Department of Psychiatry and Behavioral Sciences (SLW, YZ, KD, PM, WC, SQ, S-NB, AP, VGC, VM), Stanford School of Medicine, Stanford; and Department of Psychology (SLW), Palo Alto University, Palo Alto, California; and the State Key Laboratory of Cognitive Neuroscience and Learning (SQ), IDG/McGovern Institute for Brain Research, Beijing Normal University, Beijing, China.

SLW and YZ contributed equally to this work.

Address correspondence to Vinod Menon, Ph.D., Department of Psychiatry and Behavioral Sciences, Stanford School of Medicine, Stanford, CA 94305; E-mail: menon@stanford.edu.

Received Aug 13, 2019; revised Jan 22, 2020; accepted Feb 10, 2020.

Supplementary material cited in this article is available online at https://doi.org/10.1016/j.biopsych.2020.02.011.

REFERENCES

Anxiety and Stress Alter Emotion Dynamics and Circuits