Emotion

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BRIEF REPORT

Microstructural Integrity of a Pathway Connecting the Prefrontal Cortex and Amygdala Moderates the Association Between Cognitive Reappraisal and Negative Emotions

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Duke University

Cognitive reappraisal is a commonly used form of emotion regulation that utilizes frontal-executive control to reframe an approaching emotional event to moderate its potential psychological impact. Use of cognitive reappraisal has been associated with diminished experience of anxiety and depressive symptoms, as well as greater overall well-being. Using data from a study of 647 healthy young adults, we provide initial evidence that an association between typical use of cognitive reappraisal in daily life and the experience of anxiety and depressive symptoms is moderated by the microstructural integrity of the uncinate fasciculus, which provides a major anatomical link between the amygdala and prefrontal cortex. Our findings are consistent with the nature of top-down regulation of bottom-up negative emotions and suggest the uncinate fasciculus may be a useful target in the search for biomarkers predicting not only disorder risk but also response to psychotherapy utilizing cognitive reappraisal.

Keywords: cognitive reappraisal, depression, anxiety, uncinate fasciculus, emotion regulation

Cognitive reappraisal is a common form of negative emotion regulation that requires top-down executive control to reframe a potentially aversive experience to mitigate its emotional impact (Gross, 1999; Gross & John, 2003). Individuals who typically employ reappraisal techniques in daily life report fewer depressive symptoms, less anticipatory anxiety, and greater overall well-being (Gross & John, 2003). Accordingly, poor emotion regulation and infrequent or ineffective use of cognitive reappraisal is observed across mental disorders, especially depression and anxiety (Gross & John, 2003; Joormann & Gotlib, 2010; Sloan et al., 2017).

The neural mechanisms supporting cognitive reappraisal parallel its core features of top-down executive control over negative emotion. Specifically, successful downregulation of negative emotion through cognitive reappraisal involves concurrent increases in the activity of the dorsolateral prefrontal cortex (dLPFC), ventrolateral prefrontal cortex (vLPFC), and dorsomedial prefrontal cortex (DMpFC), as well as decreases in amygdala activity (Buhle et al., 2014; Drabant, McRae, Manuck, Hariri, & Gross, 2009; Ochsner, Bunge, Gross, & Gabrieli, 2002). As there is no evidence for direct structural connections between these regions and the amygdala, this functional interaction is likely mediated through the ventromedial prefrontal cortex (vmPFC), which has direct connections with the dLPFC, vLPFC, DMpFC, and with the amygdala, acting as a gateway between these regions (Delgado, Nearing, LeDoux, & Phelps, 2008). Principal among these connections is the uncinate fasciculus (UF), a large bidirectional ipsilateral white matter tract that broadly connects the vmPFC and limbic regions (Von Der Heide, Skipper, Klobusicky, & Olson, 2013; Figure 1.).

We have previously reported that the microstructural integrity of the UF, assayed using noninvasive diffusion-weighted MRI, is correlated with the typical use of cognitive reappraisal (Zuurbier, Nikolova, Åhs, & Hariri, 2013). Prior research has also implicated diminished UF microstructural integrity in depression and anxiety.

Tracy C. d’Arbeloff, M. Justin Kim, Annchen R. Knodt, Spenser R. Radtke, Bartholomew D. Brigidi, and Ahmad R. Hariri, Laboratory of NeuroGenetics, Department of Psychology and Neuroscience, Duke University.

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Tracy C. d’Arbeloff developed the study concept. Data analyses were performed by Tracy C. d’Arbeloff and Annchen R. Knodt under the supervision of Ahmad R. Hariri. Bartholomew D. Brigidi designed the clinical interview and neuropsychological testing protocols. Spenser R. Radtke and Annchen R. Knodt collected the data. Tracy C. d’Arbeloff, M. Justin Kim, and Ahmad R. Hariri drafted the manuscript. Ahmad R. Hariri conceived and designed the parent Duke Neurogenetics Study. All authors reviewed and approved the final version of the manuscript.

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Correspondence concerning this article should be addressed to Tracy C. d’Arbeloff, Department of Psychology and Neuroscience, Duke University, Durham, NC 27708. E-mail: tracy.darbeloff@duke.edu
specific self-report questionnaires. All 647 participants were free of the following conditions: (1) medical diagnoses of cancer, stroke, head injury with loss of consciousness, untreated migraine headaches, diabetes requiring insulin treatment, chronic kidney, or liver disease; (2) use of psychotropic, glucocorticoid, or hypolipidemic medication; and (3) conditions affecting cerebral blood flow and metabolism (e.g., hypertension).

**Self-Report Questionnaires**

The Mood and Anxiety Symptoms Questionnaire (MASQ) Short Form is a 62-item self-report questionnaire used to assess current anxious and depressive symptoms over the past week. Symptoms of depression and anxiety are scored along four subscales: two General Distress factors (22 items, 11 assessing symptoms relative to depression and 12 assessing symptoms relative to anxiety); an Anxious Arousal factor (17 items), which is specific to anxious symptoms; and an Anhedonic Depression factor (22 items) specific to depression (Watson, Clark et al., 1995). A total MASQ score is generated by summing all items across the four subscales, which have high internal consistency across multiple samples (Watson, Weber et al., 1995). Items are rated on a 5-point scale ranging from 1 (not at all) to 5 (extremely). To comply with IRB protocol, one item on the Anhedonic Depression subscale relating to suicidality was removed.

The Emotion Regulation Questionnaire (ERQ) is a 10-item self-report questionnaire used to measure individual differences in two divergent emotion regulation strategies: Cognitive Reappraisal (6 items) and Expressive Suppression (4 items), (Gross & John, 2003). Items are rated on a 7-point scale from 1 (strongly disagree) to 7 (strongly agree).

**MRI Data Acquisition and Processing**

Each participant was scanned using one of two identical research-dedicated GE MR750 3T scanners at the Duke-UNC Brain Imaging and Analysis Center. Each identical scanner was equipped with high-power, high-duty cycle 50-mT/m gradients at 200 T/m/s slew rate and an eight-channel head coil for parallel imaging at high bandwidth up to 1 MHz. Following an ASSET calibration scan, two 2-min 50-s high angular resolution diffusion imaging acquisitions were collected, providing full brain coverage with 2-mm isotropic resolution and 15 diffusion-weighted directions (10-s repetition time, 84.9-ms echo time, b value 1,000 s/mm², 240-mm field of view, 90° flip angle, 128 × 128 acquisition matrix, slice thickness = 2 mm). High-resolution T1-weighted images were also obtained using a 3D Ax FSPGR BRAVO sequence with the following parameters: Repetition Time (TR) = 8.148 s; Echo Time (TE) = 3.22 ms; 162 sagittal slices; flip angle, 12°; Field of View (FOV), 240 mm; matrix = 256 × 256; slice thickness = 1 mm with no gap; and total scan time = 4 min and 13 s.

Diffusion tensor images were processed according to the protocol developed by the Enhancing Neuro Imaging Genetics Through Meta-Analysis consortium (Jahanshad et al., 2013). In brief, raw diffusion-weighted images underwent eddy current correction and linear registration to the non-diffusion-weighted image in order to correct for head motion. These images were skull-stripped and diffusion tensor models were fit at each voxel using the (Functional Magnetic Resonance Imaging of the Brain’s FMRIB’s) Diffusion Toolbox (http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/FDTI). This produced a whole-brain fractional anisotropy (FA) image for each participant, which was next processed using tract-based spatial statistics in FMRIB’s Software Library (Smith et al., 2006). FA
images were realigned to the FMRIB standard-space FA image and transformed into MN1 standard space. A mean FA skeleton was created and thresholded at .2, and each participant’s FA data were projected onto the skeleton. Regions of interest were then created using the Johns Hopkins University White Matter Tractography Atlas (Mori, Wakana, Van Zijl, & Nagae-Poetscher, 2005). The left and right UF regions were binarized and skeletonized in order to extract mean FA values from each hemisphere.

**Moderation Analysis**

PROCESS for SPSS (Hayes, 2013) was used within SPSS 21 (IBM Corp., Armonk, NY) to test whether average FA of the UF moderated the association between ERQ reappraisal and MASQ total scores (independent and dependent variables, respectively). Age, sex, and mean head motion were included as additional covariates to control for any potential confounds within the model.

**Results**

**Behavioral Measures**

Means and standard deviations for MASQ total score and the ERQ reappraisal subscale are as follows: MASQ total score: 109 ± 24.18, range = 60 and ERQ reappraisal: 5.23 ± 8.3, range = 5.33, n = 646; ERQ scores were missing for one participant. As expected, MASQ total and ERQ reappraisal were negatively correlated, $r(646) = -0.147, p < .001$. There were no significant sex differences for MASQ total scores; however, there were significant sex differences, $t(493.01) = -4.015, p < .001$, in ERQ reappraisal subscale scores, with women (5.34 ± .04) scoring higher than men (5.07 ± .06). ERQ suppression subscale scores were positively correlated with MASQ total scores, $r(646) = 0.308, p < .001$. There were significant sex differences, $t(598.84) = 4.48, p < .001$, in ERQ suppression subscale scores as well, with men (4.03 ± 1.05) scoring higher than women (3.63 ± 1.17).

Post hoc analyses revealed that the association between ERQ reappraisal and MASQ total scores was primarily driven by the two MASQ depression symptom subscales. Specifically, the Depression General Distress subscale, $r(646) = -0.14, p = .011$, and Anhedonic Depression subscale, $r(646) = -0.23, p < .001$, were both significantly correlated with ERQ reappraisal scores and survived correction for multiple comparisons. Neither the Anxiety General Distress subscale, $r(646) = -0.012, p = .762$, nor the Anxious Arousal subscale, $r(646) = 0.011, p = .782$, were significantly correlated with ERQ reappraisal scores.

**Moderation Analysis**

The overall model was significant in predicting MASQ total scores, $R = 0.22, F(5, 640) = 6.58, p < .001$. Additionally, a significant two-way interaction between average FA values of the UF and ERQ reappraisal scores predicted MASQ ($b = -151.01, 95\% CI [-249.03, -52.99], \Delta R^2 = .014, p = .026$; see Figure 2). Conditional effects of ERQ reappraisal scores on MASQ total scores at three separate levels (below −1 SD, between −1 SD and +1 SD, and above +1 SD) of average FA values of the UF are summarized in Figure 2. In contrast to these patterns with ERQ reappraisal scores, the interaction between ERQ suppression scores and average FA values of the UF did not significantly predict MASQ total scores ($b = -71.29, 95\% CI [-143.92, 1.35], \Delta R^2 = .0051, p = .55$). Post hoc analyses again showed that the significant two-way interaction between average UF FA values and ERQ reappraisal scores predicting MASQ scores was primarily driven by the two depression subscales ($b = -103.62, 95\% CI [-176.33, -30.9], \Delta R^2 = .012, p = .0053$). Additional analyses using the average FA of the cingulum bundle and the sagittal stratum, which were selected as control pathways, did not significantly interact with ERQ to predict MASQ symptoms. A post hoc analysis controlling for scanner revealed no significant difference in our results.

**Discussion**

Here, we provide initial evidence that the microstructural integrity of a pathway linking the dIPFC and amygdala moderates an association between typical use of cognitive reappraisal to downregulate negative emotions and the experience of anxiety and depressive symptoms. A significant negative correlation was observed between cognitive reappraisal and symptoms for individuals with relatively high and average but not low FA values in the UF. This pattern was specific to cognitive reappraisal as no significant moderation was observed for suppression, an alternative and less effective form of emotion regulation (Gross & John, 2003).

Our current findings unite prior research demonstrating that the microstructural integrity of the UF is associated with both depression and anxiety (Baur et al., 2012; Tronn et al., 2012), as well as cognitive reappraisal (Zuurbier et al., 2013). Moreover, our findings provide a possible mechanistic link between typical use of cognitive reappraisal and the experience of depressive and anxiety symptoms (Gross & John, 2003; Joormann & Gotlib, 2010; Sloan et al., 2017). Our findings are further consistent with recent work.
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2017).

It is important to note that our cross-sectional data cannot determine if higher microstructural integrity of the UF supports more typical use of cognitive reappraisal or vice versa. Longitudinal designs can address this question in future research, which can also employ strategies such as ecological momentary assessment to overcome the limitations of self-report. In addition, given that our DTI scans were limited to 15 directions, future studies could improve on the resolution of our findings by using tractography to define the UF pathway. Another limitation is that the ERQ assesses spontaneous or natural use of cognitive reappraisal in daily life (Gross, 1999). Thus, while the patterns observed herein suggest that the effectiveness of cognitive reappraisal in regulating the experience of depression and anxiety is, at least in part, shaped by the microstructural integrity of a pathway linking the dlPFC and amygdala, it is possible that specific targeting of cognitive reappraisal strategies within a clinical context are not similarly limited. Nevertheless, our findings suggest that FA values of the UF may represent a useful target in the search for biomarkers predicting not only disorder risk (e.g., Swartz, Knodt, Radtke, & Hariri, 2015) but also response to cognitive–behavioral therapy, which often emphasizes cognitive reappraisal of negative emotions (Sloan et al., 2017).

References


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