Behavioral/Cognitive

Anxiety and the Neurobiology of Temporally Uncertain Threat Anticipation

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When extreme, anxiety—a state of distress and arousal prototypically evoked by uncertain danger—can be debilitating. Uncertain anticipation is a shared feature of situations that elicit signs and symptoms of anxiety across psychiatric disorders, species, and assays. Despite the profound significance of anxiety for human health and wellbeing, the neurobiology of uncertain-threat anticipation remains unsettled. Leveraging a paradigm adapted from animal research and optimized for fMRI signal decomposition, we examined the neural circuits engaged during the anticipation of temporally uncertain and certain threat in 99 men and women. Results revealed that the neural systems recruited by uncertain and certain threat anticipation are anatomically colocalized in frontocortical regions, extended amygdala, and periaqueductal gray. Comparison of the threat conditions demonstrated that this circuitry can be fractionated, with frontocortical regions showing relatively stronger engagement during the anticipation of uncertain threat, and the extended amygdala showing the reverse pattern. Although there is widespread agreement that the bed nucleus of the stria terminalis and dorsal amygdala—the two major subdivisions of the extended amygdala—play a critical role in orchestrating adaptive responses to potential danger, their precise contributions to human anxiety have remained contentious. Follow-up analyses demonstrated that these regions show statistically indistinguishable responses to temporally uncertain and certain threat anticipation. These observations provide a framework for conceptualizing anxiety and fear, for understanding the functional neuroanatomy of threat anticipation in humans, and for accelerating the development of more effective intervention strategies for pathological anxiety.

Key words: affective neuroscience; anxiety and fear; bed nucleus of the stria terminalis; emotion; extended amygdala; Research Domain Criteria (RDoC)

Significance Statement

Anxiety—an emotion prototypically associated with the anticipation of uncertain harm—has profound significance for public health, yet the underlying neurobiology remains unclear. Leveraging a novel neuroimaging paradigm in a relatively large sample, we identify a core circuit responsive to both uncertain and certain threat anticipation, and show that this circuitry can be fractionated into subdivisions with a bias for one kind of threat or the other. The extended amygdala occupies center stage in neuropsychiatric models of anxiety, but its functional architecture has remained contentious. Here we demonstrate that its major subdivisions show statistically indistinguishable responses to temporally uncertain and certain threat. Collectively, these observations indicate the need to revise how we think about the neurobiology of anxiety and fear.

Received Mar. 26, 2020; revised July 31, 2020; accepted Aug. 5, 2020.

Author contributions: A.J.S., J.F.S., and K.A.D. designed research. J.F.S. contributed new software. A.J.S., A.S.F., H.C.K., J.F.S., J.H., J.K., K.A.D., and R.M.T. performed research. A.J.S., J.F.S., and J.H. analyzed data. A.J.S., A.S.F., H.C.K., J.F.S., J.H., and M.K. wrote the paper. A.J.S. funded and supervised the study. All authors edited the paper.

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This work was supported by the California National Primate Center; National Institutes of Health DAO40717, MH107444, and MH121409; University of California, Davis; and University of Maryland, College Park. Raw data are available at the National Institute of Mental Health's Data Archive. Key statistical maps are publicly available at www.NeuroVault.org. We thank the following for assistance and critical feedback:

A. Antonacci, M. Barstead, D. Bradford, J. Curtin, L. Friedman, J. Furcolo, C. Gorgolewski, C. Grubb, D. Holley, R. Hum, C. Kaplan, C. Lejuez, D. Limon, B. Nacewicz, S. Padmala, L. Pessoa, M. Roesch, S. Rose, J. Swayambunathan, and A. Vogel; members of the Affective and Translational Neuroscience laboratory; the staff of the Maryland Neuroimaging Center; the Office of the Registrar (University of Maryland); and two anonymous reviewers.

The authors declare no competing financial interests.

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https://doi.org/10.1523/JNEUROSCI.0704-20.2020 Copyright © 2020 the authors

Introduction

Anxiety is widely conceptualized as a state of distress and arousal elicited by the anticipation of uncertain danger (Davis et al., 2010; Grupe and Nitschke, 2013). Anxiety lies on a continuum and, when extreme, can be debilitating (Salomon et al., 2015; Conway et al., 2019). Anxiety disorders are the most common family of psychiatric illnesses, and existing treatments are inconsistently effective or associated with significant adverse effects, underscoring the urgency of developing a clearer understanding of the underlying neurobiology (Griebel and Holmes, 2013; Global Burden of Disease Collaborators, 2016; Craske et al., 2017).

Perturbation and recording studies in mice have begun to reveal the specific molecules and cellular ensembles that underlie defensive responses to uncertain threat (Fadok et al., 2018; Fox and Shackman, 2019), but the relevance of these tantalizing discoveries to the complexities of human anxiety is unclear. Humans and mice diverged ~75 MYA, leading to marked behavioral, genetic, and neurobiological differences between the two species (Van Essen et al., 2019). The role of frontocortical regions that are especially well-developed in humans—including the midcingulate cortex (MCC), anterior insula (AI), and dlPFC—remains particularly opaque, reflecting equivocal or absent anatomic homologies and the use of disparate paradigms across species (Vogt and Paxinos, 2014; Shackman et al., 2016; Carlén, 2017; Roberts, 2020).

Beneath the neocortex, the role of the central extended amygdala—including the dorsal amygdala in the region of the central nucleus (Ce) and the bed nucleus of the stria terminalis (BST) remains particularly contentious (Fox and Shackman, 2019). Inspired by an earlier generation of lesion studies in rodents (Davis, 2006), it is widely believed that these regions are functionally dissociable, with the amygdala mediating phasic responses to clear-and-immediate danger ("acute threat") and the BST mediating sustained responses to uncertain-or-remote danger ("potential threat") (e.g., Sylvers et al., 2011; Somerville et al., 2013; Avery et al., 2016; LeDoux and Pine, 2016; Klumpers et al., 2017; Watson et al., 2017). This "strict-segregation" hypothesis has even been enshrined in the National Institute of Mental Health's Research Domain Criteria framework (National Institute of Mental Health, 2011, 2020a,b). Yet a growing body of optogenetic, chemogenetic, and electrophysiological work in rodents demonstrates that defensive responses elicited by the anticipation of uncertain threat (e.g., elevated-plus maze) are assembled by microcircuits encompassing both regions (Gungor and Paré, 2016; Lange et al., 2017; Ahrens et al., 2018; Griessner et al., 2018; Pomrenze et al., 2019a,b; Ressler et al., 2020), motivating the competing hypothesis that the dorsal amygdala and BST are both important substrates for human anxiety (Shackman and Fox, 2016; Fox and Shackman, 2019).

To address these fundamental questions, we combined fMRI with a novel threat-anticipation task in 99 adults. Advanced data acquisition and processing techniques enhanced our ability to resolve small subcortical regions. Building on earlier work (e.g., Somerville et al., 2013; Grupe et al., 2016), the Maryland Threat Countdown (MTC) paradigm is an fMRI-optimized variant of assays that have been validated using fear-potentiated startle and acute pharmacological manipulations in rodents (Miles et al., 2011; Daldrup et al., 2015), and humans (Hefner et al., 2013), maximizing translational relevance. It takes the form of a 2 (Valence: Threat/Safety) × 2 (Temporal Certainty: Uncertain/Certain) randomized event-related design (Fig. 1). On Certain

Threat trials, subjects saw a descending stream of integers for 18.75 s, sufficiently long to enable the dissection of onset-evoked from sustained hemodynamic responses. To ensure robust emotion, this anticipatory epoch ("countdown") always culminated with the delivery of a multimodal reinforcer (aversive shock, photograph, and audio clip). Uncertain Threat trials were similar, but the integer stream was randomized and presented for an uncertain and variable duration (mean = 18.75 s, range = 8.75-30.00 s). Here, subjects knew the threat was coming, but they did not know when it would occur. Safety trials were similar but terminated in benign reinforcers. Comparison of the well-matched anticipatory epochs enabled us to rigorously isolate circuits recruited during uncertain-threat anticipation.

Materials and Methods

Subjects

As part of an ongoing prospective-longitudinal study focused on the emergence of anxiety disorders and depression, we used well-established measures of dispositional negativity (often termed neuroticism or negative emotionality) (Shackman et al., 2018; Hur et al., 2019a, 2019b) to screen 6594 young adults (57.1% female; 59.0% White, 19.0% Asian, 9.9% African American, 6.3% Hispanic, 5.8% Multiracial/Other; mean = 19.2 years, SD = 1.1 years). Screening data were stratified into quartiles (top quartile, middle quartiles, bottom quartile) separately for men and women. Individuals who met preliminary inclusion criteria were independently recruited from each of the resulting six strata. Given the focus of the larger study, approximately half the subjects were recruited from the top quartile, with the remainder split between the middle and bottom quartiles (i.e., 50% high, 25% medium, and 25% low), enabling us to sample a wide range of risk for the development of internalizing disorders. A total of 121 subjects were recruited. Of these, 2 withdrew during the imaging assessment because of excess distress. Of the 119 subjects who completed the imaging assessment, a total of 20 were excluded from analyses because of incidental neurological findings (n = 4), scanner problems (n = 2), insufficient fMRI data (<2 usable scans, n = 1), excessive global motion artifact (see below; n=3), or excessive taskcorrelated motion (see below, n = 10). This yielded a final sample of 99 subjects (52 females; 65.7% White, 17.2% Asian, 8.1% African American, 3.0% Hispanic, 6.1% Multiracial/Other; mean = 18.8 years, SD = 0.4 years), providing substantially greater power to detect medium-sized (0.5 < Cohen's d < 0.8) statistical effects (Geuter et al., 2018) compared with typical fMRI studies of uncertain threat anticipation (median N = 29; range = 15-108) (Chavanne and Robinson, 2020). All subjects had normal or corrected-to-normal color vision; and reported the absence of lifetime neurologic symptoms, pervasive developmental disorder, very premature birth, medical conditions that would contraindicate MRI, and prior experience with noxious electrical stimulation. All subjects were free from a lifetime history of psychotic and bipolar disorders; a current diagnosis of a mood, anxiety, or trauma disorder (past 2 months); severe substance abuse; active suicidality; and ongoing psychiatric treatment as determined by an experienced masters-level diagnostician using the Structured Clinical Interview for DSM-5 (First et al., 2015). Subjects provided informed written consent, and all procedures were approved by the Institutional Review Board at the University of Maryland, College Park.

MTC fMRI paradigm

Paradigm structure and design considerations. Building on earlier imaging work (Somerville et al., 2013; Grupe et al., 2016; Pedersen et al., 2019), the MTC paradigm is an fMRI-optimized version of temporally uncertain-threat assays that have been validated using fear-potentiated startle and acute anxiolytic administration (e.g., benzodiazepine) in mice (Daldrup et al., 2015; Lange et al., 2017), rats (Miles et al., 2011), and humans (Hefner et al., 2013), enhancing its translational relevance. The MTC paradigm takes the form of a 2 (Valence: Threat/Safety) × 2 (Temporal Certainty: Uncertain/Certain) randomized event-related

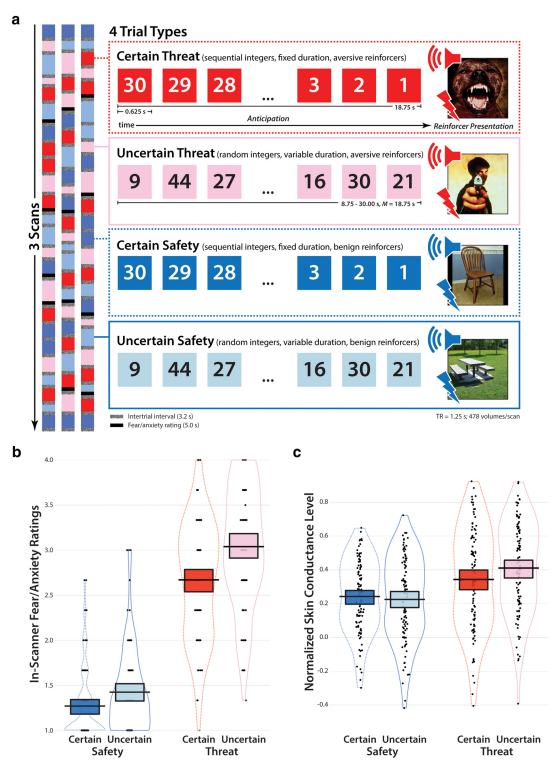


Figure 1. MTC paradigm. As shown schematically in *a*, the MTC paradigm takes the form of a 2 (Valence: Threat/Safety) × 2 (Temporal Certainty: Uncertain/Certain) repeated-measures design (for details, see Materials and Methods). Subjects provided ratings of anticipatory fear/anxiety for each trial type during each scan. Skin conductance was continuously acquired during scanning. Simulations were used to optimize the detection and deconvolution of task-related hemodynamic signals (variance inflation factors <1.54). Middle panels, Structure of each trial type. Trial valence was continuously signaled during the anticipatory epoch by the background color of the display. Safety trials were similar but terminated with the delivery of benign stimuli (e.g., just-perceptible electrical stimulation). Trial certainty was signaled by the nature of the integer stream. Certain trials always began with the presentation of 30. On Uncertain trials, integers were randomly drawn from a uniform distribution ranging from 1 to 45 to reinforce the belief that uncertain trials could be much longer than certain ones. To mitigate potential confusion and eliminate mnemonic demands, a lowercase 'c' or 'u' was presented at the lower edge of the display throughout the anticipatory epoch (not depicted). As shown in *b* and *c*, threat anticipation robustly increased subjective symptoms (in-scanner ratings) and objective signs (skin conductance) of anxiety, and this was particularly evident when the timing of aversive stimulation was uncertain (Valence × Certainty, *p* values < 0.001; Uncertain Threat > Certain Threat, *p* values < 0.001). *b*, *c*, Data (black points; individual participants), density distribution (bean plots), Bayesian 95% highest density interval (colored bands), and mean (black bars) for each condition. Highest density intervals permit population-generalizable visual inferences about mean differences and were estimated using 1000 samples from a posterior Gaussian distribution. TR, the time required to collect a single

design (3 scans; 6 trials/condition/scan). Simulations were used to optimize the detection and deconvolution of task-related hemodynamic signals (variance inflation factors <1.54). Stimulus presentation and ratings acquisition were controlled using Presentation software (version 19.0, Neurobehavioral Systems).

On Certain Threat trials, subjects saw a descending stream of integers (countdown; e.g., 30, 29, 28...3, 2, 1) for 18.75 s. To ensure robust emotion, this anticipatory epoch always culminated with the delivery of a noxious electric shock, unpleasant photographic image (e.g., mutilated body), and thematically related audio clip (e.g., scream, gunshot). Uncertain Threat trials were similar, but the integer stream was randomized and presented for an uncertain and variable duration (8.75-30.00 s; mean = 18.75 s). Here, subjects knew that something aversive was going to occur, but they had no way of knowing precisely when it would occur. Consistent with recent recommendations (Shackman and Fox, 2016), the average duration of the anticipatory epoch was identical across conditions, ensuring an equal number of measurements (TRs/condition). Mean duration was chosen to enhance detection of task-related differences in the BOLD signal (Henson, 2007), and to enable dissection of onset from genuinely sustained responses. Safety trials were similar but terminated with the delivery of benign reinforcers (see below). Valence was continuously signaled during the anticipatory epoch by the background color of the display. Temporal certainty was signaled by the nature of the integer stream. Certain trials always began with the presentation of the number 30 (Fig. 1). On uncertain trials, integers were randomly drawn from a near-uniform distribution ranging from 1 to 45 to reinforce the impression that Uncertain trials could be much longer than Certain ones and to minimize incidental temporal learning (time-keeping). To mitigate potential confusion and eliminate mnemonic demands, a lowercase 'c' or 'u' was presented at the lower edge of the display throughout the anticipatory epoch. White-noise visual masks (3.2 s) were presented between trials to minimize persistence of the visual reinforcers in iconic memory. Subjects provided ratings of anticipatory fear/anxiety for each trial type during each scan using an MRI-compatible response pad (MRA; Fig. 1). Subjects were instructed to rate the intensity of the fear/ anxiety experienced during the prior anticipatory (countdown) epoch using a 1 (minimal) to 4 (maximal) scale. Subjects were prompted to rate each trial type once per scan. A total of 6 additional EPI volumes were acquired at the beginning and end of each scan (see below).

Procedures. Before scanning, subjects practiced an abbreviated version of the paradigm, without electrical stimulation, until they indicated and staff confirmed that they understood the task. Benign and aversive electrical stimulation levels were individually titrated. Benign stimulation: Subjects were asked whether they could "reliably detect" a 20 V stimulus and whether it was "at all unpleasant." If the subject could not detect the stimulus, the voltage was increased by 4 V and the process repeated. If the subject indicated that the stimulus was unpleasant, the voltage was reduced by 4 V and the process repeated. The final level chosen served as the benign electrical stimulation during the imaging assessment (mean = 20.67, SD = 6.23). Aversive stimulation: Subjects received a 100 V stimulus and were asked whether it was "as unpleasant as you are willing to tolerate." If the subject indicated that they were willing to tolerate more intense stimulation, the voltage was increased by 10 V and the process repeated. If the subject indicated that the stimulus was too intense, the voltage was reduced by 5 V and the process repeated. The final level chosen served as the aversive electrical stimulation during the imaging assessment (mean = 115.21, SD = 25.05). Following each scan of the MTC paradigm, we reassessed whether stimulation was sufficiently intense and recalibrated as necessary. In total, 32.3% of subjects adjusted the level of benign or aversive stimulation at least once during the imaging assessment.

Electrical stimuli. Electrical stimuli (100 ms; 2 ms pulses every 10 ms) were generated using an MRI-compatible constant-voltage stimulator system (STMEPM-MRI; Biopac Systems). Stimuli were delivered using MRI-compatible, disposable carbon electrodes (Biopac) attached to the fourth and fifth phalanges of the non-dominant hand.

Visual stimuli. Visual stimuli (1.8 s) were digitally back-projected (Powerlite Pro G5550, Epson America) onto a semi-opaque screen mounted at the head-end of the scanner bore and viewed using a mirror

mounted on the head-coil. A total of 72 photographs were selected from the International Affective Picture System (IAPS identification numbers), Benign: 1670, 2026, 2038, 2102, 2190, 2381, 2393, 2397, 2411, 2850, 2870, 2890, 5390, 5471, 5510, 5740, 7000, 7003, 7004, 7014, 7020, 7026, 7032, 7035, 7050, 7059, 7080, 7090, 7100, 7140, 7187, 7217, 7233, 7235, 7300, 7950. Aversive: 1300, 3000, 3001, 3010, 3015, 3030, 3051, 3053, 3061, 3062, 3063, 3069, 3100, 3102, 3150, 3168, 3170, 3213, 3400, 3500, 6022, 6250, 6312, 6540, 8230, 9042, 9140, 9253, 9300, 9405, 9410, 9414, 9490, 9570, 9584, 9590 (Lang et al., 2008). Based on normative ratings, the aversive images were significantly more negative and arousing than the benign images, $t_{(70)} > 24.3, \, p < 0.001$. On a 1 (negative/low-arousal) to 9 (positive/high-arousal) scale, the mean valence and arousal scores were 2.2 (SD = 0.6) and 6.3 (SD = 0.6) for the aversive images, and 5.2 (SD = 0.4) and 2.8 (SD = 0.3) for the benign images.

Auditory stimuli. Auditory stimuli (0.80 s) were delivered using an amplifier (PA-1 Whirlwind) with inline noise-reducing filters and ear buds (S14; Sensimetrics) fitted with noise-reducing ear plugs (Hearing Components). A total of 72 auditory stimuli (half aversive, half benign) were adapted from open-access online sources.

Peripheral physiology data acquisition

Peripheral physiology was continuously acquired during each fMRI scan using a Biopac system (MP-150). Skin conductance (250 Hz; 0.05 Hz high-pass) was measured using MRI-compatible disposable electrodes (EL507) attached to the second and third phalanges of the nondominant hand. For imaging analyses, measures of respiration and breathing were also acquired using a respiration belt and photo-plethysmograph placed on the first phalange of the non-dominant hand.

MRI data acquisition

MRI data were acquired using a Siemens Magnetom TIM Trio 3 Tesla scanner (32-channel head-coil). Foam inserts were used to immobilize the participant's head within the head-coil and mitigate potential motion artifact. Subjects were continuously monitored from the control room using an MRI-compatible eye-tracker (Eyelink 1000; SR Research). Head motion was monitored using the AFNI real-time plugin (Cox, 1996). Sagittal T1-weighted anatomic images were acquired using a MPRAGE sequence (TR = 2400 ms; TE = 2.01 ms; inversion time = 1060 ms; flip angle = 8° ; sagittal slice thickness = 0.8 mm; in-plane = 0.8×0.8 mm; matrix = 300×320 ; FOV = 240×256). A T2-weighted image was collected coplanar to the T1-weighted image (TR = 3200 ms; TE = 564 ms; flip angle = 120°). To enhance resolution, a multiband sequence was used to collect oblique-axial EPI volumes (multiband acceleration = 6; TR = 1250 ms; TE = 39.4 ms; flip angle = 36.4° ; slice thickness = 2.2 mm, number of slices = 60; in-plane resolution = 2.1875×2.1875 mm; matrix = 96 \times 96). Images were collected in the oblique axial plane (\sim -20° relative to the AC-PC plane) to minimize potential susceptibility artifacts. Three 478-volume EPI scans were acquired. The scanner automatically discarded 7 volumes before the first recorded volume. To enable fieldmap correction, two oblique-axial spin echo (SE) images were collected in each of two opposing phase-encoding directions (rostral-to-caudal and caudal-to-rostral) at the same location and resolution as the functional volumes (i.e., coplanar; TR = 7220 ms; TE = 73 ms). Following the last scan, subjects were removed from the scanner, debriefed, compensated, and discharged.

Skin conductance data pipeline

Skin conductance data were processed using PsPM (version 4.0.2) and in-house MATLAB code (Bach and Friston, 2013; Bach et al., 2018). Data from each scan were bandpass filtered (0.01-0.25 Hz), resampled to match the TR used for fMRI data acquisition (1.25 s), and z-transformed. Using standard MATLAB functions, SCR data were modeled in a manner that approximated that used for the fMRI data. A GLM was used to estimate skin conductance levels during the anticipatory epoch of each condition of the MTC paradigm (see above) for each subject (Bach et al., 2009, 2013; Bach, 2014). Predictors from the first-level fMRI model (see below) were convolved with a canonical skin conductance response function (Bach et al., 2010; Gerster et al., 2018), bandpass filtered to match the data, and z-transformed.

MRI data pipeline

Methods were optimized to minimize spatial normalization error and other potential sources of noise. Structural MRI and fMRI data were visually inspected before and after processing for quality assurance.

Anatomical data processing

Methods were similar to those described in other recent reports by our group (Hur et al., 2018; Smith et al., 2018; Tillman et al., 2018). T1weighted images were inhomogeneity-corrected using N4 (Tustison et al., 2010) and filtered using the denoise function in ANTS (Avants et al., 2011). The brain was then extracted using a variant of the BEaST algorithm (Eskildsen et al., 2012) with brain-extracted and normalized reference brains from the IXI database (https://brain-development.org/ ixi-dataset). Brain-extracted T1 images were normalized to a version of the brain-extracted 1 mm T1-weighted MNI152 (version 6) template (Grabner et al., 2006), modified to remove extracerebral tissue. This was motivated by evidence that brain-extracted T1 images and a brainextracted template enhance the quality of spatial normalization (Fein et al., 2006; Acosta-Cabronero et al., 2008; Fischmeister et al., 2013). Normalization was performed using the diffeomorphic approach implemented in SyN (version 1.9.x0.2017-09.11; Klein et al., 2009; Avants et al., 2011). T2-weighted images were rigidly coregistered with the corresponding T1 before normalization, and the brain extraction mask from the T1 was applied. Tissue priors (Lorio et al., 2016) were unwarped to the native space of each T1 using the inverse of the diffeomorphic transformation. Brain-extracted T1 and T2 images were simultaneously segmented using native-space priors generated using FAST (FSL version 5.0.9) (Zhang et al., 2001) for use in T1-EPI coregistration (see below).

Fieldmap data processing

SE images were used to create a fieldmap in topup (Andersson et al., 2003; Smith et al., 2004; Graham et al., 2017). Fieldmaps were converted to radians, median filtered, and smoothed (2 mm). The average of the distortion-corrected SE images was inhomogeneity-corrected using N4, and brain-masked using 3dSkullStrip in AFNI (version 17.2.10) (Cox, 1996). The resulting mask was minimally eroded to further exclude extracerebral voxels.

Functional data processing

EPI files were despiked using 3dDespike and slice-time-corrected (to the center of the TR) using 3dTshift, inhomogeneity corrected using N4, and motion-corrected to the first volume using a 12-parameter affine transformation implemented in ANTs. Recent work indicates that despiking is more effective than "scrubbing" for attenuating motion-related artifacts (Jo et al., 2013; Siegel et al., 2014; Power et al., 2015). Transformations were saved in ITK-compatible format for subsequent use. The first volume was extracted for EPI-T1 coregistration. The reference EPI volume was simultaneously coregistered with the corresponding T1-weighted image in native space and corrected for geometric distortions using boundary-based registration (Greve and Fischl, 2009). This step incorporated the previously created fieldmap, undistorted SE, T1, white matter image, and masks. The spatial transformations necessary to transform each EPI volume from native space to the reference EPI, from the reference EPI to the T1, and from the T1 to the template were concatenated and applied to the processed (despiked and slicetime-corrected) EPI data in a single step to minimize incidental spatial blurring. Normalized EPI data were resampled to 2 mm isotopic voxels using fifth-order b-splines and smoothed (6 mm FWHM) using 3DblurInMask.

Data exclusions

To assess residual motion artifact, we computed the number of times the brain showed a volume-to-volume displacement >0.5 mm using the motion-corrected data. Scans with excess artifact ($\geq 7.5\%$) were discarded. Three subjects with insufficient usable data (<2 scans) were excluded from analyses, while 6 subjects with 2 usable scans were retained. To assess task-correlated motion, we computed correlations between the design matrix and the motion estimates (see above). Scans showing extreme correlations (>2 SD) were discarded. On this basis, 10

subjects with insufficient usable data (<2 scans) were excluded from analyses, while 19 subjects with 2 usable scans were retained.

Canonical first-level fMRI modeling

Modeling was performed using SPM12 (version 6678; https://www.fil. ion.ucl.ac.uk/spm). Bandpass was set to the HRF and 128 s for low and high pass, respectively. The MTC paradigm was modeled using variableduration rectangular (boxcar) regressors time-locked to the anticipatory epochs of the Uncertain Threat, Certain Threat, and Uncertain Safety trials. Certain Safety trials were treated as an unmodeled (implicit) highlevel baseline. EPI volumes collected before the first trial, during intertrial intervals, and following the final trial were also unmodeled, and contributed to the baseline estimate. Regressors were convolved with a canonical HRF and its temporal derivative. The periods corresponding to the delivery of the four reinforcers and rating trials were modeled using a similar approach (Fig. 1). Volume-to-volume displacement and motion parameters (including 1- and 2-volume lagged versions) were also included, similar to other recent work (Reddan et al., 2018). To further attenuate potential noise, CSF time-series, instantaneous pulse and respiration rates, and their estimated effect on the BOLD time-series were also included as nuisance variates. ICA-AROMA (Pruim et al., 2015) was used to model several other potential sources of noise (brainedge, CSF-edge, white matter). These and the single ICA component showing the strongest correlation with motion estimates were included as additional nuisance variates. EPI volumes with excessive volume-tovolume displacement (>0.25 mm), as well as those during and immediately following the delivery of aversive reinforcers, were censored.

Decomposing canonical effects using finite impulse response (FIR) modeling

The canonical modeling approach estimates the amplitude of anticipatory activity under the assumption that it approximates a boxcar-like square-wave function. This makes it tempting to conclude that regions showing significant activation represent sustained responses. Yet there is ample evidence that a variety of other signals are plausible (e.g., Gonzalez-Castillo et al., 2015; Gungor and Paré, 2016; Sreenivasan and D'Esposito, 2019) and, importantly, can yield similarly strong statistical effects (Fig. 2). Addressing this ambiguity necessitates a finer decomposition of the signal underlying significant "omnibus" effects revealed by canonical modeling, a surprisingly rare approach in the neuroimaging literature. To do so, we identified the most extreme peak (e.g., BST) in each of the major regions identified in our canonical analyses. These peak locations were then interrogated using a FIR analysis, which provides an estimate of the magnitude and shape of anticipatory activity (Glover, 1999; Ollinger et al., 2001; for a similar approach by our group, see Guller et al., 2012). To perform the FIR modeling, variance related to reinforcer delivery, ratings, and the nuisance variables was removed from the preprocessed data using the canonical approach described above. Residualized data were bandpass filtered (0.007813-0.2667 Hz) and normalized to the SD of the Certain Safety trials (i.e., the implicit baseline in the canonical HRF GLM). We then estimated the mean response at each TR of the anticipatory epoch for each condition of the MTC paradigm for each subject.

Experimental design and statistical analyses

Overview. Study design is described in MTC fMRI paradigm. The number of usable datasets, data exclusions, and power considerations are detailed in Subjects.

In-scanner fear/anxiety ratings and skin conductance. Data were analyzed using standard repeated-measures GLM approaches with Huynh-Feldt correction for potential nonsphericity implemented in SPSS (version 24; IBM). Significant interactions were decomposed using simple effects. Figures were created using R Studio (http://www.rstudio.com) and yarrr (version 0.1.5) for R (version 3.6.1.; https://www.R-project.org).

Canonical second-level GLM. Standard whole-brain voxelwise GLMs (random effects) were used to compare anticipatory hemodynamic activity elicited by each threat-anticipation condition and its corresponding control condition (e.g., Uncertain Threat vs Uncertain Safety). Sig-

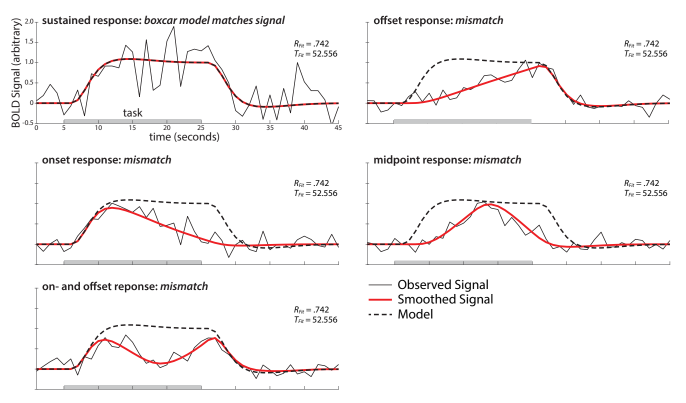


Figure 2. Interpretive ambiguities of canonical HRF modeling. The canonical approach to fMRI analysis models the amplitude of anticipatory activity (solid black line) under the assumption that it approximates a boxcar-like square-wave shape (dotted line; convolution of a canonical HRF with task duration). In some cases, such as the top left panel, the hemodynamic signal and the model will match. But in others, it will not. Importantly, a variety of physiologically plausible hemodynamic responses can produce similarly strong and statistically significant results (T = 52.556 in this example), highlighting the importance of modeling the BOLD signal on a finer temporal scale.

nificance was assessed using FDR q < 0.05, whole-brain-corrected. As in prior work by our group (Shackman et al., 2013; Shackman et al., 2017), a minimum conjunction (logical "AND") was used to identify voxels sensitive to both temporally certain and temporally uncertain threat anticipation (Nichols et al., 2005). We also directly examined potential differences in anticipatory activity between the two threat conditions (Certain Threat vs Uncertain Threat). We did not examine hemodynamic responses to reinforcer delivery given the possibility of artifact. Some figures were created using MRIcron (http://people.cas.sc.edu/rorden/mricron). Clusters and local maxima were labeled using a combination of the Allen Institute, Harvard-Oxford, and Mai atlases (Frazier et al., 2005; Desikan et al., 2006; Makris et al., 2006; Hawrylycz et al., 2012; Mai et al., 2015) and a recently established consensus nomenclature (ten Donkelaar et al., 2018).

Descriptive decomposition of canonical effects using FIR modeling To decompose the signal underlying significant canonical effects, we identified the most extreme peak in each of the major regions (e.g., amygdala) identified in our canonical analyses (indicated by a blackand-white asterisk in the accompanying figures). These peaks were then descriptively interrogated using FIR models. As shown in Figure 1, the duration of anticipatory epoch differed between certain (18.75 s) and uncertain trials (8.75-32.5 s; mean = 18.75 s), necessitating slightly different procedures for specific contrasts. For the comparison of Certain Threat to Certain Safety, responses were modeled for 15 TRs (1.25 s TR; total = 18.125 s). Given the temporal resolution and autocorrelation of the BOLD signal, data were averaged for 3 windows (TR-1 to TR-5, TR-6 to TR-10, TR-11 to TR-15). For the comparison of Uncertain Threat to Uncertain Safety, responses were modeled for 24 TRs (total = 30.00 s) and averaged for four windows, the first three corresponding to those used for certain trials and a fourth spanning TR-16 to TR-24. This choice was partially motivated by the modest number of trials with the longest anticipatory epoch (Fig. 1). For the comparison of the two threat conditions, responses were modeled for 15 TRs and averaged across three windows, as above. "Sustained" activity was operationally defined as greater mean activity across two consecutive windows. Using this criterion, descriptive tests indicated nominally significant (p < 0.05) evidence of sustained responses for most of the key regions for most of the contrasts (e.g., Uncertain Threat vs Uncertain Safety). Exceptions were the periaqueductal gray (PAG) for the Certain Threat vs Certain Safety contrast, and the PAG and BST for the Certain Threat versus Uncertain Threat contrast. Because this approach yields optimistically biased effect-size estimates (Kriegeskorte et al., 2010; Davenport and Nichols, 2020), we refrain from reporting exact p values, and instead provide the SEM as a descriptive guide to the size of observed effects. Naturally, any inferences drawn from inspection of the standard errors only apply to the peak voxels depicted in the accompanying figures (black-and-white asterisks) and not necessarily to the entire parent region (e.g., amygdala).

Testing whether the BST and amygdala show different responses to threat

To test hypothesized regional differences in threat sensitivity (see the Introduction), we used a combination of anatomic and functional criteria to independently identify BST and amygdala voxels that were most sensitive to each kind of threat. As shown in Figure 3, each region was anatomically defined using a priori probabilistic ROIs (Frazier et al., 2005; Desikan et al., 2006; Theiss et al., 2017). Next, we extracted and averaged standardized regression coefficients for voxels showing significant (FDR q < 0.05, whole-brain-corrected) activation for each of the relevant contrasts, separately for each region: Uncertain Threat > Uncertain Safety, Certain Threat > Certain Safety, and Certain Threat > Uncertain Threat. Potential regional differences (i.e., Region × Condition interactions) were then assessed using standard repeated-measures GLM approaches implemented in SPSS. Nonparametric tests (Wilcoxon signed rank) yielded identical conclusions (not reported). Inferences

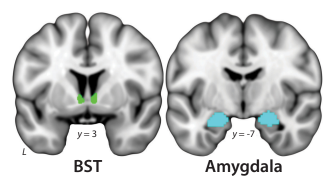


Figure 3. Amygdala and BST ROIs. BST: The probabilistic BST ROI (green) was described by Theiss et al. (2017) and was thresholded at 0%. The seed mostly encompasses the supracommissural BST, given the difficulty of reliably discriminating the borders of regions below the anterior commissure using T1-weighted images (Kruger et al., 2015). Amygdala: The Harvard-Oxford probabilistic amygdala (cyan) was described by Frazier et al. (2005) and Desikan et al. (2006) and conservatively thresholded at 50%. Analyses used ROIs decimated to the 2 mm resolution of the EPI data. For illustrative purposes, 1 mm ROIs are shown. Single-subject data were visually inspected to ensure that the ROIs were correctly aligned to the spatially normalized T1-weighted images.

necessarily apply only to the subset of BST or amygdala voxels that proved sensitive to one or more of the threat contrasts, not the entire region.

Testing whether the BST and amygdala show equivalent responses to threat

Consistent with prior work by our group (McMenamin et al., 2009, 2010), we used the well-established two one-sided tests (TOST) procedure for formally testing whether the BST and amygdala show statistically equivalent activity during threat anticipation. While it is not possible to definitively show that the true difference in regional activity is zero, TOST provides a well-established and widely used framework for testing whether mean differences are small enough to be considered equivalent (Lakens, 2017; Lakens et al., 2018). Regression coefficients were extracted and averaged using the approach detailed in the prior section. For present purposes, we considered regional differences smaller than a "medium" standardized effect for dependent means (Cohen's d_z = 0.35) to be equivalent (Lakens, 2017). TOST procedures were performed using the TOSTER package (version 0.3.4) for R.

Results

Temporally uncertain threat anticipation elicits robust symptoms and signs of anxiety

As shown in Figure 1, threat anticipation markedly increased subjective symptoms (in-scanner ratings) and objective signs (skin conductance) of anxiety, and this was particularly evident when the timing of aversive stimulation was uncertain. Anticipatory feelings: Subjects reported experiencing more intense fear/anxiety when anticipating aversive outcomes ($F_{(1,98)} = 543.27$, p < 0.001), and when anticipating outcomes with uncertain timing ($F_{(1.98)}$ = 85.46, p < 0.001). The impact of threat on fear/anxiety ratings was potentiated by temporal uncertainty (Valence × Uncertainty: $F_{(1.98)} = 13.08$, p < 0.001; Uncertain Threat > Certain Threat: $t_{(98)} = 7.58$, p < 0.001; Uncertain Safety > Certain Safety: $t_{(98)} =$ 4.90, p < 0.001; Uncertain Threat > Uncertain Safety: $t_{(98)} = 21.98$, p < 0.001; Certain Threat > Certain Safety: $t_{(98)} = 20.36$, p < 0.0010.001), consistent with prior work (Grillon et al., 2006; Nelson and Shankman, 2011; Somerville et al., 2013; Bennett et al., 2018). Anticipatory arousal: Subjects showed elevated skin conductance levels when anticipating aversive outcomes ($F_{(1.98)} = 345.31$, p < 0.001), and when anticipating outcomes with uncertain timing $(F_{(1.98)} = 85.86, p < 0.001)$. The impact of threat on skin conductance was potentiated by temporal uncertainty (Valence ×

Uncertainty: $F_{(1,98)} = 93.63$, p < 0.001; Uncertain Threat > Certain Threat: $t_{(98)} = 11.53$, p < 0.001; Uncertain Safety > Certain Safety: $t_{(98)} = -3.99$, p < 0.001; Uncertain Threat > Uncertain Safety: $t_{(98)} = 25.59$, p < 0.001; Certain Threat > Certain Safety: $t_{(98)} = 9.84$, p < 0.001). Together, these results confirm the validity of the MTC paradigm for understanding the neural circuits underpinning human anxiety.

Temporally uncertain threat anticipation recruits a distributed network of subcortical and cortical regions

Next, a voxelwise GLM was used to identify brain regions recruited during the anticipation of temporally Uncertain Threat (Uncertain Threat > Uncertain Safety; FDR q < 0.05, wholebrain-corrected). As shown in Figure 4, this highlighted a widely distributed network of regions previously implicated in the expression and regulation of human fear and anxiety (Fullana et al., 2016; Qi et al., 2018; Fox and Shackman, 2019; Chavanne and Robinson, 2020), including the MCC; AI extending into the frontal operculum (FrO); dlPFC extending to the frontal pole (FP); brainstem encompassing the PAG; basal forebrain, in the region of the BST; and dorsal amygdala, in the region of the central and medial nuclei. Heightened activity during the anticipation of Uncertain Threat was also evident in the orbitofrontal cortex, basal ganglia, hippocampus, and ventrolateral amygdala in the region of the lateral nucleus (Extended Data Fig. 4-1). Consistent with prior work (Choi et al., 2012; Grupe et al., 2016), Uncertain Threat anticipation was associated with reduced activity in a set of midline regions that resembled the default mode network (e.g., anterior rostral sulcus/ventromedial PFC, postcentral gyrus, and precuneus), as well as the posterior insula and parahippocampal gyrus (Extended Data Fig. 4-2). Reduced activity was also observed in the most rostral tip of the amygdala, underscoring the functional heterogeneity of this complex structure.

Temporally uncertain threat anticipation elicits sustained hemodynamic activity

Anxiety is widely conceptualized as a sustained state (Davis et al., 2010; Tye et al., 2011; LeDoux and Pine, 2016; Mobbs, 2018), and it is tempting to interpret clusters of enhanced activity (e.g., Fig. 4) through this lens. But do we actually see evidence of sustained responses during the anticipation of temporally Uncertain Threat? Although a wide variety of other signals are physiologically plausible (Fig. 2), the vast majority of fMRI studies never address this question; they simply assume the shape of the hemodynamic response and focus on estimates of response magnitude ("activation"). To address this ambiguity, we used an FIR approach to estimate responses elicited by the anticipation of Uncertain Threat and Uncertain Safety on a finer time-scale. Descriptively, this revealed sustained activity (see Materials and Methods) across key cortical (MCC, AI/FrO, dlPFC/FP) and subcortical (PAG, BST, dorsal amygdala) regions (Uncertain Threat > Uncertain Safety; 6.25-30 s; Fig. 5).

Temporally certain threat anticipation recruits an anatomically and functionally similar network

Having identified a distributed neural circuit sensitive to Uncertain Threat, we used a parallel approach to identify regions recruited during the anticipation of temporally Certain Threat (Certain Threat > Certain Safety; FDR q < 0.05, whole-brain-corrected). As shown in Figure 4, results were similar to those found for Uncertain Threat (Extended Data Figs. 4-3, 4-4). In fact, a minimum conjunction analysis (Logical "AND") (Nichols

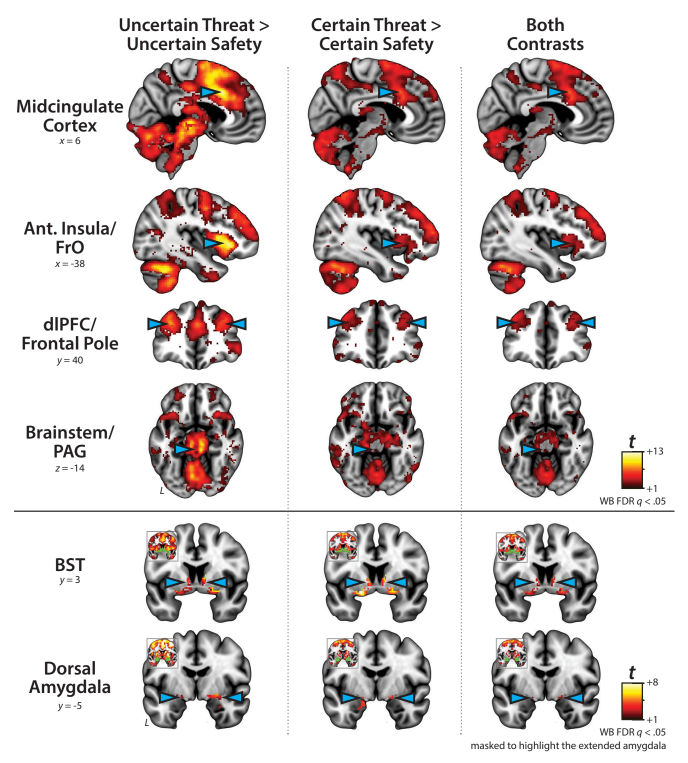


Figure 4. The anticipation of temporally Uncertain and Certain Threat recruits broadly similar neural systems. Key regions (cyan arrowheads) showing significantly elevated activity during the anticipation of Uncertain Threat (left column) and Certain Threat (middle column) compared with their respective control conditions. Right column, Voxels showing significantly increased activity in both contrasts. BST and dorsal amygdala images are masked to highlight significant voxels in extended amygdala (green). Coronal insets, The thresholded statistical parametric maps without the additional mask. Together, these observations indicate that these regions are sensitive to both temporally certain and uncertain threat. For additional details, see Extended Data Figures 4-1, 4-2, 4-3, 4-4, 4-5. Ant., Anterior; FDR, false discovery rate; WB, whole-brain-corrected.

et al., 2005) revealed voxelwise colocalization in every key cortical and subcortical region, including the BST and dorsal amygdala in the region of the central and medial nuclei (Fig. 4; Extended Data Fig. 4-5). FIR results also suggested functional convergence across conditions, with all but one of these key

regions (PAG) showing sustained levels of heightened hemodynamic activity during the anticipation of Certain Threat (see Materials and Methods; Fig. 6). Together, these results suggest that this network of subcortical and cortical regions is sensitive to multiple kinds of threat anticipation, both certain and uncertain.

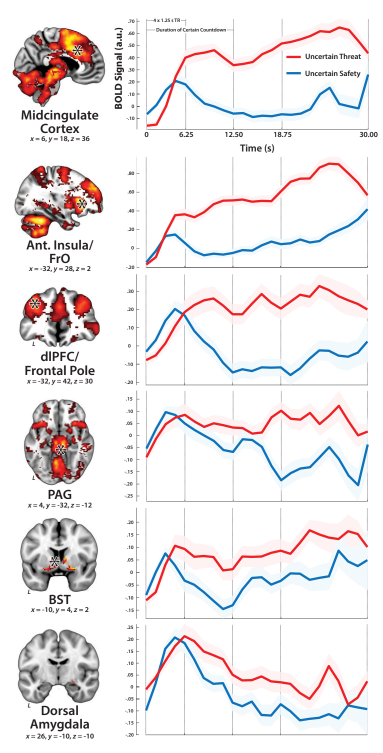


Figure 5. Regions sensitive to temporally Uncertain Threat show sustained hemodynamic activity. Mean responses to the anticipatory epoch were estimated on a TR-by-TR (1.25 s) basis for Uncertain Threat (red) and Uncertain Safety (blue) trials, using data from the local maxima of key clusters (black-and-white asterisks in the left panels) identified using a canonical analytic approach. Given the temporal resolution and autocorrelation of the hemodynamic signal, data were averaged for 4 windows (TR-1 to TR-5, TR-6 to TR-10, TR-11 to TR-15, and TR-16 to TR-24), spanning a total of 24 measurements (30 s). Broken vertical lines indicate windows. Shaded envelopes represent SEM. Ant., Anterior; TR, the time needed to acquire a single volume of fMRI data.

The threat anticipation network can be fractionated into subdivisions

To determine whether regions recruited during threat anticipation are sensitive to temporal uncertainty, we directly compared the Uncertain and Certain Threat conditions (FDR q < 0.05,

whole-brain-corrected). This indicated that the threat anticipation network can be fractionated into subdivisions. As shown in Figure 7, key cortical regions (MCC, AI/FrO, and dlPFC/FP) showed a further increase in activity during the anticipation of Uncertain Threat (Extended Data Fig. 7-1). In contrast, the BST and dorsal amygdala (adjacent to the Ce, in the region of the cortical nucleus and amygdala-hippocampal transition area) showed the reverse pattern, with relatively greater activity during the anticipation of Certain Threat (Extended Data Fig. 7-2). The PAG did not discriminate the two threat conditions. FIR results suggest a similar conclusion (Fig. 7).

The anticipation of temporally uncertain and certain threat elicits statistically indistinguishable responses in the extended amygdala

Our results indicate that the BST and dorsal amygdala—the two major subdivisions of the extended amygdala—respond similarly to threat anticipation. Both regions show signs of elevated activity during threat anticipation, and this is evident whether or not the timing of aversive stimulation is uncertain (Fig. 4). Furthermore, both regions showed parallel increases in activity during the anticipation of Certain Threat (Figs. 6, 7). Yet it remains possible that the BST and the amygdala exhibit subtler differences in threat sensitivity. To rigorously test this, we directly compared regional responses for each of the threat contrasts (e.g., Uncertain Threat vs Uncertain Safety), equivalent to testing the Region × Condition interactions. As shown in Figure 8, mean differences were small to very small ($d_Z < 0.17$) and all nonsignificant (Extended Data Fig. 8-1). Likewise, the proportion of subjects showing numerically greater activity in one region or the other never exceeded 55% (Fig. 8). Naturally, these results do not license strong claims of regional equivalence. While it is impossible to demonstrate that the true difference in regional activity is zero, the TOST procedure provides a well-established and widely used framework for testing whether mean differences-here, in regional activity-are small enough to be considered statistically equivalent (Lakens, 2017; Lakens et al., 2018). For present purposes, we considered differences smaller than a "medium"

standardized effect (Cohen's $d_z = 0.35$) to be statistically equivalent. Using the voxels that were most sensitive to each threat contrast (see Materials and Methods), our results revealed significant equivalence for all three contrasts (p values = 0.001-0.03; Fig. 8;

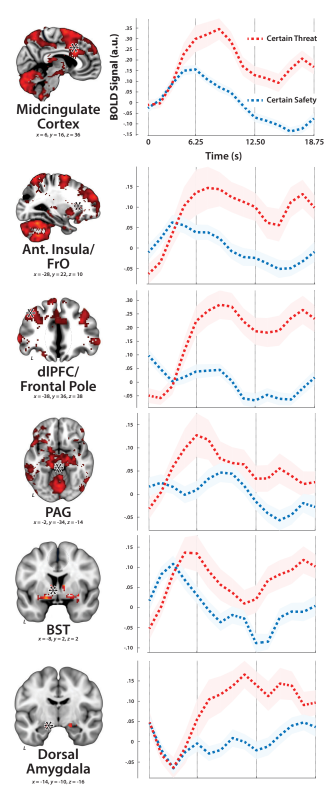


Figure 6. Regions sensitive to temporally Certain Threat show sustained hemodynamic activity. Mean responses to the anticipatory epoch were estimated on a TR-by-TR (1.25 s) basis for Certain Threat (red) and Certain Safety (blue) trials, using data from the local maxima of key clusters (black-and-white asterisks in the left panels) identified using a canonical HRF GLM approach. Given the temporal resolution and autocorrelation of the hemodynamic signal, data were averaged for 3 windows (TR-1 to TR-5, TR-6 to TR-10, and TR-11 to TR-15), spanning a total of 15 measurements (18.75 s). Broken vertical lines indicate windows. Shaded envelopes represent SEM. Ant., Anterior.

Extended Data Fig. 8-1). Although these statistical findings do not demonstrate that the amygdala and the BST are functionally interchangeable ("the same"), they do enable us to decisively reject claims of strict functional segregation (i.e., that the BST is sensitive to uncertain danger, whereas the amygdala is not) for the subset of these regions engaged by the MTC paradigm.

Discussion

Uncertain-threat anticipation is the prototypical trigger of anxiety, a core theme that cuts across psychiatric disorders, species, and assays, including novelty, darkness, and other "diffuse" threats. Despite the immense significance of anxiety for public health, the neural systems recruited by uncertain threat have remained unclear. Leveraging a translationally relevant paradigm optimized for fMRI signal decomposition (Fig. 1), our results reveal that the anticipation of temporally uncertain aversive stimulation recruits a distributed network of frontocortical (MCC, AI/FrO, and dlPFC/FP) and subcortical (PAG, BST, and dorsal amygdala) regions (Fig. 4), mirroring robust changes in experience and psychophysiology (Fig. 1). Closer inspection of signal dynamics in these regions provided descriptive support for sustained activity during the anticipation of Uncertain Threat (Fig. 5). Analyses focused on the anticipation of temporally Certain Threat revealed a similar pattern, with voxels sensitive to both kinds of threat evident in key cortical and subcortical regions (Fig. 4), suggesting that this circuitry is sensitive to both certain and uncertain threat. Direct comparison of the two threat conditions demonstrated that this network can be fractionated: cortical regions showed relatively greater activity during the anticipation of Uncertain Threat, whereas the extended amygdala showed relatively greater activity during the anticipation of Certain Threat (Fig. 7). While there is consensus that the BST and dorsal amygdala play a critical role in orchestrating adaptive responses to danger, their precise contributions to human anxiety have remained contentious. Our results suggest that these regions respond similarly to different kinds of threat anticipation. Indeed, we show that the BST and dorsal amygdala exhibit statistically indistinguishable responses to threat anticipation across a variety of comparisons (Fig. 8), reinforcing the possibility that they make broadly similar contributions to human anxiety (Gungor and Paré, 2016; Fox and Shackman, 2019).

Since the time of Freud (Freud, 1920), the distinction between certain ("fear") and uncertain ("anxiety") danger has been a key feature of neuropsychiatric models of emotion (Davis et al., 2010; LeDoux and Pine, 2016; Mobbs, 2018). Our findings show that the regions recruited during the anticipation of Certain and Uncertain Threat are colocalized in several key regions (Fig. 4). This common threat anticipation network encompasses subcortical regions that are critical for assembling defensive responses to uncertain threat in animals (Fox and Shackman, 2019). But it also includes frontocortical regions—such as the MCC, AI/FrO, and dlPFC/FP—that have received less empirical attention and are challenging to study in rodents (e.g., Carlén, 2017). These regions have traditionally been associated with the controlled processing and regulation of emotion and cognition (Shackman et al., 2011; Morawetz et al., 2017, 2020; Langner et al., 2018; Kroes et al., 2019; Picó-Pérez et al., 2019) and more recently implicated in the conscious experience of emotion (LeDoux, 2020). As shown in Figure 9, the present results are well aligned with recent meta-analyses of neuroimaging studies of "fear" (Fullana et al., 2016) and "anxiety" (Chavanne and Robinson, 2020). Across studies encompassing tens of studies and hundreds

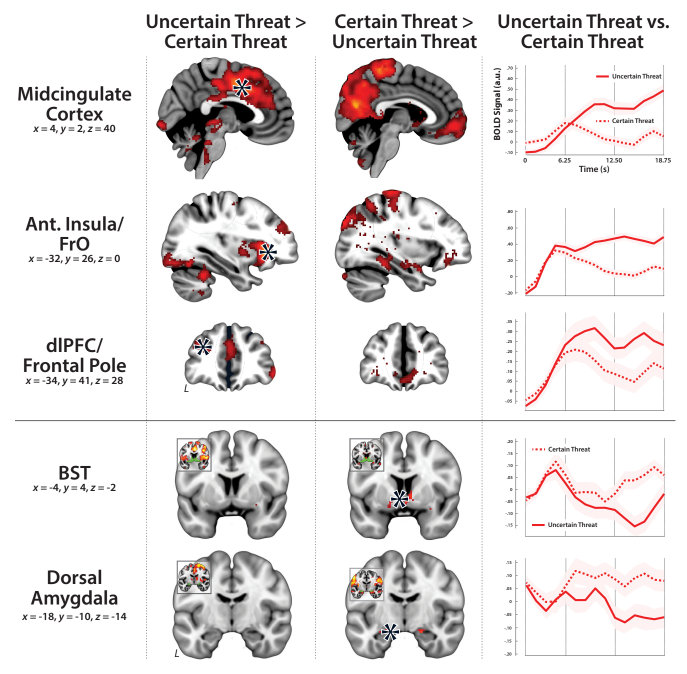


Figure 7. The threat anticipation network can be fractionated into subdivisions. The MCC, Al/FrO, and dlPFC showed greater activity during the anticipation of Uncertain Threat (left column), whereas the BST and dorsal amygdala showed greater activity during the anticipation of Certain Threat (middle column). Thresholds and other conventions are identical to Figure 4. For additional details, see Extended Data Figures 7-1 and 7-2. Right column, TR-by-TR (1.25 s) hemodynamic responses during the anticipation of Uncertain Threat (solid red line) and Certain Threat (broken red line). Data were extracted from the local maxima of key clusters (black-and-white asterisks in the left and center columns) identified using a canonical HRF GLM approach. Given the temporal resolution and autocorrelation of the hemodynamic signal, data were averaged for 3 windows (TR-1 to TR-5, TR-6 to TR-10, and TR-11 to TR-15), spanning a total of 15 measurements (18.75 s). Broken vertical lines indicate windows. Shaded envelopes represent SEM. Ant., Anterior.

of subjects, this work demonstrates that the anticipation of certain-threat (Pavlovian threat cues; the prototypical "fear" stimulus in laboratory studies) and uncertain-threat (instructed "threat-of-shock") recruit an overlapping network of core regions, including the BST (but not the Ce; see below). This similarity cannot be dismissed as an artifact of neuroimagers' penchant for partial-reinforcement Pavlovian paradigms, which render "certain" threat uncertain (Fullana et al., 2016; median threat probability = 63%; Picó-Pérez et al., 2019; median threat probability = 62%). In fact, the same general pattern—including elevated activity in the region of the BST—is evident in large-

scale studies of certain-threat (Sjouwerman et al., 2020) (Study 2, n = 113, threat probability = 100%; https://neurovault. org/collections/6031) and uncertain-threat anticipation (Klumpers et al., 2017) (Sample 1: n = 108, threat probability = 33%), consistent with our results.

Our observations provide insight into the functional architecture of the threat anticipation network, demonstrating that frontocortical regions prefer Uncertain over Certain Threat, whereas the BST and dorsal amygdala show the reverse preference—a difference in degree, not in kind. Trivial differences cannot account for this nuance; the two threat conditions were pseudo-randomly

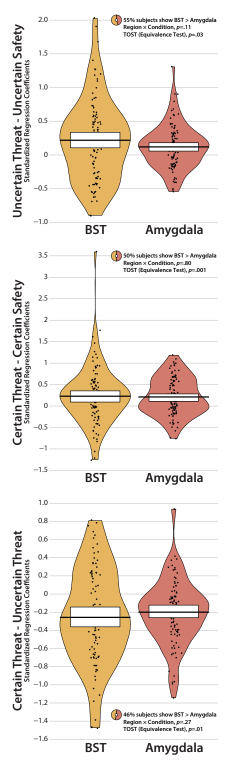


Figure 8. The BST and dorsal amygdala regions recruited by the MTC paradigm show statistically indistinguishable responses during threat anticipation. While it is impossible to demonstrate that the true difference in regional hemodynamic activity is zero, the TOST procedure provides a well-established and widely used statistical framework for testing whether mean differences in regional activity are small enough to be considered equivalent (Lakens, 2017; Lakens et al., 2018). Using the subset of voxels that were most sensitive to each threat contrast (see Materials and Methods), results revealed significant equivalence for all contrasts (Extended Data Fig. 8-1). Figure represents the data (black points; individual subjects), density distribution (bean plots), Bayesian 95% highest density interval (colored bands), and mean (black bars) for each condition. Highest density intervals permit population-generalizable visual inferences about mean differences and were estimated using 1000 samples from a posterior Gaussian distribution. Inset ring plots, Percentage of subjects showing greater activity in the BST compared with the dorsal amygdala for each contrast.

intermixed and nearly identical in terms of their perceptual, nociceptive, motor, and statistical features (Fig. 1). What might explain the observed regional preferences? Aside from temporal certainty, the most conspicuous difference between the conditions is the degree of cognitive scaffolding. On Certain Threat trials, the descending stream of integers provided a precise and predictable index of momentary changes in threat imminence, encouraging a reactive, stimulus-bound cognitive mode. On Uncertain Threat trials, this support was absent, necessitating greater reliance on the kinds of sustained, endogenous representations that are the hallmark of frontocortical regions (Badre and Nee, 2018). A second notable difference between the two threat conditions is the intensity of anxiety. Uncertain-Threat anticipation was associated with greater distress and arousal (Fig. 1). The observed increase in frontocortical activity could reflect either heightened anxiety or compensatory processes aimed at downregulating distress and arousal. Testing these nonexclusive hypotheses will require a multipronged approach that encompasses carefully optimized tasks, mechanistic interventions, and a broader assessment of the nomological network. Multivoxel classifier approaches are likely to be useful for linking specific facets of anxiety (e.g., feelings) to particular elements of the threat anticipation network, and determining whether this reflects expressive or regulatory processes (Chang et al., 2015).

The present results add to a growing body of evidence indicating that the BST and dorsal amygdala, while certainly not interchangeable, are more alike than different (Fox and Shackman, 2019). The BST and dorsal amygdala are characterized by broadly similar patterns of anatomical connectivity, cellular composition, neurochemistry, and gene expression (Fox et al., 2015), although some differences in functional connectivity have been identified (Gorka et al., 2018). Both regions are poised to trigger defensive responses via dense projections to downstream effectors (Fox et al., 2015). Neuroimaging studies have documented similar responses in the two regions to a range of anxiety-eliciting stimuli (Fox and Shackman, 2019; Hudson et al., 2020) (https://neurovault.org/collections/6237), and mechanistic work in rodents reinforces the hypothesis that the BST and dorsal amygdala (Ce) are crucial substrates for human anxiety (Fox and Shackman, 2019). Indeed, work using a variant of the present paradigm in mice shows that Ce-BST projections are necessary for mounting defensive responses during the anticipation of temporally uncertain shock (Lange et al., 2017), consistent with our general conclusions. While our understanding remains far from complete, this body of observations underscores the need to revise models of anxiety that imply a strict segregation of certain and uncertain threat processing in the extended amygdala. The present results imply that the magnitude of regional differences in hemodynamic sensitivity to threat-uncertainty is modest ($< d_z = 0.35$); conditional on perceptual confounds, collinearities, or other moderators; or simply nonexistent. An important challenge for the future will be to determine whether the type of threat uncertainty (e.g., temporal vs likelihood) is a crucial determinant of regional differences in

Our results indicate that the amygdala's response to threat anticipation is sparse, at least when compared to widely used emotional face and scene paradigms. This was not unexpected. The amygdala is a heterogeneous collection of at least 13 nuclei and cortical areas—not "a thing" (Swanson and Petrovich, 1998; Yilmazer-Hanke, 2012)—and converging lines of mechanistic and imaging evidence point to the special importance of the dorsal amygdala in the region of the Ce (Davis et al., 2010; Fox and

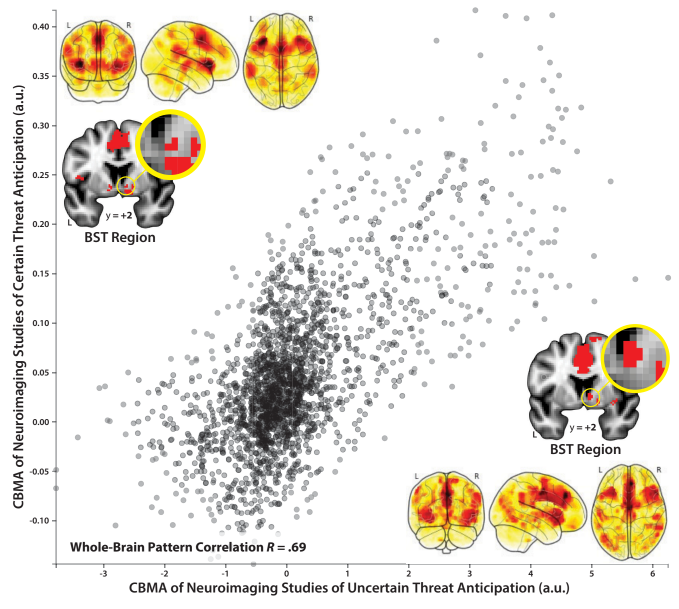


Figure 9. Certain and uncertain threat anticipation elicit broadly similar patterns of neural activity. Figure summarizes the results of two coordinate-based meta-analyses (CBMA) of functional neuroimaging studies. Top left inset, Results for 27 "fear conditioning" studies (N = 677), highlighting regions showing consistently greater activity during the anticipation of certain threat ($CS^+ > CS^-$; https://neurovault.org/collections/2472) (Fullana et al., 2016). Bottom right inset, Results for 18 "threat-of-shock" studies (N = 693), highlighting regions showing consistently greater activity during the anticipation of uncertain threat (Threat > Safe; https://neurovault.org/collections/6012) (Chavanne and Robinson, 2020). Visual inspection of the results (red clusters) suggests that the anticipation of certain and uncertain threat elicits qualitatively similar patterns, including heightened activity in the region of the BST. This impression is reinforced by the substantial correlation between the two whole-brain patterns (r = 0.69). Consistent amygdala activity was not detected in either meta-analysis. The pattern correlation was estimated in Neurovault using a brain-masked, 4 mm transformation of the publicly available, vectorized meta-analytic maps (Gorgolewski et al., 2015). For illustrative purposes, every 10th voxel is depicted in the scatter plot.

Shackman, 2019; Hur et al., 2019a). In humans, Ce represents \sim 3% of total amygdala volume (Wegiel et al., 2014; Avino et al., 2018). The dorsal amygdala clusters that we observed extend beyond the Ce to encompass neighboring dorsocaudal aspects of the medial, lateral, and cortical nuclei, and amygdala-hippocampal transition area. While meta-analyses of small-sample neuroimaging studies have failed to detect significant amygdala responses to threat anticipation (Fullana et al., 2016, median n=16; Chavanne and Robinson, 2020, median n=29), the location and extent of the dorsal amygdala clusters reported here align with more recent large-sample studies of certain-(Sjouwerman et al., 2020) and uncertain-threat anticipation (Reddan et al., 2018) (n=68, threat probability = 33%). In short,

our results are broadly aligned with amygdala anatomy, prior theory, and emerging neuroimaging evidence.

In conclusion, the neural circuits recruited by temporally uncertain and certain threat are not categorically different, at least when viewed through the macroscopic lens of fMRI. We see evidence of anatomic colocalization—not segregation—in a number of key regions, in broad accord with animal models and recent imaging meta-analyses. This shared threat-anticipation system can be fractionated, with frontocortical regions showing relatively stronger engagement during the anticipation of temporally uncertain threat, and the BST and dorsal amygdala showing the reverse pattern. In direct comparisons, the BST and dorsal amygdala exhibited statistically indistinguishable responses,

reinforcing the possibility that they make similar contributions to human anxiety. These observations provide a framework for conceptualizing fear and anxiety and for guiding mechanistic work aimed at developing more effective intervention strategies for pathologic anxiety. A large sample, well-controlled task, and advanced techniques for data acquisition and processing enhance confidence in the robustness and translational relevance of these results.

References

- Acosta-Cabronero J, Williams GB, Pereira JM, Pengas G, Nestor PJ (2008)

 The impact of skull-stripping and radio-frequency bias correction on grey-matter segmentation for voxel-based morphometry. Neuroimage 39:1654–1665.
- Ahrens S, Wu MV, Furlan A, Hwang GR, Paik R, Li H, Penzo MA, Tollkuhn J, Li B (2018) A central extended amygdala circuit that modulates anxiety. J Neurosci 38:5567–5583.
- Andersson JLR, Skare S, Ashburner J (2003) How to correct susceptibility distortions in spin-echo echo-planar images: application to diffusion tensor imaging. Neuroimage 20:870–888.
- Avants BB, Tustison NJ, Song G, Cook PA, Klein A, Gee JC (2011) A reproducible evaluation of ANTs similarity metric performance in brain image registration. Neuroimage 54:2033–2044.
- Avery SN, Clauss JA, Blackford JU (2016) The human BNST: functional role in anxiety and addiction. Neuropsychopharmacology 41:126–141.
- Avino TA, Barger N, Vargas MV, Carlson EL, Amaral DG, Bauman MD, Schumann CM (2018) Neuron numbers increase in the human amygdala from birth to adulthood, but not in autism. Proc Natl Acad Sci USA 115:3710–3715.
- Bach DR (2014) A head-to-head comparison of SCRalyze and Ledalab, two model-based methods for skin conductance analysis. Biol Psychol 103:63–68.
- Bach DR, Friston KJ (2013) Model-based analysis of skin conductance responses: towards causal models in psychophysiology. Psychophysiology 50:15–22.
- Bach DR, Flandin G, Friston KJ, Dolan RJ (2009) Time-series analysis for rapid event-related skin conductance responses. J Neurosci Methods 184:224–234.
- Bach DR, Flandin G, Friston KJ, Dolan RJ (2010) Modelling event-related skin conductance responses. Int J Psychophysiol 75:349–356.
- Bach DR, Friston KJ, Dolan RJ (2013) An improved algorithm for modelbased analysis of evoked skin conductance responses. Biol Psychol 94:490–497.
- Bach DR, Castegnetti G, Korn CW, Gerster S, Melinscak F, Moser T (2018) Psychophysiological modeling: current state and future directions. Psychophysiology 55:e13214.
- Badre D, Nee DE (2018) Frontal cortex and the hierarchical control of behavior. Trends Cogn Sci 22:170–188.
- Bennett KP, Dickmann JS, Larson CL (2018) If or when? Uncertainty's role in anxious anticipation. Psychophysiology 55:e13066.
- Carlén M (2017) What constitutes the prefrontal cortex? Science 358:478–
- Chang LJ, Gianaros PJ, Manuck SB, Krishnan A, Wager TD (2015) A sensitive and specific neural signature for picture-induced negative affect. PLoS Biol 13:e1002180.
- Chavanne AV, Robinson OJ (2020) The overlapping neurobiology of adaptive and pathological anxiety: a meta-analysis of functional neural activation. Am J Psychiatry.
- Choi JM, Padmala S, Pessoa L (2012) Impact of state anxiety on the interaction between threat monitoring and cognition. Neuroimage 59:1912–1923
- Conway CC, et al. (2019) A hierarchical taxonomy of psychopathology can reform mental health research. Perspect Psychol Sci 14:419–436.
- Cox RW (1996) AFNI: software for analysis and visualization of functional magnetic resonance neuroimages. Comput Biomed Res 29:162–173.
- Craske MG, Stein MB, Eley TC, Milad MR, Holmes A, Rapee RM, Wittchen HU (2017) Anxiety disorders. Nat Rev Dis Primers 3:17024.
- Daldrup T, Remmes J, Lesting J, Gaburro S, Fendt M, Meuth P, Kloke V, Pape HC, Seidenbecher T (2015) Expression of freezing and fear-potentiated startle during sustained fear in mice. Genes Brain Behav 14:281–291.

- Davenport S, Nichols TE (2020) Selective peak inference: unbiased estimation of raw and standardized effect size at local maxima. Neuroimage 209:116375.
- Davis M (2006) Neural systems involved in fear and anxiety measured with fear-potentiated startle. Am Psychol 61:741–756.
- Davis M, Walker DL, Miles L, Grillon C (2010) Phasic vs sustained fear in rats and humans: role of the extended amygdala in fear vs anxiety. Neuropsychopharmacology 35:105–135.
- Desikan RS, Ségonne F, Fischl B, Quinn BT, Dickerson BC, Blacker D, Buckner RL, Dale AM, Maguire RP, Hyman BT, Albert MS, Killiany RJ (2006) An automated labeling system for subdividing the human cerebral cortex on MRI scans into gyral based regions of interest. Neuroimage 31:968–980.
- Eskildsen SF, Coupé P, Fonov V, Manjón JV, Leung KK, Guizard N, Wassef SN, Østergaard LR, Collins DL, Alzheimer's Disease Neuroimaging Initiative (2012) BEaST: brain extraction based on nonlocal segmentation technique. Neuroimage 59:2362–2373.
- Fadok JP, Markovic M, Tovote P, Lüthi A (2018) New perspectives on central amygdala function. Curr Opin Neurobiol 49:141–147.
- Fein G, Landman B, Tran H, Barakos J, Moon K, Di Sclafani V, Shumway R (2006) Statistical parametric mapping of brain morphology: sensitivity is dramatically increased by using brain-extracted images as inputs. Neuroimage 30:1187–1195.
- First MB, Williams JB, Karg RS, Spitzer RL (2015) Structured clinical interview for DSM-5, Research version (SCID-5 for DSM-5, research version; SCID-5-RV). Arlington, VA: American Psychiatric Association.
- Fischmeister FP, Hollinger I, Klinger N, Geissler A, Wurnig MC, Matt E, Rath J, Robinson SD, Trattnig S, Beisteiner R (2013) The benefits of skull stripping in the normalization of clinical fMRI data. Neuroimage Clin 3:369–380.
- Fox AS, Shackman AJ (2019) The central extended amygdala in fear and anxiety: closing the gap between mechanistic and neuroimaging research. Neurosci Lett 693:58–67.
- Fox AS, Oler JA, Tromp DP, Fudge JL, Kalin NH (2015) Extending the amygdala in theories of threat processing. Trends Neurosci 38:319–329.
- Frazier JA, Chiu S, Breeze JL, Makris N, Lange N, Kennedy DN, Herbert MR, Bent EK, Koneru VK, Dieterich ME, Hodge SM, Rauch SL, Grant PE, Cohen BM, Seidman LJ, Caviness VS, Biederman J (2005) Structural brain magnetic resonance imaging of limbic and thalamic volumes in pediatric bipolar disorder. Am J Psychiatry 162:1256–1265.
- Freud S (1920) A general introduction to psychoanalysis. New York: Boni and Liveright.
- Fullana MA, Harrison BJ, Soriano-Mas C, Vervliet B, Cardoner N, Àvila-Parcet A, Radua J (2016) Neural signatures of human fear conditioning: an updated and extended meta-analysis of fMRI studies. Mol Psychiatry 21:500–508.
- Gerster S, Namer B, Elam M, Bach DR (2018) Testing a linear time invariant model for skin conductance responses by intraneural recording and stimulation. Psychophysiology 55:e12986.
- Geuter S, Qi G, Welsh RC, Wager TD, Lindquist MA (2018) Effect size and power in fMRI group analysis. bioRxiv 295048.
- Global Burden of Disease Collaborators (2016) Global, regional, and national incidence, prevalence, and years lived with disability for 310 diseases and injuries, 1990-2015: a systematic analysis for the Global Burden of Disease Study 2015. Lancet 388:1545–1602.
- Glover GH (1999) Deconvolution of impulse response in event-related BOLD fMRI. Neuroimage 9:416–429.
- Gonzalez-Castillo J, Hoy CW, Handwerker DA, Roopchansingh V, Inati SJ, Saad ZS, Cox RW, Bandettini PA (2015) Task dependence, tissue specificity, and spatial distribution of widespread activations in large single-subject functional MRI datasets at 7T. Cereb Cortex 25:4667–4677.
- Gorgolewski KJ, Varoquaux G, Rivera G, Schwarz Y, Ghosh SS, Maumet C, Sochat VV, Nichols TE, Poldrack RA, Poline JB, Yarkoni T, Margulies DS (2015) NeuroVault.org: a web-based repository for collecting and sharing unthresholded statistical maps of the human brain. Front Neuroinform 9:8.
- Gorka AX, Torrisi S, Shackman AJ, Grillon C, Ernst M (2018) Intrinsic functional connectivity of the central nucleus of the amygdala and bed nucleus of the stria terminalis. Neuroimage 168:392–402.
- Grabner G, Janke AL, Budge MM, Smith D, Pruessner J, Collins DL (2006) Symmetric atlasing and model based segmentation: an application to the

- hippocampus in older adults. Med Image Comput Comput Assist Interv 9:58–66.
- Graham MS, Drobnjak I, Jenkinson M, Zhang H (2017) Quantitative assessment of the susceptibility artefact and its interaction with motion in diffusion MRI. PLoS One 12:e0185647.
- Greve DN, Fischl B (2009) Accurate and robust brain image alignment using boundary-based registration. Neuroimage 48:63–72.
- Griebel G, Holmes A (2013) 50 years of hurdles and hope in anxiolytic drug discovery. Nat Rev Drug Discov 12:667–687.
- Griessner J, Pasieka M, Bohm V, Grossl F, Kaczanowska J, Pliota P, Kargl D, Werner B, Kaouane N, Strobelt S, Kreitz S, Hess A, Haubensak W (2018) Central amygdala circuit dynamics underlying the benzodiazepine anxiolytic effect. Mol Psychiatry.
- Grillon C, Baas JM, Cornwell B, Johnson L (2006) Context conditioning and behavioral avoidance in a virtual reality environment: effect of predictability. Biol Psychiatry 60:752–759.
- Grupe DW, Nitschke JB (2013) Uncertainty and anticipation in anxiety: an integrated neurobiological and psychological perspective. Nat Rev Neurosci 14:488–501.
- Grupe DW, Wielgosz J, Davidson RJ, Nitschke JB (2016) Neurobiological correlates of distinct post-traumatic stress disorder symptom profiles during threat anticipation in combat veterans. Psychol Med 46:1885– 1895.
- Guller Y, Ferrarelli F, Shackman AJ, Sarasso S, Peterson MJ, Langheim FJ, Meyerand ME, Tononi G, Postle BR (2012) Probing thalamic integrity in schizophrenia using concurrent transcranial magnetic stimulation and functional magnetic resonance imaging. Arch Gen Psychiatry 69:662– 671
- Gungor NZ, Paré D (2016) Functional heterogeneity in the bed nucleus of the stria terminalis. J Neurosci 36:8038–8049.
- Hawrylycz MJ, Lein ES, Guillozet-Bongaarts AL, Shen EH, Ng L, Miller JA, van de Lagemaat LN, Smith KA, Ebbert A, Riley ZL, Abajian C, Beckmann CF, Bernard A, Bertagnolli D, Boe AF, Cartagena PM, Chakravarty MM, Chapin M, Chong J, Dalley RA, et al. (2012) An anatomically comprehensive atlas of the adult human brain transcriptome. Nature 489:391–399.
- Hefner KR, Moberg CA, Hachiya LY, Curtin JJ (2013) Alcohol stress response dampening during imminent versus distal, uncertain threat. J Abnorm Psychol 122:756–769.
- Henson R (2007) Efficient experimental design for fMRI. In: Statistical parametric mapping: the analysis of functional brain images (Friston K, Ashburner J, Kiebel S, Nichols T, Penny W, eds), pp 193–210. San Diego: Academic.
- Hudson M, Seppälä K, Putkinen V, Sun L, Glerean E, Karjalainen T, Karlsson HK, Hirvonen J, Nummenmaa L (2020) Dissociable neural systems for unconditioned acute and sustained fear. Neuroimage 216: 116522
- Hur J, Kaplan CM, Smith JF, Bradford DE, Fox AS, Curtin JJ, Shackman AJ (2018) Acute alcohol administration dampens central extended amygdala reactivity. Sci Rep 8:16702.
- Hur J, Stockbridge MD, Fox AS, Shackman AJ (2019a) Dispositional negativity, cognition, and anxiety disorders: an integrative translational neuroscience framework. Prog Brain Res 247:375–436.
- Hur J, DeYoung KA, Islam S, Anderson AS, Barstead MG, Shackman AJ (2019b) Social context and the real-world consequences of social anxiety. Psychol Med 1–12.
- Jo HJ, Gotts SJ, Reynolds RC, Bandettini PA, Martin A, Cox RW, Saad ZS (2013) Effective preprocessing procedures virtually eliminate distance-dependent motion artifacts in resting state FMRI. J Appl Math 2013:1–9.
- Klein A, Andersson J, Ardekani BA, Ashburner J, Avants B, Chiang MC, Christensen GE, Collins DL, Gee J, Hellier P, Song JH, Jenkinson M, Lepage C, Rueckert D, Thompson P, Vercauteren T, Woods RP, Mann JJ, Parsey RV (2009) Evaluation of 14 nonlinear deformation algorithms applied to human brain MRI registration. Neuroimage 46:786–802.
- Klumpers F, Kroes MC, Baas J, Fernandez G (2017) How human amygdala and bed nucleus of the stria terminalis may drive distinct defensive responses. J Neurosci 37:9645–9656.
- Kriegeskorte N, Lindquist MA, Nichols TE, Poldrack RA, Vul E (2010) Everything you never wanted to know about circular analysis, but were afraid to ask. J Cereb Blood Flow Metab 30:1551–1557.
- Kroes MC, Dunsmoor JE, Hakimi M, Oosterwaal S, NYU PROSPEC Collaboration, Meager MR, Phelps EA (2019) Patients with dorsolateral

- prefrontal cortex lesions are capable of discriminatory threat learning but appear impaired in cognitive regulation of subjective fear. Soc Cogn Affect Neurosci 14:601–612.
- Kruger O, Shiozawa T, Kreifelts B, Scheffler K, Ethofer T (2015) Three distinct fiber pathways of the bed nucleus of the stria terminalis to the amygdala and prefrontal cortex. Cortex 66:60–68.
- Lakens D (2017) Equivalence tests: a practical primer for *t* tests, correlations, and meta-analyses. Soc Psychol Personal Sci 8:355–362.
- Lakens D, Scheel AM, Isager PM (2018) Equivalence testing for psychological research: a tutorial. Adv Methods Pract Psychol Sci 1:259–269.
- Lang PJ, Bradley MM, Cuthbert BN (2008) International Affective Picture System (IAPS): affective ratings of pictures and instruction manual. Technical Report A-8. Gainesville, FL: University of Florida.
- Lange MD, Daldrup T, Remmers F, Szkudlarek HJ, Lesting J, Guggenhuber S, Ruehle S, Jungling K, Seidenbecher T, Lutz B, Pape HC (2017) Cannabinoid CB1 receptors in distinct circuits of the extended amygdala determine fear responsiveness to unpredictable threat. Mol Psychiatry 22:1422–1430.
- Langner R, Leiberg S, Hoffstaedter F, Eickhoff SB (2018) Towards a human self-regulation system: common and distinct neural signatures of emotional and behavioural control. Neurosci Biobehav Rev 90:400–410.
- LeDoux JE (2020) Thoughtful feelings. Curr Biol 30:R619-R623.
- LeDoux JE, Pine DS (2016) Using neuroscience to help understand fear and anxiety: a two-system framework. Am J Psychiatry 173:1083–1093.
- Lorio S, Fresard S, Adaszewski S, Kherif F, Chowdhury R, Frackowiak RS, Ashburner J, Helms G, Weiskopf N, Lutti A, Draganski B (2016) New tissue priors for improved automated classification of subcortical brain structures on MRI. Neuroimage 130:157–166.
- Mai JK, Majtanik M, Paxinos G (2015) Atlas of the human brain, Ed 4. San Diego: Academic.
- Makris N, Goldstein JM, Kennedy D, Hodge SM, Caviness VS, Faraone SV, Tsuang MT, Seidman LJ (2006) Decreased volume of left and total anterior insular lobule in schizophrenia. Schizophr Res 83:155–171.
- McMenamin BW, Shackman AJ, Maxwell JS, Greischar LL, Davidson RJ (2009) Validation of regression-based myogenic correction techniques for scalp and source-localized EEG. Psychophysiology 46:578–592.
- McMenamin BW, Shackman AJ, Maxwell JS, Bachhuber DR, Koppenhaver AM, Greischar LL, Davidson RJ (2010) Validation of ICA-based myogenic artifact correction for scalp and source-localized EEG. Neuroimage 49:2416–2432.
- Miles L, Davis M, Walker D (2011) Phasic and sustained fear are pharmacologically dissociable in rats. Neuropsychopharmacology 36:1563–1574.
- Mobbs D (2018) The ethological deconstruction of fear(s). Curr Opin Behav Sci 24:32–37.
- Morawetz C, Bode S, Derntl B, Heekeren HR (2017) The effect of strategies, goals and stimulus material on the neural mechanisms of emotion regulation: a meta-analysis of fMRI studies. Neurosci Biobehav Rev 72:111–128.
- Morawetz C, Riedel MC, Salo T, Berboth S, Eickhoff S, Laird AR, Kohn N (2020) Multiple large-scale neural networks underlying emotion regulation. Neurosci Biobehav Rev 116:111–128.
- National Institute of Mental Health (2011) Negative valence systems: Workshop proceedings. Rockville, MD.
- National Institute of Mental Health (2020a) Construct: potential threat ("anxiety"). Bethesda, MD: National Institute of Mental Health.
- National Institute of Mental Health (2020b) Construct: acute threat ("fear"). Bethesda, MD: National Institute of Mental Health.
- Nelson BD, Shankman SA (2011) Does intolerance of uncertainty predict anticipatory startle responses to uncertain threat? Int J Psychophysiol 81:107–115.
- Nichols T, Brett M, Andersson J, Wager T, Poline JB (2005) Valid conjunction inference with the minimum statistic. Neuroimage 25:653–660.
- Ollinger JM, Shulman GL, Corbetta M (2001) Separating processes within a trial in event-related functional MRI. Neuroimage 13:210–217.
- Pedersen WS, Muftuler LT, Larson CL (2019) A high-resolution fMRI investigation of BNST and centromedial amygdala activity as a function of affective stimulus predictability, anticipation, and duration. Soc Cogn Affect Neurosci 14:1167–1177.
- Picó-Pérez M, Alemany-Navarro M, Dunsmoor JE, Radua J, Albajes-Eizagirre A, Vervliet B, Cardoner N, Benet O, Harrison BJ, Soriano-Mas C, Fullana MA (2019) Common and distinct neural correlates of fear

- extinction and cognitive reappraisal: a meta-analysis of fMRI studies. Neurosci Biobehav Rev 104:102–115.
- Pomrenze MB, Giovanetti SM, Maiya R, Gordon AG, Kreeger LJ, Messing RO (2019a) Dissecting the roles of GABA and neuropeptides from rat central amygdala CRF neurons in anxiety and fear learning. Cell Rep 29:13–21.e14.
- Pomrenze MB, Tovar-Diaz J, Blasio A, Maiya R, Giovanetti SM, Lei K, Morikawa H, Hopf FW, Messing RO (2019b) A corticotropin releasing factor network in the extended amygdala for anxiety. J Neurosci 39:1030– 1043.
- Power JD, Schlaggar BL, Petersen SE (2015) Recent progress and outstanding issues in motion correction in resting state fMRI. Neuroimage 105:536–551.
- Pruim RH, Mennes M, van Rooij D, Llera A, Buitelaar JK, Beckmann CF (2015) ICA-AROMA: a robust ICA-based strategy for removing motion artifacts from fMRI data. Neuroimage 112:267–277.
- Qi S, Hassabis D, Sun J, Guo F, Daw N, Mobbs D (2018) How cognitive and reactive fear circuits optimize escape decisions in humans. Proc Natl Acad Sci USA 115:3186–3191.
- Reddan MC, Wager TD, Schiller D (2018) Attenuating neural threat expression with imagination. Neuron 100:994–1005.e1004.
- Ressler RL, Goode TD, Evemy C, Maren S (2020) NMDA receptors in the CeA and BNST differentially regulate fear conditioning to predictable and unpredictable threats. Neurobiol Learn Mem 174:107281.
- Roberts A (2020) Prefrontal regulation of threat-elicited behaviors: a pathway to translation. Annu Rev Psychol 71:357–387.
- Salomon JA, Haagsma JA, Davis A, de Noordhout CM, Polinder S, Havelaar AH, Cassini A, Devleesschauwer B, Kretzschmar M, Speybroeck N, Murray CJ, Vos T (2015) Disability weights for the Global Burden of Disease 2013 study. Lancet Glob Health 3:e712-723.e723.
- Shackman AJ, Fox AS (2016) Contributions of the central extended amygdala to fear and anxiety. J Neurosci 36:8050–8063.
- Shackman AJ, Salomons TV, Slagter HA, Fox AS, Winter JJ, Davidson RJ (2011) The integration of negative affect, pain and cognitive control in the cingulate cortex. Nat Rev Neurosci 12:154–167.
- Shackman AJ, Fox AS, Oler JA, Shelton SE, Davidson RJ, Kalin NH (2013) Neural mechanisms underlying heterogeneity in the presentation of anxious temperament. Proc Natl Acad Sci USA 110:6145–6150.
- Shackman AJ, Tromp DP, Stockbridge MD, Kaplan CM, Tillman RM, Fox AS (2016) Dispositional negativity: an integrative psychological and neurobiological perspective. Psychol Bull 142:1275–1314.
- Shackman AJ, Fox AS, Oler JA, Shelton SE, Oakes TR, Davidson RJ, Kalin NH (2017) Heightened extended amygdala metabolism following threat characterizes the early phenotypic risk to develop anxiety-related psychopathology. Mol Psychiatry 22:724–732.
- Shackman AJ, Weinstein JS, Hudja SN, Bloomer CD, Barstead MG, Fox AS, Lemay EP Jr (2018) Dispositional negativity in the wild: social environment governs momentary emotional experience. Emotion 18:707–724.
- Siegel JS, Power JD, Dubis JW, Vogel AC, Church JA, Schlaggar BL, Petersen SE (2014) Statistical improvements in functional magnetic resonance imaging analyses produced by censoring high-motion data points. Hum Brain Mapp 35:1981–1996.
- Sjouwerman R, Scharfenort R, Lonsdorf TB (2020) Individual differences in fear acquisition: multivariate analyses of different Emotional Negativity scales, physiological responding, subjective measures, and neural activation. bioRxiv 233528.

- Smith JF, Hur J, Kaplan CM, Shackman AJ (2018) The impact of spatial normalization for functional magnetic resonance imaging data analyses revisited. BioRxiv.
- Smith SM, Jenkinson M, Woolrich MW, Beckmann CF, Behrens TE, Johansen-Berg H, Bannister PR, De Luca M, Drobnjak I, Flitney DE, Niazy R, Saunders J, Vickers J, Zhang Y, De Stefano N, Brady JM, Matthews PM (2004) Advances in functional and structural MR image analysis and implementation as FSL. Neuroimage 23:S208–S219.
- Somerville LH, Wagner DD, Wig GS, Moran JM, Whalen PJ, Kelley WM (2013) Interactions between transient and sustained neural signals support the generation and regulation of anxious emotion. Cereb Cortex 23:49–60.
- Sreenivasan KK, D'Esposito M (2019) The what, where and how of delay activity. Nat Rev Neurosci 20:466–481.
- Swanson LW, Petrovich GD (1998) What is the amygdala? Trends Neurosci 21:323–331.
- Sylvers P, Lilienfeld SO, LaPrairie JL (2011) Differences between trait fear and trait anxiety: implications for psychopathology. Clin Psychol Rev 31:122–137
- ten Donkelaar HJ, Tzourio-Mazoyer N, Mai JK (2018) Toward a common terminology for the gyri and sulci of the human cerebral cortex. Front Neuroanat 12:93.
- Theiss JD, Ridgewell C, McHugo M, Heckers S, Blackford JU (2017) Manual segmentation of the human bed nucleus of the stria terminalis using 3T MRI. Neuroimage 146:288–292.
- Tillman RM, Stockbridge MD, Nacewicz BM, Torrisi S, Fox AS, Smith JF, Shackman AJ (2018) Intrinsic functional connectivity of the central extended amygdala. Hum Brain Mapp 39:1291–1312.
- Tustison NJ, Avants BB, Cook PA, Zheng YJ, Egan A, Yushkevich PA, Gee JC (2010) N4ITK: improved N3 bias correction. IEEE Trans Med Imaging 29:1310–1320.
- Tye KM, Prakash R, Kim SY, Fenno LE, Grosenick L, Zarabi H, Thompson KR, Gradinaru V, Ramakrishnan C, Deisseroth K (2011) Amygdala circuitry mediating reversible and bidirectional control of anxiety. Nature 471:358–362.
- Van Essen DC, Donahue CJ, Coalson TS, Kennedy H, Hayashi T, Glasser MF (2019) Cerebral cortical folding, parcellation, and connectivity in humans, nonhuman primates, and mice. Proc Natl Acad Sci USA 116:26173–26180.
- Vogt BA, Paxinos G (2014) Cytoarchitecture of mouse and rat cingulate cortex with human homologies. Brain Struct Funct 219:185–192.
- Watson D, Stanton K, Clark LA (2017) Self-report indicators of negative valence constructs within the research domain criteria (RDoC): a critical review. J Affect Disord 216:58–69.
- Wegiel J, Flory M, Kuchna I, Nowicki K, Ma SY, Imaki H, Wegiel J, Cohen IL, London E, Wisniewski T, Brown WT (2014) Stereological study of the neuronal number and volume of 38 brain subdivisions of subjects diagnosed with autism reveals significant alterations restricted to the striatum, amygdala and cerebellum. Acta Neuropathol Commun 2:141.
- Yilmazer-Hanke DM (2012) Amygdala. In: The human nervous system (Mai JK, Paxinos G, eds), pp 759–834. San Diego: Academic.
- Zhang Y, Brady M, Smith S (2001) Segmentation of brain MR images through a hidden Markov random field model and the expectation-maximization algorithm. IEEE Trans Med Imaging 20:45–57.

Anxiety and the neurobiology of temporally uncertain threat anticipation: **Extended Data**

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Descriptive statistics for clusters and local extrema showing greater activity during the anticipation of Uncertain Threat compared to Uncertain Safety (FDR q<.05, whole-brain corrected).

	mm ³	t	X	y	Z
Cluster 1	685,144				
L Anterior Orbital Gyrus		4.26	-26	50	-14
R Anterior Orbital Gyrus		3.19	26	46	-14
R Frontal Pole		7.59	28	46	26
L Middle Frontal Gyrus		6.92	-36	34	38
L Paracingulate Gyrus		8.13	0	34	30
R Paracingulate Gyrus		8.33	6	32	30
R Frontal Orbital Cortex		11.39	32	30	2
L Frontal Orbital Cortex		13.68	-34	26	0
R Inferior Frontal Gyrus, pars triangularis		5.20	54	24	20
L Cingulate Gyrus, anterior division		10.49	-4	20	34
R Cingulate Gyrus, anterior division		11.95	6	18	36
R Frontal Operculum Cortex		13.29	48	16	2
L Frontal Operculum Cortex		11.56	-42	14	2
R Inferior Frontal Gyrus, pars opercularis		6.58	38	12	26
L Inferior Frontal Gyrus, pars opercularis		10.00	-54	10	2
R Juxtapositional Lobule Cortex		10.97	4	8	56
R Superior Frontal Gyrus		10.18	14	8	64
R Central Opercular Cortex		11.24	50	8	2
R Temporal Pole		6.51	44	6	-3
L Temporal Pole		4.64	-40	4	-4

	L Superior Frontal Gyrus	9.23	-16	4	66
	L Accumbens R Putamen L Bed Nucleus of the Stria Terminalis L Central Opercular Cortex L Putamen R Pallidum R Precentral Gyrus L Inferior Temporal Gyrus, anterior division L Insular Cortex L Precentral Gyrus L Parahippocampal Gyrus, anterior division L Caudate R Caudate R Caudate R Amygdala (in the region of the lateral nucleus)a-b R Middle Frontal Gyrus R Planum Polare L Pallidum L Temporal Fusiform Cortex, anterior division L Juxtapositional Lobule Cortex	3.68	-10	4	-6
	R Putamen	10.89	30	4	-2
	L Bed Nucleus of the Stria Terminalis	4.59	-10	4	2
	L Central Opercular Cortex	8.73	-50	2	4
	L Putamen	9.05	-28	2	-2
	R Pallidum	8.38	16	2	0
	R Precentral Gyrus	10.92	46	2	50
	L Inferior Temporal Gyrus, anterior division	5.12	-44	0	-34
	L Insular Cortex	5.05	-32	0	14
	L Precentral Gyrus	7.56	-42	-2	44
	L Parahippocampal Gyrus, anterior division	3.21	-32	-2	-32
	L Caudate	7.62	-16	-2	20
	R Caudate	8.87	16	-2	18
	R Amygdala (in the region of the lateral nucleus) ^{a,b}	4.39	30	-2	-26
	R Middle Frontal Gyrus	7.52	32	-2	44
	R Putamen L Bed Nucleus of the Stria Terminalis L Central Opercular Cortex L Putamen R Pallidum R Precentral Gyrus L Inferior Temporal Gyrus, anterior division L Insular Cortex L Precentral Gyrus L Parahippocampal Gyrus, anterior division L Caudate R Caudate R Caudate R Amygdala (in the region of the lateral nucleus) ^{a,b} R Middle Frontal Gyrus R Planum Polare L Pallidum L Temporal Fusiform Cortex, anterior division	4.36	42	-2	-18
		6.19	-14	-4	0
	L Temporal Fusiform Cortex, anterior division	4.38	-32	-6	-38
	L Juxtapositional Lobule Cortex	6.43	-8	-6	60
	R Insular Cortex	4.32	44	-6	-4
	R Amygdala (in the region of the central nucleus) ^{a,c}	4.85	26	-10	-10
	L Planum Polare	3.48	-44	-10	-12
	R Parahippocampal Gyrus, anterior division	3.38	20	-12	-32
	L Bed Nucleus of the Stria Terminalis L Central Opercular Cortex L Putamen R Pallidum R Precentral Gyrus L Inferior Temporal Gyrus, anterior division L Insular Cortex L Precentral Gyrus L Parahippocampal Gyrus, anterior division L Caudate R Caudate R Amygdala (in the region of the lateral nucleus) ^{a,b} R Middle Frontal Gyrus R Planum Polare L Pallidum L Temporal Fusiform Cortex, anterior division L Juxtapositional Lobule Cortex R Insular Cortex R Amygdala (in the region of the central nucleus) ^{a,c} L Planum Polare R Parahippocampal Gyrus, anterior division R Hippocampus	5.12	30	-14	-12
	R Inferior Temporal Gyrus, posterior division	4.39	54	-16	-32

L Thalamus	6.75	-4	-20	-2
R Thalamus R Temporal Fusiform Cortex, posterior division L Postcentral Gyrus L Cingulate Gyrus, posterior division R Cingulate Gyrus, posterior division L Supramarginal Gyrus, anterior division L Hippocampus R Supramarginal Gyrus, anterior division L Hippocampus R Supramarginal Gyrus, anterior division L Temporal Fusiform Cortex, posterior division L Parahippocampal Gyrus, posterior division L Superior Temporal Gyrus, posterior division R Middle Temporal Gyrus, posterior division L Brainstem/Periaqueductal Gray L Inferior Temporal Gyrus, posterior division R Middle Temporal Gyrus, posterior division L Middle Temporal Gyrus, posterior division L Parietal Operculum Cortex R Parietal Operculum Cortex R Supramarginal Gyrus, posterior division	10.10	6	-20	-2
R Temporal Fusiform Cortex, posterior division	2.18	36	-20	-30
L Postcentral Gyrus	6.13	-56	-24	24
L Cingulate Gyrus, posterior division	8.51	-8	-24	40
R Cingulate Gyrus, posterior division	8.00	10	-24	40
L Supramarginal Gyrus, anterior division	6.30	-60	-26	22
L Hippocampus	3.66	-34	-26	-10
R Supramarginal Gyrus, anterior division	8.42	56	-26	26
R Cingulate Gyrus, posterior division	8.00	10	-24	40
L Supramarginal Gyrus, anterior division	6.30	-60	-26	22
L Hippocampus	3.66	-34	-26	-10
R Supramarginal Gyrus, anterior division	8.42	56	-26	26
L Temporal Fusiform Cortex, posterior division	3.71	-38	-30	-24
L Parahippocampal Gyrus, posterior division	3.63	-26	-30	-28
L Superior Temporal Gyrus, posterior division	3.51	-58	-32	4
R Middle Temporal Gyrus, posterior division	8.34	52	-32	-6
L Brainstem/Periaqueductal Gray	6.23	4	-32	-12
L Inferior Temporal Gyrus, posterior division	4.12	-56	-34	-26
L Middle Temporal Gyrus, posterior division	4.62	-54	-34	-8
R Postcentral Gyrus	5.78	42	-34	54
L Parietal Operculum Cortex	7.45	-58	-36	26
R Parietal Operculum Cortex	2.50	42	-36	22
R Supramarginal Gyrus, posterior division	9.66	56	-42	32
L Supramarginal Gyrus, posterior division	8.73	-62	-44	32
R Middle Temporal Gyrus, temporooccipital part	7.68	58	-44	6

R Superior Parietal Lobule		7.51	36	-46	
R Inferior Temporal Gyrus, temporooccipital part		4.19	56	-46	
L Temporal Occipital Fusiform Cortex		2.67	-36	-48	
L Superior Parietal Lobule		6.98	-30	-50	
R Precuneus Cortex		5.11	10	-50	
L Angular Gyrus		5.71	-60	-54	
L Precuneus Cortex		5.42	-10	-54	
L Middle Temporal Gyrus, temporooccipital part		6.37	-48	-60	
L Lateral Occipital Cortex, superior division		5.18	-16	-60	
R Temporal Occipital Fusiform Cortex		2.70	42	-60	
L Lateral Occipital Cortex, inferior division		4.74	-38	-64	
R Lateral Occipital Cortex, inferior division		4.32	44	-64	
R Angular Gyrus		3.63	-18	-66	
R Lateral Occipital Cortex, inferior division		4.55	42	-70	
R Occipital Fusiform Gyrus		6.51	20	-76	
R Lingual Gyrus		5.62	6	-82	
L Cuneal Cortex		3.48	-8	-84	
L Lingual Gyrus		4.45	-12	-86	
L Occipital Fusiform Gyrus		3.27	-12	-88	
R Occipital Pole		5.40	26	-94	-
L Occipital Pole		3.96	-14	-100	
Cluster 2	368				
L Subcallosal Cortex		3.05	-2	26	
R Subcallosal Cortex		4.60	0	22	
Cluster 3	72				
L Inferior Temporal Gyrus, temporooccipital part		2.48	-60	-56	

Cluster 4	32				
R Superior Temporal Gyrus, anterior division		2.48	54	4	-12

^a Within the Harvard-Oxford amygdala (p>.25), a total of 72 voxels (576 mm³) exceeded threshold. ^b 91% probability of lying within the Harvard-Oxford amygdala. ^c 39% probability of lying within the Harvard-Oxford amygdala.

Descriptive statistics for clusters and local extrema showing greater activity during the anticipation of Uncertain Safety compared to Uncertain Threat (FDR q<.05, whole-brain corrected).

	mm³	t	X	У	Z
Cluster 1	51,072				
R Hippocampus		3.80	24	-20	-20
R Parahippocampal Gyrus, anterior division		3.16	32	-22	-24
R Parahippocampal Gyrus, posterior division		3.18	28	-28	-22
R Temporal Fusiform Cortex, posterior division		5.15	28	-36	-16
L Cingulate Gyrus, posterior division		6.46	-6	-44	4
R Cingulate Gyrus, posterior division		8.04	6	-46	4
R Lingual Gyrus		7.59	14	-50	2
R Precuneus Cortex		10.10	4	-60	20
L Lingual Gyrus		7.62	-4	-62	6
L Precuneous Cortex		7.64	0	-62	34
R Supracalcarine Cortex		11.20	2	-68	18
R Cuneal Cortex		10.66	6	-70	22
R Intracalcarine Cortex		10.17	10	-78	8
L Intracalcarine Cortex		11.11	-14	-84	4
Cluster 2	41,664				
L Postcentral Gyrus		9.59	-52	-10	26
L Planum Polare		3.42	-50	-10	-2
L Heschls Gyrus (includes H1 and H2)		4.57	-52	-16	6
L Central Opercular Cortex		9.06	-38	-16	18
L Parietal Operculum Cortex		5.64	-34	-28	18

R Precentral Gyrus		10.36	2	-30	
L Precentral Gyrus		10.58	-2	-32	
R Postcentral Gyrus		10.32	4	-34	
Cluster 3	8,840				
R Insular Cortex		7.67	36	-10	
Cluster 4	8,096				
L Frontal Pole		4.23	-6	68	
R Inferior Rostral Sulcus		5.62	4	54	
R Frontal Medial Cortex		4.30	6	34	
R Subcallosal Cortex		3.67	2	30	
L Subcallosal Cortex		4.57	-2	20	
Cluster 5	2,648				
R Lateral Occipital Cortex, superior division		7.20	48	-72	
Cluster 6	1,456				
L Lateral Occipital Cortex, superior division		3.70	-40	-80	
Cluster 7	1,432				
L Parahippocampal Gyrus, posterior division		5.64	-26	-38	
Cluster 8	920				
R Superior Frontal Gyrus		4.80	22	32	
Cluster 9	392				
R Planum Temporale		3.78	60	-10	
Cluster 10	352				
L Middle Temporal Gyrus, posterior division		3.97	-60	-12	
Cluster 11	288				
L Hippocampus		3.91	-18	-12	
Cluster 12	264				

R Middle Temporal Gyrus, anterior division		3.34	64	-4	-20
Cluster 13	208				
L Thalamus		3.12	-18	-26	-2
Cluster 14	192				
L Amygdala ^{a,b}		3.92	-30	0	-16
Cluster 15	144				
L Frontal Medial Cortex		3.39	-2	44	-26
Cluster 16	96				
L Frontal Orbital Cortex		3.57	-34	36	-10
Cluster 17	96				
L Planum Temporale		3.37	-44	-30	6
Cluster 18	72				
R Amygdala ^{a,c}		3.61	28	2	-16
Cluster 19	64				
L Paracingulate Gyrus		3.07	-8	42	-4
Cluster 20	64				
L Middle Frontal Gyrus		3.25	-32	24	56
Cluster 21	24				
L Juxtapositional Lobule Cortex		2.99	-10	-14	46
Cluster 22	8				
R Inferior Temporal Gyrus, posterior division		2.81	54	-32	-30
Cluster 23	8				
R Planum Polare		2.87	52	-4	-6
Cluster 24	8				
L Superior Temporal Gyrus, anterior division		3.02	-50	-14	-4
Cluster 25	8				

R Thalamus		2.81	16	-28	-2
Cluster 26	8				
R Heschls Gyrus (includes H1 and H2)		2.96	50	-14	2

^a Within the Harvard-Oxford amygdala (*p*>.25), a total of 22 voxels (176 mm³) exceeded threshold. ^b 60% probability of lying within the Harvard-Oxford amygdala. ^c 61% probability of lying within the Harvard-Oxford amygdala.

$Descriptive\ statistics\ for\ clusters\ and\ local\ extrema\ showing\ greater\ activity\ during\ the\ anticipation$ of Certain Threat compared to Certain Safety (FDR q<.05, whole-brain corrected).

	mm³	t	X	У	Z
Cluster 1	494,696				
L Frontal Pole		7.08	-38	54	14
L Anterior Orbital Gyrus/Lateral Orbital Sulcus		3.08	-32	48	-12
(OP11)					
L Middle Frontal Gyrus		5.94	-38	36	38
R Frontal Pole		5.03	32	36	28
L Inferior Frontal Gyrus, pars triangularis		5.64	-54	34	4
L Medial Orbital Gyrus		4.07	-18	34	-1
R Middle Frontal Gyrus		4.43	34	34	34
R Frontal Orbital Cortex		3.76	40	32	-2
R Inferior Frontal Gyrus, pars triangularis		3.79	44	32	2
R Paracingulate Gyrus		4.06	14	30	26
L Frontal Orbital Cortex		4.96	-36	28	-4
L Cingulate Gyrus, anterior division		6.64	-6	20	34
L Frontal Operculum Cortex		4.59	-38	18	10
R Frontal Operculum Cortex		6.47	50	18	0
R Cingulate Gyrus, anterior division		7.07	6	16	30
R Inferior Frontal Gyrus, pars opercularis		6.53	52	16	-2
L Inferior Frontal Gyrus, pars opercularis		3.09	-52	14	14
L Temporal Pole		7.83	-52	12	-6
L Paracingulate Gyrus		6.18	-2	12	40
L Caudate		5.84	-16	10	16

L Central Opercular Cortex	2.68	-32	8	16
R Accumbens	3.63	12	8	-6
R Caudate	6.59	18	8	20
R Putamen	6.29	22	8	-8
L Putamen	6.51	-18	6	-10
R Temporal Pole	5.03	50	6	-40
L Bed Nucleus of the Stria Terminalis	3.75	-8	2	2
R Bed Nucleus of the Stria Terminalis	3.75	10	2	2
L Superior Frontal Gyrus	6.47	-18	0	70
R Juxtapositional Lobule Cortex	7.02	4	0	68
R Superior Frontal Gyrus	6.44	16	0	68
L Superior Temporal Gyrus, anterior division	4.28	-64	-2	0
L Middle Temporal Gyrus, anterior division	4.95	-56	-2	-36
L Parahippocampal Gyrus, anterior division / Amygdala	3.66	-22	-2	-28
L Pallidum	4.62	-16	-2	4
L Juxtapositional Lobule Cortex	6.76	-2	-2	66
L Precentral Gyrus	6.86	-30	-4	64
L Temporal Fusiform Cortex, anterior division	3.41	-30	-6	-34
R Inferior Temporal Gyrus, anterior division	3.16	44	-6	-36
R Pallidum	5.13	18	-8	-2
L Amygdala (in the region of the Cortical nucleus and Amygdala-Hippocampal Transition Area) ^{a,b}	2.86	-32	0	-22
L Inferior Temporal Gyrus, anterior division	2.88	-44	-10	-34
R Precentral Gyrus	7.20	30	-10	66
R Amygdala/Hippocampus ^{a,c}	5.33	18	-12	-14
L Inferior Temporal Gyrus, posterior division	4.77	-56	-14	-30

R Hippocampus	4.97	30	-16	-12
L Superior Temporal Gyrus, posterior division	3.11	-64	-18	2
L Hippocampus	4.61	-20	-22	-16
R Cingulate Gyrus, posterior division	4.34	6	-24	42
R Inferior Temporal Gyrus, posterior division	4.14	52	-24	-22
L Postcentral Gyrus	2.67	-46	-26	46
L Middle Temporal Gyrus, posterior division	4.47	-66	-28	-6
L Cingulate Gyrus, posterior division	4.59	-12	-30	36
L Supramarginal Gyrus, anterior division	3.56	-54	-32	34
L Parahippocampal Gyrus, posterior division	5.35	-14	-32	-12
R Middle Temporal Gyrus, posterior division	5.27	48	-32	-6
R Supramarginal Gyrus, anterior division	4.68	50	-32	56
R Superior Temporal Gyrus, posterior division	3.36	58	-32	10
L Brainstem/Periaqueductal Gray	2.66	-2	-34	-14
L Temporal Fusiform Cortex, posterior division	4.16	-30	-34	-28
R Thalamus	5.65	10	-34	10
R Temporal Fusiform Cortex, posterior division	2.94	40	-34	-18
R Parietal Operculum Cortex	3.20	56	-34	26
L Thalamus	5.55	-22	-36	-2
R Planum Temporale	3.30	50	-36	16
L Planum Temporale	3.29	-64	-38	18
R Postcentral Gyrus	5.85	38	-38	66
R Middle Temporal Gyrus, temporooccipital part	6.14	50	-38	2
R Superior Parietal Lobule	5.91	34	-44	62
R Supramarginal Gyrus, posterior division	7.25	56	-44	34
L Supramarginal Gyrus, posterior division	6.17	-62	-46	32

L Precuneus Cortex		3.94	-8	-48	48
R Temporal Occipital Fusiform Cortex		2.79	42	-52	-10
L Angular Gyrus		7.51	-56	-54	44
L Inferior Temporal Gyrus, temporooccipital part		5.05	-56	-54	-2
		6.90		-54	62
L Superior Parietal Lobule			-40		
L Middle Temporal Gyrus, temporooccipital part		5.28	-48	-56	6
R Angular Gyrus		3.76	38	-56	30
R Inferior Temporal Gyrus, temporooccipital part		4.18	54	-58	-1
L Lateral Occipital Cortex, superior division		6.36	-30	-60	6
R Precuneus Cortex		5.93	8	-62	5
R Lateral Occipital Cortex, superior division		6.59	14	-68	6
R Lateral Occipital Cortex, inferior division		4.48	48	-76	-1
L Cuneal Cortex		3.81	-18	-78	3
R Lingual Gyrus		5.82	4	-78	-1
L Occipital Fusiform Gyrus		5.76	-8	-84	-2
R Occipital Pole		4.36	2	-94	-1
Cluster 2	600				
L Lateral Occipital Cortex, inferior division		3.64	-48	-80	-:
Cluster 3	280				
R Planum Polare		3.20	42	-8	-1
Cluster 4	160				
	100	204	20	1.4	
R Parahippocampal Gyrus, anterior division		2.94	20	-14	-3
Cluster 5	160				
R Subcallosal Cortex		2.86	10	16	-2
Cluster 6	48				
R Middle Temporal Gyrus, anterior division		3.12	50	-2	-2

Cluster 7	48				
					_
L Lingual Gyrus		2.46	-20	-52	0
Cluster 8	40				
L Frontal Medial Cortex		2.67	-4	52	-28
Cluster 9	40				
R Central Opercular Cortex		2.61	38	-10	22
Cluster 10	32				
L Planum Polare		2.51	-40	-4	-16
Cluster 11	16				
L Parietal Operculum Cortex		2.80	-54	-24	20
Cluster 12	16				
R Cuneal Cortex		2.34	16	-80	28

^a Within the Harvard-Oxford amygdala (p>.25), a total of 148 voxels (1,184 mm³) exceeded threshold. ^b 41% probability of lying within the Harvard-Oxford amygdala. c 57% probability of lying within the Harvard-Oxford amygdala.

Continued...

$Descriptive\ statistics\ for\ clusters\ and\ local\ extrema\ showing\ greater\ activity\ during\ the\ anticipation$ of Certain Safety compared to Certain Threat (FDR q<.05, whole-brain corrected).

	mm ³	t	X	у	Z
Cluster 1	63,216				
R Parahippocampal Gyrus, posterior division		4.98	22	-36	-12
R Lingual Gyrus		6.70	20	-46	-10
R Cingulate Gyrus, posterior division		4.44	2	-48	20
L Cingulate Gyrus, posterior division		3.07	-6	-50	10
L Precuneus Cortex		3.39	-2	-54	6
R Temporal Occipital Fusiform Cortex		5.42	34	-56	-14
L Supracalcarine Cortex		10.57	0	-68	14
R Cuneal Cortex		9.02	8	-68	20
R Occipital Fusiform Gyrus		6.75	30	-74	-8
R Intracalcarine Cortex		11.55	12	-80	4
L Occipital Fusiform Gyrus		5.24	-26	-86	-16
L Cuneal Cortex		4.00	0	-86	28
L Intracalcarine Cortex		11.84	-12	-88	4
R Occipital Pole		11.16	6	-90	6
L Occipital Pole		9.10	-8	-100	-2
Cluster 2	3,864				
R Precentral Gyrus		5.25	4	-28	54
L Precentral Gyrus		5.27	-4	-30	58
Cluster 3	2,688				
L Postcentral Gyrus		5.39	-52	-10	26
Cluster 5	2,240				

L Insular Cortex		4.29	-36	-12	1
L Central Opercular Cortex		6.39	-38	-18	1
L Heschl's Gyrus (includes H1 and H2)		3.44	-36	-24	8
L Parietal Operculum Cortex		5.97	-34	-28	1
Cluster 6	1,096				
R Insular Cortex		5.97	34	-26	1
Cluster 8	864				
L Lingual Gyrus		4.28	-18	-50	-1
Cluster 12	328				
R Central Opercular Cortex		3.35	50	-10	1
R Heschls Gyrus (includes H1 and H2)		3.98	46	-14	(
Cluster 13	328				
L Juxtapositional Lobule Cortex		4.33	-10	-10	4
Cluster 14	256				
R Cingulate Gyrus, anterior division		3.87	2	34	-
Cluster 16	232				
R Postcentral Gyrus		4.58	54	-14	5
Cluster 19	144				
L Superior Temporal Gyrus, anterior division		3.75	-50	-4	-:
Cluster 21	56				
R Subcallosal Cortex		3.35	0	24	-:
Cluster 22	56				
R Superior Frontal Gyrus		3.42	24	18	4
Cluster 23	48				
L Frontal Orbital Cortex		3.47	-28	10	-;
Cluster 24	40				

L Parahippocampal Gyrus, posterior division		3.22	-32	-34	-14
Cluster 26	24				
R Frontal Pole		3.17	8	62	0
Cluster 27	24				
R Precuneus Cortex		3.29	24	-58	10
Cluster 29	16				
L Temporal Occipital Fusiform Cortex		3.16	-34	-48	-16

Descriptive statistics for clusters and local extrema identified by the minimum conjunction (logical 'AND') of the Uncertain Threat greater than Uncertain Safety and Certain Threat greater than Certain Safety contrasts (FDR *q*<.05, whole-brain corrected).

	mm³	$t_{Minimum}$	X	у	Z
Cluster 1	259,760				
L Frontal Pole		6.83	-30	54	28
L Anterior Orbital Gyrus/Lateral Orbital Sulcus		2.89	-30	48	-12
(OP11)					
L Brainstem/Periaqueductal Gray		2.66	-2	34	-14
R Inferior Frontal Gyrus, pars triangularis		2.99	48	30	4
L Frontal Orbital Cortex		4.96	-36	28	-4
R Frontal Orbital Cortex		3.64	38	24	-12
L Insular Cortex		5.06	-28	22	10
L Cingulate Gyrus, anterior division		6.64	-6	20	34
L Frontal Operculum Cortex		4.59	-38	18	10
R Frontal Operculum Cortex		6.47	50	18	0
R Cingulate Gyrus, anterior division		7.07	6	16	36
R Inferior Frontal Gyrus, pars opercularis		6.53	52	16	-2
L Inferior Frontal Gyrus, pars opercularis		3.09	-52	14	14
L Temporal Pole		7.78	-52	12	-4
L Caudate		5.12	-18	12	14
L Paracingulate Gyrus		6.18	-2	12	46
L Putamen		5.86	-24	10	-4
R Accumbens		2.80	10	8	-4
R Caudate		6.11	18	8	18

R Putamen		6.18	22	8	-
L Accumbens		2.62	-8	6	-
L Bed Nucleus of the Stria Terminalis		3.70	-8	4	
L Superior Frontal Gyrus		6.47	-18	0	7
R Juxtapositional Lobule Cortex		7.02	4	0	6
R Superior Frontal Gyrus		6.44	16	0	6
L Pallidum		4.50	-16	-2	
L Juxtapositional Lobule Cortex		6.49	0	-2	6
R Amygdala (in the region of the Central and Medial nuclei) ^{a,b}		2.36	20	-4	-
L Precentral Gyrus		6.38	-34	-6	(
R Pallidum		4.71	18	-6	
R Amygdala/Lentiform Nucleus ^a		2.51	30	-8	-
L Thalamus		5.12	0	-10	-
R Precentral Gyrus		7.20	30	-10	(
R Thalamus		4.36	6	-24	-
L Temporal Fusiform Cortex, posterior division		2.59	-36	-32	-:
L Parahippocampal Gyrus, posterior division		4.57	-12	-32	-
L Inferior Temporal Gyrus, temporooccipital part		4.46	-48	-46	-:
R Temporal Occipital Fusiform Cortex		2.37	42	-52	-
R Lingual Gyrus		5.82	4	-78	-
Cluster 2	79,048				
L Middle Frontal Gyrus		5.94	-38	36	3
L Cingulate Gyrus, posterior division		3.64	-4	-24	4
R Cingulate Gyrus, posterior division		4.34	6	-24	4
R Supramarginal Gyrus, anterior division		4.16	50	-30	4
R Middle Temporal Gyrus, posterior division		5.27	48	-32	

	264	E4	24	20
	3.20	56	-34	26
	3.59	-46	-36	50
	4.55	36	-36	50
	3.29	-64	-38	18
	6.14	48	-38	2
	4.73	-54	-40	38
	7.25	56	-44	34
	6.17	-62	-46	32
	5.67	32	-46	66
	2.84	58	-46	14
	5.36	-34	-50	64
	7.06	-56	-52	42
	4.54	10	-56	54
	3.83	16	-58	66
	3.92	-10	-72	34
	4.43	-12	-76	46
	3.57	-16	-80	32
10,152				
	5.03	32	36	28
	4.43	34	34	34
3,848				
	3.75	-60	-58	6
	3.15	-52	-64	12
3,296				
	4.55	48	6	-42
	3,848	4.55 3.29 6.14 4.73 7.25 6.17 5.67 2.84 5.36 7.06 4.54 3.83 3.92 4.43 3.57 10,152 5.03 4.43 3,848 3,296	3.20 56 3.59 -46 4.55 36 3.29 -64 6.14 48 4.73 -54 7.25 56 6.17 -62 5.67 32 2.84 58 5.36 -34 7.06 -56 4.54 10 3.83 16 3.92 -10 4.43 -12 3.57 -16 10,152 5.03 3,848 3.75 -60 3,296 -52	3.20 56 -34 3.59 -46 -36 4.55 36 -36 3.29 -64 -38 6.14 48 -38 4.73 -54 -40 7.25 56 -44 6.17 -62 -46 5.67 32 -46 2.84 58 -46 5.36 -34 -50 7.06 -56 -52 4.54 10 -56 3.83 16 -58 3.92 -10 -72 4.43 -12 -76 3.57 -16 -80 10,152 5.03 32 36 4.43 34 34 3,848 3.75 -60 -58 3,296 -52 -64

R Middle Temporal Gyrus, anterior division		2.37	56	-2	-3
R Temporal Fusiform Cortex, posterior division		2.48	44	-10	-3
R Inferior Temporal Gyrus, posterior division		3.26	58	-20	-3
Cluster 6	2,152				
L Inferior Temporal Gyrus, anterior division		3.58	-52	-2	-3
L Middle Temporal Gyrus anterior division		3.64	-54	-4	-3
L Inferior Temporal Gyrus, posterior division		3.19	-56	-18	-3
L Middle Temporal Gyrus, posterior division		3.79	-58	-30	-
Cluster 7	1,248				
R Cuneal Cortex		3.81	16	-78	3
Cluster 8	736				
R Right Hippocampus		3.52	30	-38	-
Cluster 9	576				
R Occipital Pole		3.52	32	-88	1
Cluster 10	280				
L Temporal Fusiform Cortex, anterior division		3.05	-32	-4	-3
L Parahippocampal Gyrus, anterior division		2.66	-26	-6	-3
Cluster 11	264				
R Insular Cortex		3.00	42	-6	-
R Planum Polare		3.20	42	-8	-1
Cluster 12	232				
R Lateral Occipital Cortex, inferior division		3.26	46	-66	1
Cluster 13	224				
L Hippocampus		3.49	-34	-24	-1
Cluster 14	112				
R Parahippocampal Gyrus, anterior division		2.94	20	-14	-3

56				
	2.29	58	-52	-16
32				
	2.40	40	-4	-42
32				
	2.27	-20	-52	-2
32				
	2.42	10	46	4
16				
	2.51	-40	-4	-16
16				
	2.56	-32	8	16
16				
	2.80	-54	-24	20
8				
	2.12	10	18	-22
	32 32 32 16 16	2.29 32 2.40 32 2.27 32 2.42 16 2.51 16 2.56 16 2.80	2.29 58 32 2.40 40 32 2.27 -20 32 2.42 10 16 2.51 -40 16 2.56 -32 16 2.80 -54	32 2.29 58 -52 32 2.40 40 -4 32 2.27 -20 -52 32 2.42 10 46 16 2.51 -40 -4 16 2.56 -32 8 16 2.80 -54 -24 8

^aWithin the Harvard-Oxford amygdala (*p*>.25), a total of 13 voxels (104 mm³) exceeded threshold. ^b 40% probability of lying within the Harvard-Oxford amygdala.

Continued...

EXTENDED DATA FIG. 7-1

Descriptive statistics for clusters and local extrema showing greater activity during the anticipation

	mm ³	t	X	y	
Cluster 1	116,856				+
R Paracingulate Gyrus		5.10	2	40	-
R Frontal Pole		4.79	52	40	+
L Paracingulate Gyrus		5.61	-6	34	-
R Inferior Frontal Gyrus, pars triangularis		2.72	46	32	+
R Frontal Orbital Cortex		9.73	38	30	+
L Cingulate Gyrus, anterior division		6.49	-4	24	-
R Frontal Operculum Cortex		9.35	36	24	-
R Superior Frontal Gyrus		6.87	8	22	-
R Inferior Frontal Gyrus, pars opercularis		8.77	52	20	+
R Middle Frontal Gyrus		4.31	46	18	+
L Superior Frontal Gyrus		6.10	-14	12	+
R Insular Cortex		6.25	34	6	+
R Cingulate Gyrus, anterior division		8.53	4	2	+
R Central Opercular Cortex		8.02	52	2	
R Precentral Gyrus		8.22	46	0	
R Putamen		5.27	28	-2	+
R Pallidum		4.09	24	-6	+
R Heschls Gyrus		5.51	46	-12	Ŧ
R Cingulate Gyrus, posterior division		5.95	2	-18	\dagger
L Cingulate Gyrus, posterior division		4.55	-4	-24	t
R Parietal Operculum Cortex		7.26	54	-26	T
R Supramarginal Gyrus, posterior division		5.62	60	-42	t

R Angular Gyrus		5.58	58	-50	3
Cluster 2	68,280				
L Temporal Occipital Fusiform Cortex		3.64	-34	-48	-:
R Temporal Occipital Fusiform Cortex		7.74	34	-52	- 1
R Occipital Fusiform Gyrus		9.60	36	-64	-
R Lateral Occipital Cortex, inferior division		12.89	38	-82	- 1
R Lingual Gyrus		4.77	10	-86	-
L Occipital Fusiform Gyrus		6.78	-22	-88	
L Lateral Occipital Cortex, inferior division		5.14	-38	-90	-:
L Occipital Pole		7.90	-14	-98	-
R Occipital Pole		10.04	18	-98	
Cluster 3	21,816				
L Frontal Orbital Cortex		10.39	-32	26	
L Inferior Frontal Gyrus, pars opercularis		6.85	-50	20	
L Precentral Gyrus		3.96	-54	2	1
L Central Opercular Cortex		5.54	-48	-2	
L Planum Polare		2.95	-42	-2	-:
L Putamen		5.36	-34	-2	
Cluster 4	8,640				
R Thalamus		5.17	12	-10	
Cluster 5	6,784				
L Supramarginal Gyrus, anterior division		3.99	-64	-26	2
L Parietal Operculum Cortex		4.67	-56	-32	2
L Supramarginal Gyrus, posterior division		4.69	-62	-52	3
L Angular Gyrus		3.97	-54	-56	3
Cluster 6	4,720				

R Superior Parietal Lobule		7.64	20	-48	66
Cluster 7	3,336				
L Frontal Pole		4.70	-36	40	30
Cluster 8	1,616				
L Middle Frontal Gyrus		3.55	-42	8	48
Cluster 9	672				
R Temporal Pole		4.01	42	6	-3
R Middle Temporal Gyrus, anterior division		2.98	46	2	-28
Cluster 10	664				
L Superior Parietal Lobule		3.79	-20	-52	64
L Precuneus Cortex		2.76	-10	-52	58
Cluster 11	472				
R Middle Temporal Gyrus, temporooccipital part		4.03	56	-44	4
Cluster 12	344				
L Thalamus		3.50	-10	-12	10
Cluster 13	224				
L Insular Cortex		3.86	-38	-22	-2
Cluster 14	192				
L Temporal Pole		2.96	-40	2	-4
Cluster 15	152				
R Planum Polare		3.66	42	0	-2
Cluster 16	144				
R Superior Temporal Gyrus, posterior division		2.88	50	-20	-4
Cluster 17	112				
R Inferior Temporal Gyrus, posterior division		3.19	52	-36	-2
Cluster 18	104				

R Inferior Temporal Gyrus, temporooccipital part		3.67	64	-40	-22
Cluster 19	80				
R Middle Temporal Gyrus, posterior division		3.05	60	-28	-6
Cluster 20	48				
L Pallidum		3.04	-14	-4	-6
Cluster 21	40				
R Hippocampus		3.87	28	-42	2
Cluster 22	32				
L Inferior Temporal Gyrus, anterior division		2.83	-46	0	-36
Cluster 23	24				
L Temporal Fusiform Cortex, posterior division		3.31	-34	-16	-40

EXTENDED DATA FIG. 7-2

Descriptive statistics for clusters and local extrema showing greater activity during the anticipation of Certain compared to Uncertain Threat (FDR q<.05, whole-brain corrected).

	mm³	t	X	у	Z
Cluster 1	202,448			_	
L Caudate		6.56	-6	16	2
L Putamen		4.84	-14	12	-(
R Putamen		5.49	18	12	-(
L Subcallosal Cortex		6.40	-2	10	-:
R Caudate		5.52	8	8	(
R Bed Nucleus of the Stria Terminalis		4.17	8	4	(
L Temporal Pole		4.44	-34	4	-2
L Bed Nucleus of the Stria Terminalis		4.39	-4	4	-:
R Amygdala ^{a,b}		4.06	30	2	-1
L Amygdala ^{a,c}		3.80	-32	-2	-1
L Amygdala (in the region of the Cortical nucleus and the Amygdala-Hippocampal Transition Area, medial to the Central nucleus) ^{a,d}		4.83	-18	-10	-1
L Central Opercular Cortex		7.21	-36	-12	1
L Parahippocampal Gyrus, anterior division		3.42	-22	-18	-2
L Hippocampus		6.17	-22	-20	-1
R Hippocampus		6.29	24	-20	-1
R Parahippocampal Gyrus, anterior division		3.55	24	-20	-2
L Precentral Gyrus		5.89	-10	-28	7
R Precentral Gyrus		6.74	4	-30	6
L Parahipppocampal Gyrus, posterior division		4.64	-26	-32	-2
L Thalamus		5.37	-8	-32	(

R Thalamus		6.40	10	-34	
R Temporal Fusiform Cortex, posterior division		4.10	24	-36	
R Parahippocampal Gyrus, posterior division		5.79	36	-36	
L Temporal Fusiform Cortex, posterior division		6.15	-26	-40	
L Postcentral Gyrus		8.51	-8	-40	
L Cingulate Gyrus, posterior division		7.47	-6	-44	
R Cingulate Gyrus, posterior division		7.48	8	-50	
L Lingual Gyrus		6.24	-16	-54	
L Middle Temporal Gyrus, temporooccipital part		3.66	-54	-56	
L Superior Parietal Lobule		4.74	-32	-56	
R Lingual Gyrus		9.19	10	-58	
R Angular Gyrus		3.41	48	-58	
R Intracalcarine Cortex		9.23	8	-60	
R Precuneus Cortex		7.43	16	-60	
L Angular Gyrus		3.61	-42	-62	
L Precuneus Cortex		7.33	-2	-62	
L Lateral Occipital Cortex, inferior division		4.24	-44	-64	
R Cuneal Cortex		7.50	20	-66	
L Cuneal Cortex		7.90	-10	-70	
L Intracalcarine Cortex		4.84	0	-76	
R Lateral Occipital Cortex, superior division		7.34	42	-82	
L Lateral Occipital Cortex, superior division		6.36	-30	-88	
R Occipital Pole		3.42	8	-90	
Cluster 2	14,480				
L Frontal Pole		5.66	-2	56	
R Frontal Pole		5.56	4	54	

L Paracingulate Gyrus		3.11	-12	50	
R Frontal Medial Cortex		4.87	0	48	-
R Paracingulate Gyrus		3.47	14	42	
L Frontal Medial Cortex		3.27	-4	36	-
Cluster 3	9,256				
R Central Opercular Cortex		4.01	44	-8	
R Insular Cortex		5.23	36	-10	
R Postcentral Gyrus		6.29	64	-10	7
R Supramarginal Gyrus, anterior division		2.84	60	-22	4
Cluster 4	2,720				
R Middle Temporal Gyrus, anterior division		5.41	56	-2	-
R Middle Temporal Gyrus, posterior division		3.23	68	-10	-
Cluster 5	2,576				
L Inferior Temporal Gyrus, temporooccipital part		3.43	-50	-58	-
Cluster 6	2,512				
R Superior Frontal Gyrus		4.80	24	30	4
Cluster 7	1,208				
L Middle Temporal Gyrus, anterior division		4.04	-58	-8	-
L Middle Temporal Gyrus, posterior division		4.29	-64	-12	-
Cluster 8	1,016				
L Superior Frontal Gyrus		3.51	-26	30	!
Cluster 9	984				
L Transverse Orbital Sulcus		4.38	-36	36	-
Cluster 10	808				
L Planum Temporale		4.21	-52	-38	
L Supramarginal Gyrus, posterior division		2.93	-54	-48	-

Cluster 11	808				
L Juxtapositional Lobule Cortex		4.68	-4	-2	64
Cluster 12	640				
R Middle Frontal Gyrus		3.43	28	2	52
Cluster 13	424				
L Superior Temporal Gyrus, posterior division		3.29	-64	-28	2
Cluster 14	312				
R Transverse Orbital Sulcus		3.09	30	36	-6
Cluster 15	296				
R Supramarginal Gyrus, posterior division		2.87	48	-36	46
Cluster 16	88				
R Inferior Temporal Gyrus, posterior division		3.45	50	-26	-20
Cluster 17	88				
R Subcallosal Cortex/Inferior Rostral Sulcus		2.72	6	22	-14
Cluster 18	88				
L Supramarginal Gyrus, anterior division		3.02	-50	-32	34
Cluster 19	56				
L Inferior Temporal Gyrus, posterior division		2.82	-60	-22	-24
Cluster 20	48				
L Middle Frontal Gyrus		3.00	-40	22	26
Cluster 21	32				
R Planum Temporale		3.00	40	-34	12
Cluster 22	24				
R Temporal Pole		3.12	54	14	-34
Cluster 23	16				
R Planum Polare		2.80	52	-6	-8

Cluster 24	16				
L Insular Cortex		2.65	-32	-26	4
Cluster 25	16				
L Inferior Frontal Gyrus, pars triangularis		2.74	-38	32	12
Cluster 26	8				
L Pallidum		2.75	-26	-12	-6
Cluster 27	8				
L Occipital Pole		3.03	-12	-90	12

^aWithin the Harvard-Oxford amygdala (p>.25), a total of 251 voxels (2,008 mm³) exceeded threshold. ^b 57% probability of lying within the Harvard-Oxford amygdala. c 42% probability of lying within the Harvard-Oxford amygdala. d 80% probability of lying within the Harvard-Oxford amygdala.

Continued...

EXTENDED DATA FIG. 8-1

Detailed inferential statistics for the direct comparison of the BST to the Amygdala.

	BST		Amygdala					TOST for a 'Medium' Effect
Contrast ^a	М	SD	M	SD	R	Region × Condition	Cohen's dz	$d_Z = .35^a$
Uncertain Threat vs. Uncertain Safety	.2186	.60067	.1200	.32312	.273	t(98)=1.637, p=.105	.16	t(98)=1.85, p=.034
Certain Threat vs. Certain Safety	.2290	.67035	.2118	.43576	.300	t(98)=.251, p=.802	.03	t(98)=3.23, p=.001
Certain Threat vs. Uncertain Threat	2569	.52219	1997	.35992	.369	t(98)=1.109, p=.270	.11	t(98)=2.37, p=.010

^a (Lakens, 2017).

EXTENDED REFERENCE

Lakens D (2017) Equivalence tests: A practical primer for t tests, correlations, and meta-analyses. Social Psychological and Personality Science 8:355-362.