Hippocampal representation of threat features and behavior in a human approach-avoidance conflict anxiety task

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Abstract

Decisions under threat are crucial to survival and require integration of distinct situational features such as threat probability and magnitude. Recent evidence from human lesion and neuroimaging studies implicated anterior hippocampus (aHC) and amygdala in approach/avoidance decisions under threat, and linked their integrity to cautious behavior. Here we sought to elucidate how threat dimensions and behavior are represented in these structures.

Twenty human participants (11 female) completed an approach-avoidance conflict task during high-resolution functional MRI. Participants could gather tokens under threat of capture by a virtual predator, which would lead to token loss. Threat probability (predator wake-up rate) and magnitude (amount of token loss) varied on each trial. To disentangle effects of threat features, and ensuing behavior, we performed a multifold parametric analysis.

We found that high threat probability and magnitude related to BOLD signal in left anterior hippocampus/entorhinal cortex. However BOLD signal in this region was better explained by avoidance behavior than by these threat features. A priori region-of-interest analysis confirmed the relation of anterior hippocampus BOLD response with avoidance. Exploratory subfield analysis revealed that this relation was specific to anterior CA2/3 but not CA1. Left lateral amygdala responded to low and high, but not intermediate threat probability.

Our results suggest that anterior hippocampus BOLD signal is better explained by avoidance behavior than by threat features in approach-avoidance conflict. Rather than representing threat features in a monotonic manner, it appears that anterior hippocampus may compute approach/avoidance decisions based on integration of situational threat features represented in other neural structures.

Significance statement

An effective threat anticipation system is crucial to survival across species. Natural threats, however, are diverse and have distinct features. To be able to adapt to different modes of danger, the brain needs to recognize these features, integrate them and use them to modify behavior. Our results disclose the human anterior hippocampus as a likely arbiter of approach/avoidance decisions harnessing compound environmental information while partially replicating previous findings and blending into recent efforts to illuminate the neural basis of approach-avoidance conflict in humans.
**Introduction**

Integrating divergent situational demands is critical to survival; in particular when predatory or metabolic threat is involved (Korn and Bach, 2015, 2018, 2019). A standard laboratory model of this situation is provided by approach-avoidance conflict (AAC) tests, e.g. open-field test and elevated plus-maze (Calhoon and Tye, 2015), which are thought to reflect aspects of human clinical anxiety disorders (Aupperle and Paulus, 2010). Situational threat features are manifold and distinct in these tests, and even more so in biological scenarios (Evans et al., 2019). For a human during wintertime, there is a low probability of being attacked when encountering a hibernating bear and a higher probability when coming across wolves, who are short on food. The metabolic loss incurred by a bear chase, however, may be much higher than when being charged by a single wolf. How the neural system represents and integrates such different threat dimensions, and how they influence behavior, e.g., the decision to approach food under threat or passively avoid threat, remains unknown.

In rodent AAC tests, cautious ("anxiety-like") behavior is consistently reduced by anxiolytic drugs such as benzodiazepines (Gray and McNaughton, 2000). Ventral hippocampus lesions have a similar impact (Kjelstrup et al., 2002; Bannerman et al., 2003; McHugh et al., 2004; Ito and Lee, 2016; Kirlic et al., 2017), and it has been suggested that behavioral control requires interplay of hippocampal subfields (Schumacher et al., 2018). Theta oscillations of hippocampal local field potential (Gordon et al., 2005), and synchronization with prefrontal cortex (Adhikari et al., 2010; Padilla-Coreano et al., 2016), are increased in AAC, while area-specific circuits influence decisions (Wallis et al., 2019). In a human computer game resembling open-field test, benzodiazepines (Korn et al., 2017) and other anxiolytics (Bach et al., 2018) reduced cautious behavior similar to hippocampus (Bach et al., 2014) and amygdala (Korn et al., 2017) lesions in humans and non-human primates (Chudasama et al., 2008; Machado et al., 2009). Amygdala contribution is inconsistently reported in rodents (Kirlic et al., 2017); in humans it appears to be specifically required for retreat from threat after reward collection, rather than for the decision to approach (Bach et al., 2019).

While this suggests involvement of hippocampus and amygdala in behavioral control, it remains elusive how different threat features, ultimately determining behavior, are represented and integrated. Features such as magnitude and probability of threat are not experimentally controlled in many tests that build on innate anxiety, or that are extended in time. For example, we have shown using functional magnetic resonance imaging (fMRI) that neural mass activity of anterior hippocampus (aHC) increases with threat probability in continuous-time AAC (Bach et al., 2014). However, fMRI studies with more abstract AAC...
tests not requiring immediate behavior have yielded conflicting results, some suggesting the same relation of aHC activity with threat probability (Korn and Bach, 2019); others a relation of aHC activity (Loh et al., 2017) or multivoxel patterns (O'Neil et al., 2015) with behavior.

Operant conflict tests provide the opportunity to more precisely control threat features as demonstrated in rodents (Evenden et al., 2009; Oberrauch et al., 2019) and humans (Bach, 2015, 2017; Bach et al., 2019). Here, we capitalized on this latter operant AAC test to disambiguate representation of attack probability, its metabolic cost, and behavior, in aHC and amygdala. We previously used the same task to show that putative hippocampal gamma oscillations, and hippocampal theta synchronization with prefrontal cortex, increased with threat probability (Khemka et al., 2017). Presently, we gained from the superior spatial resolution of fMRI collecting 1.5 mm isotropic blood-oxygen-level-dependent (BOLD) images focused on amygdala and hippocampus while participants played the game. On each trial, they could either collect, or forgo, a monetary token under threat of capture by a predator. Threat probability was defined by the predator wake-up rate and learned by experience; threat magnitude by potential token loss and explicitly signaled.

Materials and Methods

Participants

Twenty participants were recruited from general and student population in Zurich (mean age ± SD, 23.10 ± 3.34 years; 11 female). Participants had no prior history of neurological or psychiatric disease and reported normal or corrected-to-normal vision. One participant was excluded from fMRI analysis due to a technical fault in MRI recordings, but included in behavioral analysis. Behavioral results remained consistent after removal of this participant. All participants gave their written informed consent before participation. The study protocol was in full accordance with the Declaration of Helsinki and approved by the governmental ethics committee (Kantonale Ethikkommission Zürich).

Experimental procedure

Participants performed an AAC computer game as previously used in Khemka et al. (2017), which was modified from Bach (2015). At the beginning of each trial, the human player was
located in a "safe place" in the bottom block of a 2 x 2 diamond grid (Fig 1) opposite of a sleeping predator, and was given the opportunity to collect a monetary token that would appear in the left or right grid block. Red diamonds underneath the grid explicitly signaled the number of tokens that would be lost (0-5) if captured by the predator. Threat probability was implicitly signaled through frame color (blue, pink and orange). Threat probability was implemented by setting the wake-up rate per time unit to result in catch probabilities of 0.1, 0.2, or 0.3 per 100 ms spent outside of the safe grid block. These probabilities were learned from experience during 36 preceding training trials without token loss that did not count towards ultimate earnings.

A token appeared after a random time interval drawn from a truncated gamma distribution ($k = 2, \theta = 1$; mean = 2 s, $t \leq 6$ s). If the player chose not to collect the token, it would disappear after another time interval drawn from the same distribution, and the trial would end one second later. If the player went to acquire the token and successfully returned to the safe place, the trial would proceed until the same predetermined end time. Finally, if the predator caught the player, the predator changed its color from gray to red and remained on the screen until the predetermined end time of the trial. After a random intertrial interval (ITI) also drawn from a gamma distribution truncated at $t \leq 4$ s, during which a blank screen was presented, the next trial would start. Participants completed 648 trials in random order, balanced for each combination of experimental factors, i.e. threat magnitude, and threat probability. Participants were instructed beforehand that their payment depended on performance in six trials randomly drawn from the experiment excluding training trials. The experiment was programmed in Cogent (Version 2000v1.25; www.vislab.ucl.ac.uk/Cogent) and MATLAB (Version 7.14; MathWorks).

**Acquisition of MRI data**

Data was recorded in a 3.0-Tesla MRI scanner (Phillips Achieva; Phillips Medical Systems, Best Netherlands) using a 32-channel head coil. Anatomical images were acquired using a 0.76 mm isotropic resolution T1-weighted scan (TR = 7.37 ms, TE = 3.29 ms, flip angle = 8°, field of view (FOV) = 255x255x180 mm, matrix = 336x336, thickness = 0.76 mm, in-plane resolution = 0.76 x 0.76 mm², slice tilt = 0°, 237 sagittal slices) and a 1.0 x 0.5 x 0.5 mm resolution T2-weighted scan centered on hippocampus (TR = 3200 ms, TE = 353 ms, flip angle = 90°, FOV = 200x52x200 mm, matrix = 400x400, thickness = 1 mm, in-plane resolution = 0.5 x 0.5 mm², slice tilt = 22°, 104 transverse slices). B0 Field maps were acquired with a double-echo fast gradient echo sequence (TR = 698.22 ms, TE = 4.10 and 7.10, flip angle = 44°, FOV = 240x224x240 mm, matrix = 80 x 80, thickness = 3 mm, in-plane...
resolution = 3 x 3 mm$^2$, slice tilt = 0°, 2x64 sagittal slices). Functional images during the approach-avoidance paradigm were recorded with 1.5 mm isotropic resolution T2* -weighted echo-planar imaging (EPI) sequence (TR = 2800 ms, TE = 30 ms, flip angle = 85°, in-plane resolution = 1.5 x 1.5 mm$^2$, FOV = 216x54x216 mm, matrix = 144 x 144; 36 transverse slices with thickness = 1.5 mm; slice order = interleaved ascending; slice tilt = -40°). Field of view (FOV) was centered on amygdala/hippocampus, but also encompassed striatum, thalamus, prefrontal cortices with exclusion of orbitofrontal cortex and cranio-posterior segments of frontal lobe, greater parts of temporal lobes and cerebellum as well as complete coverage of insular cortices and brainstem (Fig 1).

Preprocessing of MRI data

Preprocessing of functional images was performed using a standard pipeline in SPM12 (Statistical Parametric Mapping; Wellcome Centre for Human Neuroimaging, London, UK; http://www.fil.ion.ucl.ac.uk/spm/software/spm12). In a first step, slice time correction was performed to account for differences in acquisition time of individual brain slices (Sladky et al., 2011). Geometric distortions due to susceptibility-induced field inhomogeneities were addressed using a combined approach, which takes static distortions as well as changes in distortion due to head motion into account (Andersson et al., 2001; Hutton et al., 2002). Static distortions were derived for each subject individually from a B0 field map using the FieldMap toolbox in SPM12. Echo-planar images were subsequently realigned and unwarped integrating the measured static distortion and the estimation of distortion caused by head motion, as well as head motion itself. EPI-images as well as T2w-images were then coregistered to the individual T1w whole brain image using a 12-parameter affine transformation. Finally, EPI-images were normalized into Montreal Neurological Institute (MNI) space and smoothed using an isotropic 8 mm full-width at half maximum (FWHM) Gaussian kernel for primary mass-univariate analysis, and a 4 mm FWHM Gaussian kernel for a secondary analysis to improve localization of effects in amygdala and hippocampus. We note that the smoothing kernel must strike a balance between anatomical intersubject variability, and regional specificity (Mikl et al., 2008). Thus, the larger smoothing kernel is expected to be more sensitive in detecting activations, but the smaller kernel can provide additional information on the localization of clusters. Unsmoothed EPI images in native space were used for region-of-interest (ROI) analysis.
In a primary analysis (P1), we defined a general linear model consisting of a delta function at token appearance (consistent with Khemka et al. 2017), convolved with a canonical hemodynamic response function (HRF). Parametric modulators for linear and quadratic effect of threat probability (1-3), linear and quadratic effect of threat magnitude (0-5), and linear interaction effect of threat probability x magnitude, were also convolved with the HRF. All parametric modulators were serially orthogonalized. Motion correction parameters were included as six additional regressors of no interest.

To distinguish effects of behavior from threat features, we ran a second parametric analysis (P2) with approach or avoidance as a first parametric modulator, followed by linear and quadratic effect of threat probability, linear and quadratic effect of threat magnitude, linear combination of threat probability x magnitude, linear combinations of approach x probability, approach x magnitude and finally approach x probability x magnitude.

To extricate effects of threat probability and magnitude that were specific to ensuing behavior, we computed a third general linear model with separate trial regressors for approach trials and avoidance trials (P3), each with parametric modulators for linear and quadratic effects of threat probability and magnitude as well as for effect of linear combination of probability x magnitude. To assess a potential relation of neural activity with response latencies, we defined three further models in an analogous manner with parametric regressors for linear and quadratic effects of approach and withdrawal latency during approach trials. We controlled for threat features in all models using serial orthogonalization.

Since the player was often caught by the predator during attempts to obtain a token, data for withdrawal latency was available on fewer trials than for approach latency. Thus, we defined one model for approach latency over all trials without control for withdrawal latency, and two models over trials without capture, where approach and withdrawal latencies were orthogonalized in respect to each other.

Region-of-interest definition

Subcortical and cortical structures including hippocampus and amygdala were identified in native subject space using the "recon-all" pipeline in FreeSurfer Version 6.0 (http://surfer.nmr.mgh.harvard.edu/) (Dale et al., 1999; Fischl et al., 1999a; Fischl et al., 1999b; Fischl et al., 2002; Segonne et al., 2005; Desikan et al., 2006; Fischl et al., 2008). Individual voxels were assigned neuroanatomical labels in an automated volumetric subcortical parcellation based on a probabilistic atlas from a manual training set (Fischl et al.,
The hippocampus segmentation was then further parcellated into anterior and mid-to-posterior hippocampus by automatically splitting the mask at one-third length along the anterior-posterior axis of the image in MATLAB (Strange et al., 2014). For exploratory purposes, CA1 and CA2/3 subfields as well as a mask for dentate gyrus of the hippocampus were obtained from FreeSurfer 6.0, which uses a statistical atlas based on ultra-high resolution ex vivo data and combines T1w- and T2w-images for multispectral segmentation (Iglesias et al., 2015). CA1, CA2/3 and dentate gyrus images were then multiplied with the binary anterior hippocampus mask to focus only on the anterior segments.

For small volume correction of group-level analysis, a group-level bilateral hippocampus mask was generated by warping the individual bilateral hippocampus masks into MNI space using the deformation fields acquired during normalization of whole-brain T1w images in SPM12. These were then averaged, thresholded at 0.1 and binarized using the SPM12 function ImCalc. For visualization, group-level masks in MNI space for all significant clusters were extracted using SPM12 Results.

**Region-of-interest fMRI analysis**

For analysis of estimated condition-by-condition BOLD response, averaged within regions of interest (ROI), we defined a first-level general linear model with separate regressors for 36 possible distinct combinations of threat probability (1-3), magnitude (0-5) and behavioral response (0/1). We extracted estimated condition-by-condition BOLD response for anterior hippocampus, anterior subfields CA1 and CA2/3, anterior dentate gyrus, entire amygdala, centrocortical and basolateral amygdala subnucleus groups, and, for visualization, for significant clusters from focused brain analysis.

**Statistical Analysis**

Image-based statistical tests for fMRI analysis were performed with SPM group level analysis using cluster-level family-wise error (FWE) correction for multiple comparisons at a voxel-inclusion threshold of p < 0.001 (correction for whole field of view, or small volume corrected for hippocampus) and applying a random-field theory based approach as implemented in SPM (Worsley et al., 1992).

For a priori ROIs amygdala and hippocampus, we implemented a mixed effects analysis in R 3.4.3 (www.r-project.org) using function lmer (lme4 package) with the following fixed effects...
that followed the definition of the voxel-wise analysis while adding a hemispheric difference:
linear and quadratic effects of threat probability and magnitude, behavioral response and
hemisphere, and ensuing interactions. We added a random intercept for subject. This
resulted in the R formula (where all predictors are numerical rather than factors):

\[ Y \sim 1 + (\text{threat probability} \times \text{threat magnitude} + \text{threat probability}^2 + \text{threat magnitude}^2) \times \text{behavioral response} \times \text{hemisphere} + (1|\text{subject}) \]

Exploratory analysis in anterior hippocampus subfields (CA1, CA2/3) and amygdala
subnuclei groups (basolateral and centrocortical) was then performed using the same
formula. Significance level $\alpha$ was adjusted for multiple comparisons across two regions of
interest for a priori tests, and four regions of interest for exploratory analysis using the Holm-
Bonferroni method (Holm, 1979). To further differentiate for region- and subfield-specific
effects in an exploratory analysis, ROI was included as a fixed effect in one combined model
for amygdala vs. aHC and another for anterior CA1 vs. CA2/3. Lastly, post-hoc ROI analysis
was performed in anterior dentate gyrus using the initial model without ROI as factor.
Statistical analysis of behavioral data was likewise performed in R using a linear mixed-
effects model (lme4 package), which can deal with the inherently unbalanced data (see for
details: (Bach, 2015; Khemka et al., 2017)), using Satterthwaite approximation to degrees of
freedom to appropriately control the false positive rate (Luke, 2017).

Data availability

A repository of unthresholded SPM activations maps for parametric analyses P1-3 (group-
level; 4 and 8 mm kernel) is publicly available https://github.com/a-abivardi/neural-threat-
behavior-AAC-fMRI (Abivardi et al., 2020).

Results

Behavioral results

We first interrogated whether behavior was comparable to previous findings. Passive
avoidance (i.e. the proportion of avoidance over approach decisions) increased with higher
threat probability and magnitude. Behavioral inhibition, measured as approach latency,
increased with higher threat probability and magnitude, while the opposite pattern was
observed for withdrawal latency (Fig 2 / Table 1). These results replicate previous reports 
(Bach, 2015, 2017; Khemka et al., 2017; Bach et al., 2019) and are in concordance with 
known behavior from rodent studies.

Mass-univariate FMRI results

As threat features and approach/avoidance behavior are strongly related, we chose a 
threefold parametric design (P1-3) to disentangle distinct effects using serial 
orthogonalization as implemented in SPM12. In a primary analysis (P1; Table 2), we 
analyzed how BOLD signal related to linear and quadratic components of the two threat 
dimensions and their interactions, by including them as parametric modulators. A second 
analysis (P2; Table 3) prepended these modulators by behavioral response 
(approach/avoidance), making use of serial orthogonalization in SPM12, and further 
examined interactions between threat dimensions and behavior. Lastly (P3; Table 4), 
approach and avoidance trials were analyzed separately to account for behavior-specific 
effects of threat dimensions on brain activation. All results were corrected for family-wise 
error (FWE) within the FOV. For bilateral hippocampus, additional FWE small volume 
correction was performed using a group-level bihemispheric mask, as we had strong a priori 
hypotheses for this region. Mass-univariate results are reported for images smoothed with an 
8 mm Gaussian kernel, unless otherwise specified.

In analysis P1, we observed higher BOLD signal with a combination of higher threat 
probability and higher threat magnitude in left anterior hippocampus (specifically subiculum) 
and entorhinal cortex (linear x linear interaction, FOV-corrected corrected, Fig 3, Table 2). 
This effect was not reproduced in a secondary analysis with a narrower smoothing kernel 
size of 4 mm. There were no mass-univariate effects in the amygdala. Exploratory analysis of 
the remaining brain coverage (Table 2) revealed higher BOLD signal with lower threat 
probability (linear negative effect of threat probability) in left dorsolateral prefrontal cortex 
(dIPFC), a cluster extending into left putamen and anterior insula, and in the posterior lobe of 
the right cerebellum. Low threat magnitude was related to higher BOLD signal (linear 
negative effect of threat magnitude) in left internal capsule/putamen, posterior short gyrus of 
left insula, ventrolateral PFC (vLPFC), left inferior temporal gyrus and multiple clusters in 
bilateral cerebellum and vermis.

In P2, there were no significant hippocampus or amygdala clusters at FOV correction. After 
small volume correction (SVC) in bilateral hippocampus, we observed a cluster in which 
avoidance behavior related to higher BOLD activity. This cluster in left anterior hippocampus
combined threat magnitude and probability in P1 (Fig 3, Table 3). This result was replicated
in a secondary analysis using 4 mm kernel smoothed images for higher localization
accuracy. In this analysis, the cluster was in adjacent location, but more superior in the
anterior CA3/dentate gyrus area (Fig 3). In a distinct cluster in left middle hippocampus (8
mm kernel only), high BOLD signal related to high threat magnitude. Exploratory analysis of
the remaining brain coverage (Table 3) revealed that approach behavior related to BOLD
signal in two large clusters encompassing bilateral cerebellum and extending from bilateral
thalamus to striatum and midbrain structures. Furthermore, approach behavior related to
activation in left substantia nigra, bilateral anterior cingulate cortex and dorsomedial PFC
dmPFC, anterior short gyrus of right insula, opercular part of right inferior frontal gyrus and
precentral cortex. These clusters showed partial overlap with impact of low threat magnitude
as shown in P1, in bilateral cerebellum, left putamen and anterior insula, as well as with
impact of low threat probability in right cerebellum and left putamen. After controlling for
behavior in P2, no linear effects of threat probability were observed. A quadratic modulating
effect of threat probability emerged in the left lateral amygdaloid nucleus (8 mm kernel only),
i.e. high activation for low and for high, but not for medium threat probability (Fig 3). High
threat magnitude was related to high BOLD signal in right anterior insula (anterior and middle
short gyrus) and frontal operculum. A second adjacent cluster in the right frontal operculum
showed a linear relation of BOLD signal with threat magnitude specifically in combination
with approach (interaction threat magnitude x behavior). This effect, however, was only
estimable in 16 subjects.

In P3, there were no hippocampus or amygdala clusters at whole-brain or small-volume
correction. Exploratory analysis of the remaining brain coverage (Table 4) revealed that for
approach trials, high threat magnitude was associated with high BOLD signal in bilateral
anterior short gyrus of insula, opercular part of inferior frontal gyrus and bilateral anterior
cingulate. Overlap with activation related to approach behavior in P2 was seen primarily in
anterior cingulate while overlap with activation related to high threat magnitude in P2 was
seen in right anterior insula (replicating the previous finding). Linear interaction of high threat
probability and magnitude in approach trials (estimable in 17 subjects) furthermore related to
BOLD signal in right superior colliculus (partial overlap with approach related activation from
P2) and a cluster extending from left brachium of inferior colliculus into the medial geniculate
nucleus. Specifically, BOLD response increased with threat magnitude for medium and high
threat probabilities, but not for low probability.

Effects in avoidance trials were only partially estimable due to unequal distribution (i.e.
relative scarcity of avoidance trials across participants) and yielded no significant results.
Finally, we found a positive relation of approach latency with BOLD activation in left anterior cingulate cortex and right anterior insula over all approach trials. When controlling for withdrawal latency in the subset of trials where the player was not caught, neither this nor any other relation was seen. There were no significant clusters in relation to withdrawal latency independent of approach latency or threat features.

Region-of-interest analysis results

A priori ROI analysis was carried out across both anterior hippocampi, and across both amygdalae. Results were corrected for multiple comparisons across the two ROIs using Holm-Bonferroni adjusted significance level (Table 5).

In the anterior hippocampus ROI, we observed a linear main effect of behavioral response and quadratic main effects for threat probability and magnitude. BOLD signal was higher for avoidance than for approach trials. Similar to left lateral amygdala in parametric analysis P2, anterior hippocampus also responded to low and high threat probability (positive quadratic effect). Strikingly, this effect seemed to be behavior-dependent and lateralized as left anterior hippocampus responded to high threat probability and right hippocampus activation related to low threat probability; both during avoidance only (quadratic x linear interaction of threat probability and behavior, and linear interaction of threat probability x behavior x hemisphere) (Fig 4). Moreover, for zero threat magnitude, hippocampus BOLD signal was low, while increasing to peak levels for low to intermediate levels and falling again with higher magnitude, resulting in a significant negative quadratic pattern. Finally, anterior hippocampus exhibited a complex linear interaction of threat features and hemisphere: BOLD response showed a negative linear relation with threat magnitude for high threat probability and for left hemisphere only.

The response to low and high threat probability seen in lateral amygdala after control for behavior was replicated in the amygdala ROI analysis (positive quadratic main effect), while interactions between threat probability and behavior were not detected. Left hemisphere showed overall higher BOLD responses in the amygdala.

A combined analysis of amygdala and anterior hippocampus revealed distinct activation patterns in relation to behavior (Fig 4 / Table 5). While aHC was clearly more active during avoidant behavior amygdala exhibited a slightly higher BOLD response during approach (behavior x ROI interaction). The quadratic response to threat magnitude appeared to be specific to aHC; moreover aHC was different from amygdala in its lateralized response to threat probability during avoidance (linear threat probability x magnitude x behavior x ROI
In addition to planned ROI analysis, exploratory follow-up analyses were carried out in bilateral anterior hippocampus subfields CA1 and combined CA2/3 as well as basolateral and centrocortical amygdala ROIs. Results were corrected for multiple comparisons across four ROIs using Holm-Bonferroni method (Table 5).

Subfield analysis in anterior CA1 revealed complex and interacting effects of threat dimensions with distinct activation patterns for approach and avoidant behavior and depending on hemisphere. As in entire aHC, a relation to low and high threat probabilities was seen during avoidance only (quadratic x linear interaction). A complex linear interaction effect of threat features and hemisphere also similar to entire aHC was observed.

Activation in combined hippocampal subfield CA2/3 was higher for avoidance than approach behavior, reflecting the main effect found in the combined anterior hippocampus ROI. The difference (or increase) in BOLD response for avoidance compared to approach conditions, was furthermore higher for anterior CA2/3 (M=1.31, SD=1.28) than for subfield CA1 (M=0.22, SD=1.17) in a post-hoc paired sample t-test (t(17) = -3.31, p = 0.004), underlining the difference between the two subfields. A combined model for the subfields confirmed this distinction with a significant behavior x ROI interaction effect (Fig 4 / Table 5). At the suggestion of a reviewer, we analyzed BOLD responses in anterior dentate gyrus, based on findings that this area may have a role similar to that of CA3. However, we did not find a significant relation with avoidant behavior here.

Further exploratory analysis in amygdala subnuclei using a probabilistic amygdala mask from a previous study (Abivardi and Bach, 2017) revealed activation of basolateral amygdala with increasing threat magnitude (linear main effect). This effect was not seen for entire amygdala. Left basolateral and centrocortical amygdala were more active than amygdala of the right hemisphere as also seen for entire amygdala. Also, centrocortical amygdala exhibited heightened BOLD response to intermediate threat magnitudes, especially during avoidance (quadratic main effect + quadratic x linear interaction). We note that the centrocortical amygdala parcellation was defined by structural connectivity with lateral orbitofrontal cortex (Bach et al., 2011) based on preferred projections to central, medial and cortical amygdala in rodents and non-human primates (Carmichael and Price, 1995; McDonald, 1998; Pitkänen, 2000). Morphologically, this group parcellation (resulting from a sample with size of n=50; Abivardi and Bach, 2017) probably includes central, medial, cortical, as well as basomedial nuclei.
Discussion

Rodent and human ventral or anterior hippocampus are crucial to cautious behavior in AAC tests (Ito and Lee, 2016). However how distinct threat features are represented and integrated has only recently received attention (Korn and Bach, 2019). Harnessing a human operant AAC computer game during high-resolution fMRI, we investigated representation of threat probability and threat magnitude, and of approach or avoidance behavior, in anterior hippocampus (aHC) and amygdala. Two key findings emerged. First, aHC BOLD activity was related to behavioral avoidance, particularly for CA2/3 but not for CA1. Secondly, there was no evidence that aHC unambiguously represents elementary threat features in a linear manner. Similarly, exploratory analyses of further brain areas within our limited coverage did not reveal a coherent linear representation of threat probability or magnitude.

In mass-univariate analysis we observed that BOLD signal in left aHC/entorhinal cortex, specifically the subiculum-entorhinal area, was related to the combination of high probability and magnitude of threat (analysis P1), both of which result in more avoidant behavior. After controlling for behavior (P2), no such relation was found. Instead, neural activity in a slightly more posterior cluster was related to avoidant behavior. Using a smaller smoothing kernel to fully harness high spatial resolution, we localized this second cluster to the anterior CA3/dentate gyrus area. A priori ROI analysis confirmed these findings: averaged aHC BOLD signal was increased during avoidance. Follow-up analysis of anterior subfields revealed that this avoidance-related increase occurred in CA2/3 but not CA1. This finding resonates with a rat experiment by Schumacher et al. (2018) who demonstrated that selective pharmacological inactivation of ventral CA3 increased approach behavior. A role paralleling CA3 has been recently described for rodent ventral dentate gyrus (Yeates et al., 2019). We note that it remains possible that our CA2/3 parcellations contain individual voxels belonging to bordering dentate gyrus. Nonetheless, exploratory ROI analysis in dentate gyrus did not detect a similar effect here. On the other hand, CA1 activity in our study showed no simple relationship with threat features or behavior, whereas selective pharmacological ventral CA1 inactivation increased avoidance in a previous rat experiment (Schumacher et al., 2018).

Our finding of aHC activity relating to avoidance are in keeping with a previous human fMRI study involving abstract AAC decisions, which reported inferior aHC BOLD activity during avoidance (Loh et al., 2017), in proximity to the left aHC cluster relating to avoidance here. We note that in this previous study, most voxels in this cluster were labeled as belonging to CA1; however, the authors noted that anatomical specificity might have been limited due to lower spatial resolution (3 mm), as opposed to the present approach.
In a lesion study with the same paradigm as used here, we found that hippocampus lesions impaired approach-avoidance decisions, whereas impact of threat on other behaviors remained intact (Bach et al., 2019), further suggesting a specific role of aHC in generating avoidance behavior. Selective amygdala and hippocampus lesions were moreover associated with shorter approach latency, but not with a different relationship between threat and approach latency. This may suggest that these regions do not contribute to parametric variation in approach latency. In keeping with this, we presently found that variation in approach latency did not relate to hippocampus or amygdala signal. A previous magnetoencephalography study reported a relation between approach latency and posterior hippocampus activity (Khemka et al., 2017), not observed here.

Regarding threat feature representation, ROI analysis revealed a more complicated picture than previously assumed. Though we observed significant responses of aHC to low and intermediate threat magnitude levels, forming a quadratic pattern, BOLD signal also depended on interactions between threat features and behavior, with some effects strikingly different between hemispheres. Specifically, left aHC responded to high threat probability while right aHC related to low probability during avoidance. In humans, left hippocampus has been implicated in contextual and spatial memory encoding while right hippocampus has been linked to navigation accuracy (Maguire et al., 1998; Spiers et al., 2001). Hemisphere-specific connectivity profiles in human aHC (Robinson et al., 2016) and task-related activity in rat ventral HC (Sakaguchi and Sakurai, 2017) have been reported. However, we note the historical and ongoing debate on lateralization of emotional functions, which is based on partly contradicting observations (Gainotti, 2019). It would therefore appear useful to replicate our findings in an independent sample.

In contrast, a previous fMRI study (Bach et al., 2014) using a more ethological paradigm reported linearly increasing activity in left aHC with higher threat probability. Accounting for the influence of behavior in this temporally extended paradigm was, however, difficult. Furthermore, previous threat probabilities were higher (0.2/0.5/0.8) than the current ones (0.1/0.2/0.3). Also, this previous study did not explicitly control threat magnitude, which we achieved here. Another fMRI study involving more abstract foraging decisions under predation (Korn and Bach, 2019) found a cluster in which aHC signal increased with threat probability (0.1-0.4) but a partly overlapping cluster in which aHC signal decreased from 0.1-0.3 and increased from 0.3-0.4, yielding an overall quadratic pattern. To reconcile these findings, it appears necessary to cover a larger probability range.
As a further finding, BOLD signal in left lateral amygdala related to low and high, but not intermediate threat probability independent of behavior (P2). ROI analysis in entire amygdala exhibited replicated this behavior-independent activation pattern. The role of amygdala in AAC is reported more controversially than for aHC (Kirlic et al., 2017); nevertheless a recent human lesion study suggested specific involvement in controlling vigor of return to safety (Bach et al., 2019).

Results from exploratory focused brain analysis revealed several clusters with complex and differential relation with threat features and behavior. Left dorsolateral PFC (dlPFC) response was related to low threat probability (P1), resonating with reports that anxiety is inversely correlated with dlPFC activity (Balderston et al., 2017). Right ventrolateral PFC (vlPFC) has been implicated in motor inhibition and characterized as a "brake", which however has been debated (Aron et al., 2004, 2014; Swick and Chatham, 2014). Here, right vlPFC activity related to low threat magnitude (P1), approach behavior (P2), and high magnitude during approach (P3). While we did observe behavioral inhibition during approach trials relating to threat, the relation to approach behavior seems at odds with pure motor inhibition. Swick and Chatham (2014) propose that vlPFC monitors action-relevant situational changes, compatible with response to threat magnitude here.

In a recent optogenetic study, anterior cingulate cortex activation decreased rodent freezing behavior via input to basolateral amygdala (Jhang et al., 2018). Anterior cingulate also appears to signal value predictions of rewards and punishments (Monosov, 2017). Conceptually, dorsal anterior cingulate has been theorized to monitor conflict (or expected value of top-down control) (Botvinick et al., 1999; Shenhav et al., 2016) or to adaptively track context-relevant and action-guiding variables. (Heilbronner and Hayden, 2016). Here, dorsal anterior cingulate related to approach behavior (P2) while also relating to rises in threat magnitude during approach trials (P3). The former finding matches anterior cingulate role in freezing in mice and supports a more active role arbitrating behavior. The latter finding may equally well constitute measurement of conflict, context-relevant variable tracking or punishment-related value predictions.

Anterior insular cortex activity was related to approach decisions and both threat features. Left insula related to low threat magnitude and probability before accounting for behavior (P1), right anterior insula activation was related to approach (P2) and bilateral insula to high threat magnitude in separated approach trials (P3). This contrasts reports from a study implicating anterior insula activation in avoidance decisions (Aupperle et al., 2015). Overall, insula showed similar responses to anterior cingulate; adding to evidence of their close functional link (Medford and Critchley, 2010).
Limitations of our study include the use of a limited field-of-view as a necessary compromise for higher resolution imaging of regions-of-interest. Furthermore a relative scarcity of avoidance decisions across participants, compared to previous studies with cumulative token collection (Bach, 2015, 2017; Bach et al., 2019) hindered analysis of threat representation during avoidance and reduced power to detect brain areas involved in avoidant decision-making. A focus on single-stage decisions in the present study precludes analyzing to what extent assumptions about future foraging attempts may prompt avoidance on the current one (Korn and Bach, 2019; Zorowitz et al., 2020). Lastly, orthogonalization in SPM12 penalizes parametric modulators in a serial manner along the design matrix, which demands careful interpretation of results (Mumford et al., 2015).

To summarize, in this study we disambiguated a relation of neural tissue activity with behavior and situational threat features. Anterior hippocampus BOLD signal, in particular in CA2/3, increased when participants avoided threat. Representation of threat features showed a complicated pattern, and for threat probability depended on behavior. This is in line with a notion that hippocampus does not linearly represent threat features but retrieves them, possibly in a manner that changes over time, in order to compute decisions. It would be useful to increase the range of these threat features, as well as improve both spatial and temporal precision of recording, for example using electrophysiology, to understand how these computations emerge over time in different hippocampal subfields.

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Legends

Table 1. Analysis of Variance (ANOVA) of effects of threat features on behavioral measures with Satterthwaite’s approximation. Abbreviations: TP = threat probability, TM = threat magnitude.

Table 2. Parametric modulating effects of threat probability, magnitude and their interaction on brain activation (Analysis P1). FWE-corrected results (p < .05) at cluster level (whole-brain + whole-brain/small volume corrected (SVC) for hippocampus), at a voxel-inclusion level inclusion threshold of p < 0.001. Manual labeling in comparison with schematic brain atlas (Mai et al., 2016). Automated labeling shows AAL (Tzourio-Mazoyer et al., 2002) peak labels verbatim.
Table 3. Parametric modulating effects of behavioral response (approach/avoidance), followed by threat probability, magnitude and their interactions on brain activation (Analysis P2). FWE-corrected results ($p < .05$) at cluster level (whole-brain + small volume corrected (SVC) for hippocampus), at a voxel-inclusion level inclusion threshold of $p < 0.001$. Manual labeling in comparison with schematic brain atlas (Mai et al., 2016).

Table 4. Parametric modulating effects of threat probability, magnitude and their interactions on brain activation in separated approach and avoidance trials (Analysis P3). FWE-corrected results ($p < .05$) at cluster level (whole-brain), at a voxel-inclusion level inclusion threshold of $p < 0.001$. Manual labeling in comparison with schematic brain atlas (Mai et al., 2016).

Table 5. Main and interaction effects significant after Holm-Bonferroni correction (* $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$) from Analysis of Variance (ANOVA) of mixed effects model of estimated condition-by-condition BOLD response averaged across region-of-interest (entire/subregional amygdala and anterior hippocampus). Significant main and interaction effects from ANOVA of combined mixed effects model for amygdala vs. anterior hippocampus and anterior CA1 vs. anterior C2/3. Abbreviations: TP = threat probability, TM = threat magnitude, A = action, H = hemisphere, ROI = region-of-interest.

Figure 1. A: Field-of-view focused on amygdala/hippocampus. The image shows the EPI coverage across participants (thresholded at $p=.5$), overlaid on a mean T1 image in MNI space. B: Approach/avoidance conflict task. In each trial, the human participant (green triangle) started out in a safe (dark gray) grid block opposite a sleeping predator (gray circle) and was presented with a reward token (yellow rhombus) on the left or right side. Threat probability was signaled by frame color (blue/magenta/orange). The player then had the choice to collect the token using left/right keys to move out of, and return to, the safe place. If caught whilst outside, the amount of tokens signaled in red below the frame (here two) was lost, thus constituting the magnitude of threat.

Figure 2. A: Proportion of approach/avoidance decisions ± SEM defined as SD of generalized linear mixed-effects model residuals divided by square root of number of data points. B: Approach and withdrawal latency, estimated from linear mixed-effects model ± SEM (defined as SD of model residuals divided by square root of number of data points).

Figure 3. Cluster-level significant anterior hippocampus (aHC) and amygdala clusters from parametric analysis with, for purposes of illustration, extracted estimated condition-by-condition BOLD response ± SEM as defined by SD of BOLD response amplitude estimates divided by square root of number of data points. Primary analysis clusters using 8 mm
FWHM smoothing kernel are displayed in red, secondary analysis cluster (B) using 4 mm kernel is overlaid in blue. **A:** Left anterior subiculum-entorhinal cortex cluster modulated by combined threat probability and magnitude (linear positive interaction effect, analysis P1). **B:** Left anterior subiculum-entorhinal cortex area relating to avoidance (P2; small volume corrected). Secondary analysis localized this cluster to the left anterior CA3/dentate gyrus area. BOLD estimates ± SEM are displayed for the 4 mm cluster. **C:** Left lateral amygdala cluster quadratically modulated by threat probability (P1). All results are FWE corrected at cluster level (p < .05; voxel inclusion threshold: p < .001).

**Figure 4.** Region-of-interest analyses for anterior hippocampus (aHC) (A), amygdala vs. aHC (B) and anterior CA1 vs. anterior CA2/3 (B). **A:** Interaction effect of threat probability, approach and hemisphere: i.e. estimated condition-by-condition BOLD response amplitudes ± SEM defined as SD of mixed effects model residuals divided by square root of number of data points. **B:** Interaction effect of behavior x ROI for amygdala vs. aHC and anterior CA1 vs. anterior CA2/3 (condition-by-condition BOLD response ± SEM).
Table 1.  
Linear and omnibus effects of threat features on behavioral responses

| Action (proportion approach) | Approach latency |
|-----------------------------|------------------|
|                             | F    | df  | p     | F    | df  | p     |
| TP: omnibus                 | 61.36 | 2, 12923.0 | < .001*** | 12.57 | 2, 8883.2 | < .001*** |
| TP: linear                  | 214.74 | 1, 12923.0 | < .001*** | 14.14 | 1, 8883.9 | < .001*** |
| TM: omnibus                 | 463.61 | 5, 12923.0 | < .001*** | 37.59 | 5, 8884.8 | < .001*** |
| TM: linear                  | 1945.61 | 1, 12923.0 | < .001*** | 173.95 | 1, 8892.3 | < .001*** |
| TP x TM: omnibus            | 3.58  | 10, 12923.0 | < .001*** | 3.60  | 10, 8881.6 | < .001*** |
| TP x TM: linear             | 15.73 | 1, 12923.0 | < .001*** | 3.52  | 1, 8883.1 | 0.061 |

| Withdrawal latency | Movement into correct direction |
|--------------------|---------------------------------|
| F    | df  | p     | F    | df  | p     |
| TP: omnibus         | 13.74 | 2, 7070.0 | < .001*** | 0.56  | 2, 8939.0 | 0.570 |
| TP: linear          | 18.18 | 1, 7070.4 | < .001*** | 1.00  | 1, 8939.0 | 0.318 |
| TM: omnibus         | 10.04 | 5, 7070.5 | < .001*** | 1.53  | 5, 8939.0 | 0.176 |
| TM: linear          | 37.90 | 1, 7074.0 | < .001*** | 5.57  | 1, 8939.0 | 0.018* |
| TP x TM: omnibus    | 0.92  | 10, 7069.1 | 0.514 | 0.97  | 10, 8939.0 | 0.467 |
| TP x TM: linear     | 0.80  | 1, 7069.9 | 0.371 | 2.14  | 1, 8939.0 | 0.144 |
Table 2.
Parametric analysis (P1): effects of threat features on brain activation

| Cluster anatomy (manual label.) | Cluster size | FWE p-value (cluster level) | Peak z-score | Peak coordinates (MNI - mm) | Peak label (AAL) |
|---------------------------------|--------------|----------------------------|--------------|----------------------------|-----------------|
| **Threat probability - negative linear effect** |             |                           |              |                            |                 |
| L middle frontal gyrus [dIPFC]  | 14           | 0.049                      | 4.02         | -33 45 30                  | Frontal_Mid_L   |
| L putamen; L insula [anterior short gyrus] | 17           | 0.015                      | 3.97; 3.80   | -20 15 -3; -30 18 -8      | Putamen_L; Insula_L |
| R cerebellum                    | 25           | 0.015                      | 4.23         | 44 -47 -33                 | Cerebelum_Crus1_R |
| **Threat magnitude - negative linear effect** |             |                           |              |                            |                 |
| L anterior limb of internal capsule / putamen | 36           | <0.001                     | 4.41         | -23 15 8                  | Putamen_L       |
| L insula [posterior short gyrus] | 16           | 0.042                      | 4.29         | -36 -2 6                  | NA              |
| R inferior frontal gyrus, opercular part [vIPFC] | 16           | 0.042                      | 4.28         | 42 9 27                   | Frontal_Inf_Oper_R |
| R cerebellum                    | 784          | <0.001                     | 4.76         | 33 -53 -23                | Cerebelum_6_R   |
| 82                              | <0.001       | 4.50                       | 18 -47 -18   | Cerebelum_4_5_R           |
| 56                              | <0.001       | 4.30                       | 26 -62 -57   | Cerebelum_8_R             |
| 59                              | <0.001       | 4.12                       | 9 -72 -26    | Cerebelum_6_R             |
| 53                              | <0.001       | 4.08                       | 15 -66 -45   | Cerebelum_8_R             |
| 34                              | <0.001       | 3.92                       | 29 -66 -27   | Cerebelum_6_R             |
| L cerebellum                    | 33           | <0.001                     | 3.93         | -30 -51 -23               | Cerebelum_6_L   |
| 16                              | 0.008        | 3.67                       | -35 -63 -26  | Cerebelum_6_L             |
| Cerebellar vermis               | 18           | 0.021                      | 3.76         | -3 -59 -32                | Vermis_9        |
| L inferior temporal gyrus       | 17           | 0.030                      | 3.88         | -51 -62 -20               | Temporal_Inf_L  |
| **Threat probability x magnitude - positive linear effect** |             |                           |              |                            |                 |
| L entorhinal cortex; L pre- and parasubiculum extending into CA1 [of anterior hippocampus] | 14           | 0.043                      | 3.50         | -16 -9 -27                | ParaHippocampal_L |
| 12                              | 0.002 (SVC)  | 3.50; 3.29                | -17 -9 -27; -18 -14 -21 | ParaHippocampal_L; Hippocampus_L |
Table 3.
Parametric analysis (P2): effects of approach/avoidance behavior and serially orthogonalized threat features on brain activation

| Cluster anatomy (manual labeling) | Cluster size | FWE p-value (cluster) | Peak z-score | Peak coordinates (MNI - mm) | Peak label (AAL) |
|----------------------------------|-------------|-----------------------|--------------|-----------------------------|-----------------|
| **Effect of approach**           |             |                       |              |                             |                 |
| LR Cerebellum                    | 14178       | <0.001                | 6.74; 6.33; 6.00 | 14 -63 -52; 18 -50 -21; 27 -56 -20 | Cerebelum_8_R; Cerebelum_4_5_R; Cerebelum_6_R |
| LR ventral anterior, mediodorsal and ventral lateral thalamic nuclei; L ventral posterior lateral thalamic nucleus; LR caudate; L putamen; L insula [posterior short gyrus]; L frontal operculum; R habenular nucleus and habenular commissure; periaqueductal grey; R medial geniculate nucleus, L substantia nigra | 4677         | <0.001                | 5.50; 5.50; 5.49 | -4 -20 12; -15 -15 6; 14 -12 10 | Thalamus_L; Thalamus_L; Thalamus_R |
| R inferior frontal gyrus, opercular part [vIPFC] | 220          | <0.001                | 5.03; 4.51; 4.06 | 58 15 0; 62 14 14; 57 10 8 | Frontal_Inf_Oper_R; Frontal_Inf_Oper_R; Frontal_Inf_Oper_R |
| L substantia nigra               | 35          | <0.001                | 4.76 | -4 -12 -14 | NA |
| LR superior frontal gyrus, medial part [dIPFC / ACC]; LR cingulate gyrus [ACC] | 600          | <0.001                | 4.53; 4.51; 4.49 | 0 42 26; 0 22 32; 4 40 18 | Frontal_Sup_Medial_ LCingulum_Mid_L; Cingulum_Ant_L |
| L cerebellum                     | 40          | <0.001                | 4.41; 3.28 | -10 -54 -36; -2 -52 -39 | Cerebelum_9_L; Cerebelum_9_L |
| R insula [anterior short gyrus]  | 32          | <0.001                | 4.40 | 42 8 2 | Insula_R |
| R inferior frontal gyrus, opercular part [vIPFC]; R precentral gyrus | 187          | <0.001                | 4.24; 4.20; 4.15 | 48 10 22; 60 10 28; 57 6 20 | Frontal_Inf_Oper_R; Precentral_R; Precentral_R |
| **Effect of avoidance**          |             |                       |              |                             |                 |
| L pre- and parasubiculum [anterior hippocampus]/L entorhinal cortex | 10 (SVC)    | 0.012                  | 3.99; 3.81 | -20 -20 -21; -24 -21 -20 | ParaHippocampal_L; NA |
| L anterior CA3/dentate gyrus [anterior hippocampus] (4 mm smoothing kernel) | 12 (SVC)    | 0.039                  | 3.66 | -21 -18 -18 | |
|                                    | L lateral amygdaloid nucleus | 25 | 0.001 | 4.30 | -33 -4 -22 | NA |
|------------------------------------|-----------------------------|----|-------|------|------------|----|

**Threat magnitude - positive linear effect**

| R insula / area orbitoinsularis [anterior and middle short gyrus]; R frontal operculum, R basal operculum | 344 | <0.001 | 4.53; 4.39; 4.29 | 42 18 -2; 36 21 -10; 51 15 -4 | Insula_R; Frontal_Inf_Orb_R; NA |
|---------------------------------------------------------------------------------------------------|-----|---------|----------------|-------------------------------|--------------------------------|
| L middle Hippocampus                                                                           | 11  | 0.004 (SVC) | 4.39 | -27 -26 -12 | Hippocampus_L                  |

**Behavioral response x threat magnitude - positive linear effect (n = 16)**

| R frontal operculum | 21 | 0.002 | 3.63 | 50 20 -3 | Frontal_Inf_Oper_R |
|---------------------|----|-------|------|----------|-------------------|
Table 4.
Parametric analysis (P3): effects of threat features on brain activation, separately for approach and for avoidance trials

| Cluster anatomy (manual lab.) | Cluster size | FWE p-value (cluster) | Peak z-score | Peak coordinates (MNI - mm) | Peak label (AAL) |
|------------------------------|--------------|------------------------|--------------|-----------------------------|------------------|
| Threat magnitude - positive linear effect (separated approach trials) | | |  |  |  |
| R insula [anterior short gyrus] / R inferior frontal gyrus, opercular part | 183 | <0.001 | 4.75; 4.46; 3.66 | 48 -10; 46 -3; 42 14 3 | Frontal_Inf_Orb_R; Insula_R; Frontal_Inf_Oper_R |
| L insula [anterior short gyrus] / L inferior frontal gyrus, opercular part | 30 | 0.001 | 4.26 | -46 16 -4 | Frontal_Inf_Orb_L |
| | 22 | 0.012 | 3.90; 3.82 | -39 20 -6; -33 20 2 | Insula_L |
| LR superior frontal gyrus, medial part [ACC], LR cingulate gyrus [ACC] | 158 | <0.001 | 4.25; 4.23; 4.15 | 0 28; -2 26; 6 38 24 | Cingulum_Ant_L; Cingulum_Ant_R |
| | 39 | <0.001 | 4.06; 3.79 | -2 39 15; 3 44 21 | Cingulum_Ant_L; Cingulum_Ant_R |
| Threat probability x magnitude - linear positive effect (separated approach trials; n = 17) | | |  |  |  |
| L brachium of the inferior colliculus extending into medial geniculate nucleus | 19 | 0.005 | 4.43 | -8 -33 -9 | NA |
| R superior colliculus | 20 | 0.004 | 4.07 | 4 -30 -4 | NA |
Table 5. Region-of-interest analyses in anterior hippocampus and amygdala

| Region          | Effect                | F     | df     | p     |
|-----------------|-----------------------|-------|--------|-------|
| Anterior Hippocampus | A                     | 8.76  | 1, 1214.6 | .003** |
|                  | TP^2                  | 5.46  | 1, 1208.2 | .020*  |
|                  | TM^2                  | 14.22 | 1, 1210.1 | < .001*** |
|                  | TP x H                | 5.65  | 1, 1208.0 | .018*  |
|                  | TP x A                | 5.50  | 1, 1208.9 | .019*  |
|                  | TP x TM x H           | 7.30  | 1, 1208.0 | .007** |
|                  | TP x A x H            | 8.38  | 1, 1208.0 | .004** |
| Amygdala         | H                     | 8.64  | 1, 1207.9 | .003*  |
|                  | TP^2                  | 4.45  | 1, 1208.2 | .035*  |
| Anterior CA1     | A                     | 10.14 | 1, 1222.6 | .001** |
|                  | TP^2                  | 8.88  | 1, 1208.8 | .003** |
|                  | TP x TM x H           | 8.65  | 1, 1208.8 | .003** |
| Anterior CA2/3   | A                     | 10.14 | 1, 1222.6 | .001** |
| Basolateral Amygdala | TM                   | 6.26  | 1, 1210.9 | .012*  |
|                  | H                     | 11.13 | 1, 1207.5 | < .001*** |
| Centrocortical Amygdala | TM^2     | 6.81  | 1, 1208.1 | .009** |
|                  | TM^2 x A              | 9.28  | 1, 1208.4 | .002** |
|                  | TM^2 x H              | 6.77  | 1, 1208.1 | .009** |
| Combined model: Anterior Hippocampus + Amygdala | A | 9.91  | 1, 2434.9 | .002** |
|                  | ROI                   | 13.61 | 1, 2433.9 | < .001*** |
|                  | TM^2                  | 10.23 | 1, 2434.2 | .001** |
|                  | TP x H                | 6.05  | 1, 2433.9 | .014*  |
|                  | A x ROI               | 7.87  | 1, 2433.9 | .005** |
|                  | TM^2 x ROI            | 4.98  | 1, 2433.9 | .026*  |
|                  | TP x TM x H           | 6.82  | 1, 2433.9 | .009** |
|                  | TP x A x H            | 8.50  | 1, 2433.9 | .004** |
|                  | TP x H x ROI          | 4.39  | 1, 2433.9 | .036*  |
|                  | TP x TM x H x ROI     | 4.37  | 1, 2433.9 | .037*  |
|                  | TP x A x H x ROI      | 5.87  | 1, 2433.9 | .015*  |
| Combined model: Anterior CA1 + Anterior CA2/3 | A x ROI | 4.95  | 1, 2670 | .026* |
Figure A: Plot showing the approach proportion as a function of threat magnitude. The data is stratified by probability of threat (P(threat): low, medium, high) across different threat magnitudes.

Figure B: Three subplots illustrating approach latency (middle) and withdrawal latency (right) as a function of threat magnitude, with data points for 500, 250, and 125 trials.
