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Visuocortical changes during a freezing-like state in humans

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Abstract

An adaptive response to threat requires optimized detection of critical sensory cues. This optimization is thought to be aided by freezing - an evolutionarily preserved defensive state of immobility characterized by parasympathetically mediated fear bradycardia and regulated by the amygdala-periaqueductal grey (PAG) circuit. Behavioral observations in humans and animals have suggested that freezing is also a state of enhanced visual sensitivity, particularly for coarse visual information, but the underlying neural mechanisms remain unclear. We induced a freezing-like state in healthy volunteers using threat of electrical shock and measured threat-related changes in both stimulus-independent (baseline) and stimulus-evoked visuocortical activity to low- vs. high-spatial frequency gratings, using functional MRI. As measuring immobility is not feasible in MRI environments, we used fear bradycardia and amygdala-PAG coupling in inferring a freezing-like state. An independent functional localizer and retinotopic mapping were used to assess the retinotopic specificity of visuocortical modulations. We found a threatinduced increase in baseline (stimulus-independent) visuocortical activity that was retinotopically nonspecific, which was accompanied by increased connectivity with the amygdala. A positive correlation between visuocortical activity and fear bradycardia (while controlling for sympathetic activation), and a concomitant increase in amygdala-PAG connectivity, suggest the specificity of these findings for the parasympathetically dominated freezing-like state. Visuocortical responses to gratings were retinotopically specific but did not differ between threat and safe conditions across participants. However, individuals who exhibited better discrimination of low-spatial frequency stimuli showed reduced stimulus-evoked V1 responses under threat. Our findings suggest that a defensive state of freezing involves an integration of preparatory defensive and perceptual changes that is regulated by a common mechanism involving the amygdala.

Keywords: Freezing, functional MRI, visual perception, spatial frequency, fear bradycardia

Highlights

- This study investigated visuocortical changes during freezing in humans
- Threat-induced bradycardia and amygdala-PAG coupling indicated a freezing-like state
- Threat also increased baseline V1 activity and amygdala-V1 coupling
- Reduced V1 responses linked to better perception of coarse features under threat
- Freezing involves an integration of preparatory defensive and perceptual responses

Introduction

Optimal perception of relevant sensory cues is crucial to avoid harm in threatening situations. This perceptual optimization has been suggested to be aided by *freezing* – an evolutionarily preserved defensive response that prepares the body for a successful countering of threat (Blanchard, Griebel, Pobbe, & Blanchard, 2011; Campbell, Wood, & McBride, 1997; Fanselow, 1994; Hagenaars, Oitzl, & Roelofs, 2014; Lojowska, Gladwin, Hermans, & Roelofs, 2015; Öhman & Wiens, 2003). Freezing commonly takes place during threat anticipation and is characterized by bodily immobility and a parasympathetically dominated decrease in heart rate, or *fear bradycardia*, in both animals and humans (Fanselow & Lester, 1988; Gladwin, Hashemi, van Ast, & Roelofs, 2016; Obrist, Wood, & Perez-Reyes, 1965; Roelofs, 2017; Walker & Carrive, 2003). Behavioral observations suggest that freezing is also a state of enhanced visual sensitivity, specifically for coarse visual information (Blanchard et al., 2011; Lojowska et al., 2015). However, despite the potential relevance of these perceptual changes for threat coping, the underlying neural mechanism remains unclear.

The anticipatory state of freezing is characterized by a specific pattern of sympathetic and parasympathetic autonomic nervous system activity. Whereas concurrent sympathetic activation during freezing, as observed for instance in increased pupil size and skin conductance, prepares the body for potential actions (fight-or-flight), parasympathetic dominance serves as a break on the motor system, allowing for threat assessment and decision-making about the most adaptive behavior in given circumstances (Blanchard et al., 2011; Kozlowska, Walker, McLean, & Carrive 2015). Crucially, because these decisions largely depend on the visual characteristics of a threat (e.g., its ambiguity or magnitude), freezing has been conceptualized as a state of enhanced sensory processing during which an animal gathers relevant visual information

(Blanchard et al., 2011; Eilam, 2005; Roelofs, 2017). This has been supported by behavioral observations in rodents which, despite total bodily immobility, show slight head and ear movements suggestive of environmental scanning and increased visual sensitivity (Blanchard et al., 2011). However, whereas much human and animal research has focused on how visual input (e.g., angry faces, conditioned cues) shapes the expression of freezing behavior (Azevedo et al., 2005; Blanchard, Griebel, & Blanchard, 2001; Fanselow & Lester, 1988; Gladwin et al., 2016; Hermans, Henckens, Roelofs, & Fernandez, 2013; Maren, 2001; Roelofs, Hagenaars, & Stins, 2010; Tovote, Fadok, & Luthi, 2015), the neural mechanism underlying the opposite, i.e., how visual processing is altered during the anticipatory state of freezing, remains unclear.

Enhanced visual processing during freezing may be neurally regulated by the amygdala, which is also involved in the expression of freezing. In particular, fear bradycardia and immobility are regulated through amygdalar efferents to (ventral) periaqueductal grey (PAG) and medulla (Koba, Inoue, & Watanabe, 2016; Kozlowska et al., 2015; Tovote et al., 2016; Walker & Carrive, 2003). PAG activation and its functional connectivity with the amygdala have furthermore been linked to fear bradycardia in humans (Hermans et al., 2013). Crucially, non-human primate work has also demonstrated widespread projections from amygdala to early and ventral visual areas (Amaral, Behniea, & Kelly, 2003; Freese & Amaral, 2005), whereas in humans the amygdala drives enhanced visuocortical responses to emotionally salient stimuli (Anderson, 2001; Vuilleumier, Richardson, Armony, Driver, & Dolan, 2004). Based on these findings, we hypothesized that the anticipatory state of freezing should be characterized by a preparatory amygdala-driven enhancement in visuocortical activity, even in the absence of visual input.

In addition, experimental work in humans has shown that freezing is associated with enhanced visual perception specifically of coarse visual information, i.e., low spatial frequencies (Lojowska et al., 2015). This finding is consistent with the observation that the majority of amygdalar projections to the visual cortex are magnocellular and therefore biased toward processing of lower spatial frequency information (Amaral et al., 2003; Freese & Amaral, 2005; Leonova, Pokorny, & Smith, 2003). These anatomical findings are supported by functional evidence where enhanced neural responses to low-relative to high-spatial frequency fearful faces were found in inferior temporal cortex (Vuilleumier, Armony, Driver, & Dolan, 2003), and larger steady-state visual potentials in response to low-spatial frequency stimuli were observed over occipito-temporal areas following presentation of aversive versus neutral images (Song & Keil, 2013). We therefore hypothesized that a state of freezing should also be associated with enhanced early visuocortical responses to coarse visual information.

We conducted a functional magnetic resonance imaging (fMRI) study combined with autonomic nervous system measures in which a transient freezing-like state was induced by means of threat of electric shock during trials with 7-second duration. An equal number of safe trials was included as control. Threat of electrical shock is a commonly used procedure during which the main concomitants of freezing, i.e., bodily immobility, fear bradycardia and amygdala-PAG recruitment have been observed in both animals and humans (Fanselow, 1994; Fanselow & Lester, 1988; Gladwin et al., 2016; Kapp, Frysinger, Gallagher, & Haselton, 1979; LeDoux, Iwata, Cicchetti, & Reis, 1988; Liebman, Mayer, & Liebeskind, 1970; Obrist, 1968; Schneiderman, Smith, Smith, & Gormezano, 1966). As measuring physical immobility in an MRI scanner is not feasible, we relied on fear bradycardia and amygdala-PAG coupling (using psychophysiological interaction models) in validating a freezing-like state in the current study.

To test our hypotheses regarding the effect of threat anticipation on stimulus-independent and stimulus-dependent activity in early visual cortices, we recorded visuocortical activity which we recorded activity in early visual cortices in the absence of visual stimulation (during threat versus safe *Omission* trials, in which no grating stimuli were presented), as well as in response to peripherally presented gratings (during threat versus safe Grating trials, in which maximally three gratings were presented). Omission trials were used to assess an increase in baseline visuocortical activity without the confounding effect of stimulus presentation, and its modulation by the amygdala. Grating trials were used to examine transient neural responses to relatively low (3 cpd) compared to relatively high (6 cpd) spatial frequencies. We carried out an independent localizer and retinotopic mapping to allow for quantification of retinotopically specific (i.e., stimulus location-selective) and nonspecific activity, within V1, V2, and V3. We first predicted that if freezing is associated with an amygdala-driven upregulation of baseline activity, we should observe a threat-related increase in visuocortical activity and its functional coupling with the amygdala during Omission trials. Second, if freezing is associated with enhanced early visuocortical responses to coarse visual information, larger stimulus-evoked activity should be observed to low-spatial frequency gratings on threat versus safe conditions within retinotopically specific voxels in early visual areas.

Materials and Methods

Participants

Forty-eight participants participated in this study. Inclusion criteria were: normal or corrected-tonormal vision, no color blindness, no history of psychiatric or neurological treatment, no cardiovascular conditions and no smoking, right-handedness. Fourteen participants were excluded because of not completing the full task (4), inability to tolerate shocks (3, comparable with our previous study with shocks; Lojowska et al., 2015), falling asleep (2, during low demanding tasks of retinotopy and localizer), feeling uncomfortable inside the scanner (2, firsttime MRI participants), arrhythmia detected during data acquisition (1), and performance accuracy below 75% during the tilt titration task before the actual experiment (2, comparable with the previous behavioral study; Lojowska et al., 2015). The final sample of 34 participants consisted of 19 males and 15 females with the mean age of 23.15 years (range 19-32). Additionally, we removed behavioral outliers, i.e., >3 SD from the mean (maximum of two participants) or tilt offset >20 degrees (which is unlikely to reflect visual sensitivity; maximum of two participants), for all relevant statistical tests. Before participating in the experiment, all participants were required to grant written informed consent. Participants were reimbursed with 40 euros for their participation in both sessions of the study. This study was approved by the local ethical review board (CMO Region Arnhem-Nijmegen, The Netherlands).

Sample size calculation (G*Pwer; Faul, Erdfelder, Lang, & Buchner, 2007) revealed that 34 participants were sufficient to detect a medium effect size (for interaction effect threat x SF, $\eta_p^2 = 0.06$) at 80% power. Although only 12 participants were necessary to replicate a relatively large effect size ($\eta_p^2 = .17$) found previously for this interaction effect in a similar behavioral

study (Lojowska et al., 2015), we nevertheless used the same sample size as in this study (34 participants) in order to increase the chance of detecting possibly smaller effects arising from differences between the tasks (e.g., in testing environment).

Stimuli and apparatus

Stimuli. Gabor grating stimuli (Gaussian-enveloped sinusoidal gratings) were generated using MATLAB R2010a, in combination with the Psychophysics Toolbox Stimuli (Brainard, 1997) were displayed on a rear-projection screen using a luminance-calibrated EIKI projector (1024 x 768 resolution, 60 Hz refresh rate) located 75 cm behind the participant's head. Gratings (4.5° x 4.5° size) were presented at 100% Michelson contrast and in two spatial frequencies: 3 cycles per degree (cpd) and 6 cpd. These frequencies were used because they fall within the sensitive range of magnocellular and parvocellular cells (Leonova et al., 2003), and because the modulatory effect of threat anticipation was previously found for those frequencies (Lojowska et al., 2015). The gratings were displayed at 4° eccentricity from the central fixation. A fixation point (size: 0.15°, luminance: 116 cd/m²) was displayed in the center of a uniform gray background screen (luminance: 76 cd/m2), and participants were asked to maintain fixation through the course of the experiment. During the visual task, this fixation point could change from grey to orange (luminance: 116 cd/m²) or blue (luminance: 117 cd/m²), signaling threat or safe conditions. Stimulus parameters were chosen based on the following considerations. Peripheral display was used to avoid ceiling effects, to allow for delineation of modulation across visual areas, and to increase ecological validity of the task, as peripheral detection may be particularly adaptive in threatening situations, where not all relevant information (e.g., predator approaching from behind) is immediately accessible to overt attention. 4° eccentricity was used to minimize the bias in spatial frequency processing due to uneven distribution of cones and rods

in retina (Purves et al., 2001) and was based on previous psychophysiological work in which discrimination performance for 3 and 6 cpd was comparable at this eccentricity (Rovamo, Virsu, & Näsänen, 1978). Furthermore, horizontal presentation was used to avoid a potential difference in spatial frequency processing between upper and lower parts of the visual field (Christman, 1993). Finally, by using the same stimulus parameters (eccentricity, peripheral horizontal presentation, and spatial frequencies) as in previous studies in which threat-induced modulation on spatial frequency processing was found (Bocanegra & Zeelenberg, 2009; Lojowska et al., 2015; Nicol, Perrotta, Caliciuri, & Wachowiak, 2013), we hoped to replicate these behavioral effects in the current study. In the visual task, the maximal tilt offset the gratings could deviate from the vertical orientation was set to 40 degrees. Contrast, size and presentation time of the gratings (100 ms) were chosen to ensure their full visibility while rear-projected in the scanner during the visual task, and to prevent participants from making saccadic eye movements towards the stimuli, which would require about 160 ms (Walker, Deubel, Schneider, & Findlay, 1997).

Shock administration and physiological measurements. Electric shocks were delivered transcutaneously through the participant's fourth and fifth distal phalanges of the right hand using a 9V battery-operated MAXTENS 2000 (Bio-Protech, Inc., Wonju, Korea) and standard Ag/AgCl electrodes. The duration of the electrical stimulation was 200 ms, with a 150 Hz repetition of 250 μ s pulses. The intensity varied between 0-40V/0-80mA across 500 Ω with 10 steps in total. Shock intensity was adjusted at the individual level to ensure the shocks were unpleasant but not painful. Shock calibration was performed using a standardized staircase procedure comprising 5 shock presentations after each of which the shock intensity was adjusted according to the participant's verbal reports of its unpleasantness on a scale from 1 (not

unpleasant) to 5 (very unpleasant). The final shock intensity obtained with this method (mean score: 4.36, SD = .65) was used in the visual orientation discrimination task.

The following physiological measures were recorded: heart rate (HR), pupil size and respiration in both sessions, and skin conductance in the second session when the visual task was carried out. HR was used for offline assessment of the parasympathetically driven decelerative HR responses, whereas pupil size and skin conductance index sympathetic activity during the task (Hermans et al., 2013). Heart rate and respiration were additionally used for retrospective image-based correction (RETROICOR) of physiological noise artifacts in BOLD signal in fMRI data (Glover, Li, & Ress, 2000). HR, skin conductance and respiration were acquired using an MR-compatible BrainAmp EXG MR 16 channel recording system (Brain Products, Gilching, Germany). HR was measured using a pulse oximeter that was attached to a fourth distal phalange of the participants' left hand. SCR data were collected with two standard Ag/AgCl electrodes attached to the second and third distal phalanges of the participant's left hand. Respiration data were collected using a respiration belt positioned around the participant's abdomen. Pupil size was measured using a 50 Hz iView system with an MR-compatible eye tracker (MEye Track-LR camera unit, SMI, SensoMotoric Instruments).

MRI data acquisition. MRI data were acquired at the Donders Centre for Cognitive Neuroimaging using a 3T Siemens MAGNETOM Skyra MRI scanner (Siemens, Erlangen, Germany). Functional MRI data were acquired using a T2*-weighted gradient-echo EPI sequence and parallel multiband excitation with an acceleration factor of four (TR = 909 ms, TE = 24.6 ms, 60 axial slices acquired in interleaved order, flip angle = 59°, field of view = 212*212 mm, slice thickness = 2 mm with no gap between slices, voxel size = 2 mm isotropic, GRAPPA in-plane acceleration factor of 2). Furthermore, field maps were acquired for each participant to

allow for correction of spatial distortions in the BOLD-fMRI images due to magnetic field inhomogeneity (TR = 1020 ms; TE1 = 10 ms; TE2 = 12.46 ms; flip angle = 90° ; slice matrix size = 64*64; field of view = 224*224 mm). A high-resolution whole-brain anatomical image was acquired using a T1-weighted Magnetization-Prepared RApid Gradient Echo (MP-RAGE) sequence (TR = 2300 ms; TE = 315 ms; flip angle = 8° ; slice matrix size = 256*256; field of view = 256*256*180 mm; voxel size = 0.8 mm isotropic).

Experimental procedure

The entire experiment was divided over two sessions, scheduled 3-4 days apart. The first session consisted of the functional localizer task, retinotopic mapping task, as well as resting state and susceptibility-weighted imaging sequence not used in the current study. The second session consisted of questionnaires, a tilt titration task, a shock calibration procedure and the visual task with electric shocks (i.e., the task of primary interest). Participants lay in the scanner in a head-first supine position and viewed the back-projection screen via a mirror mounted on top of the head coil.

The first session started with the functional localizer task. This task comprised flickering gratings presented at 100% contrast in 16 blocks of 38 s each. In each block, a grating (either 3 cpd or 6 cpd) were presented at 4° eccentricity for 9.5 s and with a random phase change between the blocks. The orientation of the gratings was fixed within, but varied between the blocks. We included the following tilt offsets: +/-3, +/-6, +/-10, +/-14. Blocks with each orientation were repeated twice. Blocks were separated by a 9.5s grey screen containing only a fixation at the center of the screen. To ensure central fixation during the task, the white fixation in the center of the screen turned black at pseudorandom time points, which prompted a button-press response from participants. The localizer task lasted approximately 14 minutes.

During the retinotopic mapping task, participants viewed a wedge consisting of a flashing black-and-white checkerboard pattern (3 Hz), first rotating clockwise for 6 cycles and then anticlockwise for another 6 cycles (at a rotation speed of 36s/cycle). Also during this task, the central fixation changed color from white to black, prompting a button-press response from participants to ensure central fixation.

The second session started with participants completing questionnaires, which were included to enable post-hoc assessment of whether participant samples of our previous behavioral and current fMRI study were comparable (Lojowska, 2015). During the tilt titration task, tilt offset was titrated separately for the 3 cpd and 6 cpd stimuli in the same visual orientation task, but without shocks and omission trials. The final staircase thresholds obtained in this task were subsequently used as a starting tilt offset for gratings in the actual experiment (one threshold for 3 cpd stimuli under threat and safe conditions, and one threshold for 6 cpd stimuli under threat and safe conditions). The tilt titration task consisted of 72 gratings in each spatial frequency, taking about 10 minutes, and was administered in the scanner during the acquisition of the anatomical scan. Subsequently, to adjust shock intensity to individual sensitivity thresholds, participants underwent a shock calibration procedure. This was followed by visual task lasting about 50 - 60 minutes, with short breaks in between.

Visual task. Each trial started with a 7 s presentation of the central fixation point, the color of which – blue or orange - indicated either threat or safe conditions (Figure 1). Color fixation was followed by an ITI of 2-4 s, during which the fixation color changed to grey. Freezing was evoked in the threat condition by a chance of receiving an electric shock, administered to the participant's fourth and fifth distal phalanges of the right hand. No shocks were administered during the safe condition. Participants received verbal instructions about the

association between fixation color and shocks before the start of the experiment. The association between fixation color and threat condition was counterbalanced across participants. 25% of all trials were omission trials, during which only the central fixation was presented. These trials were used to quantify threat-related changes in baseline visual activity. The remaining trials were used to probe threat-related changes in neural visual responses to visual information. To this end, participants performed a visual orientation discrimination task (Lojowska et al., 2015) that required discrimination of the orientation of the gratings under orthogonalized threat and safe conditions. These trials were the same as omission trials with the exception of gratings presented either to the left or right side of the fixation. In total, there were four conditions: threat 3 cpd, threat 6 cpd, safe 3 cpd, and safe 6 cpd. Gratings were presented only during the display of a threat cue, i.e., orange or blue fixation, for the duration of 100 ms. In a single trial, up to 3 gratings could be presented. The minimum interstimulus interval (ISI) was set to 2 s, allowing for separation of BOLD responses to each grating (Wager & Nichols, 2003). Participants were required to indicate through button press whether the orientation of the grating was tilted to the left or right, relative to vertical. The presentation side, spatial frequency of the gratings and their orientation (left versus right) were counterbalanced between sessions and participants. Participants were instructed to fixate on the central dot during the whole task and avoid excessive blinking when the orange or blue fixation was presented.

To ensure that neural responses to the gratings reflected threat-related manipulations, rather than task difficulty known to increase activity in the visual cortex (Chen et al., 2008), we used an adaptive staircase procedure (QUEST; Watson & Pelli, 1983) to maintain an overall level of 75% accuracy while varying tilt offset in each of the four conditions. This procedure was

applied separately for each of the four conditions and the resulting changes in the tilt offsets were used as measures of behavioral performance.

The task was divided into three runs, each consisting of 102 trials (36 safe/threat trials with gratings, 12 safe/threat omission trials and 6 threat trials with shocks). In each run, 36 gratings belonging to each of the four conditions were presented in grating trials. Trials with shock presentation were excluded from the analyses. The trial sequence was fully randomized for each participant to ensure that potential carry-over effects between trial types could not result in a systematic bias. Total duration of the second session was about 2 hours. After MRI scanning, participants were asked debriefing questions to assure they understood all instructions (e.g., regarding the association between the fixation color and chance of receiving a shock), and to verify the experimental manipulations (e.g., whether they received shocks and how many, and about the perceived averseness of the shocks).

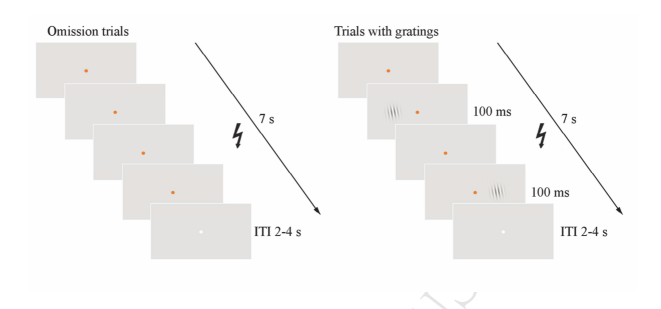


Figure 1. Visual task. The task consisted of omission trials and trials with gratings. Threat and safe conditions were signaled by fixation color, i.e., blue or orange, counterbalanced across participants. The figure illustrates an example of a threat condition trial, here signaled by an orange fixation, during which electric shocks could be delivered.

Data analysis

Behavioral and physiological analyses. Behavioral performance was defined as the mean tilt offset required to perform the orientation discrimination task at 75% accuracy, for each of the 4 conditions. Data was calculated for each of the runs and analyzed using SPSS (IBM SPSS Statistics for Windows, Version 21.0), with run number (1-3), threat condition and spatial frequency as within subject factors. Data of four participants were excluded from the behavioral analysis because of tilt offsets exceeding outlier thresholds (larger than 3 SD when tested separately for 3 cpd and 6 cpd).

The analysis of physiological data was performed offline using in-house software implemented in MATLAB (Matlab, R2013a, The MathWorks, Inc), allowing for visual assessment and removal of signal artifacts. Blinks in pupil data were removed from the signal using linear interpolation. HR data for 6 subjects were discarded due to extensive amount of noise and artifacts (e.g., pulse oximeter fell off the finger during scanning, excessive hand movement, scanner artifacts, poor signal). For similar reasons, skin conductance data of two participants were also removed from the analysis. To assess parasympathetic and sympathetic activity during the visual task, pre-trial baseline-corrected HR responses, pupil size responses, and skin conductance levels (SCL) were calculated for each trial. HR responses were quantified by calculating the mean HR between 2-7s following trial onsets corrected for the baseline (mean HR during 25 s prior to stimulus onset). This relatively long baseline window was chosen to account for fluctuations of HR with the respiratory cycle due to respiratory sinus arrhythmia (De Geus, Willemsen, Klaver, & Van Doornen, 1995). Baseline-corrected pupil size responses were quantified by subtracting the averaged pupil diameter within a 1 s period prior to trial onset (baseline) from the averaged pupil diameter within a period from 2-7 s following trial onset. Changes in SCLs were calculated using the same time windows. We reasoned that SCL change is a more appropriate measure of sympathetic arousal in the current block design than conventional scoring of skin conductance response magnitudes commonly used in event-related designs (Phelps et al., 2001). The difference between HR responses on threat vs. safe conditions (across both grating and omission trials) were analyzed using paired sample t-test. As SCLs and pupil responses did not have a normal distribution (Shapiro-Wilk test p-value < .001), the difference between threat and safe conditions for these measures were assessed using a nonparametric permutation t-test (with 50,000 permutations).

Because of expected physiological changes during our task (due to threat modulation), we used physiological noise correction in all our fMRI analyses to remove variance in BOLD signal originating from these fluctuations. In this way, any differences between threat and safe trials are more likely to reflect the underlying changes in task-related neural responses rather than physiological fluctuations. We included 10 nuisance regressors for cardiac noise and 10 for respiratory noise, which were specified by calculating fifth-order Fourier models of the cardiac and respiratory phase-related modulation of the BOLD signal (Glover et al., 2000). In addition to these RETROICOR regressors, we calculated regressors for heart rate frequency, heart rate variability, (raw) abdominal circumference, respiratory frequency, respiratory amplitude, and respiration volume per unit time (Birn, Diamond, Smith, & Bandettini, 2006; van Buuren et al., 2009), yielding a total of 26 nuisance regressors that were subsequently used as regressors of no interests in GLM models.

fMRI data preprocessing. The fMRI data were spatially preprocessed using SPM8 (www.fil.ion.ucl.ac.uk/spm) for analysis in native space. The first five volumes of each session were discarded to allow for T1 equilibration. Field maps were subsequently used to calculate a voxel displacement map (Hutton et al., 2002) for each session which was used for the realignment and dynamic unwarping of functional images. Functional images were realigned to the first image within and across all three runs, yielding head movement parameters (translations and rotations) that were subsequently used as nuisance regressors in the general linear models (GLMs). The mean realigned and unwarped functional image was used to coregister functional images with the T1 image for each participant. Furthermore, inverse normalization parameters were created by segmenting the T1 image into grey matter, white matter and CSF images using a unified probabilistic template registration and tissue classification method (Ashburner & Friston,

2005). These inverse normalization parameters were used to warp MNI-space masks of regions of interest into native space. All subsequent statistical analyses were performed in native space (i.e., without spatial normalization) and without spatial smoothing.

Functional localizer and regions of interest. To find retinotopically specific voxels in the left and right hemispheres of V1, block regressors for right and left stimulus presentations were constructed and convolved with a canonical hemodynamic response function (HRF). Additional HRF-convolved stick function regressors modeled pseudorandom flickering of the fixation point, the aim of which was to prevented participants from looking at the gratings. Left and right hemisphere responses were found by respective contrasting of beta maps for right vs. left, and left vs. right stimulus presentation. Activations within V1 were defined using the retinotopic boundaries of V1, delineated based on the retinotopy task and well-established methods (DeYoe et al., 1996; Engel, Glover, & Wandell, 1997; Sereno et al., 1995) implemented in Freesurfer (http://surfer.nmr.mgh.harvard.edu/). 140 voxels that responded maximally to the contralateral stimulus presentation and belonged to the same cluster (i.e., showing spatial contiguity) were selected for left and right V1, and subsequently used as retinotopically specific ROIs in the analysis of the visual task. 140 voxels corresponded approximately to an activation threshold of p < 0.05, uncorrected, using a one-tailed t-test. To examine the retinotopic specificity on neural responses in V1, retinotopically nonspecific ROIs were created by subtracting retinotopically specific ROIs created at a lenient threshold of p < 0.2 (uncorrected) from the V1 masks created based on the retinotopy task. Left and right amygdala were defined in native space using automated anatomical segmentation of T1-weighted images in FSL FIRST (http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/FIRST). PAG voxels were delineated on each participant's high-resolution T1 scan using fslview and according to previously described guidelines (Satpute

et al., 2013). The resulting PAG ROI had a shape of a hollow cylinder with length of about 10 mm, external diameter of about 6 mm and internal diameter of about 2 mm. The masks were then converted from 4D to 3D space, resliced to each participant's native space of functional images.

Visual task. For analyses of neural responses in the visual task, two separate finite impulse response (FIR) models were created for omission and grating trials. In the model for omission trials, we modeled block responses to threat and safe trials. This model was used to test changes in baseline visuocortical activity on threat relative to safe conditions without the confounding effects of visual stimulation. For each run, regressors for threat and omission trials with 14 time bins (TRs) were created starting with the trial onset. Additional 3 regressors of no interest were created for grating threat and safe trials, and threat trials with shocks. For each participant and condition of interest, parameter estimates were extracted from an a priori defined period between 4.5 and 9 s where the peak BOLD response was expected (5 – 10 TR post-trial onset) for the following visual ROIs: retinotopically specific and nonspecific V1 voxels contralateral and ipsilateral to stimulus presentation. This period allowed for capturing trialspecific BOLD responses without interfering with subsequent trials which - with minimum ITI of 2 s - could start at 9 s following the onset of the previous trial. Visual inspection of the BOLD signal averaged across threat and safe conditions confirmed that this period captured the BOLD response peak occurring at around 6.5 s following stimulus onset. Changes in baseline V1 activity were tested in a three-way repeated-measures ANOVA with threat condition (threat, safe), retinotopic space (retinotopically specific versus retinotopically nonspecific voxels in V1) and run number (1-3) as within-subject factors. We additionally analyzed the difference in BOLD responses between threat and safe trials for the amygdala and PAG to check whether any

threat-induced changes in functional connectivity (see below) are associated with a concurrent overall increase in activation of these regions.

In the second FIR model, we estimated neural responses time-locked to the presentation of 3 cpd and 6 cpd gratings under threat and safe conditions. For each run, eight regressors of interest were created representing the following conditions: threat 3 cpd, threat 6 cpd, safe 3 cpd, and safe 6 cpd, for left and right presentations. Gratings to which no responses were given and shock trials were modeled by additional FIR regressors. For each participant and condition of interest, parameter estimates in a period from 2.7 to 5.4 s following stimulus onset were extracted. This period allowed for capturing stimulus-evoked BOLD responses peaking around 3-5 s following stimulus onset. This selection was based on visual inspection of the average of BOLD responses across all conditions (retinotopically specific and nonspecific voxels, ipsilateral and contralateral to stimulus presentation), and therefore was orthogonal to the results of interests. To validate localizer procedure (i.e., larger BOLD responses were expected in the retinotopically specific compared to nonspecific voxels contralateral to stimulus presentation side) and to assess threat-related changed in V1 responses to 3 cpd gratings, averaged parameter estimates were analyzed with a repeated measured ANOVA with stimulated hemisphere (contralateral versus ipsilateral relative to stimulus presentation), retinotopic space (retinotopically specific versus nonspecific voxels in V1), run number (1-3), threat condition (threat, safe) and SF (3 and 6 cpd) as within-subject factors. In each of these models, we included head motion parameters (3 translations, 3 rotations) from realignment, 26 physiological noise regressors, high-pass filtering (1/128 Hz cut-off) to remove low-frequency signal drifts, and AR (1) serial correlation correction.

Effective connectivity analysis. To test whether threat of shock leads to the predicted changes in connectivity between the amygdala with V1 and PAG, we performed a psychophysical interaction (PPI) analysis (Gitelman, Penny, Ashburner, & Friston, 2003). We used voxels in the left and right amygdala separately as seed regions, which were defined by individually created masks in native space (see above). We calculated the first eigenvariates of the time series within left and right amygdala for each participant and run. After deconvolution, these were multiplied by a psychological component (threat vs. safe omission trial regressor) to obtain a run-specific PPI term, which was subsequently re-convolved with the HRF. A GLM with three regressors for each session (PPI term, seed-region eigenvariate, and psychological component) was conducted to calculate participant-specific parameter estimate maps for the PPI term. These parameter estimates were averaged across retinotopically specific and retinotopically nonspecific voxels within V1, as well as within PAG. The resulting values, which represent participant-specific estimates of threat-related changes in connectivity, were tested for statistical significance across participants using a repeated measures ANOVA for each ROI.

Results

Peripheral physiological responses

We first analyzed peripheral physiological responses to verify the effectiveness of our threat manipulation. First, we found a significantly lower heart rate on threat versus safe conditions, t(1,27) = 5.56, p < 001, $\eta_p^2 = .54$, verifying that our threat-of-shock manipulation evoked fear bradycardia. Furthermore, we also found a significant increase in pupil dilation, t(1,33) = 5.09, p < .001, $\eta_p^2 = .44$, as well as SCL, t(1,31) = 3.81, p = .001, $\eta_p^2 = .32$, indicating the concurrent presence of sympathetic activation during threat conditions. However, given the long duration of the task (approximately an hour), there was a risk of habituation effects with parasympathetic and sympathetic responses decreasing over time. Indeed, a repeated-measures ANOVA with run number as a within-subject factor showed a main effect of run number on differential scores (threat – safe) of HR responses, F(2,54) = 9.12, p < .001, $\eta_p^2 = .25$, pupil dilation responses, $F(2,66) = 27.08, p < .001, \eta_p^2 = .45, \text{ and SCLs}, F(2,62) = 24.67, p < .001, \eta_p^2 = .44 \text{ (Figure 2)}. A$ further planned linear contrast revealed a significant linear trend for all physiological responses, namely HR responses F(1,27) = 10.04, p = .004, $\eta_p^2 = .27$, pupil dilation responses, F(1,33) =61.90, p < .001, $\eta_p^2 = .65$, and SCRs, F(1,31) = 41.31, p < .001, $\eta_p^2 = .57$, confirming that the magnitude of both sympathetic and parasympathetic responses to threat relative to safe conditions decreased over time. Habituation of physiological responses to threat of shock is accompanied by a reduction of subjectively perceived shock aversiveness toward the end of the task (average rating of shock aversiveness = 2.88 on a scale from 1 to 5) compared to during shock calibration (M=4.38). Because these manipulation checks indicate that threat of shock was successful and had the strongest effect in the first run, we included run number as a factor in all further analyses.

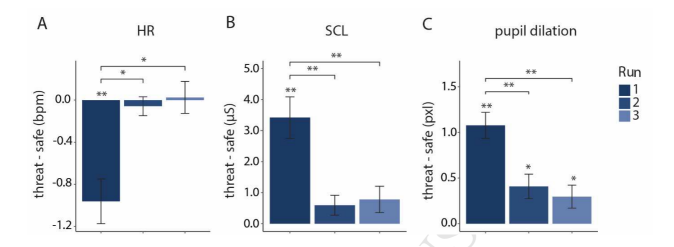


Figure 2. Physiological responses represented by the difference in baseline-corrected responses on threat versus safe trials for each of the three runs of the task. Differences between threat and safe conditions were strongest in run 1, with a (parasympathetically controlled) reduction in heart rate (HR) and (sympathetically controlled) increases in skin conductance levels (SCL) and pupil dilation. bpm, beats per minute; μ S, microSiemens, pxl, pixels; **, p < .001; *, p < .05

Behavioral results

To test if discrimination of the gratings differed as a function of threat condition, we conducted a 2 (threat condition: threat, safe) x 2 (spatial frequency: 3 cpd, 6 cpd) x 3 (run number, 1-3) repeated-measures ANOVA with tilt offset as dependent variable. First, we found a main effect of spatial frequency, F(1,29) = 12.18, p = .002, $\eta_p^2 = .30$, with 3 cpd gratings (mean tilt offset = 2.12) discriminated better than the 6 cpd gratings (mean tilt offset = 3.28) which conforms with generally higher sensitivity to lower spatial frequencies when presented in the periphery (Díez-Ajenjo & Capilla, 2010; Rovamo et al., 1978). There was no main effect of threat, F(1,29) = 2.06,

p=.16, nor an interaction between threat condition and spatial frequency, F(1,29)=3.08, p=.09. However, there was a significant interaction between threat condition, spatial frequency, and run number, F(2,58)=4.02, p=.023, $\eta_p^2=.12$, indicating time-dependent differences in the effects of threat on tilt offset between the gratings.

Given that we used an adaptive staircase procedure (QUEST) that started at the same tilt offset for both threat conditions within each spatial frequency, threat-related changes could become evident as a difference in tilt offset between the two threat conditions building up over time within each spatial frequency (Table 1). We tested this by examining the linear trend across runs within the interaction between threat condition and run number for each spatial frequency. Indeed, a steeper linear trend for threat versus safe conditions was found for 3 cpd stimulus, F(1,29) = 6.14, p = .019, $\eta_p^2 = .17$, but not for the 6 cpd stimulus, F(1,29) = 2.00, p = .17, $\eta_p^2 = .065$, indicating that, as expected, the decrease in tilt offset (i.e., increase in performance) for the 3 cpd stimulus was steeper in the threat condition than in the safe condition.

Table 1. Means and standard deviation (in parentheses) of tilt offset for 3 cpd (LSF) and 6 cpd (HSF) gratings under threat and safe conditions across runs.

SF	Threat condition	Run 1	Run 2	Run 3
3 cpd (LSF)	threat	2.22(1.10)	2.09(.96)	2.03(.92)
	safe	2.21(1.22)	2.11(1.05)	2.08(1.06)
6 cpd (HSF)	threat	3.32(2.57)	3.11(2.29)	3.10(2.31)
	safe	3.43(2.92)	3.42(2.90)	3.31(2.82)

Neuroimaging results

Threat effects on baseline V1 activity. To test our first (stimulus-independent) hypothesis that freezing is a state associated with an increase in baseline V1 activity, we compared the amplitude of BOLD responses in V1 on threat vs. safe trials during omission trials where no gratings were presented. To assess retinotopic specificity of these changes, we further compared the response of retinotopically specific (i.e., stimulus-responsive) and nonspecific voxels within V1. A three-way repeated-measures ANOVA with threat condition (threat, safe), retinotopic space (retinotopically specific versus retinotopically nonspecific voxels in V1) and run number (1-3) as within-subject factors revealed no significant main effect of threat condition, F(1,33) = 1.25, p = .27, $\eta_p^2 = .19$. However, the effect of threat on the activation of V1 differed between runs, as indicated by a significant interaction between threat condition and run number, F(2,66) = 3.28, p = .044, $\eta_p^2 = .60$ (Figure 3 A). There was a significantly higher baseline V1 activity in threat versus safe conditions in run 1, F(1,33) = 6.50, p = .016, $\eta_p^2 = .16$, but not in run 2 and run 3 (ps > .05), conforming to the pattern of physiological habituation. A follow-up analysis for run 1 revealed that significantly higher baseline activity was observed in both retinotopically specific, t(33) = 2.34, p = .026, $\eta_p^2 = .14$, and nonspecific voxels, t(33) = 2.73, p= .010, η_p^2 = .18, and that the difference in threat effects between the two retinotopic areas was nonsignificant [interaction between threat condition and retinotopic areas: F(1,33) = 2.08, p = .15, $\eta_p^2 = .06$]. Thus, the observed threat-related increase in baseline V1 activity appears to be retinotopically nonspecific. A summary of the brain regions from a whole-brain analysis for run 1 (where threat-related effects were found in the above native-space FIR-model analysis) are reported in the Supplementary Material.

Next, to test if the threat-induced increase in baseline V1 activity is associated with functional coupling with the amygdala, we performed a PPI analysis between these two regions. Threat-related changes in connectivity strength were assessed by averaging parameter estimates for a PPI term expressing the threat-related change in connectivity with the amygdala for two regions of interest: retinotopically specific V1 voxels and retinotopically nonspecific V1 voxels. The resulting participant-specific averages were tested using a repeated-measures ANOVA with retinotopic space (retinotopically specific versus nonspecific V1 voxels), run number (1-3), seed region (left versus right amygdala), and hemisphere (left, right V1) as within-subject factors. As predicted, we found that threat of shock led to an increase in connectivity between the amygdala and V1, F(1,33) = 19.33, p < .001, $\eta_p^2 = .37$ (Figure 4). This threat-induced increase in connectivity did not differ between retinotopically specific and nonspecific voxels, F(1,33) = $3.02, p = .092, \eta_p^2 = .084, \text{ or between runs}, F(2,66) = .83, p = .44, \eta_p^2 = .025. \text{ A follow-up}$ analysis confirmed that the threat-related increase in connectivity was indeed observed for both retinotopically specific [one sample t-test: t(33) = 4.40, p < .001, $\eta_p^2 = .37$], and nonspecific voxels [one-sample t-test: t(33) = 4.16, p < .001, $\eta_p^2 = .34$]. We also found no interaction between these two factors, F(2,66) = 1.92, p = .15, $\eta_p^2 = .055$. These findings indicate that the observed connectivity increases were neither retinotopically specific nor time dependent. Taken together, threat of shock resulted in a retinotopically nonspecific increase of baseline activity in early visuocortical areas which was associated with an increase in functional coupling with the amygdala.

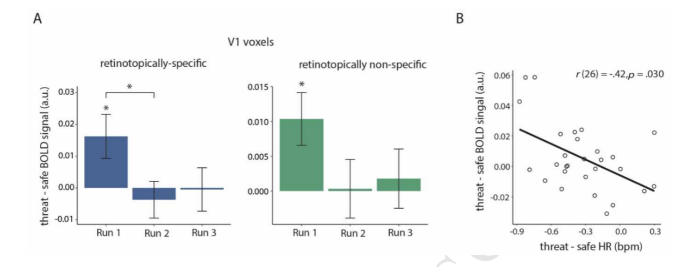


Figure 3. A. Neural responses in V1 in threat versus safe omission trials as a function of run number. Results show averaged BOLD responses across left and right V1, within retinotopically specific (blue) and nonspecific (green) V1 voxels. B. Correlation between fear bradycardia, threat-related reduction in heart rate (HR) and BOLD responses (retinotopically specific and nonspecific) in V1 in threat versus safe conditions. Error bars represent standard errors of the mean. bpm, beats per minutes; *, p < .05.

Relationship between visuocortical changes and defensive responses to threat. To test the hypothesis that physiological and perceptual changes are part of an integrated defensive response, we first tested whether the magnitude of change in BOLD responses between threat and safe trials correlated with threat-induced changes in heart rate (combined across both Grating and Omission trials). We found a significant negative correlation between threat-induced heart rate change and change in baseline activity in V1 (r(26) = -.41, p = .030; Figure 3 B). Crucially, this correlation remained significant when controlling for variance associated with sympathetic responses, i.e., threat-induced changes in SCLs and pupil dilation, r(22) = -.42, p = .033 (tested

across all runs). Furthermore, there was no significant correlation between threat-related responses in V1 and either SCL, $\rho(30) = -.15$, p = .40, or pupil dilation responses, $\rho(32) = -.19$, p = .26. Thus, threat-induced increases in V1 activity were stronger in participants who demonstrated stronger fear bradycardia, and these associations remained when controlling for concurrent sympathetic activation.

Second, we tested whether the observed increase in functional connectivity between the amygdala and visuocortical areas was accompanied by an increase in functional connectivity between the amygdala and PAG (Hermans et al., 2013). Threat-related changes in connectivity strength between the amygdala and PAG were assessed by averaging the parameter estimates for a PPI term for the PAG. In line with previous reports (Hermans et al., 2013), a repeated-measures ANOVA with threat condition, run number and seed region (left, right amygdala) as within subject factors revealed a threat-related increase in connectivity between the amygdala and PAG, F(1,33) = 7.66, p = .009, $\eta_p^2 = .18$, which did not differ between runs, F(1,33) = .93, p = .40, $\eta_p^2 = .027$ (Figure 4). Together, the correlation between threat-related changes in baseline V1 activity and fear bradycardia, and the increase in functional connectivity of the amygdala with both V1 and PAG supports the notion of integrated visual and defensive responses.

Additional control analyses revealed a significant interaction between threat condition and run number for amygdala activation, F(2,66) = 4.87, p = .011, $\eta_p^2 = .13$, with a significant difference present only in the first run [run1: t(33) = 3.43, p = .002; run2: t(33) = -.26, p = .79; run 3: t(33) = -.35, p = .73]. No main effect of threat condition, F(1,33) = .072, p = .79, $\eta_p^2 = .002$, or an interaction between threat and run number, F(2,66) = .20, p = .82, $\eta_p^2 = .006$, was observed for PAG activation.

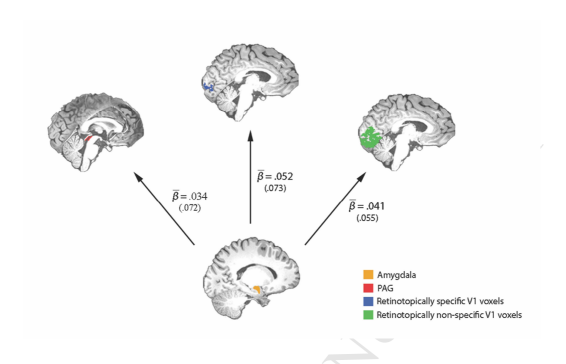


Figure 4. Amygdala (depicted in yellow) and its functional coupling with retinotopically specific (blue) and nonspecific (green) V1 voxels, and periaqueductal grey (PAG, in red). Beta values represent average parameter estimates for a PPI term expressing threat-related changes in connectivity with the amygdala for the three regions of interests. Values in parentheses represent standard deviations. All values were significantly different from zero (all p < .05).

Transient V1 responses to low and high spatial frequency gratings under threat. In order to address our second main hypothesis concerning stimulus-dependent visuocortical activity, we tested whether low-spatial frequency gratings were associated with increased stimulus-evoked activity within retinotopically specific voxels under threat in V1. To this end, we first validated our localizer procedure. A repeated-measures ANOVA with stimulated hemisphere (contralateral versus ipsilateral relative to stimulus presentation), retinotopic space (retinotopically specific versus nonspecific voxels in V1), run number (1-3), threat condition

(threat, safe) and spatial frequency (3 and 6 cpd) as within-subject factors revealed a significant interaction between stimulated hemisphere and retinotopic space, F(1,29) = 37.18, p < .001, $\eta_p^2 = .56$. This interaction was driven by significantly larger BOLD responses in retinotopically specific versus retinotopically nonspecific voxels contralateral to stimulus presentation, F(1,29) = 24.12, p < .001, $\eta_p^2 = .45$, but not on the ipsilateral side, F(1,29) = 3.46, p = .073, $\eta_p^2 = .11$ (Figure 5). These results confirm that the localizer procedure was successful.

Next, we examined the amplitude of BOLD responses to low- and high-spatial frequency gratings in retinotopically specific V1 voxels. A repeated-measures ANOVA with threat condition, spatial frequency and run number revealed a significant main effect of SF, F(1,29) = 4.93, p = .034, $\eta_p^2 = .014$, with 3 cpd gratings evoking higher responses than 6 cpd gratings. However, for both spatial frequencies, the difference in amplitude of BOLD responses between threat and safe trials was not significantly different, F(1,29) = .46, p = .50, $\eta_p^2 = .016$, and did not change as a function of run number F(2,58) = 2.29, p = .11, $\eta_p^2 = .073$.

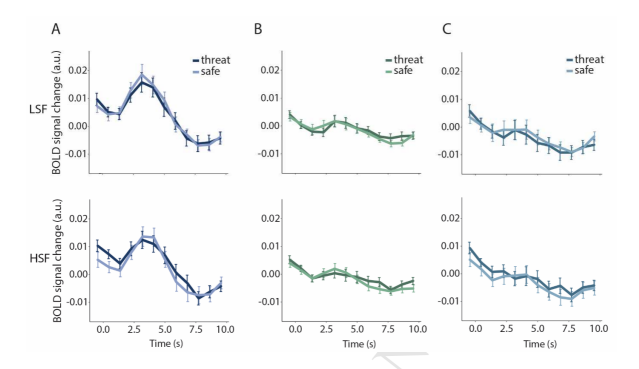


Figure 5. BOLD responses time-locked to the presentation of LSF and HSF gratings in A) retinotopically specific voxels contralateral to stimulus presentation, B) retinotopically nonspecific voxels contralateral to stimulus presentation, and C) retinotopically specific voxels ipsilateral to stimulus presentation. As expected, gratings evoked larger BOLD responses in contralateral retinotopically specific voxels compared to both other ROIs. Within the contralateral, retinotopically specific V1 voxels, the amplitude of BOLD responses to low spatial frequency (3 cpd) and high spatial frequency (6 cpd) gratings did not differ significantly between threat and safe conditions.

In spite of the lack differences between threat and safe conditions, we were still interested in whether the magnitude of threat-induced change in event-related responses would be related to visual discrimination performance. We therefore tested the correlation between threat-related changes in performance (threat – safe conditions across all trials) with the underlying stimulus-

evoked responses in retinotopically specific V1 voxels, for both low (3 cpd) - and high (6 cpd) - spatial frequency gratings. We found that threat-related improvements in performance on the 3 cpd stimulus (larger difference in tilt offset between threat and safe condition) was associated with smaller stimulus-evoked activity in retinotopically specific voxels, r(30) = .49, p = .008 (Figure 6). No significant correlation was found for the 6 cpd stimulus, $\rho(29) = .064, p = .73$. These two correlations differed at trend level (z = 1.69, p = .091), implying that the effect of threat on the relationship between behavioral performance and visual activity was marginally stronger for LSF than HSF gratings. Together, these results show that although V1 responses to low-spatial frequency gratings were on average comparable on threat and safe trials, threat-induced reductions in neural responses were associated with improved discrimination of low-spatial frequency gratings.

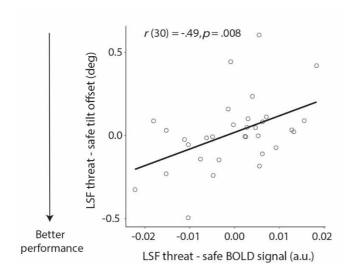


Figure 6. Correlation between threat-evoked responses in retinotopically specific voxels and behavioral performance to low spatial frequency gratings on threat versus safe trials. The facilitating effect of threat on discrimination of low-spatial frequency gratings was associated with smaller stimulus-evoked BOLD responses.

Threat modulation of stimulus-independent and stimulus-evoked responses in V2 and V3. To examine whether the observed effects of threat of shock are specific to V1, we extended our analyses to include retinotopically defined areas V2 and V3. Similar threat-induced effects were observed across V1, V2 and V3, i.e., a retinotopically nonspecific increase in baseline activity in Run 1, F(1,33) = 5.81, p = .022, $\eta_p^2 = .15$, and increased functional connectivity with the amygdala, F(1,33) = 19.20, p < .001, $\eta_p^2 = .37$, neither of which differed between visual areas (all F < 1). We also found positive relationships between stronger fear bradycardia and threat-induced increases in baseline activity, similar to V1, in both V2 [r(24) = .49, p = .010] and V3 [r(24) = -.45, p = .022], both controlling for SCL changes and pupil dilation. Furthermore, no threat-related modulation of event-related responses to the gratings,

and their relation with behavioral performance (p > .05), were observed in V2 and V3, F(2,66) = 1.41, p = .25, $\eta_p^2 = .041$. As in V1, we did find clear evidence of retinotopic specificity of event-related responses across all three visual areas, as evidenced by a retinotopic space by hemisphere interaction, F(1,33) = 50.5, p < .001, $\eta_p^2 = .60$, which did not differ between regions (F < 1). In sum, findings for V2 and V3 were largely identical to V1.

Discussion

The goal of the present study was to identify the neural mechanisms underlying behavioral observations of enhanced visual sensitivity during an anticipatory state of freezing. Our observation of threat-induced fear bradycardia, a parasympathetically dominated response characteristic of freezing (Fredericks, Moore, Metcalf, Schwaber, & Schneiderman, 1974; Koba et al., 2016; Kozlowska et al., 2015; Obrist, 1968), as well as an increase in functional connectivity between amygdala and PAG, core regions involved in expression of freezing in animals (Fanselow, 1994; Fanselow & Lester, 1988; Kapp et al., 1979; LeDoux et al., 1988; Liebman et al., 1970; Schneiderman et al., 1966), indicated the successful induction of a freezing-like defensive state, although directly measuring immobility was not possible given the constraints of an MRI environment. First, we found that threat of shock resulted in a retinotopically nonspecific increase in baseline visuocortical activity accompanied by a concurrent increase in functional connectivity with the amygdala. The magnitude of the threat-related increase in baseline visuocortical activity correlated with fear bradycardia. Second, better discrimination of low-spatial frequency (3 cpd) gratings under threat was associated with reduced responses in V1 voxels retinotopically specific to the location of the gratings.

Threat of shock resulted in an increase of baseline visuocortical activity. This finding is in line with previous studies using a threat anticipation procedure in which increased early visual responses were observed during phasic (Keil, Stolarova, Moratti, & Ray, 2007; Song & Keil, 2013; Stolarova, Keil, & Moratti, 2006; Thigpen, Bartsch, & Keil, 2017) but also prolonged (Herrmann et al., 2016; Sege, Bradley, Weymar, & Lang, 2017; Straube, Mentzel, & Miltner, 2007) presentation of conditioned cues predictive of aversive outcomes. However, although threat anticipation commonly evokes freezing (Fanselow & Lester, 1988), the lack of

concurrently acquired state-specific autonomic and neural measures makes it difficult to generalize these findings to freezing. In addition, because the anticipatory stimuli (e.g., signs) were constantly displayed during anticipatory periods in the studies so far, the contribution of anticipatory versus stimulus-driven processing to enhanced visuocortical responses remains unclear. Our study extends these findings by linking an anticipatory increase in baseline visuocortical activity to defensive responses characteristic of freezing, and by showing that this process is independent of bottom-up visual stimulation.

The upregulation of baseline visuocortical activity under threat was associated with concurrent recruitment of defensive reactions characteristic of a state of freezing. Two findings support this notion. First, enhanced activation of early visual areas (V1, V2, and V3) under threat was associated with stronger fear bradycardia, a parasympathetically dominated response observed during freezing (Fredericks et al., 1974; Koba et al., 2016; Kozlowska et al., 2015; Lang, Bradley, & Cuthbert, 1997; Obrist et al., 1965). In agreement with earlier work showing neural correlates of fear bradycardia (Hermans et al., 2013) and a correlation between low-spatial frequency perception and fear bradycardia (Lojowska et al., 2015), this association was statistically independent of simultaneously observed sympathetic activation indexed by larger skin conductance and pupil dilation responses under threat. Second, a threat-related increase in functional connectivity between the amygdala and PAG - main regions involved in expression of freezing in animals (Fanselow, 1994; Fanselow & Lester, 1988; Kapp et al., 1979; LeDoux et al., 1988; Liebman et al., 1970; Schneiderman et al., 1966), was associated with concomitant increase in functional coupling between the amygdala and early visual areas. A limitation of the current study is that, due to inherent limitations of an MRI environment, it was not possible to measure physical immobility. Such a measure would have allowed for additional validation of

the current results for the state of freezing. However, the close association of fear bradycardia and amygdala-PAG coupling with physical immobility in previous work in the context of threat anticipation, in both animals and humans (Applegate, Kapp, Underwood, & McNall, 1983; Fanselow, 1994; Gladwin et al., 2016; Gozzi et al., 2010; Kozlowska et al., 2015; Niermann, Figner, Tyborowska, Cillessen, & Roelofs, 2018; Roelofs, 2017; Roelofs et al., 2010; Tovote et al., 2016) indicates the presence of a similar defensive state. Future neuroimaging studies on visuocortical changes during freezing may make use of EEG, which may be feasible to combine with direct measurements of mobility using a stabilometric platform (Gladwin et al., 2016).

The amygdala may be a core region involved in the coordination of visual and defensive processes during freezing. Based on anatomical evidence in non-human primates, the observed increase in stimulus-independent visual cortex activity may be driven by modulatory projections from the basal nucleus of the amygdala to early visual areas (Amaral et al., 2003; Freese & Amaral, 2005, 2006). The existence of amygdala-visual connections in humans has been evidenced by morphological changes in the visual areas (Boes et al., 2012), and the absence of otherwise increased responses in the visual cortex to fearful and arousing stimuli in patients with amygdala damage (Anderson, 2001; Vuilleumier et al., 2004). Simultaneous projections of the central nucleus of the amygdala to PAG, on the other hand, are crucial for the expression of fear bradycardia and bodily immobility during animal freezing (Applegate et al., 1983; Gozzi et al., 2010; Kapp et al., 1979; Kim, Rison, & Fanselow, 1993; LeDoux et al., 1988; Liebman et al., 1970; Tovote et al., 2016; Vianna, Graeff, Landeira-Fernandez, & Brandao, 2001; Walker & Carrive, 2003). In particular, lesions to the amygdala or ventral PAG reduced bodily immobility and fear bradycardia during threat anticipation (e.g., fear conditioning) in animals (Carrive, 1993; Kapp et al., 1979; Kim et al., 1993; LeDoux et al., 1988; Liebman et al., 1970; Vianna et

al., 2001), and the activation of the PAG was associated with stronger fear bradycardia in humans (Hermans et al., 2013). In line with these findings, our results support the view that the anticipatory visual processes under threat may be integrated with defensive responses specific for the state of freezing through a common neural mechanism involving the amygdala.

What could be the functional relevance of a spatially nonspecific increase in baseline visuocortical activity during freezing? One possibility is that this increase reflects a preparatory state that may facilitate visual processing of relevant stimuli across the entire visual field. This notion is consistent with the observation that freezing commonly takes place during anticipation of potential threats or in ambiguous environments (Eilam, 2005) where the uncertainty of threat requires higher sensitivity to all spatial locations. In our study, this facilitation may be evidenced by the striking association between reduced neural responses to low-spatial frequency gratings in retinotopically specific regions and enhanced discrimination performance for these stimuli. This correlation cannot be taken to indicate that attention was directed away from the peripheral gratings by the central threat cue (fixation color), because participants with a smaller threatinduced increase in BOLD-responses would then be expected to perform worse, not better. While entirely speculative at this point, a possible explanation for this finding could be that the reduced activity may reflect a smaller prediction error occurring when a bottom-up visual input matches stimulus expectations, which are thought to sharpen stimulus representations and suppress neural responses inconsistent with them (Keller, Bonhoeffer, & Hubener, 2012; Kok, Jehee, & de Lange, 2012; Lee & Mumford, 2003). Accordingly, a threat-related increase in baseline activity found here may reflect a magnocellular input from the amygdala, which would enhance visual representations of coarse features, resulting in reduced visual responses upon their presentation. The modulatory effect of amygdala projections on neural responses is

supported by their physiological properties, i.e., their termination in superficial and deep layers of the visual cortex, which is also characteristic for feedback projections from higher to lower visual areas (Freese & Amaral, 2005, 2006). Such a mechanism would also explain an otherwise contradictory observation of increased early visuocortical responses to lower spatial frequency stimuli following a brief presentation of an arousing image where no threat anticipation was induced(Song & Keil, 2013). The prediction error explanation, however, cannot be directly supported by the current findings. Future studies utilizing, for instance, multi-voxel pattern analyses or electrophysiological recordings may be able to establish whether the observed threat-related increase in baseline visuocortical activity indeed reflects preparatory activity in lower-frequency sensitive voxels (Henriksson, Nurminen, Hyvarinen, & Vanni, 2008; Issa, Trepel, & Stryker, 2000).

One may argue that increased visuocortical activation observed independently of stimulus presentation reflects noradrenergic modulation by arousal. Indeed, increased noradrenergic activity was previously observed during shock anticipation and indexed by pupil dilation (Joshi, Li, Kalwani, & Gold, 2016; Tsuda, Ida, Satoh, Tsujimaru, & Tanaka, 1989). Animal studies have shown that these state-dependent elevations of norepinephrine increase the signal-to-noise ratio of visuocortical neurons (Bennett, Arroyo, & Hestrin, 2013; Polack, Friedman, & Golshani, 2013; Vinck, Batista-Brito, Knoblich, & Cardin, 2015), and their sensitivity to preferred visual input (Polack et al., 2013). However, although pupil dilation was also observed in the current study, it was not related to threat-related changes in visuocortical activity. Instead, our data show that a threat-related increase in baseline activity was associated with parasympathetically controlled fear bradycardia. As we found in previous work showing an association between parasympathetic activity and PAG activation (Hermans et al., 2013), this correlation remained

significant when partialing out variance associated with pupil dilation. Thus, our stimulus-independent effects within visual cortices suggest that threat anticipation evokes a pattern of modulation that is qualitatively different from pure noradrenergic activity, perhaps involving cholinergic activity that is strongly associated with the parasympathetic nervous system and is also known to modulate early visual cortices (Soma, Shimegi, Suematsu, & Sato, 2013).

We cannot exclude the possibility that the initial part of the observed threat-induced fear bradycardia involves an orienting response being also characterized by a decelerative heart rate response (Kapp et al., 1979). However, while the orienting response is typically induced by a sudden change in the environment (e.g., cue onset), freezing commonly occurs during threat anticipation after the cue has been evaluated as threatening, and is often associated with a relatively stronger and more prolonged heart rate deceleration than the orienting response (Bradley, Codispoti, Cuthbert, & Lang, 2001; Gabrielsen, Blix, & Ursin, 1985; Hagenaars et al., 2014; Lang et al., 1997; Pavlov, 1927). Because our findings concern the entire anticipatory phase following onset of the threat cue, they are unlikely to reflect only orienting.

In contrast to previous observations (Bocanegra & Zeelenberg, 2009; Song & Keil, 2013), neither an impairment in higher spatial frequency perception nor differences in neural responses to these stimuli under threat were observed in the current study. Although a threat-related impairment of perception for higher spatial frequency stimuli has been proposed to rely on a neural mechanism involving a cross-inhibition between magnocellular and parvocellular cells in the visual system (Bocanegra & Zeelenberg, 2009), no direct psychophysiological evidence currently exists to support this claim. Alternatively, the fact that current and other findings show specific improvement in lower spatial frequencies in the absence of a higher spatial frequency impairment (Lee, Baek, Lu, & Mather, 2014; Vuilleumier et al., 2003) may

suggest that these two processes are rather independent and may therefore not always co-occur. Future studies should establish what additional factors, specifically those contributing to specific impairment in higher spatial frequencies, such as pupil dilation (Campbell & Green, 1965), contribute to threat-related shifts in spatial frequency perception.

To conclude, our data demonstrate that a freezing-like defensive state is associated with visuocortical changes that may underlie the observed change in visual sensitivity during this state. These visuocortical changes coincide with simultaneously recruited defensive responses which may be regulated through a common neural mechanism involving the amygdala. These results also emphasize the importance of anticipatory threat states in visual perception, and open the way for further research on their role in modulating visual sensitivity to specific visual input such as coarse visual information.

Authors' Contribution

ML, EJH, KR designed research. ML performed research. ML analyzed data. ML, SL, EJH, KR wrote the paper. All authors approved the final version of the manuscript for submission.

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References

- Amaral, D. G., Behniea, H., & Kelly, J. L. (2003). Topographic organization of projections from the amygdala to the visual cortex in the macaque monkey. *Neuroscience*, *118*(4), 1099-1120. doi:10.1016/s0306-4522(02)01001-1
- Anderson, A. K. P., E. A. (2001). Lesions of the human amygdala impair enhanced perception of emotionally salient events. . *Nature*, *411*, 305-309.
- Applegate, C. D., Kapp, B. S., Underwood, M. D., & McNall, C. L. (1983). Autonomic and somatomotor effects of amygdala central N. stimulation in awake rabbits. *Physiology & Behavior*, 31(3), 353-360.
- Ashburner, J., & Friston, K. J. (2005). Unified segmentation. *Neuroimage*, 26(3), 839-851.
- Azevedo, T. M., Volchan, E., Imbiriba, L. A., Rodrigues, E. C., Oliveira, J. M., Oliveira, L. F., . .

 . Vargas, C. D. (2005). A freezing-like posture to pictures of mutilation.

 Psychophysiology, 42(3), 255-260. doi:10.1111/j.1469-8986.2005.00287.x
- Bennett, C., Arroyo, S., & Hestrin, S. (2013). Subthreshold mechanisms underlying state-dependent modulation of visual responses. *Neuron*, 80(2), 350-357. doi:10.1016/j.neuron.2013.08.007
- Birn, R. M., Diamond, J. B., Smith, M. A., & Bandettini, P. A. (2006). Separating respiratory-variation-related fluctuations from neuronal-activity-related fluctuations in fMRI.

 Neuroimage, 31(4), 1536-1548. doi:10.1016/j.neuroimage.2006.02.048
- Blanchard, Griebel, G., Pobbe, R., & Blanchard, R. J. (2011). Risk assessment as an evolved threat detection and analysis process. *Neurosci Biobehav Rev*, *35*(4), 991-998. doi:10.1016/j.neubiorev.2010.10.016

- Blanchard, D. C., Griebel, G., & Blanchard, R. J. (2001). Mouse defensive behaviors: pharmacological and behavioral assays for anxiety and panic. . *Neuroscience & Biobehavioral Reviews*, 25, 205-218.
- Bocanegra, & Zeelenberg, R. (2009). Emotion improves and impairs early vision. *Psychological Science*, 20(6), 707-713. doi:10.1111/j.1467-9280.2009.02354.x
- Boes, A. D., Mehta, S., Rudrauf, D., Van Der Plas, E., Grabowski, T., Adolphs, R., & Nopoulos,
 P. (2012). Changes in cortical morphology resulting from long-term amygdala damage.
 Soc Cogn Affect Neurosci, 7(5), 588-595. doi:10.1093/scan/nsr047
- Bradley, M. M., Codispoti, M., Cuthbert, B. N., & Lang, P. J. (2001). Emotion and motivation I: Defensive and appetitive reactions in picture processing. *Emotion*, 1(3), 276-298. doi:10.1037/1528-3542.1.3.276
- Brainard, D. H. (1997). The Psychophysics Toolbox. *Spat Vis*, *10*(4), 433-436. doi:10.1163/156856897X00357
- Campbell, B. A., Wood, G., & McBride, T. (1997). Origins of orienting and defensive responses:

 An evolutionary perspective. In P.J.Lang, R.F. Simons, & M.T. Balaban (Eds.), Attention and orienting: Sensory and motivational processes (pp. 41–67).
- Campbell, F. W., & Green, D. G. (1965). Optical and retinal factors affecting visual resolution. *J Physiol*, 181(3), 576-593.
- Carrive, P. (1993). The periaqueductal gray and defensive behavior: functional representation and neuronal organization. *Behav Brain Res*, 58(1-2), 27-47.
- Chen, Y., Martinez-Conde, S., Macknik, S. L., Bereshpolova, Y., Swadlow, H. A., & Alonso, J.
 M. (2008). Task difficulty modulates the activity of specific neuronal populations in primary visual cortex. *Nat Neurosci*, 11(8), 974-982. doi:10.1038/nn.2147

- Christman, S. D. (1993). Local-global processing in the upper versus lower visual fields. *Bulletin* of the Psychonomic Society, 31(4), 275-278.
- De Geus, E. J. C., Willemsen, G. H., Klaver, C. H., & Van Doornen, L. J. (1995). Ambulatory measure- ment of respiratory sinus arrhythmia and respiration rate. *Biol. Psychol.*, *41*, 205–227.
- DeYoe, A. E., Carman, G. J., Bandettini, P., Glickman, S., Wieser, J., Cox, R., . . . Neitz, J. (1996). Mapping striate and extrastriate visual areas in human cerebral cortex. *Proc Natl Acad Sci U S A*, *93*(6), 2382–2386.
- Díez-Ajenjo, M. A., & Capilla, P. (2010). Spatio-temporal Contrast Sensitivity in the Cardinal Directions of the Colour Space. A Review. *Journal of Optometry*, *3*(1), 2-19. doi:10.3921/joptom.2010.2
- Eilam, D. (2005). Die hard: a blend of freezing and fleeing as a dynamic defense--implications for the control of defensive behavior. *Neurosci Biobehav Rev*, 29(8), 1181-1191. doi:10.1016/j.neubiorev.2005.03.027
- Engel, S. A., Glover, G. H., & Wandell, B. A. (1997). Retinotopic organization in human visual cortex and the spatial precision of functional MRI. *Cereb Cortex.*, 7(2), 181-192.
- Fanselow, M. S. (1994). Neural organization of the defensive behavior system responsible for fear. *Psychonomic Bulletin & Review*, *1*(4), 429-438. doi:DOI: 10.3758/BF03210947
- Fanselow, M. S., & Lester, L. S. (1988). A functional behavioristic approach to aversively motivated behavior: Predatory imminence as a determinant of the topography of defensive behavior. In R. C. Bolles & M. D. Beecher (Eds.), *Evolution and learning* (pp. 185-212). Hillsdale: NJ: Lawrence Erlbaum Associates.

- Faul, F., Erdfelder, E., Lang, A. G., & Buchner, A. (2007). G*Power 3: A flexible statistical power analysis program for the social, behavioral, and biomedical sciences. *Behavior Research Methods*, 39, 175-191.
- Fredericks, A., Moore, J. W., Metcalf, F. U., Schwaber, J. S., & Schneiderman, N. (1974).

 Selective autonomic blockade of conditioned and unconditioned heart rate changes in rabbits. *Pharmacology Biochemistry and Behavior*, 2(4), 493-501.
- Freese, J. L., & Amaral, D. G. (2005). The organization of projections from the amygdala to visual cortical areas TE and V1 in the macaque monkey. *J Comp Neurol*, 486(4), 295-317. doi:10.1002/cne.20520
- Freese, J. L., & Amaral, D. G. (2006). Synaptic organization of projections from the amygdala to visual cortical areas TE and V1 in the macaque monkey. *J Comp Neurol*, 496(5), 655-667. doi:10.1002/cne.20945
- Gabrielsen, G. W., Blix, A. S., & Ursin, H. (1985). Orienting and freezing responses in incubating ptarmigan hens. *Physiology & Behavior*, *34*, 925-934.
- Gitelman, D. R., Penny, W. D., Ashburner, J., & Friston, K. J. (2003). Modeling regional and psychophysiologic interactions in fMRI: the importance of hemodynamic deconvolution.

 *Neuroimage, 19(1), 200-207. doi:10.1016/s1053-8119(03)00058-2
- Gladwin, T. E., Hashemi, M. M., van Ast, V., & Roelofs, K. (2016). Ready and waiting: Freezing as active action preparation under threat. *Neurosci Lett*, 619, 182-188. doi:10.1016/j.neulet.2016.03.027
- Glover, G. H., Li, T. Q., & Ress, D. (2000). Image-based method for retrospective correction of physiological motion effects in fMRI: RETROICOR. *Magn Reson Med*, 44(1), 162-167.

- Gozzi, A., Jain, A., Giovannelli, A., Bertollini, C., Crestan, V., Schwarz, A. J., . . . Bifone, A. (2010). A neural switch for active and passive fear. *Neuron*, 67(4), 656-666. doi:10.1016/j.neuron.2010.07.008
- Hagenaars, M. A., Oitzl, M., & Roelofs, K. (2014). Updating freeze: Aligning animal and human research. *Neurosci Biobehav Rev*, *47C*, 165-176. doi:10.1016/j.neubiorev.2014.07.021
- Henriksson, L., Nurminen, L., Hyvarinen, A., & Vanni, S. (2008). Spatial frequency tuning in human retinotopic visual areas. *J Vis*, 8(10), 5 1-13. doi:10.1167/8.10.5
- Hermans, E. J., Henckens, M. J., Roelofs, K., & Fernandez, G. (2013). Fear bradycardia and activation of the human periaqueductal grey. *Neuroimage*, 66C, 278-287. doi:10.1016/j.neuroimage.2012.10.063
- Herrmann, M. J., Boehme, S., Becker, M. P., Tupak, S. V., Guhn, A., Schmidt, B., . . . Straube, T. (2016). Phasic and sustained brain responses in the amygdala and the bed nucleus of the stria terminalis during threat anticipation. *Hum Brain Mapp*, *37*(3), 1091-1102. doi:10.1002/hbm.23088
- Hutton, C., Bork, A., Josephs, O., Deichmann, R., Ashburner, J., & Turner, R. (2002). Image distortion correction in fMRI: A quantitative evaluation. *Neuroimage*, 16(1), 217-240. doi:10.1006/nimg.2001.1054
- Issa, N. P., Trepel, C., & Stryker, M. P. (2000). Spatial frequency maps in cat visual cortex. *J Neurosci.*, 20(22), 8504-8514.
- Joshi, S., Li, Y., Kalwani, R. M., & Gold, J. I. (2016). Relationships between Pupil Diameter and Neuronal Activity in the Locus Coeruleus, Colliculi, and Cingulate Cortex. *Neuron*, 89(1), 221-234. doi:10.1016/j.neuron.2015.11.028

- Kapp, B. S., Frysinger, R. C., Gallagher, M., & Haselton, J. R. (1979). Amygdala central nucleus lesions: Effect on heart rate conditioning in the rabbit. *Physiology & Behavior*, 23(6), 1109-1117.
- Keil, A., Stolarova, M., Moratti, S., & Ray, W. J. (2007). Adaptation in human visual cortex as a mechanism for rapid discrimination of aversive stimuli. *Neuroimage*, 36(2), 472-479. doi:10.1016/j.neuroimage.2007.02.048
- Keller, G. B., Bonhoeffer, T., & Hubener, M. (2012). Sensorimotor mismatch signals in primary visual cortex of the behaving mouse. *Neuron*, 74(5), 809-815. doi:10.1016/j.neuron.2012.03.040
- Kim, J. J., Rison, R. A., & Fanselow, M. S. (1993). Effects of amygdala, hippocampus, and periaqueductal gray lesions on short- and long-term contextual fear. *Behav Neurosci*, 107(6), 1093-1098. doi:10.1037/0735-7044.107.6.1093
- Koba, S., Inoue, R., & Watanabe, T. (2016). Role played by periaqueductal gray neurons in parasympathetically mediated fear bradycardia in conscious rats. *Physiol Rep*, *4*(12). doi:10.14814/phy2.12831
- Kok, P., Jehee, J. F., & de Lange, F. P. (2012). Less is more: expectation sharpens representations in the primary visual cortex. *Neuron*, 75(2), 265-270. doi:10.1016/j.neuron.2012.04.034
- Kozlowska, K., Walker, P., McLean, L., & Carrive, P. (2015). Fear and the defense cascade: clinical implications and management. *Harv. Rev. Psychiatry*, 23, 263 287. doi:doi:10.1097/HRP.000000000000000065

- Lang, P. J., Bradley, M. M., & Cuthbert, B. N. (1997). Motivated attention: Affect, activation and action. In P.J. Lang, R.F. Simons, & M.T. Balaban (Eds.), Attention and orienting: Sensory and motivational processes (pp. 97–135). Hillsdale, NJ: Erlbaum.
- LeDoux, J. E., Iwata, J., Cicchetti, P., & Reis, D. J. (1988). Different projections of the central amygdaloid nucleus mediate autonomic and behavioral correlates of conditioned fear.

 LeDoux, J. E., Iwata, J., Cicchetti, P., & Reis, D. J. (1988). Different projections of the central amygdaloid nucleus mediate autonomic and behavioral correlates of conditioned fear. J Neurosci, 8(7), 2517-2529., 8(7), 2517-2529.
- Lee, Baek, J., Lu, Z. L., & Mather, M. (2014). How arousal modulates the visual contrast sensitivity function. *Emotion*, 14(5), 978-984. doi:10.1037/a0037047
- Lee, T. S., & Mumford, D. (2003). Hierarchical Bayesian inference in the visual cortex. *J. Opt. Soc. Am. A Opt. Image Sci. Vis.*, 20(7), 1434-1448.
- Leonova, A., Pokorny, J., & Smith, V. C. (2003). Spatial frequency processing in inferred PC-and MC-pathways. *Vision Research*, 43(20), 2133-2139. doi:10.1016/s0042-6989(03)00333-x
- Liebman, J. M., Mayer, D. J., & Liebeskind, J. C. (1970). Mesencephalic central gray lesions and fear-motivated behavior in rats. *Brain Research*, 23(3), 353-370.
- Lojowska, M., Gladwin, T. E., Hermans, E. J., & Roelofs, K. (2015). Freezing promotes perception of coarse visual features. *J Exp Psychol Gen, 144*(6), 1080-1088. doi: 10.1037/xge0000117
- Maren, S. (2001). Neurobiology of Pavlovian fear conditioning. *Annu Rev Neurosci*, 24, 897-931.

- Nicol, J. R., Perrotta, S., Caliciuri, S., & Wachowiak, M. P. (2013). Emotion-specific modulation of early visual perception. *Cogn Emot*, 27(8), 1478-1485. doi:10.1080/02699931.2013.793654
- Niermann, H. C. M., Figner, B., Tyborowska, A., Cillessen, A. H. N., & Roelofs, K. (2018).

 Investigation of the Stability of Human Freezing-Like Responses to Social Threat From

 Mid to Late Adolescence. *Front Behav Neurosci*, 12. doi:10.3389/fnbeh.2018.00097
- Obrist, P. A. (1968). Heart rate and somatic-motor coupling during classical aversive conditioning in humans. . *Journal of Experimental Psychology*, 77(2), 180-193.
- Obrist, P. A., Wood, D. M., & Perez-Reyes, M. (1965). Heart rate during conditioning in humans: Effects of UCS intensity, vagal blockade, and adrenergic block of vasomotor activity. *Journal of Experimental Psychology*, 70(1), 32-42.
- Öhman, A., & Wiens, S. (2003). On the automaticity of autonomic responses in emotion: an evolutionary perspective. In: Davidson, R.J., Scherer, K., Hill, H.H. (Eds.). *Handbook of affective sciences. Oxford University Press, New York.*, 256–275.
- Pavlov, J. P. (1927). Conditioned Reflexes. London: Oxford University Press.
- Phelps, E. A., O'Connor, K. J., Gatenby, J. C., Gore, J. C., Grillon, C., & Davis, M. (2001).

 Activation of the left amygdala to a cognitive representation of fear. . *Nat Neurosci*, 4(4), 437-441. doi:10.1038/86110
- Polack, P. O., Friedman, J., & Golshani, P. (2013). Cellular mechanisms of brain state-dependent gain modulation in visual cortex. *Nat Neurosci*, *16*(9), 1331-1339. doi:10.1038/nn.3464
- Purves, D., Augustine, G. J., Fitzpatrick, D., Katz, L. C., LaMantia, A.-S., McNamara, J., & Williams, M. (2001). *Neuroscience* (2 ed.). Sunderland (MA): Sinauer Associates.

- Roelofs, K. (2017). Freeze for action: neurobiological mechanisms in animal and human freezing. *Philos Trans R Soc Lond B Biol Sci*, *372*(1718). doi:10.1098/rstb.2016.0206
- Roelofs, K., Hagenaars, M. A., & Stins, J. (2010). Facing freeze: social threat induces bodily freeze in humans. *Psychol Sci*, *21*(11), 1575-1581. doi:10.1177/0956797610384746
- Rovamo, J., Virsu, V., & Näsänen, R. (1978). Cortical magnification factor predicts the photopic contrast sensitivity of peripheral vision. *Nature*, *271*(5640), 54-56.
- Satpute, A. B., Wager, T. D., Cohen-Adad, J., Bianciardi, M., Choi, J. K., Buhle, J. T., . . . Barrett, L. F. (2013). Identification of discrete functional subregions of the human periaqueductal gray. *Proc Natl Acad Sci U S A*, *110*(42), 17101-17106. doi:10.1073/pnas.1306095110
- Schneiderman, N., Smith, M. C., Smith, A. C., & Gormezano, I. (1966). Heart rate classical conditioning in rabbits. *Psychonomic Science*, 6(5), 241–242.
- Sege, C. T., Bradley, M. M., Weymar, M., & Lang, P. J. (2017). A direct comparison of appetitive and aversive anticipation: Overlapping and distinct neural activation. *Behav Brain Res*, 326, 96-102. doi:10.1016/j.bbr.2017.03.005
- Sereno, M. I., Dale, A. M., Reppas, J. B., Kwong, K. K., Belliveau, J. W., Brady, T. J., . . . Tootell, R. B. (1995). Borders of multiple visual areas in humans revealed by functional magnetic resonance imaging. *Science*, *268*(5212), 889-893.
- Soma, S., Shimegi, S., Suematsu, N., & Sato, H. (2013). Cholinergic modulation of response gain in the rat primary visual cortex. *Sci Rep, 3*, 1138. doi:10.1038/srep01138
- Song, I., & Keil, A. (2013). Affective engagement and subsequent visual processing: effects of contrast and spatial frequency. *Emotion*, *13*(4), 748-757. doi:10.1037/a0031553

- Stolarova, M., Keil, A., & Moratti, S. (2006). Modulation of the C1 visual event-related component by conditioned stimuli: evidence for sensory plasticity in early affective perception. *Cereb Cortex*, 16(6), 876-887. doi:10.1093/cercor/bhj031
- Straube, T., Mentzel, H. J., & Miltner, W. H. (2007). Waiting for spiders: brain activation during anticipatory anxiety in spider phobics. *Neuroimage*, *37*(4), 1427-1436. doi:10.1016/j.neuroimage.2007.06.023
- Thigpen, N. N., Bartsch, F., & Keil, A. (2017). The malleability of emotional perception: Short-term plasticity in retinotopic neurons accompanies the formation of perceptual biases to threat. *J Exp Psychol Gen*, *146*(4), 464-471. doi:10.1037/xge0000283
- Tovote, P., Esposito, M. S., Botta, P., Chaudun, F., Fadok, J. P., Markovic, M., . . . Luthi, A. (2016). Midbrain circuits for defensive behaviour. *Nature*, *534*(7606), 206-212. doi:10.1038/nature17996
- Tovote, P., Fadok, J. P., & Luthi, A. (2015). Neuronal circuits for fear and anxiety. *Nat Rev Neurosci*, 16(6), 317-331. doi:10.1038/nrn3945
- Tsuda, A., Ida, Y., Satoh, H., Tsujimaru, S., & Tanaka, M. (1989). Stressor predictability and rat brain noradrenaline metabolism. *Pharmacol Biochem Behav*, 32(2), 569-572.
- van Buuren, M., Gladwin, T. E., Zandbelt, B. B., van den Heuvel, M., Ramsey, N. F., Kahn, R. S., & Vink, M. (2009). Cardiorespiratory effects on default-mode network activity as measured with fMRI. *Hum Brain Mapp*, *30*(9), 3031-3042. doi:10.1002/hbm.20729
- Vianna, D. M., Graeff, F. G., Landeira-Fernandez, J., & Brandao, M. L. (2001). Lesion of the ventral periaqueductal gray reduces conditioned fear but does not change freezing induced by stimulation of the dorsal periaqueductal gray. *Learn Mem*, 8(3), 164-169. doi:10.1101/lm.36101

- Vinck, M., Batista-Brito, R., Knoblich, U., & Cardin, J. A. (2015). Arousal and locomotion make distinct contributions to cortical activity patterns and visual encoding. *Neuron*, 86(3), 740-754. doi:10.1016/j.neuron.2015.03.028
- Vuilleumier, P., Armony, J. L., Driver, J., & Dolan, R. J. (2003). Distinct spatial frequency sensitivities for processing faces and emotional expressions. *Nat Neurosci*, 6(6), 624-631.
- Vuilleumier, P., Richardson, M. P., Armony, J. L., Driver, J., & Dolan, R. J. (2004). Distant influences of amygdala lesion on visual cortical activation during emotional face processing. *Nat Neurosci*, 7(11), 1271-1278. doi:10.1038/nn1341
- Wager, T. D., & Nichols, T. E. (2003). Optimization of experimental design in fMRI: a general framework using a genetic algorithm. *Neuroimage*, 18(2), 293-309. doi:10.1016/s1053-8119(02)00046-0
- Walker, P., & Carrive, P. (2003). Role of the ventrolateral periaqueductal gray neuronsin the behavioral and cardiovascular responses to contextual conditioned fear and poststress recovery. *Neuroscience*, *116*, 897–912. doi:doi:10.1016/S0306-4522(02)00744
- Walker, R., Deubel, H., Schneider, W. X., & Findlay, J. M. (1997). Effect of remote distractors on saccade programming: evidence for an extended fixation zone. *J Neurophysiol*, 78(2), 1108–1119.
- Watson, A. B., & Pelli, D. G. (1983). QUEST: a Bayesian adaptive psychometric method. *Percept Psychophys*, 33(2), 113-120. doi:10.3758/BF03202828