Neural responses to emotional involuntary memories in posttraumatic stress disorder: Differences in timing and activity

Shana A. Hallb,c,⁎, Kaitlyn E. Brodarb,c, Kevin S. LaBarc, Dorthe Berntsend, David C. Rubinc,d

aDepartment of Psychology & Neuroscience, University of North Carolina, Chapel Hill, United States
bDepartment of Psychology, University of Miami, United States
cDepartment of Psychology & Neuroscience, Duke University, United States
dCenter on Autobiographical Memory Research, Aarhus University, Denmark

ARTICLE INFO

Keywords:
Posttraumatic stress disorder (PTSD)
Involuntary memory
Functional magnetic resonance imaging (fMRI)
Finite impulse response (FIR)
Memory network
Ventromedial prefrontal cortex (vmPFC)

ABSTRACT

Background: Involuntary memories are a hallmark symptom of posttraumatic stress disorder (PTSD), but studies of the neural basis of involuntary memory retrieval in posttraumatic stress disorder (PTSD) are sparse. The study of the neural correlates of involuntary memories of stressful events in PTSD focuses on the voluntary retrieval of memories that are sometimes recalled as intrusive involuntary memories, not on involuntary retrieval while being scanned. Involuntary memory retrieval in controls has been shown to elicit activity in the para-hippocampal gyrus, precuneus, inferior parietal cortex, and posterior midline regions. However, it is unknown whether involuntary memories are supported by the same mechanisms in PTSD. Because previous work has shown that both behavioral and neural responsivity is slowed in PTSD, we examined the spatiotemporal dynamics of the neural activity underlying negative and neutral involuntary memory retrieval.

Methods: Twenty-one individuals with PTSD and 21 non-PTSD, trauma-exposed controls performed an involuntary memory task, while undergoing a functional magnetic resonance imaging scan. Environmental sounds served as cues for well-associated pictures of negative and neutral scenes. We used a finite impulse response model to analyze temporal differences between groups in neural responses.

Results: Compared with controls, participants with PTSD reported more involuntary memories, which were more emotional and more vivid, but which activated a similar network of regions. However, compared to controls, individuals with PTSD showed delayed neural responsivity in this network and increased vmPFC/ACC activity for negative > neutral stimuli.

Conclusions: The similarity between PTSD and controls in neural substrates underlying involuntary memories suggests that, unlike voluntary memories, involuntary memories elicit similar activity in regions critical for memory retrieval. Further, the delayed neural responsivity for involuntary memories in PTSD suggests that factors affecting cognition in PTSD, like increased fatigue, or avoidance behaviors could do so by delaying activity in regions necessary for cognitive processing. Finally, compared to neutral memories, negative involuntary memories elicit hyperactivity in the vmPFC, whereas the vmPFC is typically shown to be hypoactive in PTSD during voluntary memory retrieval. These patterns suggest that considering both the temporal dynamics of cognitive processes as well as involuntary cognitive processes would improve existing neurobiological models of PTSD.

1. Introduction

Involuntary episodic memories are explicit memories of past events that can be described by the rememberer, but are not preceded by a retrieval attempt (Berntsen, 1996). Involuntary memories are assumed to be critical for maintaining a sense of continuity of self across time and for giving fast access to memories that may have functional relevance in novel situations (Rasmussen and Berntsen, 2009). They occur at least as frequently as their voluntary counterparts (Rasmussen and Berntsen, 2011; Rubin and Berntsen, 2009), and they are viewed as

Abbreviations: PTSD, posttraumatic stress disorder; ACC, anterior cingulate cortex; PCC, posterior cingulate cortex; IPC, inferior parietal cortex; vmPFC, ventromedial prefrontal cortex; MTL, medial temporal lobes; IAPS, International Affective Picture System; SPGR, spoiled gradient recalled; TR, repetition time; TE, echo time; TI, inverse recovery time; SPM, Statistical Parametric Mapping; FIR, finite impulse response; FDR, false detection rate; FWE, family-wise error

⁎ Corresponding author at: Department of Psychology & Neuroscience, University of North Carolina, Chapel Hill, 235 E. Cameron Ave., Campus Box #3270, Chapel Hill, NC 27599-3270, United States.
E-mail address: shanahall@unc.edu (S.A. Hall).

https://doi.org/10.1016/j.nicl.2018.05.009
Received 3 February 2018; Received in revised form 30 April 2018; Accepted 8 May 2018
Available online 10 May 2018
2213-1582/ © 2018 The Authors. Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/BY-NC-ND/4.0/).
a basic mode of remembering, operating on the same system as voluntary memories (Berntsen, 2010).

Involuntary memories of a traumatic event are a hallmark symptom of posttraumatic stress disorder (PTSD; American Psychiatric Association, 2013), with intrusive memories being present in almost 90% of individuals with PTSD (Rozzell et al., 1991). Past research indicates the frequency of these intrusive memories—and especially associated responses such as negative reinterpretations, a strong sense of reliving, and maladaptive emotional reactions to the memories—correlates highly with PTSD severity (e.g., Michael et al., 2005). Little is known about the neural underpinnings of the retrieval of involuntary memory. This paucity of research stands in marked contrast with the fact that in clinical disorders, such as PTSD, involuntary memories often become disruptive and distressing. Despite their centrality to the disorder, the basic cognitive and neural mechanisms underlying both emotional and non-emotional involuntary memories in PTSD are not well understood.

We were able to identify four studies that specifically addressed the neural basis of intrusive involuntary memories of stressful events. Two of these studies used a prospective design with healthy individuals to examine whether neural activity during the encoding of stressful stimuli would predict the reporting of involuntary intrusive memories of this material during a subsequent diary study. Both reported increased activity in limbic regions, including the amygdala, anterior cingulate cortex (ACC), striatum, thalamus, and parahippocampal gyrus, with such increased activity during the encoding of stressful films subsequently reported as flashbacks (Bourne et al., 2013), and during encoding in individuals reporting a high level of intrusions (Battaglini et al., 2016). Neither of these studies examined neural activation during retrieval.

Whalley et al. (2013) had PTSD patients and controls provide a narrative account of their trauma or most distressing event before the scan. During the scan they conducted a recognition task in which they were asked to identify keywords culled from their own trauma narrative versus keywords derived from other participants’ narratives. Outside the scanner, the participants subsequently identified which of the keywords had led to flashback experiences during the scan and which led to ordinary episodic memories. Whalley et al. (2013) found that flashbacks versus episodic recollections were associated with increased neural activity in the sensory and motor areas. However, since a voluntary autobiographical recognition task generated both types of memories, the neural activity did not target involuntary retrieval per se.

Clark et al. (2016) examined involuntary retrieval of emotional film clips in healthy participants while they were at rest. The participants indicated that they had an involuntary memory from the film clip by pressing a button. In contrast to other work examining the neural underpinnings of involuntary memories in healthy individuals (e.g., Hall et al., 2014), Clark et al. (2016) found that involuntary retrieval elicited activity in frontal regions. This finding may reflect that the memory recording necessarily involved a source monitoring judgment to decide whether a memory ‘popping into mind’ matched what had been seen in the film clip or not, since only involuntary memories related to the film clips should be reported. Thus, the frontal activity may be due to making this judgment, which might involve a voluntary search process and related cognitive control processes. We designed our task to address this issue by using retrospective report to determine when involuntary memories occur (Hall et al., 2014). We also expand on this work by including individuals with PTSD and comparing negative to neutral involuntary memories.

Outside the clinical literature, a few studies examined brain activity during involuntary memory retrieval. Hall et al. (2014) compared voluntary and involuntary memories of scenes cued by environmental sounds to mimic the kinds of cues and memories that often occur during everyday life (Berntsen et al., 2013; Rubin et al., 2008; Schlagman and Kvavilashvili, 2008). Both voluntary and involuntary memories elicited activity in regions typically found to be active during visual voluntary memory retrieval, including the parahippocampal gyrus, posterior midline regions like the posterior cingulate cortex (PCC) and retrosplenial cortex, precuneus, and inferior parietal cortex (IPC). However, involuntary memory retrieval did not elicit activity in frontal cortices, likely reflecting the lack of effort involved in such retrieval (Hall et al., 2014).

To our knowledge, no studies addressing the neural underpinnings of involuntary memories in PTSD have used a methodology that does not include the voluntary retrieval of a memory during the scan either to rate directly or to compare with a memory that has come involuntarily. The goal of the present study was to fill this gap in the literature. We wanted to examine the neural activations associated with involuntary memory processing in PTSD in general and not specifically for traumatic events. Developing a deeper understanding of such general processing deficits is relevant because, among other things, diagnostic inclusion criteria for PTSD include a general enhancement of negative affect that is not limited to trauma-relevant material. In particular, this includes difficulty experiencing positive affect, decreased interest in activities, and overly negative thoughts or assumptions about oneself or the world (American Psychiatric Association, 2013).

Involuntary memories in PTSD compared to non-PTSD participants, irrespective of their trauma-relevance, involve greater emotionality and are more central to the life story (Berntsen et al., 2015; Rubin et al., 2008, 2011). Also, there is evidence of broad emotion regulation deficits in PTSD beyond traumatic material. Individuals with PTSD have abnormal responses to non-trauma-related negative stimuli, including increased self-reported emotion (Amdu et al., 2000; Orsillo et al., 2004; Wolf et al., 2009) and enhanced neural activity to non-trauma-related stimuli (Offringa et al., 2013; Rauch et al., 2000; Shin et al., 2005). Our focus on the broad emotion processing deficit in PTSD is consistent with an increasing emphasis in the PTSD literature on dysfunctional emotion regulation processes across a range of contexts, and not simply in response to the traumatic event per se (e.g., Bonanno et al., 2004; Brohawn et al., 2010; Orcutt et al., 2014; Shin et al., 2005). Because of this aim we used the same paradigm as Hall et al. (2014, described in detail below), and developed in our previous work (Berntsen et al., 2013; Staagaard and Berntsen, 2014), but varied the emotionality of the scenes.

Voluntary retrieval of scenes, such as those we used here, is commonly used in studies of involuntary memories in healthy individuals (Berntsen, 2009; Berntsen et al., 2013). They elicit activity in a well-characterized network of regions including the parahippocampal gyrus, the inferior and superior parietal cortices, precuneus, posterior cingulate cortex, and dorsomedial and dorsolateral prefrontal cortex (for reviews, see Kim, 2013; Spaniol et al., 2009). In previous work, we showed activity in a subset of these regions for involuntary memory retrieval (Hall et al., 2014): the parahippocampal gyrus, thought to underlie successful retrieval, making contributions to scene and context memory (Ranganath and Ritchey, 2014), the inferior parietal cortex, thought to be involved in bottom-up (i.e., non-goal-driven, or source-driven) attention to memory (Cabeza et al., 2008; Hutchinson et al., 2009) and posterior midline regions like the posterior cingulate cortex, a key region in the default mode network, which has been associated with self-referential thought (Buckner and Carroll, 2007; Greicius et al., 2003; Gusnard et al., 2001; Wicker et al., 2003) and the precuneus, which has been associated with visual imagery (Cavanna and Trimble, 2006). There may be an additional anterior/posterior dissociation within the inferior parietal cortex, with anterior regions being involved in attention reorienting and posterior regions being associated with the default mode network (Hutchinson et al., 2009). Other regions not found to be active during involuntary memory retrieval (Hall et al., 2014) are regions thought to be involved in top-down processes that are less likely to be invoked when retrieval is involuntary. The dorsolateral PFC is thought to be involved in memory search and maintenance (Svoboda et al., 2006). The superior parietal cortex underlies top-down, goal-driven attention to memory (Cabeza et al., 2008; Hutchinson et al., 2009).
The differential and common activity in the superior and inferior parietal cortices, respectively, in voluntary and involuntary retrieval illustrates the dissociation in top-down versus bottom-up cognitive processes used during these two types of retrieval. These top-down processes allow for the maintenance of retrieval goals and may be more engaged during low-confidence memory retrieval in an attempt to retrieve additional details of a weak memory, whereas bottom-up processes are particularly engaged when the memory is salient (Cabeza et al., 2008). Though these processes are thought to exist in two dimensions, they interact. Bottom-up processes can occur during voluntary memories and top-down processes can occur during involuntary memories.

To examine the neural activations associated with involuntary memory processing in PTSD generally, we did not intentionally select scenes that were trauma-relevant for each participant. Critically, participants did not indicate when they had an involuntary memory during the task because doing so would likely invoke top-down processes that would not typically occur during naturalistic involuntary remembering (Berntsen, 2009) and would produce neural activity common to voluntary retrieval tasks. We also examined the specific time course for the neural activations, following up on previous work. Because retrieval in our task and in autobiographical memory can take seconds, we used fMRI to study the time course of retrieval, which allowed us to track the time course of neural processes (Daseelaar et al., 2008; Hall et al., 2014). Investigating differences in the time course of neural processes is critical in PTSD because previous work indicates such differences in neural activity during memory retrieval (St Jacques et al., 2011a) and in response time during reflexive, involuntary processing in PTSD (Clark et al., 2003; Veltmeyer et al., 2005). Below, we present whole-brain results but with a theoretical focus on what we refer to as the involuntary memory network, derived from previous work on non-emotional involuntary memories, which includes the MTI, the precuneus, IPC, and posterior midline regions, as well as on emotion-processing regions consisting of the amygdala and the vmPFC.

2. Materials & methods

2.1. Participants

Fifty-seven participants (35 women, mean age = 21.9 years, range = 18–37 years) completed the scan session. Participants gave written informed consent for a protocol approved by the Duke University Institutional Review Board and the Duke University Medical Center Institutional Review Board. All participants had an A1 trauma in which they were exposed to death, threatened death, actual or threatened serious injury, or actual or threatened sexual violence (American Psychiatric Association, 2013). After excluding participants who did not meet inclusion criteria (see Supplemental materials) there were 21 participants in the PTSD group (15 women, mean age = 21.5 years, range = 18–31 years) and 21 in the trauma-exposed control group (14 women, mean age = 22.1 years, range = 18–37 years). Additional demographic and clinical data are shown in Table 1 and described in the Supplemental materials.

2.2. Materials

The stimuli included 55 scenes and 95 environmental sounds (e.g., woman sneezing, lawn mower starting). We chose these stimuli because scene memories are typical of autobiographical memories and sounds are common cues for such memories (Berntsen et al., 2013). We obtained the images from the International Affective Picture System (Lang et al., 2008) and used normative IAPS valence ratings. We considered ratings from 1.0–3.5 to be negative, 3.6–5.5 to be neutral, and 5.6–9.0 to be positive. We only included negative and neutral scenes and did not choose images to be related to a specific trauma. The negative pictures depicted poverty, death/illness, violence, gore, filth, an accident, and hate groups. The neutral pictures depicted animals sometimes looked threatening, daily life activities, objects, and locations, military people and equipment, modes of transportation, locations, and portraits of people. While we did not intentionally choose scenes to be relevant to any individual participant’s index trauma, some images may have been relevant to particular traumas. We did not obtain trauma-relevance ratings from the participants; however, a rater judged the trauma-relevance of each image for each trauma and found no more than nine images to be relevant to any participant’s index trauma.

2.3. Experimental design and procedure

The experimental design and procedure is nearly identical to that described in a previous publication in healthy subjects (Hall et al., 2014). Here we report a short summary of the experimental design and procedures modified for the goals of the current study. For a full description, see the Supplemental materials and Fig. 1.

Two days before MRI scanning, participants encoded the 55 paired and 40 unpaired sounds. To ensure strong memory associations, there were four encoding runs, during which participants were presented with the sound-picture pairs and wrote a sentence or a short story linking the sound and the picture. Two days after encoding, they participated in an fMRI session during which they re-encoded the paired and unpaired sounds and then completed the critical sound localization/memory recall runs. During the re-encoding runs, each sound-picture pair and unpaired sound was presented. Participants rated on a 7-point scale how emotional the stimuli were. During the recall runs, the 55 paired and 40 unpaired sounds (randomly intermixed) were presented, panned 15° to either the left or the right using specialized audio software (Audacity, audacity.sourceforge.net). Participants indicated the side on which the sound was louder. We presented each sound for 4 s, followed by a 1.5-second response period, during which the screen was blank and participants completed the sound localization judgment. A fixation period followed (jittered with a mean of 4 s, range: 1.5–10.4 s). At the beginning of the run, we instructed participants to wait until after the sound had ended to respond. Immediately following the scanning session, participants completed a post-scan questionnaire to assess their memory for the scenes. Participants heard all 95 sounds. After the presentation of each sound, participants reported: (1) whether they had remembered a picture when the sound was played during the sound localization task (yes/no), (2) how hard they tried to perform the sound localization task correctly (1 = did not try at all, 7 = tried very hard), (3) how hard they tried to recall a picture during the scan (1 = did not try at all, 7 = tried very hard), and (4) how vivid the memory of the picture was during the scan (1 = not at all vivid, 7 = very vivid). Lastly, we presented the 55 paired sounds a final time and participants provided a description of the picture originally paired

| Table 1: Participant characteristics. |
|-------------------------------------|
| Demographics | PTSD mean (SD) | Control mean (SD) | p |
| Age | 21.48 (3.34) | 22.10 (4.75) | 0.640 |
| CAPS | 54.29 (15.88) | 11.29 (10.78) | < 0.001 |
| BDI | 13.55 (9.10) | 3.33 (3.69) | < 0.001 |
| CES | 25.60 (5.23) | 15.00 (8.77) | < 0.001 |
| PCL-S | 32.48 (12.42) | 8.67 (6.17) | < 0.001 |
| TLEQ | 6.14 (3.53) | 5.19 (4.48) | 0.474 |

| Type of trauma exposure | PTSD total (n) | Control total (n) |
|-------------------------|---------------|------------------|
| Sexual assault | 6 | 5 |
| Physical assault | 4 | 2 |
| Injury, illness, or accident | 4 | 6 |
| Family/friend injury or death | 7 | 6 |
| Natural disaster | 0 | 2 |
with the sound. Two independent judges rated these descriptions for accuracy. In the case of disagreement, a third judge broke the tie. As described below, only paired sounds that elicited a memory without effort and unpaired sounds that did not elicit a memory and had no retrieval effort were analyzed.

2.4. Image acquisition and preprocessing

Imaging was conducted on a 3 T GE Signa Excite MRI scanner (GE Healthcare, Waukesha, WI) with an eight-channel head coil. Following a localizer scan was a T1-weighted structural image, (96 contiguous slices acquired with a high-resolution, 3D fast inverse recovery-prepared spoiled gradient recalled (SPGR) sequence, with a repetition time (TR) = 3.22 ms, echo time (TE) = 8.2 ms, inversion recovery time (TI) = 450 ms, 1 mm slice thickness, and parallel imaging with a selection factor of 2). During the task, T2* - weighted echo-planar, functional images were acquired using a spiral-in sequence with SENSE imaging (34 contiguous slices, TR = 2000 ms, TE = 30 ms, FOV = 240 mm, 3.8 mm slice thickness with no gaps, flip angle = 70°, voxel size = 3.75 × 3.75 × 3.8 mm, 64 × 64 matrix, SENSE factor = 2). Preprocessing and analyses of functional imaging data were conducted with Statistical Parametric Mapping software (SPM12; Wellcome Department of Cognitive Neurology, London, UK), along with locally developed Matlab (Mathworks, Natick, MA) scripts. We discarded the first three volumes of each run, including the first trial of each run, for scanner stability. We also excluded the discarded trials for each run from behavioral analyses. Images were corrected for slice-timing and head motion, spatially normalized to the Montreal Neurological Institute template using a 12-parameter affine model, and then spatially smoothed with an 8-mm Gaussian kernel. A high-pass filter was included in every model to correct for scanner drift.

2.5. fMRI data analysis

We created two general linear models: a Memory model comparing paired and unpaired sound trials and an Emotion model comparing high and low emotional valence trials. In the Memory model, the first-level model included regressors for paired sound trials in which there was a reported memory and a low effort rating (1 or 2 out of 7), and unpaired sound trials for which there was no reported memory and a low effort rating. We split these trials according to sound lateralization (paired left, paired right, unpaired left, and unpaired right) to account for this additional variance. In both models, we included all high-effort trials, paired sound trials that did not elicit a memory, and unpaired sound trials that did elicit a memory as trials of no interest, as well as regressors accounting for motion and run-level effects.

In the Emotion model, at the first level, we separately included regressors for high and low emotion paired sounds and an Emotion model comparing high and low emotional valence trials. In the Memory model, the first-level model included regressors for paired sound trials in which there was a reported memory and a low effort rating (1 or 2 out of 7), and unpaired sound trials for which there was no reported memory and a low effort rating. We split these trials according to sound lateralization (paired left, paired right, unpaired left, and unpaired right) to account for this additional variance. We collapsed the sound lateralization factor at the second level. In both models, we included all high-effort trials, paired sound trials that did not elicit a memory, and unpaired sound trials that did elicit a memory as trials of no interest, as well as regressors accounting for motion and run-level effects.
we included unpaired sounds as trials of no interest.

We initially identified activations using a finite impulse response (FIR) model (Goutte et al., 2000; Henson et al., 2001), which we preferred over a standard canonical hemodynamic response model to better understand the temporal dynamics of involuntary memory retrieval. We initially modeled seven TRs, or 14 s, starting at the beginning of the trial, not adjusted for the hemodynamic lag, but included only the 3rd to 6th TRs in analyses. The 1st and 2nd TRs were not analyzed because the sound corresponded to the beginning of the subsequent trial. The 3rd and 4th TRs corresponded to the time period during which peak activity associated with the sound would be expected to occur, accounting for hemodynamic lag. The 5th and the 6th TRs corresponded to the time period during which involuntary memories could continue to be formed. Additionally, exploratory analyses revealed that activity was similar in the 3rd and 4th TRs and in the 5th and the 6th TRs. Therefore, to maintain a consistent length of time represented by each time point we collapsed these pairs of TRs to create an early and a late time point. The late time point included the 1.5 s response period and 2.5 s of the jitter. The late time point was not contaminated by the cognitive processes involved in the subsequent trial because the hemodynamic response for the subsequent trial does not peak until approximately 6 s after that trial begins. We report whole-brain results at a significance threshold of \( p < 0.05 \) using a false discovery rate (FDR) correction and a cluster correction of \( p < 0.05 \), using a family-wise error (FWE) correction.

### Table 3

Memory model behavioral results. Note, the number of trials modeled, shown in parantheses in the % recalled column, were not included in tests of behavioral differences.

| % recalled | Emotion | Vividness | Reaction time |
|------------|---------|-----------|---------------|
|            | Avg.    | SD        | Avg.          | SD        | Avg. | SD        |
| Paired sounds |         |           |               |           |      |           |
| Control    | 56.19% (30.00 trials modeled) | 24.24% (12.89) | 4.09 | 1.20 | 4.55 | 1.4 | 689.22 | 231.53 |
| PTSD       | 68.84% (36.95 trials modeled) | 22.52% (12.11) | 4.24 | 0.68 | 5.32 | 1.3 | 713.16 | 138.59 |
| Unpaired sounds |         |           |               |           |      |           |
| Control    | 4.08% (36.43 trials modeled) | 5.18% (4.34) | 2.51 | 1.18 | 1.10 | 0.40 | 751.06 | 223.20 |
| PTSD       | 6.66% (31.67 trials modeled) | 5.41% (9.60) | 2.47 | 0.77 | 1.06 | 0.22 | 765.60 | 157.02 |
| ANOVA      | F       | p         | F             | p         | F    | p         |   | |
| Main effect of group | 4.23 | < 0.05 | 0.05 | > 0.05 | 2.79 | > 0.05 | 0.32 | > 0.05 |
| Main effect of pairing | 238.29 | < 0.001 | 60.23 | < 0.001 | 309.54 | < 0.001 | 0.68 | > 0.05 |
| Interaction | 1.85 | > 0.05 | 0.20 | > 0.05 | 3.41 | > 0.05 | 0.25 | > 0.05 |

* \( p < 0.05 \).

** \( p < 0.001 \).

### 3. Results

#### 3.1. Behavioral results

##### 3.1.1. Memory model: paired > unpaired

We report behavioral results for paired > unpaired sounds in Table 2. PTSD participants reported more involuntary memories (those retrieved with low effort) than controls, and paired sounds elicited more involuntary memories than unpaired sounds. Additionally, focusing only on trials included in the fMRI analyses (paired sounds that elicited a memory and unpaired sounds that did not), an ANOVA revealed higher emotion ratings (gathered during encoding) and vividness (gathered during the post-scan retrieval run) for paired sounds relative to unpaired sounds (see Table 2).

##### 3.1.2. Emotion model: high > low emotion

We report behavioral analyses for high > low emotion sound-picture pairs in Table 3. Notably, the PTSD group had more involuntary memories to both high and low emotion trials than controls. There was no effect of emotion on the number of memories recalled.

There were no reaction time differences for either model. Behavioral performance on the localization task for both models is reported in the Supplemental materials.

##### 3.1.3. Post-scan recall

We quantified the number of pictures that were accurately recalled voluntarily after the scan. Technical error resulted in missing data for one PTSD participant. There was an effect of Group (F(1, 82) = 5.80, \( p < 0.05 \)), with the PTSD group accurately recalling a greater number of pictures than controls.
3.2. Neuroimaging results

Because we could not be certain when the involuntary memories would occur, we probed neural activity at two time points: an early time point including the 4 s when the sound played, and a late time point including the 4 s after the sound stopped. We first discuss the results of a Memory model in which the neural underpinnings of involuntary memories can be probed in both groups. The Memory model is a 2 (Group) × 2 (Time) model. Then we discuss an Emotion model in which we contrast high and low emotion trials. Since all results shown are for the negative > neutral trials, this model is also a 2 (Group) × 2 (Time) design.

3.2.1. Memory model: paired sounds > unpaired sounds

3.2.1.1. Group × time interaction. Results from the Group × Time interaction revealed timing differences in a similar set of regions for controls (early > late) and PTSD (late > early): bilateral parahippocampal gyrus, precuneus, IPC, and PCC exhibited delayed activity in the PTSD group compared to the control group (Fig. 2, Table 4). There was no activity for the reverse contrast (control late > early, PTSD early > late).

3.2.1.1.1. Group × time interaction follow-up analyses. To further explore the change in neural activity between the early and the late time points in the control and PTSD groups, we test the observation that our results may be due to a difference between groups at one time window but not the other. We first calculated the average BOLD signal estimates for each time point for each of the six regions shown in Fig. 2. We then subtracted the early and late BOLD signal estimates to determine the change in activity between the early minus the late time points in the control group and the late minus the early time points in the PTSD group. Finally, we calculated the ratio of the change in activity between time points. The closer the ratio is to one, the more similar the change in activity between time points. The average ratio across all six regions is 0.659, suggesting that the change in activity between the early and late time points was similar across groups. See Table S1 for more details.

Additionally, we examined the paired > unpaired sound contrast (i.e. sounds that elicited a memory > sounds that did not elicit a memory) during the early time point for the control group combined with the late time point for the PTSD group. This analysis revealed whether activity at the early time point for the control group and the late time point for the PTSD group was greater than zero. There was significant activity in posterior midline regions and inferior parietal cortex. When the threshold was raised slightly to \( p < 0.055 \), FDR corrected for multiple comparisons, our a priori hypothesized region, the parahippocampal gyrus, was also active (see Fig. S1 and Table S2). There was no activity for the control group late combined with the PTSD group early. This supports our claim that involuntary memories elicit activity in a similar network of regions at different times for the two groups.

3.2.1.2. Main effect of group. There was no significant main effect of group, suggesting that the regions associated with memory retrieval are similar between groups. To test for similarities within the memory network between groups, we conducted two additional analyses. First, we extracted estimates of activity in these regions for both groups in both time points and ran t-tests to compare the groups. We compared control early to PTSD late and compared PTSD early to control late since the Group × Time analysis suggested that the memory network was similar between groups but peaked at different times. There were no differences in activity between the early time point in the control group and the late time point in the PTSD group in any region and the absolute value of the t-scores was lower than one for five out of the six regions of interest. Turning to differences between the late time point in the control group and the early time point in the PTSD group, there were no differences in activity and t-scores were lower than 1.2 for the all regions (see Table 5). This suggests that memory network activity is similar between groups but with different peak times.

Second, we lowered the threshold for the test of the Main Effects of Group to determine whether there were differences between groups at a slightly more lenient threshold. There was no activity for control > PTSD at a threshold as high as \( p < 0.45 \), FDR corrected for multiple comparisons. Even raising the threshold as high as \( p < 0.65 \) only revealed a 16-voxel cluster in the postcentral gyrus. There was no activity for PTSD > control at a threshold as high as \( p < 0.95 \). The lack of activity in regions thought to be involved in memory retrieval even at these extremely lenient thresholds supports our claim that activity in these regions is similar between groups but with peaks at different times.

3.2.1.3. Main effect of time. There was no significant main effect of time.

3.2.2. Emotion model: high versus low emotion

Having established that high and low emotion involuntary memories elicit activity in regions that we predicted, but at different times for the control and PTSD groups, we turn to examining the differences between high and low emotion involuntary memories.

3.2.2.1. Comparing high > low emotion involuntary memories across group and time: group × time interaction. There was no significant Group × Time interaction.

3.2.2.2. Comparing high > low emotion involuntary memories across group and time: main effect of group. Across both time points, the PTSD group exhibited greater activity in vmPFC and ACC. (Fig. 3, Table 6).

3.2.2.3. Comparing high > low emotion involuntary memories across group and time: main effect of time. There was no significant main effect of Time.

4. Discussion

We introduce three novel findings on the neural basis of emotional and neutral involuntary memories in PTSD. First, a similar network of regions is active during involuntary memory retrieval for PTSD and controls. Second, compared to controls, individuals with PTSD showed delayed neural responsibility in regions typically associated with memory retrieval (Kim, 2013; Spaniol et al., 2009) and previously shown to be associated with involuntary memory retrieval in healthy adults (Hall et al., 2014). Third, compared to controls, individuals with PTSD showed increased vmPFC/ACC activity for high > low emotion stimuli.

4.1. Finding 1: similarities in the memory network between PTSD and controls

Involuntary memories elicited similar activity between controls and PTSD in regions involved in the memory network, including the MTL, precuneus, PCC, and IPC. This is in contrast with neurobiological models of PTSD based on voluntary memory retrieval that emphasize dysfunctional activity in the medial temporal lobes and related memory circuits (Carrion et al., 2010; Geuze et al., 2008; Rauch et al., 2006; Werner et al., 2009). Also, in contrast with the voluntary memory literature, we show that the PTSD group had more involuntary memories...
than the control group. This is consistent with work showing a positive correlation between the frequency of involuntary memories and PTSD symptoms (Berntsen et al., 2015) whereas voluntary memory retrieval, as measured by neuropsychological testing, is impaired in PTSD (Samuelson, 2011) and may even be a risk factor for PTSD (Vasterling et al., 2018). One reason for these deficits could be that individuals with PTSD have decreased executive control functioning (Hayes et al., 2011; Johnsen and Asbjørnsen, 2009; for reviews, see Aupperle et al., 2012;
Table 4
Memory model. Top peaks from the Group × Time interaction. All results are significant at $p < 0.05$, FDR corrected for multiple comparisons, cluster corrected at $p < 0.05$, FWE-corrected. Note: peaks from the ROIs were not among the top peaks in this table but are shown in Fig. 2.

| Region                        | Cluster size | Side   | x    | y    | z    | t-Score |
|-------------------------------|--------------|--------|------|------|------|---------|
| Control early ≥ late, PTSD late ≥ early |              |        |      |      |      |         |
| Orbifrontal cortex            | 73,321       | Right  | 26   | 22   | −16  | 5.04    |
| Cerebellum                    |              | Left   | −10  | −72  | −36  | 5.01    |
| Insula                        |              | Right  | 38   | −2   | −2   | 4.67    |
| Claustrum                     |              | Right  | 30   | −20  | 14   | 4.52    |
| Fusiform gyrus                |              | Right  | 30   | −48  | −18  | 4.49    |
| Lenticular nucleus            |              | Right  | 20   | −12  | 4    | 4.35    |
| Postcentral gyrus             |              | Right  | 40   | −30  | 54   | 4.27    |
| Orbifrontal cortex            |              | Left   | −20  | 26   | −16  | 4.24    |
| Precentral gyrus              |              | Right  | 34   | −16  | 56   | 4.21    |
| Cuneus                        |              | Left   | −16  | −94  | 0    | 4.19    |
| Mid cingulate cortex          |              | Right  | 6    | −8   | 44   | 4.17    |
| vlPFC (Inferior frontal gyrus)|              | Right  | 38   | 24   | −2   | 4.16    |
| Anterior cingulate cortex     |              | Right  | 18   | 28   | 30   | 4.15    |
| Inferior parietal cortex      |              | Right  | 58   | −30  | 28   | 4.14    |
| Lenticular nucleus            |              | Right  | 16   | 10   | −8   | 4.13    |

Control late ≥ early, PTSD early ≥ late
None

Table 5
A between-group, between-time point comparison of activity associated with involuntary memory retrieval. The regions are shown in Fig. 2. There is similar activity between the early time point in the control group and the late time point in the PTSD group in all regions. There is also similar activity between the late time point in the control group and the early time point in the PTSD group in all regions. This suggests that a similar neural network underlies involuntary memory retrieval in PTSD and controls with the difference being the time that it peaks.

| Region                        | Cluster size | Side   | x    | y    | z    | t-Score |
|-------------------------------|--------------|--------|------|------|------|---------|
| Control early, PTSD late      |              |        |      |      |      |         |
| Orbitofrontal cortex          | 73,321       | Right  | 26   | 22   | −16  | 5.04    |
| Cerebellum                    |              | Left   | −10  | −72  | −36  | 5.01    |
| Insula                        |              | Right  | 38   | −2   | −2   | 4.67    |
| Claustrum                     |              | Right  | 30   | −20  | 14   | 4.52    |
| Fusiform gyrus                |              | Right  | 30   | −48  | −18  | 4.49    |
| Lenticular nucleus            |              | Right  | 20   | −12  | 4    | 4.35    |
| Postcentral gyrus             |              | Right  | 40   | −30  | 54   | 4.27    |
| Orbifrontal cortex            |              | Left   | −20  | 26   | −16  | 4.24    |
| Precentral gyrus              |              | Right  | 34   | −16  | 56   | 4.21    |
| Cuneus                        |              | Left   | −16  | −94  | 0    | 4.19    |
| Mid cingulate cortex          |              | Right  | 6    | −8   | 44   | 4.17    |
| vlPFC (Inferior frontal gyrus)|              | Right  | 38   | 24   | −2   | 4.16    |
| Anterior cingulate cortex     |              | Right  | 18   | 28   | 30   | 4.15    |
| Inferior parietal cortex      |              | Right  | 58   | −30  | 28   | 4.14    |
| Lenticular nucleus            |              | Right  | 16   | 10   | −8   | 4.13    |

Control late ≥ early, PTSD early ≥ late
None

Table 6
Emotion model. Main effect of Group (PTSD versus Control). All activity is significant at $p < 0.05$, FDR corrected, cluster-corrected at $p < 0.05$, FWE corrected. A. Activity in the mPFC/ACC. Activity in this region is greater for high > low emotion stimuli in the PTSD group compared to the control group. B. Time courses plotting high > low emotion differences at each TR. Each TR is 2 s. The first, second, and seventh TRs were not included in any analyses and are included for illustrative purposes only. Activity relating to the early and late time points is indicated with light and dark gray boxes over the respective time points. This graph indicates that the high > low emotion difference is greater for the PTSD group than the control group across both time points. C. Time courses for high and low emotion stimuli plotted separately.

| Region                        | Cluster size | Side   | x    | y    | z    | t-Score |
|-------------------------------|--------------|--------|------|------|------|---------|
| PTSD ≥ control                |              |        |      |      |      |         |
| Anterior cingulate cortex     | 646          | Left   | −8   | 30   | −4   | 4.80    |
| Ventromedial prefrontal cortex|              | Right  | 8    | 62   | −2   | 4.58    |
| Control ≥ PTSD                |              |        |      |      |      |         |
None

4.2. Finding 2: delayed activity in the memory network in PTSD

Analyses indicated a lag in activity in the memory network in PTSD compared with controls. This timing effect is extremely strong, with over 70,000 voxels in 15 peaks showing earlier activity in the control group compared to no significant activity peaking earlier in the PTSD group. This effect has negligible odds of occurring by chance. Moreover, the activity includes locations we predicted would be active during involuntary memory retrieval based on the existing literature. These findings align with both behavioral work showing delayed response times in PTSD during voluntary memory retrieval (Werner et al., 2009) and neuroimaging work showing slow neural responsivity in PTSD in similar regions during voluntary memory retrieval (St Jacques et al., 2011a). Our results indicate that this processing delay is not limited to effortful retrieval.
the necessity of investigating the difference in the timing of the neural response to such memories. In the current study, when activity was compared between groups in the early time point, there appeared to be hypoactivity in the controls and hyperactivity in the PTSD group in similar regions. It was not until later activity was examined that activity in these regions was revealed in the PTSD group. This suggests that hypoactivity found in Whalley et al. (2013) and other studies (Carrion et al., 2010; Geuze et al., 2008; Rauch et al., 2006; Werner et al., 2009) could be reexamined in light of these results.

We offer four possible explanations for this delayed activity in the memory network in PTSD. The first explanation is that involuntary memories take longer to construct in PTSD. The involuntary memories measured here were more vivid in PTSD than controls. This finding is consistent with other work showing highly vivid intrusive memories in PTSD (Bernsten et al., 2003; Michael et al., 2005) that decrease in vividness after therapy (Hackmann et al., 2004). Memories become more specific and likely more vivid as more time passes to construct the memory (Daselaar et al., 2008; Haque and Conway, 2001). It is possible that more time is needed to construct highly vivid memories in PTSD. A second possible explanation is that the slowed processing speed is driven by factors that affect both effortful and non-effortful cognitive processes. Not only do PTSD patients have slowed reaction times to tasks that require cognitive control (Barrett et al., 1996; Vasterling et al., 2002; for a review, see Aupperle et al., 2012) but they have slower reaction times during reflexive attention tasks (Clark et al., 2003; Veltmeyer et al., 2005), which may reflect a generalized cognitive slowing. This could be caused by increased fatigue or decreased cognitive capacity. PTSD participants in our study did not have slowed reaction times to the localization task. This is likely because participants were given 4 s to respond and speeded responding was not emphasized in the instructions. Other work has shown that when speed is not emphasized, reaction time is not slowed in PTSD (e.g. Dickie et al., 2008; St Jacques et al., 2011a). Third, this processing delay is the result of increased cognitive demand caused by coping with internal psychological distress or monitoring the environment for threats (Jennings, 1986). Fourth is that people with PTSD try to avoid having memories. Using the think/no-think paradigm (Anderson and Green, 2001) activity during memory retrieval can be modulated by instructing participants to not think of a memory (Depue et al., 2007; Detre et al., 2013). One of the required symptoms of PTSD is avoidance (American Psychiatric Association, 2013), which involves trying to avoid having emotional memories or to suppress the emotional response. People with PTSD avoid thinking about their trauma (Bernsten et al., 2003; Moulds et al., 2007) and avoid emotionally processing memories (Williams and Moulds, 2007). In contrast, people without PTSD are less likely to suppress or distract themselves after an involuntary memory (Reynolds and Brewin, 1998). However, suppression is generally unsuccessful in PTSD (Catarino et al., 2015), so it is possible that the attempt at suppression only delays the onset of the memory. Future work controlling for these factors and investigating different types of involuntary memories, like trauma-specific memories, will shed light on this issue.

4.3. Finding 3: hyperactivity in vmPFC/ACC

There was also increased vmPFC/ACC activity in PTSD compared to controls for high versus low emotion involuntary memories. Fear regulation models have posited that the vmPFC/ACC modulates fear extinction and emotion regulation by downregulating activity in the amygdala (Koenigs and Grafman, 2009; Rauch et al., 2006); the vmPFC/ACC may be involved in regulating other emotions as well (Bush et al., 2000; Etkin et al., 2011). In PTSD, the vmPFC/ACC is typically hypoactive, putatively leading to a hyperresponsive amygdala (Shin et al., 2006). This has also been shown in voluntary recall of traumatic events and emotional scenes (Frewen et al., 2008; Lanius et al., 2007; Protopopescu et al., 2005; Shin et al., 1997, 2004; Whalley et al., 2009) but it is not clear whether this hypoactivity could be expected to generalize to involuntary memories. While there is no previous work on the neural basis of involuntary memories in PTSD to inform this question, involuntary memories do share traits with studied phenomena, like responses to unexpected stimuli. Models discussing this vmPFC/ACC hypoactivity are based on task paradigms in which the emotional items are the focal point (Bremner et al., 1999; Britton et al., 2005; Hou et al., 2007; Lanius et al., 2001, 2003; Mazza et al., 2013; Phan et al., 2006; Shin et al., 1999, 2004, 2005; Williams et al., 2006), and are thus expected. However, investigating responses to unexpected stimuli is critical because many key symptoms revolve around such responses (American Psychiatric Association, 2013). In fact, unexpected stimuli appear to elicit different neural responses than do expected stimuli. For example, stimuli presented quickly, below the threshold of conscious detection, elicited heightened vmPFC/ACC activity in PTSD compared to controls (Bryant et al., 2008a, 2008b, 2010), as do emotional stimuli presented as distractors (Bruce et al., 2013; Fani et al., 2012; Hayes et al., 2009; Kim et al., 2008, but see Shin et al., 2007). Our results suggest that the unexpected nature of involuntary memories may elicit hypervigilant activity not predicted by prevailing PTSD models. Hyperactivity in these contexts may reflect the generation of non-conscious fear signals that reorient attention to threatening stimuli (Bryant et al., 2008b; Liddell et al., 2005).

Consistent with the idea that vmPFC/ACC hypovigilance is not a canonical PTSD correlate, as implied by prevailing PTSD models, is a lesion study showing that vmPFC lesions decrease the likelihood of developing PTSD (Koenigs and Grafman, 2009). If vmPFC/ACC hypoactivity results in increased amygdala activity, then a lesion here should cause increased emotional responsivity and increased PTSD symptomatology. The role of the vmPFC is multifaceted. It has been shown to be involved in self-referential thought (Motzkin et al., 2015; St Jacques et al., 2011b), and the retrieval of contextual details and initiating subsequent action based on those details (Keveraga et al., 2011; Panichello et al., 2012). Understanding the circumstances under which there is hypo- versus hyperactivity here and integrating what is known about the role of this region in other cognitive processes is vital to understanding emotion dysfunction in PTSD.

It can also be noted that the control group had greater activity in the vmPFC/ACC for the neutral stimuli than the negative. It is possible that this relates to attention reorienting (Persaud et al., 2002) or safety signaling (Abs et al., 2015) in controls but not PTSD, but these speculative ideas warrant direct investigation in future work.

4.4. Limitations

This study has several limitations. First, we used the post-scan self-report of memory retrieval and effort to classify trials as successful, low effort memory trials or not. We classified trials as successful based on participants’ self-report of memory retrieval and effort. Though the use of self-report to classify trials may lead to the misclassification of some trials, there is both behavioral and neural evidence that this paradigm does elicit involuntary memories with high frequency. When unique cues are used to cue memories of unique scenes, similar to the cues and scenes used in this paradigm, and participants report the retrieval of involuntary memories as they occur in real time, almost 60% of the cues elicit involuntary memories (Bernsten et al., 2013). This rate is similar to the frequency with which involuntary memories are elicited in our control group, shown above. Additionally, involuntary memories occur more frequently in PTSD than in controls (Rubin et al., 2011), similar to the pattern that we find. Finally, the similarity in neural activity for voluntary and involuntary memory retrieval (Hall et al., 2014) suggests that the use of post-hoc self-report to classify trials is reasonably accurate.

Second, our claim that there is similar activity between groups in regions critical for memory retrieval, but at different times, is derived from the Group x Time interaction effect in the Memory Model, and the lack of a Main Effect of Group. To ensure that the similar activity did

S.A. Hall et al.
NeuroImage: Clinical 19 (2018) 793–804
not rely solely on a null result, we conducted four additional analyses to make four additional claims. In the first analysis, we focused on the six key regions of interest for memory retrieval. We conducted t-tests comparing activity between the early time point in the control group and the late time point in the PTSD group, as well as between the late time point in the control group and the early time point in the PTSD group. The t-scores were all smaller than 1.5 and for 9 of the 12 they were smaller than one. In the second analysis, we showed that there was no Main Effect of Group even at extremely lenient thresholds. In the third analysis, we showed that there was significant activity in these critical memory regions for the control group early and the PTSD group late, and that there was no activity for the control group late and the PTSD group early. This shows that activity in these regions was not driven solely by the difference between the early and late time points, but that the activity in both groups is significantly greater than zero.

In the fourth analysis, we compared the change in activity between the early and late time points in the six regions of interest between groups and showed that it was similar between groups for most regions. Together, this set of supplemental analyses lend significant support to our claim that the activity in this involuntary memory network is similar between groups but that it peaks at different times between groups.

Third, encoding processes are sometimes disrupted in PTSD (Samuelson, 2011), suggesting that the neural differences we find may be due to differences in encoding. However, impaired encoding should result in impaired voluntary retrieval. The voluntary post-scan recall scores were similar between groups, suggesting that in our paradigm, encoding was preserved in PTSD. This could be due to having several encoding sessions.

Fourth, we did not directly measure the relatedness of the stimuli to our participants’ trauma. However, using trauma descriptions, we determined the relatedness of each picture to each individual’s trauma and found that very few pictures were trauma-relevant for anyone. Therefore, we do not think this issue will have a large impact on results. Again, it should be underscored that the aim of the study was to examine the neural underpinning of involuntary memory processing generally in PTSD, and not specifically in relation to personally traumatic material as this overgeneralization to less traumatic situations is what produces many of the symptoms and dysfunctions of the disorder.

However, since the present study was not designed to examine neural activity underlying trauma-related involuntary memories in PTSD, it is therefore not clear whether the findings of delayed activity in the memory network in PTSD will generalize to involuntary, intrusive memories of real-life traumatic events. This is an important question for future research to address.

Fifth, we used a relatively high-functioning population; all participants were students or were in the workforce. It is possible that people with more debilitating PTSD symptoms could have different neural responses to involuntary memories.

4.5. Conclusions

Understanding the neural mechanisms underlying emotional involuntary memories can pave the way toward the development of a treatment that could alleviate this key symptom of PTSD and can also add to the understanding of the basic emotional processing deficits seen in PTSD. The importance of involuntary memories to PTSD is widely recognized with work seeking to understand the mechanisms underlying the encoding of such memories (Clark et al., 2014, 2016). However, the current study fills a gap in understanding the mechanisms underlying their retrieval. We show that existing models of PTSD do not predict spatiotemporal neural patterning of involuntary memories particularly well. Given that many of the symptoms of PTSD revolve around responses to the unexpected (involuntary memories, startle responses, hypervigilance toward unknown threats), our innovative approach opens up additional lines of inquiry that could be key to understanding the fundamental mechanisms underlying these symptoms.

Acknowledgments

The authors wish to thank the Danish National Research Foundation (DNRF89) and the National Institute of Mental Health (grant R01 MH066079) for funding.

Financial disclosures

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

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.nicl.2018.05.009.

References

Ahs, F., Kragel, P.A., Zielinski, D.J., Brady, R., LaBar, K.S., 2015. Medial prefrontal pathways for the contextual regulation of extinguished fear in humans. NeuroImage 122, 262–271. http://dx.doi.org/10.1016/j.neuroimage.2015.07.051.

Amour, R.L., Larsen, R., Liberzon, I., 2000. Emotional processing in combat-related posttraumatic stress disorder: a comparison with traumatized and normal controls. J. Anxiety Disord. 14, 219–238.

American Psychiatric Association, 2013. Diagnostic and Statistical Manual of Mental Disorders: DSM-5. American Psychiatric Association, Washington, D.C.

Anderson, M.C., Green, C., 2001. Suppressing unwanted memories by executive control. Nature 410, 366–369. http://dx.doi.org/10.1038/35066572.

Appperle, R.L., Melrose, A.J., Stein, M.B., Paulus, M.P., 2012. Executive function and PTSD: disengaging from trauma. Neuropharmacology 62, 686–694. http://dx.doi.org/10.1016/j.n薪orpharm.2011.02.008.

Barrett, D.H., Green, M.L., Morris, R., Giles, W.H., Croft, J.B., 1996. Cognitive functioning and posttraumatic stress disorder. Am. J. Psychiatry 153, 1492–1494.

Battaglini, E., Liddell, B., Das, P., Malhi, G., Felmingham, K., Bryant, R.A., 2016. Intrusive memories of distressing information: an fMRI study. PLoS ONE 11. http://dx.doi.org/10.1371/journal.pone.0140871.

Berntsen, D., 2009. Intrusive autobiographical memories: An Introduction to the Unbidden past. Cambridge University Press, New York.

Berntsen, D., 2010. The unbidden past: involuntary autobiographical memories as a basic mode of remembering. Curr. Dir. Psychol. Sci. 19, 138–142. http://dx.doi.org/10.1177/0963721410370391.

Berntsen, D., Willert, M., Rubin, D.C., 2003. Splintered memories or vivid landmarks? Qualities and organization of traumatic memories with and without PTSD. Appl. Cogn. Psychol. 17, 675–693. http://dx.doi.org/10.1002/acp.894.

Berntsen, D., Staugaard, S.R., Svennen, L.M., 2013. Why am I remembering this now? Predicting the occurrence of involuntary (spontaneous) episodic memories. J. Exp. Psychol.-Gen. 142, 426–444. http://dx.doi.org/10.1037/a0029128.

Berntsen, D., Rubin, D.C., Salgado, S., 2015. The frequency of involuntary autobiographical memories and future thoughts in relation to daydreaming, emotional distress, and age. Conscious. Cogn. 36, 352–372. http://dx.doi.org/10.1016/j.concog.2015.07.007.

Bonanno, G.A., Papa, A., Lalande, K., Westphal, M., Coullman, K., 2004. The importance of being flexible - the ability to both enhance and suppress emotional expression predicts long-term adjustment. Psychol. Sci. 15, 482–487. http://dx.doi.org/10.1111/j.1467-9280.2004.01705.x.

Bourne, C., Mackay, C.E., Holmes, E.A., 2013. The neural basis of flashback formation: the impact of viewing trauma. Psychol. Med. 43, 1521–1532. http://dx.doi.org/10.1017/S0033291712002358.

Bremner, J.D., Staib, L.H., Kaloupek, D., Southwick, S.M., Soufer, R., Charney, D.S., 1999. Neural correlates of exposure to traumatic pictures and sound in Vietnam combat veterans with and without posttraumatic stress disorder: a positron emission tomography study. Biol. Psychiatry 45, 806–816. http://dx.doi.org/10.1016/S0006-3223(98)00297-2.

Britton, J.C., Phan, K.L., Taylor, S.F., Fig, L.M., Liberzon, I., 2005. Corticolimbic blood flow in posttraumatic stress disorder during script-driven imagery. Biol. Psychiatry 57, 832–840. http://dx.doi.org/10.1016/j.biopsych.2004.12.025.

Browaw, K.H., Offringa, R., Pfaff, D.L., Hughes, K.C., Shin, L.M., 2010. The neural correlates of emotional memory in posttraumatic stress disorder. Biol. Psychiatry Neuroplast. Anxiety Disord. 68, 1023–1030. http://dx.doi.org/10.1016/j.biopsych.2010.07.018.

Bruce, S.E., Buchholz, K.R., Brown, W.J., Yan, L., Durbin, A., Sheline, Y.I., 2013. Altered
emotional interference processing in the amygdala and insula in women with post-
traumatic stress disorder. Neuroimage-Clin. 2, 43–49. http://dx.doi.org/10.1016/j.
nicl.2012.11.003.
Bryant, R.A., Felmingham, K., Kemp, A., Das, P., Hughes, G., Peduto, A., Williams, L.,
2006. Amygdala and ventral anterior cingulate expression predicts treatment re-
sponse to cognitive behaviour therapy for post-traumatic stress disorder. Psychol.
Med. 38, 555–561. http://dx.doi.org/10.1017/S0033291706002211.
Bryant, R.A., Kemp, A.H., Felmingham, K.L., Liddell, B., Olivier, G., Peduto, A., Gordon, E.,
Williams, L.M., 2008b. Enhanced amygdala and medial prefrontal activation during
nonconscious processing of fear in posttraumatic stress disorder: an fMRI study.
Hum. Brain Mapp. 29, S17–S23. http://dx.doi.org/10.1002/hbm.20415.
Bryant, R.A., Kemp, A.H., Felmingham, K., Liddell, B.J., Olivier, G., Peduto, A., Gordon, E.,
Williams, L.M., 2010. Simulating emotional responses in posttraumatic stress
disorder: an fMRI study. Psychol. Inj. Law 3, 111–117.
Buckner, R.L., Carroll, D.C., 2007. Self-projection and the brain. Trends Cogn. Sci.
11, 490–497. http://dx.doi.org/10.1016/j.tics.2007.04.003.
Bush, G., Lu, P., Posner, M.I., 2000. Cognitive and emotional influences in anterior
cingulate cortex. Trends Cogn. Sci. 4, 215–222. http://dx.doi.org/10.1016/S1364-
6613(00)01483-2.
Buxton, R.B., Ugurbil, K., Dubowitz, D.J., Liu, T.T., 2004. Modeling the hemodynamic
response to brain activation. Neuroimage Math. Brain Imaging 23, S229–S233.
http://dx.doi.org/10.1016/j.neuroimage.2004.07.013.
Cabeza, R., Ciaramelli, E., Olson, I.R., Moscovitch, M., 2008. The parietal cortex and
episodic memory: an attentional account. Nat. Rev. Neurosci. 9, 613–625. http://dx.doi.
group=10.1038/nrn2459.
Carrion, V.G., Haas, B.W., Garrett, A., Song, S., Reiss, A.L., 2010. Reduced hippocam-
pal activity in youth with posttraumatic stress symptoms: an fMRI study. J. Pediatr.
Psychol. 35, 560–569. http://dx.doi.org/10.1093/jpepsy/jsp212.
Catarino, A., Kupper, C.S., Werner-Seidler, A., Dalgleish, T., Anderson, M.C., 2015.
The role of temporary memory suppression in the treatment of posttraumatic stress
symptoms: a randomized controlled trial. Behav. Res. Ther. 62, 37–46. http://dx.doi.
group=10.1016/j.brat.2014.07.019.
Clark, I.A., Holmes, E.A., Woolrich, M.W., Mackay, C.E., 2014. Intrusive memories to
traumatic footage: the neural basis of their encoding and involuntary recall. Psychol.
Meth. 46, 505–518. http://dx.doi.org/10.1177/0146327313511907.
Davelaar, S.M., Rice, H.J., Greenberg, D.L., Cabeza, R., Labal, K.S., Rubin, D.C., 2008.
The spatiotemporal dynamics of autobiographical memory: neural correlates of recall,
emotional intensity, and reliving. Cereb. Cortex 18, 217–229. http://dx.doi.
group=10.1093/cercor/bhm048.
Depue, R.E., Carman, T.B., Mathi, M.T., 2007. Prefrontal regions orchestrate suppression
of emotional memories via a two-phase process. Science 317, 215–218. http://dx.doi.
group=10.1126/science.1139560.
Dettre, G.J., Natarajan, A., Gershman, S.J., Norman, K.A., 2013. Moderate levels of acti-
vation lead to forgetting the think/no-think effect. Neuropsychologia 51, 2371–2388.
http://dx.doi.org/10.1016/j.neuropsychologia.2013.02.017.
Dickie, E.W., Brunet, A., Akerib, V., Armony, J.L., 2008. An fMRI investigation of memory
encoding in PTSD: influence of symptom severity. Neuropsychologia 46, 1522–1531.
http://dx.doi.org/10.1016/j.neuropsychologia.2007.11.020.
Etkin, A., Egner, T., Kalisch, R., 2011. Emotional processing in anterior cingulate and
medial prefrontal cortex. Trends Cogn. Sci. 15, 85–93. http://dx.doi.org/10.1016/
S1364-6613(10)70003-0.
Fani, N., Tone, E.B., Phifer, J., Norrholm, S.D., Bradley, B., Ressler, K.J., Kamkwalala, A.,
Williams, L.M., 2005. A direct brainstem-amygdala-cortical “alarm” system for sub-
liminal signals of fear. Neuroimage 24, 235–243. http://dx.doi.org/10.1016/j.
nenimage.2004.08.016.
Maia, T.V., Tempelta, D., Pizzini, G., Catalucci, A., Gallucci, M., Ferrari, M., 2013.
Regional cerebral changes and functional connectivity during the observation of
negative emotional stimuli in subjects with post-traumatic stress disorder. Eur. Arch.
Psychiatry Clin. Neurosci. 263, 575–583. http://dx.doi.org/10.1007/s00406-013-
0246-4.
Mehta, T., Ehlers, A., Halligan, S.L., Clark, D.M., 2005. Unwanted intrusions: what
intrusion characteristics are associated with PTSD? Behav. Res. Ther. 43, 131–148.
http://dx.doi.org/10.1016/j.brat.2004.04.006.
Motzkin, J.C., Philippi, C.L., Wolf, R.C., Baskaya, M.K., Koenigs, M., 2015. Ventromedial
cingulate cortex activation during trauma-unrelated emotional interference in PTSD.
J. Psychiatr. Res. 62, 37–46. http://dx.doi.org/10.1016/j.jpsychires.2014.07.003.
Orcutt, H.K., Bonanno, G.A., Hannan, S.M., Miron, L.R., 2014. Prospective trajectories of
symptom severity. Neuropsychologia 46, 1522–1531. http://dx.doi.org/10.1016/j.
nenimage.2007.11.020.
Bryant, R.A., Felmingham, K., Kemp, A., Das, P., Hughes, G., Peduto, A., Gordon, E.,
Williams, L.M., 2005. A direct brainstem-amygdala-cortical “alarm” system for sub-
liminal signals of fear. Neuroimage 24, 235–243. http://dx.doi.org/10.1016/j.
nenimage.2004.08.016.
Maia, T.V., Tempelta, D., Pizzini, G., Catalucci, A., Gallucci, M., Ferrari, M., 2013.
Regional cerebral changes and functional connectivity during the observation of
negative emotional stimuli in subjects with post-traumatic stress disorder. Eur. Arch.
Psychiatry Clin. Neurosci. 263, 575–583. http://dx.doi.org/10.1007/s00406-013-
0246-4.
Michael, T., Ehlers, A., Halligan, S.L., Clark, D.M., 2005. Unwanted memories: what
intrusion characteristics are associated with PTSD? Behav. Res. Ther. 43, 131–148.
http://dx.doi.org/10.1016/j.brat.2004.04.006.
