### **Brief Communication**

# Emotional arousal impairs association memory: roles of prefrontal cortex regions

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The brain processes underlying impairing effects of emotional arousal on associative memory were previously attributed to two dissociable routes using high-resolution fMRI of the MTL (Madan et al. 2017). Extrahippocampal MTL regions supporting associative encoding of neutral pairs suggested unitization; conversely, associative encoding of negative pairs involved compensatory hippocampal activity. Here, whole-brain fMRI revealed prefrontal contributions: dmPFC was more involved in hippocampal-dependent negative pair learning and vmPFC in extrahippocampal neutral pair learning. Successful encoding of emotional memory associations may require emotion regulation/conflict resolution (dmPFC), while neutral memory associations may be accomplished by anchoring new information to prior knowledge (vmPFC).

Emotional arousal is well known to enhance memory for individual items (Schümann and Sommer 2018), but can have impairing effects on associative memory, particularly when items cannot be easily unitized and interitem associations have to be formed (Madan et al. 2012; Murray and Kensinger 2013; Bisby and Burgess 2017). The neural substrates of the impairing effect of emotional arousal on associative memory have only begun to be explored (Bisby et al. 2016; Madan et al. 2017). Emotional arousal may disrupt hippocampal functions that are critical to promote binding and thereby lead to reduced associative memory for emotionally arousing items ("disruption hypothesis") (Bisby et al. 2016). Conversely, encoding of neutral items may engage extrahippocampal medial temporal lobe (MTL) regions, areas we interpreted to promote better incidental unitization of neutral than negative items, leading to a net-decrease in associative memory for negative items ("bypassing hypothesis") (Madan et al. 2017).

Specifically, in our previous high-resolution fMRI study focussing on MTL regions (Madan et al. 2017), extrahippocampal MTL cortex was more active during encoding of neutral than negative picture pairs, showed a subsequent memory effect (SME) for neutral picture pairs, and neutral pair encoding was accompanied by more between-picture saccadic eye movements compared with negative pairs. In line with previous findings of extrahippocampal MTL areas involved in association memory formation of merged or unitized items (Giovanello et al. 2006; Quamme et al. 2007; Diana et al. 2008; Delhaye et al. 2019), we interpreted our fMRI and eye movement findings to suggest better incidental unitization of neutral picture pairs than negative pictures pairs. A behavioral follow-up study confirmed that unitization, that is, imagining the two pictures as one ("interactive imagery"), was indeed rated as higher for neutral than negative pairs, and this advantage in interactive imagery was linked to better associative memory for neutral pairs, lending further support to the bypassing hypothesis (Caplan et al. 2019).

What would prevent emotional pairs from being as easily merged as neutral pairs? We observed that during negative pair encoding, each individual picture was fixated longer compared with neutral pictures. These longer fixation durations for negative pictures were related to greater activity during negative than neutral pair encoding in the dorsal amygdala (likely the centromedial group) (Hrybouski et al. 2016), an activation which was functionally coupled to the more ventral amygdala (likely the lateral nucleus) (Hrybouski et al. 2016). This ventral amygdala activation exhibited a subsequent forgetting effect specifically for negative pairs. Given that emotional items attract more attention to themselves and are more likely processed as individual items (Markovic et al. 2014; Mather et al. 2016), we conjectured that this may make pairs of emotional items harder to unitize and to benefit from extrahippocampal unitization-related processes such as interactive imagery. Interestingly, the hippocampus revealed a subsequent memory effect specifically for negative pairs in Madan et al. (2017). We concluded that when sufficiently arousing information precludes extrahippocampal unitization-based encoding, an alternative, compensatory, and effortful relational hippocampusdependent encoding strategy may be used.

Both findings, extrahippocampal MTL encoding for neutral pairs and compensatory hippocampal encoding for negative pairs, raise the question of which cortical areas could be involved in these two dissociable associative encoding processes. Conceivably, successful associative encoding of negative information may require participants to evaluate the emotional content, and regulate emotional arousal/conflict, functions primarily associated with dorso-medial PFC regions (dmPFC; Dixon et al. 2017), the anterior cingulate cortex (ACC) (Botvinick 2007), and posterior areas of the ventro-medial PFC (vmPFC) (Yang et al. 2020). To unitize two pictures through interactive imagery, retrieval of semantic memories and prior knowledge regarding the contents of the two pictures is likely helpful. Semantic memory processes have been attributed to the left inferior frontal gyrus (left IFG) (Binder and Desai 2011). The vmPFC (more anterior portions) could also be involved, owing to its role in relating new information during encoding to prior knowledge, that is, a "unitization-like" process (Gilboa and Marlatte 2017; Sommer 2017). Motivated by our previous discovery and interpretation of the two distinct encoding processes in the MTL (Madan et al. 2017), the potential contribution of these cortical areas in neutral and negative association memory was

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explored here by using a whole-brain scan and overcoming the limitations of our previous high-resolution fMRI sequence focused only on the MTL in Madan et al. (2017).

In the current study, we therefore acquired standardresolution whole-brain fMRI (3 Tesla Siemens Trio scanner, 3-mm thickness, TR 2.21 sec, TE 30 msec) of 22 male participants during exactly the same task as in Madan et al. (2017). Only male participants were recruited because of known sex-dependent lateralization of amygdala activity (Cahill et al. 2004; Mackiewicz et al. 2006), limiting the conclusions of the current study to males. Eye movements were assessed as a proxy for overt attention (EyeLink 1000, SR Research, 17 participants with usable eyetracking data). In each of three encoding-retrieval cycles, 25 neutral and 25 negative picture pairs were intentionally encoded. Pictures (e.g., objects, scenes, humans, and animals) were selected from the International Affective Picture System (Lang et al. 2008) and the internet, and confirmed to have different valence and arousal through independent raters (details in Madan et al. 2017). Each encoding round was followed by a two-step memory test for each pair: In a judgement of memory (JoM) task one picture served as retrieval cue and volunteers indicated their memory (yes/ no) for the other picture of the original pair. Then the same picture was centrally presented again, surrounded by five same-valence pictures (one correct target, four lures) in a five-alternative forced-choice associative recognition test. Participants chose the target picture from the array with an MR-compatible joystick.

As in our previous studies, associative recognition was less accurate for negative (NN) (M=0.47) than neutral (nn) pairs (M= 0.51; t(22)=2.49, P=0.02). Subjective memory confidence (JoM) for neutral pairs (M=0.41) was not significantly different from confidence for negative pairs (M=0.43; t(22)=1.19, P=0.25) (Fig. 1A; Madan et al. 2017; Caplan et al. 2019).

Average fixation duration (a proxy for the depth of processing of individual pictures) was longer for negative than neutral pictures ( $F_{(1,16)} = 9.59$ , P = 0.007), with no main effect of memory ( $F_{(1,16)} = 0.11$ , P = 0.75), nor emotion–memory interaction ( $F_{(1,16)} = 1.27$ , P = 0.28). The number of fixations was also higher for negative than neutral pictures ( $F_{(1,16)} = 5.56$ , P = 0.03), again with no main effect of memory ( $F_{(1)} = 1.56$ , P = 0.23) or interaction ( $F_{(1,16)} = 0.26$ , P = 0.61). The number of saccades within each picture (i.e., visual exploration within but not across items, reflecting intraitem processing) was higher for negative than neutral pairs (Fig. 1B;  $F_{(1,16)} = 33.38$ , P < 0.001), with no main effect of memory ( $F_{(1,16)} = 0.02$ , P = 0.89) nor interaction ( $F_{(1,16)} = 0.15$ , P = 0.71). However, the number of saccades between the two pictures of a pair, which may support associative processing, was substantially lower for



**Figure 1.** Behavioral and eye tracking results. (*A*) Accuracy in the associative recognition task (5-AFC) for all negative (NN) and neutral (nn) pairs. Chance level in the 5-AFC associative recognition task was 1/5 = 0.20. (*B*) Mean number of saccades between the two pictures of a pair for remembered (Hit) and forgotten (Miss) negative (NN) and neutral (nn) pairs. (C) Mean number of saccades within pictures. Error bars are 95% confidence intervals around the mean, corrected for interindividual differences (Loftus and Mason 1994).

negative than neutral pairs (Fig. 1C;  $F_{(1,16)} = 7.67$ , P = 0.01). Importantly, there were more between-picture saccades for pairs that were later remembered than forgotten, that is, a subsequent memory effect based on between-picture saccades ( $F_{(1,16)} = 8.43$ , P = 0.01). This effect did not further interact with emotion ( $F_{(1,16)} = 2.64$ , P = 0.12). Thus, association memory success was driven by interitem saccades, and these were reduced in negative trials. Participants spent more attention to individual negative than neutral pictures (fixation duration and number of within-picture saccades), but this was unrelated to association memory success.

The fMRI data were preprocessed (slice timing corrected, realigned and unwarped, normalized using DARTEL and smoothed, FWHM = 8 mm) and analyzed using SPM12. First-level models were created with four regressors that modeled the onsets of the 2 (negative and neutral) × 2 (subsequent hits and misses) conditions of interest. Results of all fMRI analyses were considered significant at P < 0.05, family-wise error (FWE) corrected for multiple comparisons across the entire scan volume or within the a priori anatomical regions of interest (ROIs). ROIs for the hippocampus, amygdala and extrahippocampal MTL were reused from our previous study (Madan et al. 2017). The prefrontal ROIs, that is, dmPFC, ACC, vmPFC and left inferior frontal gyrus ROIs, were manually traced on the mean T1 image using ITK-SNAP 3.6.0 (Yushkevich et al. 2006) following schematic drawings based on meta-analyses (Binder and Desai 2011; Dixon et al. 2017; Gilboa and Marlatte 2017).

The second-level analyses based on the resulting individual β images and subject as a random factor replicated a wellestablished network of brain areas involved in negative emotion processing (Spalek et al. 2015): greater activity during processing negative than neutral picture pairs in the amygdala, insula, right inferior frontal gyrus, mid, and anterior cingulate cortex as well as visual areas (Fig. 2A). As in our previous study, we correlated the difference in left amygdala activity with the difference in eye movements for negative minus neutral trials, showing a significant correlation with the number of within-picture saccades (r = 0.50, P = 0.018). Thus, higher left amygdala activity was associated with increased visual search within negative pictures. We conducted a psychophysiological interaction analysis (PPI) using this amygdala region as seed and contrasted functional coupling during successful versus unsuccessful negative with successful versus unsuccessful neutral pair encoding (i.e., the interaction of valence and subsequent memory success). This PPI revealed stronger coupling during successful encoding of negative compared with neutral pairs with a (nonsignificant) cluster in the dmPFC (Z=3.01, [-12, 38, 26]). Simple effects showed that the amygdala was

more strongly coupled with the dmPFC during successful than unsuccessful negative pair encoding (Z = 3.63, [-2, 16, 42]).

Neutral-pair processing was associated with greater activity than negative-pair processing in the bilateral extrahippocampal MTL cortex, ventral precuneus (vPC), retrosplenial cortex (RSC), middle occipital gyrus, and putamen (Fig. 2B). In addition, we observed a general SME irrespective of valence in the left hippocampus ([-28, -16, -24], Z=3.49, P=0.04).

An interaction between pair valence and SME with greater neutral than negative SME was observed in vmPFC (Fig. 3A), together with a (nonsignificant) cluster in right MTL cortex ([26, -24, -28], Z = 3.16, P=0.11). We conducted a PPI using this vmPFC region as seed and contrasted



**Figure 2.** Main effects of emotion—fMRI results. (*A*) Greater activity during negative than neutral pair processing irrespective of subsequent memory success. (*B*) Greater activity during neutral than negative pairs processing. *t*-maps thresholded at P < 0.001 uncorrected for visualization purposes. *t*-value color-coded.

The current findings, first, replicated the impairing effects of emotional arousal on association memory previously observed in six experiments across four studies (Madan et al. 2012, 2017; Caplan et al. 2019). We built on these previous findings here by identifying cortical, especially prefrontal areas involved in the associative memory advantage for neutral pairs and those involved in the compensatory mechanism for learning negative pairs. In particular, vmPFC activity more strongly supported successful encoding of neutral than negative pairs and during this process, showed stronger coupling with a cluster at the border between MTL cortex and hippocampus. Conversely, the dmPFC was more engaged and more strongly coupled with the hippocampus during successful negative than neutral pair encoding.

We observed more and longer fixations, as well as more within-picture saccades for individual negative pictures compared with neutral pictures, resembling previously reported eye movement findings (Bradley et al. 2011; Dietz et al. 2011). We had previously shown that increased attention (fixation duration) to individual negative pictures is linked to centromedial amygdala activity (not measurable here due to the whole-brain scan resolution), and functionally coupled with a negative pair-specific subsequent forgetting effect in the lateral amygdala (Madan et al. 2017). These findings together suggest that increased attention attracted by individual negative pictures does not support associative memory, or may even be detrimental (cf., Hockley and Cristi 1996).

functional coupling during successful vs. unsuccessful neutral with successful vs. unsuccessful negative pair encoding. This PPI revealed stronger coupling during successful encoding of neutral compared with negative pairs in a cluster at the border of the extrahippocampal MTL cortex reaching into the hippocampus ([-20, -18, -26], Z=4.61, Fig. 3B).

Conversely, an interaction between pair valence and SME showing a greater negative than neutral SME was observed in the right hippocampal region (Fig. 3C), replicating our previous finding of compensatory hippocampal encoding, and in the insula (Z=3.7, [38, 2, 8]). Within prefrontal cortex, the dorsal medial prefrontal cortex (dmPFC, Z=4.14) (Fig. 3D), also showed this effect. Neutral pairs showed a subsequent forgetting effect, that is, greater activity during unsuccessful encoding of neutral pairs, in these regions (Fig. 3 C,D).

Similar to the PPI with the vmPFC seed, we conducted a PPI with the dmPFC cluster as seed. This PPI revealed the bilateral hippocampus to be more strongly coupled with the dmPFC during successful negative than neutral pair encoding (Z=3.98, [-24, -10, -18], Z=4.71, [30, -14, -29]) (Fig. 3E). The correlational analyses of activity in the dmPFC and vmPFC (valence × encoding success interactions) with the corresponding eye-tracking measures were nonsignificant, possibly due to low reliability of difference measures (Schümann et al. 2020).



**Figure 3.** SME × Emotion interactions and PPIs. (*A*) Activity in the vmPFC revealed a SME only for neutral but not negative pairs. (*B*) This region was stronger coupled during neutral than negative pair encoding with a cluster in the border of left MTL cortex/hippocampus. (*C*,*D*) Activity in the right hippocampus and dmPFC revealed a SME only for negative pairs. (*E*) The dmPFC was stronger coupled during negative than neutral pairs encoding with the bilateral hippocampus. *t*-maps thresholded at P < 0.001 uncorrected for visualization purposes. Error bars are 95% confidence intervals around the mean, corrected for interindividual differences (Loftus and Mason 1994).

The dmPFC contributed more to negative than neutral association memory and was functionally coupled to the hippocampus, which complements our interpretation of possibly compensatory activity in the hippocampus during negative pair encoding (Madan et al. 2017). The amygdala on the other hand was stronger coupled with the dmPFC during successful encoding of negative pairs which might reflect the detection of aversive stimuli by the amygdala. The dmPFC not only plays a role in emotion regulation (Wager et al. 2008; Ochsner et al. 2012; Kohn et al. 2014; Dixon et al. 2017): It is the central node in the cognitive control network. In particular, the dmPFC regulates conflicts between goals and distracting stimuli by boosting attention toward the relevant task (Weissman 2004; Grinband et al. 2011; Ebitz and Platt 2015; Iannaccone et al. 2015). Consistent with this role in the current task, the dmPFC was functionally more strongly coupled with the bilateral hippocampus during successful negative compared with neutral pair learning. The involvement of the dmPFC during successful negative (but unsuccessful neutral) (discussed below) pair encoding may suggest that it resolves conflicts between the prepotent attention to the individual negative pictures and the current task goals, that is, their intentional associative encoding. One way to do so might involve the dmPFC's role to regulate the negative emotions elicited by the pictures in order to focus on the associative memory task.

Neutral pairs elicited more between-picture saccades than negative pairs, as in (Madan et al. 2017). The vmPFC was more strongly involved in successful associative encoding of neutral than negative pairs and more strongly coupled with the extrahippocampal MTL cortex bordering the hippocampus during successful neutral compared with negative pair encoding. Anterior vmPFC regions and their coupling with the MTL have been implicated in retrieval of consolidated memories and in anchoring new information to prior knowledge (Nieuwenhuis and Takashima 2011; van Kesteren et al. 2013; Schlichting and Preston 2015; Gilboa and Marlatte 2017; Sommer 2017; Brod and Shing 2018; Sekeres et al. 2018). We previously observed that interactive imagery (forming one instead of two images to memorize) was higher for neutral than negative pairs (Caplan et al. 2019), perhaps reflected by the increased between-picture saccades in the current study. Assuming that the anterior vmPFC subserves retrieval of prior knowledge, its engagement during successful neutral pair encoding may have supported such incidental unitization processes here as well. Negative pictures are inherently semantically more related (Barnacle et al. 2016), which implies that they may share even more prior knowledge than neutral pictures. However, the retrieval of this prior knowledge may be inhibited by the attraction of attention to individual negative pictures, not their arbitrary pairing as in the current task. Incidental unitization can occur through rather subtle manipulations (Giovanello et al. 2006; Diana et al. 2008; Bader et al. 2010; Ford et al. 2010; Li et al. 2019) or even entirely without any instruction; for example, when the items' combination is itself meaningful or familiar (Ahmad and Hockley 2014). We suggest that similar incidental unitization processes may have occurred here as well. Memory for unitized associations is independent of hippocampal memory processes and can be based solely on the extrahippocampal MTL (Quamme et al. 2007; Haskins et al. 2008; Staresina and Davachi 2010). Our previous high-resolution fMRI study supported such a bypassing hypothesis, that is, extrahippocampal MTL cortex involvement in the successful associative encoding of neutral but not negative pairs (Madan et al. 2017). Here, this interaction did not reach significance in the MTL cortex, but the P-value of 0.11 can be considered suggestive based on our strong a priorihypothesis. Notably, in our previous study using a scanning resolution of 1 mm<sup>3</sup> the cluster included only 17 voxels, which would correspond to less than one voxel here. Therefore, we assume the lower sensitivity here was due to the lower spatial resolution.

Unexpectedly, we observed greater activity during unsuccessful encoding of neutral pairs in the same regions that promoted successful encoding of negative pairs, that is, the dmPFC and hippocampal region. Hockley et al. (2016) previously observed that incidental but not intentional encoding of associations (for word pairs) improved for items with stronger pre-experimental associations. Perhaps using an effortful (dmPFC/hippocampal) learning strategy for neutral pairs, that is, pairs that are already more likely incidentally linked or linkable (e.g., through interactive imagery) may not have helped encoding. The forgotten neutral pairs underlying the SFE in these regions may then have been simply the hardest-to-learn neutral pairs; that is, pairs where both encoding strategies failed. Evidently, future studies should test such speculations directly.

Our interpretation of the dmPFC and vmPFC as signifying in emotion regulation and unitization in this task was based on previous studies. Because we did not manipulate unitization and/or emotional regulation, these processes remain hypothetical. However, within this framework, we addressed two hypotheses regarding interactions between hippocampal/extrahippocampal MTL regions and prefrontal cortex during association memory formation. The disruption hypothesis proposes that the hippocampus is equally responsible for encoding of negative and neutral association memory but that for negative memories, hippocampal activity is inhibited by the amygdala via the prefrontal cortex (Murray and Kensinger 2013; Bisby et al. 2016). The vmPFC has known involvement in negative emotion processing (Yang et al. 2020), and the observed activity pattern in the vmPFC could appear to disrupt hippocampal association memory processes for negative pairs. However, according to the bypassing hypothesis (Madan et al. 2017), successful encoding of negative (compared with neutral) pairs requires the hippocampus since fewer extrahippocampal contributions are available. Supporting the bypassing hypothesis, we observed that the vmPFC was negatively functionally coupled with extrahippocampal MTL cortex (bordering the hippocampus), suggesting that the vmPFC decreased extrahippocampal contributions to association memory for negative pairs. The bypassing hypothesis is also supported by our finding that the hippocampus was not less but more involved in negative compared with neutral pair encoding, that is, we observed no evidence for a prefrontally (e.g., vmPFC)-mediated disruption of hippocampal activity by emotion.

In conclusion, the two critical prefrontal cortex regions linked to MTL memory processes in the current study were the dmPFC, involved in successful hippocampal-dependent negative pair learning and the vmPFC, supporting successful neutral pair learning that relied on extrahippocampal MTL involvement.

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#### References

- Ahmad FN, Hockley WE. 2014. The role of familiarity in associative recognition of unitized compound word pairs. QJ Exp Psychol 67: 2301– 2324. doi:10.1080/17470218.2014.923007
- Bader R, Mecklinger A, Hoppstädter M, Meyer P. 2010. Recognition memory for one-trial-unitized word pairs: evidence from event-related potentials. *Neuroimage* 50: 772–781. doi:10.1016/j.neuroimage.2009.12 .100

Barnacle GE, Montaldi D, Talmi D, Sommer T. 2016. The list-composition effect in memory for emotional and neutral pictures: differential contribution of ventral and dorsal attention networks to successful encoding. *Neuropsychologia* **90**: 125–135. doi:10.1016/j .neuropsychologia.2016.06.023

Binder JR, Desai RH. 2011. The neurobiology of semantic memory. *Trends Cogn Sci* 15: 527–536. doi:10.1016/j.tics.2011.10.001

Bisby J, Burgess N. 2017. Differential effects of negative emotion on memory for items and associations, and their relationship to intrusive imagery. *Curr Opin Behav Sci* 17: 124–132. doi:10.1016/j.cobeha.2017 .07.012

Bisby JA, Horner AJ, Hørlyck LD, Burgess N. 2016. Opposing effects of negative emotion on amygdalar and hippocampal memory for items and associations. *Soc Cogn Affect Neurosci* **11**: 981–990. doi:10.1093/ scan/nsw028

Botvinick MM. 2007. Conflict monitoring and decision making: reconciling two perspectives on anterior cingulate function. *Cogn Affect Behav Neurosci* 7: 356–366. doi:10.3758/CABN.7.4.356

Bradley MM, Houbova P, Miccoli L, Costa VD, Lang PJ. 2011. Scan patterns when viewing natural scenes: emotion, complexity, and repetition. *Psychophysiology* **48**: 1544–1553. doi:10.1111/j.1469-8986 .2011.01223.x

Brod G, Shing YL. 2018. Specifying the role of the ventromedial prefrontal cortex in memory formation. *Neuropsychologia* **111**: 8–15. doi:10.1016/j .neuropsychologia.2018.01.005

Cahill L, Uncapher M, Kilpatrick L, Alkire MT, Turner J. 2004. Sex-related hemispheric lateralization of amygdala function in emotionally influenced memory: an FMRI investigation. *Learn Mem* **11:** 261–266. doi:10.1101/lm.70504

Caplan JB, Sommer T, Madan CR, Fujiwara E. 2019. Reduced associative memory for negative information: impact of confidence and interactive imagery during study. *Cogn Emot* **33**: 1745–1753. doi:10.1080/ 02699931.2019.1602028

Delhaye E, Mechanic-Hamilton D, Saad L, Das SR, Wisse LEM, Yushkevich PA, Wolk DA, Bastin C. 2019. Associative memory for conceptually unitized word pairs in mild cognitive impairment is related to the volume of the perirhinal cortex. *Hippocampus* **29**: 630–638. doi:10 .1002/hipo.23063

Diana RA, Yonelinas AP, Ranganath C. 2008. The effects of unitization on familiarity-based source memory: testing a behavioral prediction derived from neuroimaging data. *J Exp Psychol Learn Mem Cogn* 34: 730–740. doi:10.1037/0278-7393.34.4.730
Dietz J, Bradley MM, Okun MS, Bowers D. 2011. Emotion and ocular

 Dietz J, Bradley MM, Okun MS, Bowers D. 2011. Emotion and ocular responses in Parkinson's disease. *Neuropsychologia* 49: 3247–3253. doi:10.1016/j.neuropsychologia.2011.07.029
 Dixon ML, Thiruchselvam R, Todd R, Christoff K. 2017. Emotion and the

Dixon ML, Thiruchselvam R, Todd R, Christoff K. 2017. Emotion and the prefrontal cortex: an integrative review. *Psychol Bull* **143**: 1033–1081. doi:10.1037/bul0000096

Ebitz RB, Platt ML. 2015. Neuronal activity in primate dorsal anterior cingulate cortex signals task conflict and predicts adjustments in pupil-linked arousal. *Neuron* **85:** 628–640. doi:10.1016/j.neuron.2014 .12.053

Ford JH, Verfaellie M, Giovanello KS. 2010. Neural correlates of familiarity-based associative retrieval. *Neuropsychologia* **48:** 3019–3025. doi:10.1016/j.neuropsychologia.2010.06.010

Gilboa A, Marlatte H. 2017. Neurobiology of schemas and schemamediated memory. *Trends Cogn Sci* **21:** 618–631. doi:10.1016/j.tics.2017 .04.013

Giovanello KS, Keane MM, Verfaellie M. 2006. The contribution of familiarity to associative memory in amnesia. *Neuropsychologia* 44: 1859–1865. doi:10.1016/j.neuropsychologia.2006.03.004
Grinband J, Savitskaya J, Wager TD, Teichert T, Ferrera VP, Hirsch J. 2011.

Grinband J, Savitskaya J, Wager TD, Teichert T, Ferrera VP, Hirsch J. 2011. The dorsal medial frontal cortex is sensitive to time on task, not response conflict or error likelihood. *Neuroimage* **57:** 303–311. doi:10.1016/j .neuroimage.2010.12.027

Haskins AL, Yonelinas AP, Quamme JR, Ranganath C. 2008. Perirhinal cortex supports encoding and familiarity-based recognition of novel associations. *Neuron* **59:** 554–560. doi:10.1016/j.neuron.2008 .07.035

Hockley WE, Cristi C. 1996. Tests of encoding tradeoffs between item and associative information. *Mem Cognit* 24: 202–216. doi:10.3758/ BF03200881

Hockley WE, Ahmad FN, Nicholson R. 2016. Intentional and incidental encoding of item and associative information in the directed forgetting procedure. *Mem Cogn* **44**: 220–228. doi:10.3758/s13421-015-0557-8

Hrybouski S, Aghamohammadi-Sereshki A, Madan CR, Shafer AT, Baron CA, Seres P, Beaulieu C, Olsen F, Malykhin NV. 2016. Amygdala subnuclei response and connectivity during emotional processing. *Neuroimage* 133: 98–110. doi:10.1016/j.neuroimage.2016.02.056

Iannaccone R, Hauser TU, Staempfli P, Walitza S, Brandeis D, Brem S. 2015. Conflict monitoring and error processing: new insights from simultaneous EEG-fMRI. Neuroimage 105: 395-407. doi:10.1016/j .neuroimage.2014.10.028

Kohn N, Eickhoff SB, Scheller M, Laird AR, Fox PT, Habel U. 2014. Neural network of cognitive emotion regulation: an ALE meta-analysis and MACM analysis. *Neuroimage* 87: 345–355. doi:10.1016/j.neuroimage .2013.11.001

Lang PJ, Bradley MM, Cuthb BN. 2008. International affective picture system (IAPS): affective ratings of pictures and instruction manual. University of Florida, Gainesville, FL.

Li B, Han M, Guo C, Tibon R. 2019. Unitization modulates recognition of within-domain and cross-domain associations: evidence from event-related potentials. *Psychophysiology* **56**: e13446. doi:10.1111/psyp .13446

Loftus GR, Masson MEJ. 1994. Using confidence intervals in within-subject designs. *Psychon Bull Rev* 1: 476–490. doi:10.3758/ BF03210951

 Mackiewicz KL, Sarinopoulos I, Cleven KL, Nitschke JB. 2006. The effect of anticipation and the specificity of sex differences for amygdala and hippocampus function in emotional memory. *Proc Natl Acad Sci* 103: 14200–14205. doi:10.1073/pnas.0601648103
 Madan CR, Caplan JB, Lau CSM, Fujiwara E. 2012. Emotional arousal does

Madan CR, Caplan JB, Lau CSM, Fujiwara E. 2012. Emotional arousal does not enhance association-memory. J Mem Lang 66: 695–716. doi:10 .1016/j.jml.2012.04.001

Madan CR, Fujiwara E, Caplan JB, Sommer T. 2017. Emotional arousal impairs association-memory: roles of amygdala and hippocampus. *Neuroimage* **156**: 14–28. doi:10.1016/j.neuroimage.2017 .04.065

Markovic J, Anderson AK, Todd RM. 2014. Tuning to the significant: neural and genetic processes underlying affective enhancement of visual perception and memory. *Behav Brain Res* **259**: 229–241. doi:10.1016/j .bbr.2013.11.018

Mather M, Clewett D, Sakaki M, Harley CW. 2016. Norepinephrine ignites local hot spots of neuronal excitation: how arousal amplifies selectivity in perception and memory. *Behav Brain Sci* **39**: e200. doi:10.1017/ S0140525X15000667

Murray BD, Kensinger EA. 2013. A review of the neural and behavioral consequences for unitizing emotional and neutral information. *Front Behav Neurosci* **7:** 42. doi:10.3389/fnbeh.2013.00042

Nieuwenhuis ILC, Takashima A. 2011. The role of the ventromedial prefrontal cortex in memory consolidation. *Behav Brain Res* **218**: 325–334. doi:10.1016/j.bbr.2010.12.009

Ochsner KN, Silvers JÅ, Buhle JT. 2012. Functional imaging studies of emotion regulation: a synthetic review and evolving model of the cognitive control of emotion: functional imaging studies of emotion regulation. *Ann N Y Acad Sci* **1251:** E1–E24. doi:10.1111/j.1749-6632 .2012.06751.x

Quamme JR, Yonelinas AP, Norman KA. 2007. Effect of unitization on associative recognition in amnesia. *Hippocampus* **17:** 192–200. doi:10 .1002/hipo.20257

Schlichting ML, Preston AR. 2015. Memory integration: neural mechanisms and implications for behavior. *Curr Opin Behav Sci* 1: 1–8. doi:10.1016/j .cobeha.2014.07.005

Schümann D, Sommer T. 2018. Dissociable contributions of the amygdala to the immediate and delayed effects of emotional arousal on memory. *Learn Mem* **25**: 283–293. doi:10.1101/lm.047282.117

Schümann D, Joue G, Jordan P, Bayer J, Sommer T. 2020. Test-retest reliability of the emotional enhancement of memory. *Memory* **28:** 49– 59. doi:10.1080/09658211.2019.1679837

Sekeres MJ, Winocur G, Moscovitch M. 2018. The hippocampus and related neocortical structures in memory transformation. *Neurosci Lett* 680: 39– 53. doi:10.1016/j.neulet.2018.05.006

Sommer T. 2017. The emergence of knowledge and how it supports the memory for novel related information. *Cereb Cortex* 27: 1906–1921. doi:10.1093/cercor/bhw031

Spalek K, Fastenrath M, Ackermann S, Auschra B, Coynel D, Frey J, Gschwind L, Hartmann F, van der Maarel N, Papassotiropoulos A, et al. 2015. Sex-dependent dissociation between emotional appraisal and memory: a large-scale behavioral and fMRI study. *J Neurosci* 35: 920– 935. doi:10.1523/JNEUROSCI.2384-14.2015

Staresina BP, Davachi L. 2010. Object unitization and associative memory formation are supported by distinct brain regions. *J Neurosci* **30**: 9890– 9897. doi:10.1523/JNEUROSCI.0826-10.2010

van Kesteren MTR, Beul SF, Takashima A, Henson RN, Ruiter DJ, Fernández G. 2013. Differential roles for medial prefrontal and medial temporal cortices in schema-dependent encoding: from congruent to incongruent. *Neuropsychologia* **51**: 2352–2359. doi:10.1016/j .neuropsychologia.2013.05.027

Wager TD, Davidson ML, Hughes BL, Lindquist MA, Ochsner KN. 2008. Prefrontal-subcortical pathways mediating successful emotion regulation. *Neuron* **59:** 1037–1050. doi:10.1016/j.neuron.2008 .09.006 Weissman DH. 2004. Dorsal anterior cingulate cortex resolves conflict from distracting stimuli by boosting attention toward relevant events. *Cereb Cortex* 15: 229–237. doi:10.1093/cercor/bhh125
 Yang M, Tsai S-J, Li C-SR. 2020. Concurrent amygdalar and ventromedial

Yang M, Tsai S-J, Li C-SR. 2020. Concurrent amygdalar and ventromedial prefrontal cortical responses during emotion processing: a meta-analysis of the effects of valence of emotion and passive exposure versus active regulation. *Brain Struct Funct* 225: 345–363. doi:10.1007/ s00429-019-02007-3 Yushkevich PA, Piven J, Hazlett HC, Smith RG, Ho S, Gee JC, Gerig G. 2006. User-guided 3D active contour segmentation of anatomical structures: significantly improved efficiency and reliability. *Neuroimage* **31**: 1116– 1128. doi:10.1016/j.neuroimage.2006.01.015

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