

The slow forgetting of emotional episodic memories: an emotional binding account

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Emotional events are remembered better than neutral events, and this emotion advantage becomes particularly pronounced over time. The time-dependent effects of emotion impact upon recollection rather than on familiarity-based recognition, and they influence the recollection of item-specific details rather than contextual details. Moreover, the amygdala, but not the hippocampus, is crucial for producing these effects. Time-dependent effects of emotion have been attributed to an emotional consolidation process whereby the amygdala gradually facilitates the storage of emotional memories by other medial temporal lobe regions. However, we propose that these effects can be better understood by an emotional binding account whereby the amygdala mediates the recollection of item-emotion bindings that are forgotten more slowly than item-context bindings supported by the hippocampus.

The effects of emotion on episodic memory

The most memorable events of our lives are often those that are emotionally arousing (e.g., an encounter with a vicious dog, viewing a photograph of a gruesome murder). It is well documented that emotional materials can attract more attention or garner more elaborative encoding than neutral materials, and that this enhanced encoding can lead emotional materials to be better remembered than neutral materials (see [1–4] for review). However, the beneficial effects of emotion cannot be explained solely on the basis of enhanced encoding because, as will be described below, emotional and neutral materials can often be remembered equally well shortly after they have occurred, and it is only after a delay period that the emotion advantage begins to emerge (e.g., [5–9]).

The reason why emotional memories are so resistant to forgetting is not yet fully understood. Although important advances have been made in developing models of episodic memory (see Glossary) that incorporate findings from behavioral, lesion, and neuroimaging studies, most of these models have focused on accounting for studies of memory

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 ${\it Keywords}: {\it emotional memory}; {\it medial temporal lobes}.$

1364-6613/

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for neutral materials. In the current paper we review the behavioral and neural studies examining emotion effects on episodic memory in human subjects, and we identify several well-established empirical regularities. Based on

Glossary

Consolidation: a process of stabilizing a memory trace after it has been encoded. Synaptic consolidation is used to refer to a set of cellular/molecular processes that are engaged to support the strengthening of the synapses in a local circuit, and it is thought to occur within the first few hours after encoding. Systems consolidation is used to refer to a process whereby hippocampus-dependent memories are transferred to the cortex over a period of weeks, months, or decades. Emotional consolidation refers to the idea that after encoding, the amygdala signals the hippocampus to preferentially stabilize or protect hippocampus-dependent memories of emotional compared to neutral events.

Episodic memory: memory of a specific event that was personally experienced at a particular time or place in the past. It is typically measured using tests of recognition or recall.

Familiarity: a memory process whereby subjects discriminate between old and new items on the basis of perceived memory strength (sometimes referred to as processing fluency or a sense of recency). It is thought to be particularly useful in tests of item recognition where old items are familiar and the new items are novel, but to be somewhat less useful in relational recognition tests or recall tests 117.1001.

Recall: memory tests in which subjects are required to generate items from a previous encoding event, such as the words or images from a previous encoding liet

Receiver operating characteristic (ROC) procedure: a procedure that can be used to measure the contribution of recollection and familiarity to recognition performance [101]. The function describes the relationship between the proportion of correctly recognized studied items against the proportion of incorrectly recognized nonstudied items across variations in response criterion or confidence. A nonlinear model is fit to the observed function to estimate the probability of recollection and familiarity.

Recognition: memory tests in which subjects must discriminate between stimuli that were earlier studied from those that are new to the experimental setting. In item recognition tests, stimuli typically include words, scenes, faces, or objects. In relational recognition tests, subjects must discriminate between pairings of items or stimuli that were earlier studied from re-pairings. For example, the task may require recognizing that a particular word was studied with a particular face previously, which is sometimes referred to as associative recognition, or it may require recognizing that a particular word was encountered in a specific location, sometimes referred to as source recognition.

Recollection: a memory process whereby subjects retrieve qualitative information about a specific study event. For example, remembering that a particular object was encountered at a specific time or location, or was associated with a particular semantic and emotional state. It is expected to play a role in free recall and in tests of recognition memory, particularly relational recognition tests.

Remember/know procedure: a procedure that can be used to measure recollection and familiarity on the basis of introspective reports [102]. For each recognition response, subjects report whether they recognize items on the basis of remembering (i.e., recollection of qualitative information about the study event) or knowing (i.e., the item is familiar in the absence of recollection). Because subjects are instructed to respond 'remember' whenever they recollect a test item, the probability of a 'remember' response is used as an index of recollection, whereas the probability that an item is familiar is equal to the conditional probability that it received a 'know' response given it was not recollected [103].



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these results, we argue that emotional memories exhibit a time-dependent memory advantage because they rely on item—emotion bindings supported by the amygdala that are forgotten more slowly than item—context bindings supported by the hippocampus.

Five empirical regularities

The majority of the existing studies examining delayed emotion effects have contrasted memory for arousing negative emotional materials, such as gruesome pictures and taboo words, to memory for neutral materials, and our review will therefore focus on the effects of negative emotion as measured with these types of materials. Although this reflects a somewhat restrictive definition of emotion, later we will return to consider further whether these findings generalize to other emotional materials such as positive arousing materials, traumatic autobiographical events, as well as fear-conditioning paradigms.

The memory advantage for emotional materials increases over time

Numerous laboratory experiments have indicated that negative emotional materials are recalled and recognized better than neutral materials [1–3]. Although these effects may be due in part to enhanced encoding of emotion compared to neutral items, several studies have shown that the emotion effects are either absent or much smaller when memory is tested immediately, and they tend to increase in magnitude after a few hours [5–13].

To illustrate the delayed emotion effects we describe a study [8] in which subjects were presented with a mixture of negative and neutral images. Images were divided into two lists that were studied one day apart. Immediately after exposure to the second list, participants completed a recognition memory test for all the studied images mixed with negative and neutral nonstudied images (Figure 1). Overall recognition performance was then assessed for the emotional and neutral items that had just been studied as well as those that had been studied 24 h earlier. For the items studied and tested on the same day (i.e., the 5 minute delay condition) emotional and neutral items were recognized equally well. However, for the items studied 24 h earlier there was a recognition memory advantage for emotional materials over neutral materials. Thus, the emotional and neutral materials were equally well encoded, but after a delay an emotion advantage emerged.

Similar effects have been seen in other studies of recognition for words [6] and visual images [10,13], as well as in tests of free recall [11], and these delayed emotion effects have been shown to appear even after 2 h [10,13]. The results show that emotion effects can emerge during retention, and thus cannot be attributed to enhanced encoding. This does not preclude the possibility that emotion advantages can be seen in immediate tests. For example, encoding factors such as enhanced attention or distinctiveness can contribute to emotional memory advantages even when tested immediately (e.g., [14,15]). However, the results indicate that a simple enhanced encoding account of emotion is not sufficient to explain the long-term effects of emotion (see also [4]).

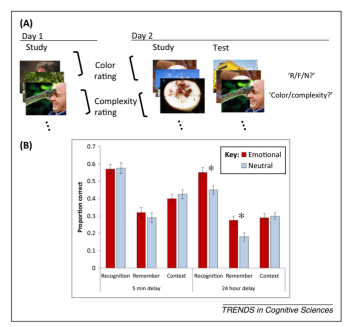


Figure 1. Procedures and results from [8]. (A) On day 1, subjects studied a mixture of negative and neutral images, half while rating visual complexity and half while rating the range of colors used in each image. On day 2, subjects studied a second list of images under similar encoding conditions, then after a 5 minute delay they received a recognition memory test containing a mixture of old items from both days and new items. For each test item subjects indicated if they could remember the occurrence of the item ('R'), if they knew it was studied on the basis of familiarity in the absence of recollection ('F'), or if they thought it was new ('N'). In addition, they indicated whether the item was encoded in the context of the color or complexity rating task. (B) The proportions of correct recognition responses are plotted for emotional and neutral materials for items tested after the 5 minute retention interval and the 24 h retention interval, Item recognition was greater for emotional than neutral materials, but only after the longer retention interval, and this effect was due to the relative increase in 'R' responses (*, statistically significant difference). By contrast, memory for the study context task was not influenced by emotion in either delay condition.

Emotion impacts recollection, rather than familiarity

Recognition memory judgments can be based either on the recollection of qualitative information about a study event or on assessments of familiarity [16–18]. Studies that have directly contrasted the contributions of recollection and familiarity to memory have indicated that emotion impacts recollection but has little or no effect on familiarity [7– 10,19–26]. Many of these studies have examined memory under conditions in which the relative increase in recollection may reflect better encoding of emotional compared to neutral items, but others have found that the recollection advantage for emotional materials is time-dependent [7-9,11,13]. For example, as seen in Figure 1, the recognition memory advantage that arose in the delay condition [8] was due to the items that were recollected. That is, in that study, a remember/know procedure was used in which subjects were required to indicate if recognition was accompanied by recollection – in the sense that they could remember some qualitative aspect of the study event, or if it was based on familiarity in the absence of recollection. The memory advantage for emotional materials that appeared in the delayed condition was entirely due to an increase in measures of recollection accuracy, and did not impact the items recognized on the basis of familiarity in the absence of recollection.

The results in this particular study were based on subjective reports of recollection and familiarity, but similar results have been observed using receiver operating characteristic methods [9,22,26]. The results show that not all forms of episodic memory benefit from emotion. Instead, emotion improves recollection, but does not benefit familiarity. Thus, any account of emotion and episodic memory needs to account for the observed selectivity of these effects.

Emotion impacts the recollection of items rather than contexts

Not all forms of recollection, however, are increased by emotion. For example, as illustrated in Figure 1, although emotional materials were more accurately remembered after the delay than were the neutral items, recollection of the encoding context was similar for emotional and neutral materials for both the immediate and delayed tests [8]. That is, in that experiment, half the images were encoded under instructions to rate the visual complexity of each image, whereas the others were encoded under instructions to rate the range of colors in each image. At test, in addition to making recognition and remember/ know responses, subjects were required to indicate which context question they had been asked about for each of the studied items. The ability to correctly recollect the encoding context was the same for emotional and neutral materials. Thus, over time, the emotional photographs themselves were more likely to be correctly recollected than neutral photos, but this did not generalize to an increase in recollection of the study context.

These effects are consistent with a growing body of research demonstrating that emotion increases recollection for details that are intrinsic to the emotional item or object [27], or to the within-item features of emotional objects [28], while having little effect, or even negative effects, on details that are extrinsic or contextual in nature. For example, emotion has been shown to increase recollection for the colors and specific visual details of emotional objects [19,29–36], but to have little, or even a negative, effect on memory for the type of encoding task, visual features of the background, or other items in the periphery [8,37–45]. These effects may be due in part to the fact that the emotional items attract attention during encoding [27,28] but, in addition, there is evidence that the emotional benefit for item compared to context information increases over time [25,46].

These results further highlight the fact that emotion effects on episodic memory are selective, and show that emotion impacts on the ability to recollect aspects of the emotional item or items rather than increasing all aspects of the emotional event such as the contextual or peripheral features of the event. Thus, emotion does not simply enhance memory for emotional events, but instead selectively improves recollection for the emotional item in the event rather than for other contextual details.

The emotion effect on episodic memory is dependent on the amygdala

Although selective amygdala lesions are rare, several cases have been reported in which bilateral amygdala damage either eliminates or severely reduces the normal advantage seen for emotional compared to neutral materials [47–50]. For example, in one study [48] individuals were presented with a set of slides along with an accompanying story that included neutral materials and negative arousing materials. In a subsequent recognition test 1 week later, patients with selective bilateral amygdala damage performed well at recognizing the neutral slides but, unlike the controls, they showed no advantage in recognizing the negative slides. A reduced memory advantage for negative materials has also been seen for verbal as well as visual materials tested with both recall and recognition [47,49]. In addition, another patient study [50] found that, in immediate tests, amygdala damage did not entirely eliminate the emotion advantage but, unlike in controls, the emotion advantage did not increase over time.

Neuroimaging results provide converging evidence that the amygdala plays a crucial role in producing the emotion effects on episodic memory. Amygdala activity during learning correlates with subsequent emotional memory (e.g., [12,51–53], see [54] for meta-analytic data), and the amygdala is more active during the retrieval of emotional relative to neutral memories [23,55-58]. Consistent with a role in recollection, amygdala activity during encoding scales with the vividness of subsequent memory [59], and its involvement during retrieval is associated with recollection rather than familiarity processes [10,23]. In addition, the amygdala is selectively involved in creating and retrieving emotional memories that carry item-specific details [36,59,60], but not necessarily other forms of contextual information, such as memory for an accompanying background scene or the decision made at encoding [39,59,60]. The amygdala seems to also be involved in promoting the time-dependent emotion advantage: amygdala activity during encoding is more strongly correlated with memory performance when memory is tested after a delay (e.g., 1-2 weeks) versus immediately ([9,61], but see [62]).

The lesion and imaging results show that the amygdala plays a central role in producing the emotion advantage in episodic memory. Moreover, the results further verify that the episodic advantage is selective to the recollection of the emotional item itself rather than influencing familiarity or the recollection of contextual or background information.

The emotion effect on episodic memory is not dependent on the hippocampus

Several studies have examined emotion and memory in patients with large medial temporal lobe (MTL) lesions that have included the amygdala, hippocampus, and surrounding perirhinal cortex, and found that these patients exhibit reduced emotion effects on memory (e.g., [11,63,64]). These impairments, however, appear to be due to amygdala damage rather than to damage elsewhere in the MTL because, in studies examining patients with MTL damage that does not include the amygdala, the emotional memory advantage is not disrupted [13,65,66]. For example, one study examining recognition memory for negative and neutral pictures found that normal controls showed no emotion advantage in an immediate test, but after a 2 h delay they showed better memory for the emotional materials than neutral materials [13]. Patients with selective hippocampal damage were tested in the delayed condition and were

found to exhibit an emotional memory advantage in recognition that was similar in magnitude to that of the controls. Another study [66] found that patients with MTL damage that did not include the amygdala exhibited a normal emotion advantage, even when controlling for overall memory performance. Similar results have been reported for story recall tests [65] and for face-emotion pair identification [64]. Finally, in temporal lobe epilepsy patients, amygdala pathology was found to be related to decreases in recollection for emotional but not neutral words, whereas hippocampal pathology was related to decreases in recollection for both emotional and neutral materials [67].

In contrast to the clear body of brain-imaging evidence linking the amygdala to emotional memory processes, neuroimaging findings have been more mixed with respect to the role of other MTL regions. Some studies have found that MTL regions contribute similarly to neutral and emotional memory formation [12,51,68], whereas others have found that the anterior hippocampus and the rhinal cortex are more involved in emotional than neutral encoding ([52], reviewed in [54]). Findings have been similarly mixed with respect to the role of the MTL during emotional retrieval (cf [10,23]). Although MTL activity on its own has not been a consistent predictor of the emotional memory advantage, several studies have found increased correlations between amygdala activity and other MTL regions during emotional than neutral memory encoding, including regions in the anterior hippocampus [12,52,68,69], anterior parahippocampal gyrus [9,12,52], and posterior parahippocampal gyrus [70]. One study found that the correlation between activation in the amygdala and rhinal cortex was stronger for items remembered after a 1 week versus 20 minute delay [9], and that this effect was related to the persistence of recollection. These findings suggest that there may be important interactions between the amygdala and hippocampus at the time of encoding, but when delay effects are taken into consideration it is the amygdala-rhinal interactions that are important for lasting memory.

In sum, the neural studies indicate that the hippocampus is not necessary for producing the delayed emotion effects. Although hippocampal damage reduces episodic memory, it reduces memory for both emotional and neutral items similarly. Moreover, hippocampal activity is related to the encoding and retrieval of both emotional and neutral materials. However, some imaging studies have suggested that the anterior hippocampus and perirhinal cortex may be more involved in emotional memory processes than are posterior regions such as the parahippocampal cortex.

The emotional binding model

How can we explain the observed effects of emotion on episodic memory? Standard models of episodic memory and the MTL (Box 1) have not typically incorporated the amygdala, which, as described above, plays a crucial role in supporting memory for emotional events and is richly connected to both the hippocampus and to the perirhinal cortex ([71], reviewed in [72]). To account for the emotion effects on memory we propose an emotional binding account whereby the amygdala supports item–emotion

Box 1. Theories of episodic memory and the MTL

Results from lesion and neuroimaging studies of humans, as well as studies of rats and nonhuman primates, have led to the development of several neuroanatomical models of memory that aim to characterize the functions of the hippocampus, the perirhinal cortex, and the parahippocampal cortex, as well as the visual processing streams that feed into those regions (see Figure 2A in main text). These models share several core assumptions and reflect what we refer to as the standard model (see Figure 2B in main text) [87-91]. In general, the hippocampus is assumed to sit at the top of a functional hierarchy, and it is involved in binding together and retrieving the items and contexts that make up an event (e.g., associating a person with a specific place or time). The perirhinal cortex is thought to process item information, which it receives from the ventral visual stream (sometimes called the 'what stream'), whereas the parahippocampal cortex is thought to process contextual information such as spatial layout, which it receives from the dorsal processing stream (sometimes called the 'where stream'). It is assumed that an event is encoded by a representation in the hippocampus that links to the various cortical ensembles involved in processing the different aspects of the event. In this way, the presentation of an item (e.g., a person's face) could lead to the reactivation of the hippocampal binding for an earlier experience with that item, which would then lead to the reactivation of the initial encoding context (e.g., the place in which the person was encountered). Conversely, re-presenting a spatial context might lead to the recollection of the individuals who were encountered in that context. Thus recollection can involve all three MTL regions. By contrast, the perirhinal cortex is thought to be sufficient to support familiarity-based recognition. That is, it is presumed to process repeated items more fluently than novel items, and thus it can support the discrimination of familiar and new items in tests of recognition memory. The evidence for the standard model is fairly extensive and has been reviewed elsewhere (see [17,88,89,104-106]). Although there are some differences in the specific focus of each of these different models, there is broad consensus about the basic process distinctions between the hippocampus, perirhinal cortex, and the parahippocampal cortex. However, a shortcoming of all of these models is that they have not incorporated predictions about emotional memories or the role of the amygdala.

bindings that are more slowly forgotten than are the item-context bindings supported by the hippocampus (Figure 2). Consistent with standard models of the MTL, we assume that the hippocampus supports the binding of item and context information, and thus plays a central role in recollection, whereas the perirhinal cortex supports item processing and is therefore capable of supporting familiarity-based recognition. We assume that this occurs for both neutral and emotional events. The important addition, however, is that we propose that the amygdala responds to emotional arousal and supports the binding of item and emotion information, and that these amygdala bindings, once formed, will support recollection and will exhibit relatively slow forgetting. Therefore, for events that elicit an emotional response, the amygdala will bind that emotional response to inputs from the cortical ensembles in the perirhinal cortex that are involved in processing the emotional item or items in the event. In this way, emotional events will be more dependent on the amygdala than will neutral events, and thus if the amygdala exhibits slower forgetting than the hippocampus then emotional materials will be more slowly forgotten.

Why would the emotional bindings supported by the amygdala be forgotten more slowly than the bindings supported by the hippocampus? As we will describe below,

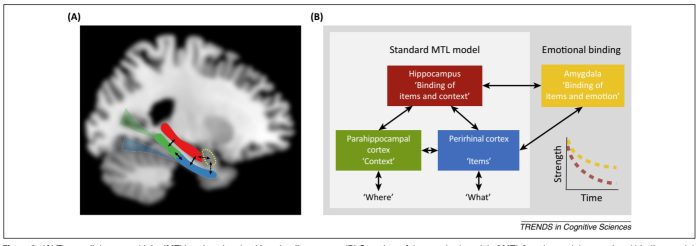


Figure 2. (A) The medial temporal lobe (MTL) regions involved in episodic memory. (B) Overview of the standard model of MTL function and the emotional binding model. In the standard model, the perirhinal cortex receives projections from the ventral 'what' stream and is thought to play a role in identifying and processing the items and objects in the environment. The parahippocampal cortex receives projections from the dorsal 'where' stream, and is thought to play a role in processing contextual information such as the ongoing spatial and temporal context. The hippocampus receives information from the perirhinal and parahippocampal cortex via entorhinal projections, and binds the item and context information together to form episodic memories. The emotional binding model subsumes the standard MTL model, but in addition assumes that the amygdala forms item-emotion bindings that are forgotten more slowly than the item-context bindings supported by the hippocampus.

this assumption is essential in accounting for the full range of empirical effects we have described above. In addition, it is consistent with results from studies of fear conditioning (Box 2), which suggest that item-emotion associations are dependent on the amygdala and are very resistant to forgetting, whereas item-context associations are dependent on the hippocampus and are forgotten more rapidly. There are also theoretical reasons why emotional bindings supported by the amygdala would be forgotten more slowly than the bindings supported by the hippocampus. First, the hippocampus is unique in the sense that it has a high rate of neurogenesis and cell death [73,74]. Relative to regions such as the amygdala that do not exhibit this high level of cell turnover, the hippocampus may be subject to accelerated forgetting due to cell death or increased interference due to new cells [75]. A related account is that the hippocampus may be more

susceptible to active decay processes than other brain regions [76,77]. Such an active decay process may remove selected memories by silencing synapses that are weakly potentiated [78]. To the extent that item—emotion bindings are dependent on the amygdala, rather than on the hippocampus, one may expect slower forgetting for emotional than neutral materials. Another important possibility is that the amygdala may simply be more resistant to interference because there are typically fewer competing emotional experiences outside of the experimental context, relative to neutral experiences.

The emotional binding model naturally accounts for each of the empirical regularities described earlier. Most obviously, the fact that emotional events are forgotten more slowly over time follows directly from the assumption that emotional events depend more on the amygdala, which exhibits slower forgetting than the hippocampus.

Box 2. Fear-conditioning studies and their relation to emotional episodic memory

Fear conditioning refers to the process by which fear responses become associated with an intrinsically neutral stimulus, such as a tone or a context, that predicts the onset of a fear-eliciting stimulus, such as a shock. Although there are several parallels between the emotion effects seen in episodic memory and results seen in fearconditioning studies, there are important differences across these paradigms - suggesting that they reflect at least partially distinct processes. Similarly to emotion effects on episodic memory, conditioned fear responses are amygdala-dependent [107-109] and extremely resistant to forgetting [110,111]. In addition, however, it has been shown that contextual fear memories are affected by post-learning manipulations of glucocorticoids in the dorsal hippocampus (posterior hippocampus in humans) and noradrenaline in the basolateral amygdala (reviewed in [81,112]. Most rodent studies linking the amygdala and hippocampus to arousal-related memory enhancements have relied on contextual fear-conditioning tasks, such as the inhibitory avoidance task, in which the fear response is bound to a context representation [81]. By contrast, in episodic memory paradigms the emotion is bound directly to the item. It may be that item-emotion bindings can be supported by the amygdala alone, whereas contextemotion bindings may additionally require the hippocampus. In fact, it has been shown that contextual fear memories require both the amygdala and hippocampus, whereas other types of fear memories

(e.g., auditorily cued fear) appear to require only the amygdala [107] – although the exact role of the amygdala has been debated [113,114]. Other work has shown that fear responses persist for a long time, but that accompanying contextual information degrades more rapidly [115,116], which is consistent with the notion that the amygdala supports bindings that are particularly resistant to forgetting.

However, the task demands of episodic memory and fear-conditioning paradigms are different. For example, in human episodic memory tasks, differential forgetting effects are observed for emotional and neutral trials that are rapidly interleaved in time (i.e., the study-lists typically contain a mixture of negative and neutral items presented in rapid succession). By contrast, rodent fear-conditioning studies focus on the protracted neurohormonal cascade that is triggered by a single, arousal-inducing event (e.g., a shock) and its influence on memory for that event. It is unlikely that the same hormonal cascade that enhances memory for a single fearful event would also selectively enhance episodic memory for a negative image without also influencing the neutral images that immediately precede or follow it. Future research directly contrasting episodic memory and fear conditioning under conditions in which the materials and task demands are closely matched should help to bridge the gap between these literatures and provide a more complete understanding of emotion and memory.





Second, the finding that the emotion effects are limited to influencing recollection, rather than familiarity-based recognition, arises because the recollection of item-emotion bindings is supported by the amygdala through its interactions with the perirhinal cortex. By contrast, familiarity of both neutral and emotional materials is expected to rely on the perirhinal cortex, and therefore familiarity should show similar forgetting rates for emotional and neutral materials. Third, the binding account also explains the finding that the emotional memory advantages are limited to recollection of item-specific information rather than generalizing to the recollection of contextual information. That is, slower forgetting should be seen for the itememotion bindings, that are expected to be dependent on the amygdala, than for the item-context bindings supported by the hippocampus. Fourth, it accounts for the finding that the emotion advantage is dependent on the amygdala. Finally, according to the emotional binding account, the hippocampus should not be crucial for the increasing advantage for emotional compared to neutral items over time. Because the hippocampus is assumed to support itemcontext associations for both emotional and neutral events, damage to this region is expected to reduce recollection, but it should do so for both emotional and neutral events.

Alternative theoretical accounts

An alternative account that has been used to explain delayed effects of emotion on memory is a modulatory emotional consolidation explanation (e.g., [2,9,13,79–81]). By this account, the amygdala plays a role in identifying and responding to emotional materials, whereas the other MTL regions are assumed to be responsible for supporting episodic memory. Crucially, the amygdala is thought to modulate the MTL after the initial encoding of emotional events to preferentially stabilize memories for these events. The consolidation account can be distinguished from the emotional binding model in two main ways. First, in the consolidation account, the amygdala facilitates long-term storage in other MTL structures but does not play a lasting role in itememotion binding. Second, in the consolidation account, the time-dependent advantage emerges because of processes that unfold after encoding. By contrast, in the emotional binding account, once item-emotion bindings are formed, there is no need to additionally assume further modulation – the slower forgetting rate will ensure that the emotional memory advantage will emerge over time.

The consolidation account can nicely account for the fact that memory for emotional and neutral materials can be comparable immediately after encoding, and that over time the emotional events will exhibit slower forgetting than neutral events. However, it does not specify which types of episodic memory should be influenced by emotion, and it therefore does not explain why the time-dependent emotion benefits are specific to item recollection. To resolve this issue, one might assume that the amygdala modulates some parts of the MTL more than others. If the amygdala were to selectively modulate the hippocampus, we would expect that emotion would benefit recollection of both item and context information, and that hippocampal damage would reduce the time-dependent advantage. The available evidence contradicts both of

these predictions. Alternatively, if the amygdala were to selectively modulate perirhinal representations, we would expect emotion to influence familiarity rather than recollection; however, this is also contradicted by the available evidence. Finally, one might assume that the amygdala modulates only the anterior portion of the hippocampus, which may be specialized for binding item and emotional information, whereas the posterior hippocampus is specialized for binding item and contextual information. However, such an account would predict that only patients with posterior hippocampal damage would exhibit normal emotion effects. By contrast, however, several of the hippocampal patients with normal emotional memory advantages suffered from ischemic events (e.g., [13,65]), which are known to cause volumetric reductions [82-84] and cell death (e.g., [85,86]) along the entire anterior-posterior extent of the hippocampus. We conclude that there do not appear to be any immediately obvious ways of bringing the emotional consolidation model into line with the existing results.

The emotional binding model is broadly consistent with several other theoretical approaches to emotion and memory. Most obviously, the approach extends current MTL models of episodic memory [87-91] by incorporating emotion and the role of the amygdala (Figure 2B). The approach is also broadly consistent with models of emotion that have highlighted the fact that the effects of emotion are not expected to impact upon memory for all aspects of the emotional event in the same way. For example, it has been proposed that emotion leads to increased memory for the central aspects of an emotional event at the cost of poorer memory for the contextual or peripheral aspects of the event [27,28]. According to the arousal biased competition model [92], this difference in memory arises because arousal biases perception and encoding toward 'high priority' information, which typically but not necessarily includes emotional items. The current approach differs by focusing on the time-dependent effects of emotion rather than on the encoding or attention processes that act on emotional materials. We emphasize, however, that although the delayed emotion effects cannot be explained by encoding processes, the emotional binding model does not preclude the possibility that other cognitive factors such as increased attention or more elaborative encoding can lead to an emotion advantage. These encoding effects would of course lead to an emotion advantage even after a brief delay.

Future directions

We believe that the emotional binding model is useful in explaining the slow forgetting of negative arousing materials seen in episodic memory, as well as some of the major functional and neural characteristics of these effects. However, we acknowledge that there are several important questions that the model leaves unanswered. For example, the model focuses on coarse neuroanatomical distinctions, such as between the amygdala and the hippocampus, but there are expected to be differences between the functions of different amygdala nuclei [93] and hippocampal subfields [94]. In addition, emerging evidence suggests that there may be functional differences

Box 3. Outstanding questions

- Can item-emotion associations be unitized? Prior studies have shown that when different features of an item or event are unitized or treated as a single object during encoding, the perirhinal cortex can support familiarity-based recognition for those associations [117,118]. If the current model is correct, then the perirhinal cortex may support unitized item-emotion associations under encoding conditions that promote unitization. In such cases, one expects that the emotion advantage should be observed for familiarity-based recognition, and, unlike the recollection advantage, it should not increase across delay.
- How does damage to the perirhinal cortex and parahippocampal cortex impact on the delayed emotion effects? The current model predicts that perirhinal damage, but not parahippocampal damage, should reduce the emergence of the emotional advantage in memory.
 Selective lesions of this type are rare in humans, but volumetric studies in human patients or analogous emotional recognition paradigms in rodents may be useful for testing these predictions.
- Are there conditions in which emotion enhances memory for context information, and are those effects time-dependent? The current model predicts that context bindings supported by the hippocampus should be forgotten more quickly than the itememotion bindings supported by the amygdala. However, it is an

- open question whether there are other types of context associations (e.g., internal context) that can be supported by a non-hippocampal pathway, such as the medial prefrontal cortex.
- What hormonal mechanisms are involved in producing the emotion effects on episodic memory, and are they specific to particular neuroanatomical regions of the medial temporal lobes?
- Do the item-emotion bindings supported by the amygdala also support memory for emotionally positive materials such as images of babies or smiling faces? Imaging studies have shown memory-related amygdala activity for both positive and negative arousing materials [119,120]. To our knowledge, however, no studies have examined emotional memory for positive materials in either patients with selective hippocampal or selective amygdala lesions, nor is it yet clear whether positive emotion effects show the same delay-dependent effects seen with negative emotion.
- How do these laboratory results relate to memory for traumatic real-life experiences? In the existing lab studies, emotion has been shown to benefit recollection rather than familiarity, but more traumatic experiences may impact on both processes. For example, people may be more likely to be reminded of real-life traumatic experiences and thus re-encode those items, which would be expected to slow the forgetting of both recollection and familiarity.

within the hippocampus along the long axis of this structure [95,96]. High-resolution fMRI may prove useful in further differentiating the functions of these subregions and in assessing whether they differentially contribute to episodic memory for emotional materials.

Another important question concerns how the emerging effects of emotion on episodic memory are related to postencoding manipulations such as stress. For example, a growing body of research indicates that when encoding is immediately followed by a stressful event this can lead to slower forgetting compared to a no-stress control condition [22,26,97–99]. Post-encoding stress effects have often been attributed to a consolidation process similar to the one originally thought to produce the delayed emotion effects. However, some findings suggest that these effects are distinct. For example, the emotion effects do not seem to require stress to emerge because they can be observed even after a 2 h no-stress delay [26]. In addition, stress can impact memory for both emotional and neutral materials, and it therefore does not seem to preferentially slow the forgetting of emotional materials [22,26,97] (but see [98,99]).

We have argued that although the MTL regions might interact during encoding, to account for the emerging emotion advantage in memory, there is no need to assume that there are any additional interactions between the amygdala and other MTL regions across the delay period. However, it is, of course, possible that under particular conditions there may be additional interactions. For example, if subjects preferentially rehearse or are reminded of the emotional events, this could lead to additional encodings of those items, slowing forgetting even further. However, in this case one would expect to see both recollection and familiarity advantages emerge over time because both the items and the item-emotion bindings are repeated. The existing literature shows that the time-dependent emotion advantage is limited to recollection, arguing against this type of reminding. We suspect, however, that with truly traumatic real-life events, this type of reminding may occur more often. Future work could test whether

traumatic events are characterized by increased reminding and whether this is related to the emergence of recollection and familiarity based emotional advantages. Outstanding questions are listed in ${\tt Box}\ 3$.

Concluding remarks

The current literature indicates that emotional memories are forgotten more slowly than neutral memories. Moreover, these effects are specific to recollection of item-related information rather than to familiarity or contextual recollection, and they are dependent on the amygdala but not the hippocampus. The results are consistent with an emotional binding model in which item-emotion bindings supported by the amygdala are particularly resistant to forgetting compared to the item-context bindings supported by the hippocampus. Although the emotional binding account appears to explain the emerging emotional memory effects, there are several questions about these effects that have not yet been fully addressed. We hope that future studies designed to test this approach will prove useful in furthering our understanding of the impact of emotion on episodic memory.

References

- 1 Hamann, S. (2001) Cognitive and neural mechanisms of emotional memory. *Trends Cogn. Sci.* 5, 394–400
- 2 LaBar, K.S. and Cabeza, R. (2006) Cognitive neuroscience of emotional memory. Nat. Rev. Neurosci. 7, 54–64
- 3 Phelps, E.A. (2004) Human emotion and memory: interactions of the amygdala and hippocampal complex. Curr. Opin. Neurobiol. 14, 198–202
- 4 Talmi, D. (2013) Enhanced emotional memory: cognitive and neural mechanisms. *Curr. Dir. Psychol. Sci.* 22, 430–436
- 5 Kleinsmith, L.J. and Kaplan, S. (1963) Paired-associate learning as a function of arousal and interpolated interval. J. Exp. Psychol. 65, 190–193
- 6 Sharot, T. and Phelps, E.A. (2004) How arousal modulates memory: disentangling the effects of attention and retention. Cogn. Affect. Behav. Neurosci. 4, 294–306
- 7 Anderson, A.K. et al. (2006) Emotional memories are not all created equal: evidence for selective memory enhancement. Learn. Mem. 13, 711–718

- 8 Sharot, T. and Yonelinas, A.P. (2008) Differential time-dependent effects of emotion on recollective experience and memory for contextual information. *Cognition* 106, 538–547
- 9 Ritchey, M. et al. (2008) Role of amygdala connectivity in the persistence of emotional memories over time: an event-related fMRI investigation. Cereb. Cortex 18, 2494–2504
- 10 Sharot, T. et al. (2004) How emotion enhances the feeling of remembering. Nat. Neurosci. 7, 1376–1380
- 11 LaBar, K.S. and Phelps, E.A. (1998) Arousal-mediated memory consolidation: role of the medial temporal lobe in humans. *Psychol. Sci.* 9, 490–493
- 12 Hamann, S.B. et al. (1999) Amygdala activity related to enhanced memory for pleasant and aversive stimuli. Nat. Neurosci. 2, 289–293
- 13 Sharot, T. et al. (2007) How emotion strengthens the recollective experience: a time-dependent hippocampal process. PLoS ONE 2, e1068
- 14 Talmi, D. and McGarry, L.M. (2012) Accounting for immediate emotional memory enhancement. J. Mem. Lang. 66, 93–108
- 15 Talmi, D. et al. (2007) The role of attention and relatedness in emotionally enhanced memory. Emotion 7, 89–102
- 16 Jacoby, L.L. (1991) A process dissociation framework: separating automatic from intentional uses of memory. J. Mem. Lang. 30, 513–541
- 17 Yonelinas, A.P. (2002) The nature of recollection and familiarity: a review of 30 years of research. J. Mem. Lang. 46, 441–517
- 18 Mandler, G. (1980) Recognizing: the judgment of previous occurrence. Psychol. Rev. 87, 252–271
- 19 Kensinger, E.A. and Corkin, S. (2003) Memory enhancement for emotional words: Are emotional words more vividly remembered than neutral words? *Mem. Cogn.* 31, 1169–1180
- 20 Dewhurst, S.A. and Parry, L.A. (2000) Emotionality, distinctiveness, and recollective experience. Eur. J. Cogn. Psychol. 12, 541–551
- 21 Atienza, M. and Cantero, J.L. (2008) Modulatory effects of emotion and sleep on recollection and familiarity. J. Sleep Res. 17, 285–294
- 22 Yonelinas, A.P. *et al.* (2011) The effects of post-encoding stress on recognition memory: examining the impact of skydiving in young men and women. *Stress* 14, 136–144
- 23 Dolcos, F. et al. (2005) Remembering one year later: role of the amygdala and the medial temporal lobe memory system in retrieving emotional memories. Proc. Natl. Acad. Sci. U.S.A. 102, 2626–2631
- 24 Ochsner, K.N. (2000) Are affective events richly recollected or simply familiar? The experience and process of recognizing feelings past. J. Exp. Psychol. Gen. 129, 242–261
- 25 Pierce, B.H. and Kensinger, E.A. (2011) Effects of emotion on associative recognition: valence and retention interval matter. $Emotion\ 11,\ 139-144$
- 26 McCullough, A.M. and Yonelinas, A.P. (2013) Cold-pressor stress after learning enhances familiarity-based recognition memory in men. Neurobiol. Learn. Mem. 106, 11–17
- 27 Kensinger, E.A. (2007) Negative emotion enhances memory accuracy: behavioral and neuroimaging evidence. Curr. Dir. Psychol. Sci. 16, 213–218
- 28 Mather, M. (2007) Emotional arousal and memory binding: an object-based framework. Perspect. Psychol. Sci. 2, 33–52
- 29 Mather, M. and Nesmith, K. (2008) Arousal-enhanced location memory for pictures. J. Mem. Lang. 58, 449–464
- 30 MacKay, D. et al. (2004) Relations between emotion, memory, and attention: evidence from taboo stroop, lexical decision, and immediate memory tasks. Mem. Cognit. 32, 474–488
- 31 MacKay, D.G. and Ahmetzanov, M.V. (2005) Emotion, memory, and attention in the taboo Stroop paradigm. Psychol. Sci. 16, 25–32
- 32 D'Argembeau, A. and Van der Linden, M. (2004) Influence of affective meaning on memory for contextual information. *Emotion* 4, 173–188
- 33 Doerksen, S. and Shimamura, A. (2001) Source memory enhancement for emotional words. $Emotion\ 1,\ 5-11$
- 34 Dougal, S. et al. (2007) The role of medial temporal lobe in item recognition and source recollection of emotional stimuli. Cogn. Affect. Behav. Neurosci. 7, 233–242
- 35 Kensinger, E.A. et al. (2006) Memory for specific visual details can be enhanced by negative arousing content. J. Mem. Lang. 54, 99–112
- 36 Kensinger, E.A. et al. (2007) How negative emotion enhances the visual specificity of a memory. J. Cogn. Neurosci. 19, 1872–1887

- 37 Mather, M. and Knight, M. (2008) The emotional harbinger effect: poor context memory for cues that previously predicted something arousing. *Emotion* 8, 850–860
- 38 Kensinger, E.A. et al. (2007) Effects of emotion on memory specificity: memory trade-offs elicited by negative visually arousing stimuli. J. Mem. Lang. 56, 575–591
- 39 Kensinger, E.A. and Schacter, D.L. (2006) Amygdala activity is associated with the successful encoding of item, but not source, information for positive and negative stimuli. J. Neurosci. 26, 2564–2570
- 40 Yegiyan, N.S. and Yonelinas, A.P. (2011) Encoding details: positive emotion leads to memory broadening. PCEM 25, 1255–1262
- 41 Bergmann, H.C. et al. (2012) The effects of valence and arousal on associative working memory and long-term memory. PLoS ONE 7, e52616
- 42 Cook, G.I. et al. (2007) Source monitoring is not always enhanced for valenced material. Mem. Cognit. 35, 222–230
- 43 Rimmele, U. et al. (2011) Emotion enhances the subjective feeling of remembering, despite lower accuracy for contextual details. Emotion 11, 553–562
- 44 Touryan, S. et al. (2007) Effect of negative emotional pictures on associative memory for peripheral information. Memory 15, 154–166
- 45 Burke, A. et al. (1992) Remembering emotional events. Mem. Cognit. 20, 277–290
- 46 Waring, J.D. and Kensinger, E.A. (2009) Effects of emotional valence and arousal upon memory trade-offs with aging. *Psychol. Aging* 24, 412–422
- 47 Adolphs, R. et al. (1997) Impaired declarative memory for emotional material following bilateral amygdala damage in humans. Learn. Mem. 4, 291–300
- 48 Cahill, L. et al. (1995) The amygdala and emotional memory. Nature 377, 295–296
- 49 Markowitsch, H.J. et al. (1994) The amygdala's contribution to memory – a study on two patients with Urbach–Wiethe disease. Neuroreport 5, 1349–1352
- 50 Phelps, E.A. et al. (1998) Specifying the contributions of the human amygdala to emotional memory: a case study. Neurocase 4, 527–540
- 51 Cahill, L. et al. (1996) Amygdala activity at encoding correlated with long-term, free recall of emotional information. Proc. Natl. Acad. Sci. U.S.A. 93, 8016–8021
- 52 Dolcos, F. et al. (2004) Interaction between the amygdala and the medial temporal lobe memory system predicts better memory for emotional events. Neuron 42, 855–863
- 53 Canli, T. et al. (2000) Event-related activation in the human amygdala associates with later memory for individual emotional experience. J. Neurosci. 20, 5
- 54 Murty, V. et al. (2010) fMRI studies of successful emotional memory encoding: A quantitative meta-analysis. Neuropsychologia 48, 3459– 3469
- 55 Smith, A.P. et al. (2006) Task and content modulate amygdalahippocampal connectivity in emotional retrieval. Neuron 49, 631–638
- 56 Smith, A. et al. (2004) fMRI correlates of the episodic retrieval of emotional contexts. Neuroimage 22, 868–878
- 57 Maratos, E.J. et al. (2001) Neural activity associated with episodic memory for emotional context. Neuropsychologia 39, 910–920
- 58 Somerville, L.H. et al. (2006) Dissociable medial temporal lobe contributions to social memory. J. Cogn. Neurosci. 18, 1253–1265
- 59 Kensinger, E.A. et al. (2011) Amygdala activity at encoding corresponds with memory vividness and with memory for select episodic details. Neuropsychologia 49, 663–673
- 60 Waring, J.D. and Kensinger, E.A. (2011) How emotion leads to selective memory: neuroimaging evidence. *Neuropsychologia* 49, 1831–1842
- 61 Mackiewicz, K.L. et al. (2006) The effect of anticipation and the specificity of sex differences for amygdala and hippocampus function in emotional memory. Proc. Natl. Acad. Sci. U.S.A. 103, 14200–14205
- 62 Mickley Steinmetz, K.R. et al. (2012) The effect of emotional arousal and retention delay on subsequent-memory effects. Cogn. Neurosci. 3, 150–159
- 63 Adolphs, R. et al. (2000) Impaired emotional declarative memory following unilateral amygdala damage. Learn. Mem. 7, 180–186

- 64 Todorov, A. and Olson, I.R. (2008) Robust learning of affective trait associations with faces when the hippocampus is damaged, but not when the amygdala and temporal pole are damaged. Soc. Cogn. Affect. Neurosci. 3, 195–203
- 65 Hamann, S.B. et al. (1997) Intact enhancement of declarative memory for emotional material in amnesia. Learn. Mem. 4, 301–309
- 66 Hamann, S.B. et al. (1997) Emotional perception and memory in amnesia. Neuropsychology 11, 104–113
- 67 Richardson, M.P. et al. (2004) Encoding of emotional memories depends on amygdala and hippocampus and their interactions. Nat. Neurosci. 7, 278–285
- 68 Kensinger, E.A. and Corkin, S. (2004) Two routes to emotional memory: distinct neural processes for valence and arousal. Proc. Natl. Acad. Sci. U.S.A. 101, 3310–3315
- 69 St. Jacques, P.L. et al. (2009) Effects of aging on functional connectivity of the amygdala for subsequent memory of negative pictures. Psychol. Sci. 20, 74–84
- 70 Kilpatrick, L. and Cahill, L. (2003) Amygdala modulation of parahippocampal and frontal regions during emotionally influenced memory storage. *Neuroimage* 20, 2091–2099
- 71 Amaral, D.G. and Price, J.L. (1984) Amygdalo-cortical projections in the monkey (*Macaca fascicularis*). J. Comp. Neurol. 230, 465–496
- 72 Price, J. (2003) Comparative aspects of amygdala connectivity. Ann. N. Y. Acad. Sci. 985, 50–58
- 73 Biebl, M. et al. (2000) Analysis of neurogenesis and programmed cell death reveals a self-renewing capacity in the adult rat brain. Neurosci. Lett. 291, 17–20
- 74 Eriksson, P.S. et al. (1998) Neurogenesis in the adult human hippocampus. Nat. Med. 4, 1313–1317
- 75 Akers, K.G. et al. (2014) Hippocampal neurogenesis regulates forgetting during adulthood and infancy. Science 344, 598–602
- 76 Hardt, O. et al. (2013) Decay happens: the role of active forgetting in memory. Trends Cogn. Sci. 17, 111–120
- 77 Sadeh, T. et al. (2014) How we forget may depend on how we remember. Trends Cogn. Sci. 18, 26–36
- 78 Tononi, G. and Cirelli, C. (2003) Sleep and synaptic homeostasis: a
- hypothesis. Brain Res. Bull. 62, 143–150 79 Cahill, L. and McGaugh, J.L. (1998) Mechanisms of emotional arousal
- and lasting declarative memory. *Trends Neurosci.* 21, 294–299 80 McGaugh, J.L. (2000) Memory – a century of consolidation. *Science* 287, 248–251
- 81 McGaugh, J.L. (2004) The amygdala modulates the consolidation of memories of emotionally arousing experiences. *Annu. Rev. Neurosci.*
- 27, 1–2882 Olsen, R.K. *et al.* (2013) Volumetric analysis of medial temporal lobe subregions in developmental amnesia using high-resolution magnetic
- resonance imaging. *Hippocampus* 23, 855–860
 83 Vargha-Khadem, F. et al. (1997) Differential effects of early hippocampal pathology on episodic and semantic memory. *Science*
- 277, 376–380

 84 Vargha-Khadem, F. et al. (2003) Developmental amnesia: effect of age
- at injury. Proc. Natl. Acad. Sci. U.S.A. 100, 10055–10060 85 Rempel-Clower, N.L. et al. (1996) Three cases of enduring memory
- so Rempel-Clower, N.L. et al. (1996) Three cases of enduring memory impairment after bilateral damage limited to the hippocampal formation. J. Neurosci. 16, 5233–5255
- 86 Zola-Morgan, S. et al. (1986) Human amnesia and the medial temporal region: enduring memory impairment following a bilateral lesion limited to field CA1 of the hippocampus. J. Neurosci. 6, 2950–2967
- 87 Davachi, L. (2006) Item, context and relational episodic encoding in humans. Curr. Opin. Neurobiol. 16, 693–700
- 88 Diana, R.A. et al. (2007) Imaging recollection and familiarity in the medial temporal lobe: a three-component model. Trends Cogn. Sci. 11, 379–386
- 89 Eichenbaum, H. et al. (2007) The medial temporal lobe and recognition memory. Annu. Rev. Neurosci. 30, 123–152
- 90 Mayes, A. et al. (2007) Associative memory and the medial temporal lobes. Trends Cogn. Sci. 11, 126–135
- 91 Norman, K.A. and O'Reilly, R.C. (2003) Modeling hippocampal and neocortical contributions to recognition memory: a complementary-learning-systems approach. *Psychol. Rev.* 110, 611–646

- 92 Mather, M. and Sutherland, M.R. (2011) Arousal-biased competition in perception and memory. Perspect. Psychol. Sci. 6, 114–133
- 93 Sah, P. et al. (2003) The amygdaloid complex: anatomy and physiology. Physiol. Rev. 83, 803–834
- 94 Kesner, R.P. (2013) Role of the hippocampus in mediating interference as measured by pattern separation processes. Behav. Processes 93, 148–154
- 95 Poppenk, J. et al. (2013) Long-axis specialization of the human hippocampus. Trends Cogn. Sci. 17, 230–240
- 96 Strange, B.A. et al. (2014) Functional organization of the hippocampal longitudinal axis. Nat. Rev. Neurosci. 15, 655–669
- 97 Andreano, J.M. and Cahill, L. (2006) Glucocorticoid release and memory consolidation in men and women. *Psychol. Sci.* 17, 466–470
- 98 Cahill, L. et al. (2003) Enhanced human memory consolidation with post-learning stress: interaction with the degree of arousal at encoding. Learn. Mem. 10, 270–274
- 99 Smeets, T. et al. (2008) True or false? Memory is differentially affected by stress-induced cortisol elevations and sympathetic activity at consolidation and retrieval. Psychoneuroendocrinology 33, 1378–1386
- 100 Tulving, E. (1982) Synergistic ecphory in recall and recognition. Can. J. Psychol. 36, 130–147
- 101 Yonelinas, A.P. (1994) Receiver-operating characteristics in recognition memory: evidence for a dual-process model. J. Exp. Psychol. Learn. Mem. Cogn. 20, 1341–1354
- 102 Tulving, E. (1984) Précis of elements of episodic memory. Behav. Brain Sci. 7, 223–238
- 103 Yonelinas, A.P. and Jacoby, L.L. (1995) Dissociating automatic and controlled processes in a memory-search task: beyond implicit memory. Psychol. Res. 57, 156–165
- 104 Yonelinas, A.P. et al. (2010) Recollection and familiarity: examining controversial assumptions and new directions. Hippocampus 20, 1178–1194
- 105 Aggleton, J.P. and Brown, M.W. (1999) Episodic memory, amnesia, and the hippocampal–anterior thalamic axis. *Behav. Brain Sci.* 22, 425–489
- 106 Rugg, M.D. and Yonelinas, A.P. (2003) Human recognition memory: a cognitive neuroscience perspective. Trends Cogn. Sci. 7, 313–319
- 107 Phillips, R.G. and LeDoux, J.E. (1992) Differential contribution of amygdala and hippocampus to cued and contextual fear conditioning. Behav. Neurosci. 106, 274–285
- 108 Bechara, A. et al. (1995) Double dissociation of conditioning and declarative knowledge relative to the amygdala and hippocampus in humans. Science 269, 1115–1118
- 109 LaBar, K.S. et al. (1995) Impaired fear conditioning following unilateral temporal lobectomy in humans. J. Neurosci. 15, 6846–6855
- 110 Fanselow, M.S. (1990) Factors governing one-trial contextual conditioning. *Anim. Learn. Behav.* 18, 264–270
- 111 Gale, G.D. et al. (2004) Role of the basolateral amygdala in the storage of fear memories across the adult lifetime of rats. J. Neurosci. 24, 3810–3815
- 112 McGaugh, J.L. and Roozendaal, B. (2002) Role of adrenal stress hormones in forming lasting memories in the brain. Curr. Opin. Neurobiol. 12, 205–210
- 113 Cahill, L. et al. (1999) Is the amygdala a locus of conditioned fear? Some questions and caveats. Neuron 23, 227–228
- 114 Fanselow, M.S. and LeDoux, J.E. (1999) Why we think plasticity underlying Pavlovian fear conditioning occurs in the basolateral amygdala. Neuron 23, 229–232
- 115 Wiltgen, B.J. and Silva, A.J. (2007) Memory for context becomes less specific with time. Learn. Mem. 14, 313–317
- 116 Winocur, G. et al. (2007) Memory consolidation or transformation: context manipulation and hippocampal representations of memory. Nat. Neurosci. 10, 555–557
- 117 Haskins, A.L. et al. (2008) Perirhinal cortex supports encoding and familiarity-based recognition of novel associations. Neuron 59, 554– 560
- 118 Quamme, J.R. et al. (2007) Effect of unitization on associative recognition in amnesia. Hippocampus 17, 192–200
- 119 Hamann, S.B. et al. (2002) Ecstasy and agony: activation of the human amygdala in positive and negative emotion. Psychol. Sci. 13, 135–141
- 120 Anderson, A.K. et al. (2003) Dissociated neural representations of intensity and valence in human olfaction. Nat. Neurosci. 6, 196–202