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Lucas S. Broster
University of Kentucky, lucas.broster@uky.edu

Lee X. Blonder
University of Kentucky, lxblond@uky.edu

Yang Jiang
University of Kentucky, yang.jiang@uky.edu

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Does emotional memory enhancement assist the memory-impaired?

Lucas S. Broster1, Lee X. Blonder1,2,3 and Yang Jiang1,2,4*

1Department of Behavioral Science, University of Kentucky College of Medicine, Lexington, KY, USA
2 Sanders-Brown Center on Aging, University of Kentucky College of Medicine, Lexington, KY, USA
3 Department of Neurology, University of Kentucky College of Medicine, Lexington, KY, USA
4Magnetic Resonance Imaging and Spectroscopy Center, Lexington, KY, USA

*Correspondence:
yang.jiang@uky.edu

We review recent work on emotional memory enhancement in older adults and patients with mild cognitive impairment (MCI) or Alzheimer dementia (AD) and evaluate the viability of incorporating emotional components into cognitive rehabilitation for these groups. First, we identify converging evidence regarding the effects of emotional valence on working memory in healthy aging. Second, we introduce work that suggests a more complex role for emotional memory enhancement in aging and identify a model capable of unifying disparate research findings. Third, we survey the neuroimaging literature for evidence of a special role for the amygdala in MCI and early AD in emotional memory enhancement. Finally, we assess the theoretical feasibility of incorporating emotional content into cognitive rehabilitation given all available evidence.

Keywords: aging, Alzheimer disease, mild cognitive impairment, cognitive rehabilitation, amygdala, emotional memory enhancement, executive function, working memory

INTRODUCTION

Contemporary cognitive rehabilitation, especially when delivered alongside pharmacotherapy, has been shown to improve memory outcomes and delay dementia progression in patients with mild cognitive impairment (MCI) and early Alzheimer dementia (AD) (Mimura and Komatsu, 2007). Historically, however, clinical results of cognitive dementia interventions have been somewhat mixed. In fact, the findings of trials investigating the efficacy of such practices in the 1980s led clinicians to question the validity of the entire behavioral intervention paradigm for the treatment of AD (Hopper, 2003; Mowszowski et al., 2010).

These early cognitive rehabilitation interventions focused on improving explicit memory as a means of treating the explicit memory loss typical of AD. Other forms of memory, including implicit memory, are relatively spared by AD. Implicit memory includes those aspects of memory of which the individual lacks conscious awareness. For example, procedural memory, the memory of how to perform physical tasks, is one aspect of implicit memory. Repetition learning, the passive, automatic learning of associations upon repeated exposure, is another (Kessels et al., 2011).

The advent of implicit memory-based methods represented a major theoretical shift in the development of more efficacious cognitive rehabilitation protocols. Rather than focusing only on improving explicit memory directly through rehearsal, these protocols also utilize intact implicit memory to compensate for eroding explicit capacity (Mimura and Komatsu, 2007). Pilot studies have suggested improved efficacy for implicit memory-based intervention methods (Zanetti et al., 1997; Kessels and de Haan, 2003; van Halteren-van Tilborg et al., 2007; Jean et al., 2010a,b).

The success of this approach leads us to question whether other neurocognitive systems broadly implicated in memory are relatively intact in patients with progressive dementia such that those systems may be analogously harnessed to help compensate for eroded explicit capacity. In other words, much as implicit memory treatment appears promising for ameliorating early explicit memory decline, perhaps recruiting other relatively intact memory processes in individuals with early impairment can additively help delay significant functional impairment. One candidate system that involves structures relatively spared in AD is the emotional processing system, and its integration with cognitive memory systems has garnered significant attention over the past decades (for reviews, refer to Pessoa, 2008, 2010). To evaluate whether this system might show promise in further enhancing cognitive rehabilitation protocols, we first describe how emotional memory enhancement, the tendency of emotional content to be better-remembered than non-emotional content, might change with aging and dementia.

EMOTIONAL MEMORY ENHANCEMENT IS INTACT IN OLDER ADULTS

A major behavioral basis for the belief that emotional content might improve memory outcomes is the persistent finding that, even in older adults, emotional content is remembered more accurately and/or more quickly than non-emotional content for both short and longer-term recall of visual images (Moayeri et al., 2000; Boller et al., 2002; Hiroaki, 2002; Mikels et al., 2005;
Redondo and Fernández-Rey, 2010; Borg et al., 2011; Evans-Roberts and Turnbull, 2011; Nashiro et al., 2011). This is independently demonstrated for emotional stimuli of both positive valence and negative valence relative to non-emotional control, and it is demonstrated in research participants with early AD (Boller et al., 2002; Borg et al., 2011).

In tests simultaneously evaluating positively and negatively valenced stimuli relative to non-emotional control, both types consistently increase memory performance better than non-emotional stimuli, and, for older adults, positively valenced stimuli tend to produce better results than negatively valenced stimuli (Nashiro et al., 2011; for a meta-analysis of positivity effects in younger and older adults, see Murphy and Isaacowitz, 2008). The tendency for older adults to perform better on positively valenced stimuli with regard to measures of attention, recognition, and emotional memory enhancement has been termed the age-related positivity effect (Charles et al., 2003; Mather and Knight, 2006; Löckenhoff and Carstensen, 2007; Grühn et al., 2007; Isaacowitz et al., 2009). The positivity effect phenomenon itself is sufficiently well established that separate attempts to model the phenomenon compete for legitimacy in the emotional aging literature (for separate accounts refer to Carstensen et al., 1999; Rozin and Royzman, 2001; Labouvie-Vief, 2003; Mather and Knight, 2005; Knight et al., 2007; or to Labouvie-Vief et al., 2007; Scheibe and Carstensen, 2010).

One account of the positivity effect holds that changes in valence biases reflect top-down changes in emotional regulation with aging (Carstensen et al., 1999). In other words, it appears that older adults may have unique mental processes that enable them to encode positively valenced stimuli in a manner categorically different from younger adults (Charles and Carstensen, 2007). This perspective suggests that older adults may have special valence-specific cognitive biases that may be used in the context of emotional memory enhancement to improve memory outcomes.

This positivity effect may not be observed in some subpopulations of older adults. For example, individuals with major depressive disorder or who have recovered from major depressive disorder show preferential attention to negatively valenced stimuli, which may predispose them to encode valence differently from individuals without a history of depression (Kerestes et al., 2012). Whether these effects persist in the face of the positivity effect is unknown, but, given the association between depression and cognitive impairment, the competing trends should be noted in the short-term and formally evaluated in future work (for a meta-analysis, see Burt et al., 1995; for a full review of the interaction between these conditions, see Korczyn and Halperin, 2009).

For the purposes of this review, the observations that older adults show emotional memory enhancement, particularly with positively valenced stimuli, and that research participants with AD may also show a degree of emotional memory enhancement are sufficient for motivating further inquiry. We next look at how emotional memory enhancement differs in old age and dementia.

**Dementia Changes the Emotional Realm**

While emotional memory enhancement, as prior discussed, is intact to a certain extent in older adults, individuals with MCI, and individuals with AD, all of these groups do significantly differ from younger adults on certain attention and memory measures (reviewed by Kensinger, 2009). The prior-discussed positivity effect between younger and older adults is just one example of such a difference, and the themes implicit in that effect are broadly consistent even for non-visual modalities, attention paradigms, and neurofunctional work (Leclerc and Kensinger, 2011; Orgeta, 2011; Yang and Hasher, 2011). However, differences in emotional memory enhancement between non-demented older adults and older adults with AD have been reported for a range of experimental paradigms, and some even report lack of significant emotional memory enhancement. For example, older adults with AD do not appear to benefit from emotional memory enhancement in the processing of complex information such as verbally related stories and even some short-term memory delay recognition memory paradigms (Abrisqueta-Gomez et al., 2002; Kensinger et al., 2003). These differences in patients with AD are attested when investigating both only positively and only negatively valenced stimuli (Hamann et al., 2002; Padovan et al., 2002).

Contrarily, other work indicates that even individuals with AD benefit modestly from emotional memory enhancement (Kensinger et al., 2002). For example, while emotional memory enhancement was attenuated relative to healthy, age-matched controls, patients with early AD showed improved memory for emotional items in a free recall paradigm (Nieuwenhuis-Mark et al., 2009). Indeed, some studies imply that people with AD may show enhanced flashbulb memory (i.e., vivid, visuospatially detailed memories) for events with particularly intense arousal and emotional content (Kazui et al., 2003). Other controlled work on flashbulb memories suggests that these faculties are intact in AD (Ikeda et al., 1998; Kazui et al., 2000; Moayeri et al., 2000; Budson et al., 2004). The strength of this phenomenon within individuals may be linked to amygdala mass (Mori et al., 1999; reviewed by McGaugh et al., 1996; Phelps, 2004). The work reviewed earlier in the context of valence effect differences similarly finds intact emotional memory enhancement of some form in people with AD (Boller et al., 2002).

These data demonstrate that different paradigms come to different conclusions about whether people with AD benefit from emotional memory enhancement or the positivity effect, though studies suggest these phenomena may be partially intact. To attempt to contextualize these disparate findings, we investigate a few studies in more detail.

**Can Emotional Memory Decrement Occur?**

While most studies indicate emotional memory enhancement in older adults and some further observe a form of this in people with AD, we have also cited work that fails to observe this phenomenon in the demented population. Borg et al. (2011) and colleagues have elucidated a model suggesting that emotional content inexorably monopolizes a certain share of executive function resources (Figure 1) based on the finding that memory deficits were greater for emotional than non-emotional stimuli when a task required more executive demand or when individuals had impaired executive resources (e.g., secondary to dementia).
In a simple visual recognition task, younger adults, older adults, and patients with early AD all showed emotional memory enhancement. However, older adults and patients with early AD no longer showed emotional memory enhancement in a more executive function-demanding visual binding task, and patients with early AD also showed a between-task performance difference for non-emotional stimuli. Borg et al. (2011) and colleagues also showed that emotional memory enhancement in older adults and patients with early AD no longer showed emotional memory enhancement in a more executive function-demanding visual binding task, and patients with early AD, would be hypothesized to have relatively normal emotional memory enhancement.

This contextualization helps explain the disparate findings prior reviewed. Individual differences in the degree of executive dysfunction between cohorts and differences in executive function required for a given task would affect whether emotional memory enhancement, emotional memory decrement, or neither would occur in people with AD. These findings also contextualize research reports outside the dementia literature that appear at-odds with consensus elsewhere. For example, the finding that older adults more than younger adults fail to remember details of complex emotional scenes such as robberies (Aizpurua et al., 2011) may be explained in part by such stimuli having more complex, less controlled elements than the simpler stimuli normatively employed in the psychological literature.

This model greatly resembles the “dual competition” framework articulated by Pessoa (2009). That framework, based on work with younger cohorts, suggested, namely, that interactions between cognition, motivation, and emotion could affect whether emotional memory enhancement or emotional memory decrement would occur.

It is worth noting that this model does not account for the finding that patients with AD appear to have intact flashbulb memory. Whereas the model might predict that memory of complex events requiring concurrent executive function may suffer given emotional context, flashbulb memory, tending as it does to be associated with emotionally arousing events, shows the opposite trend. One possibility is that recollection per se is what these flashbulb memory paradigms have tested, and that recollection itself does not require extensive executive resources, even though processing the event itself may well have required them. Another possibility is that the patients with AD tested in these protocols have confabulated the details of the various events in ways not accounted for by the various protocols and that this confabulation masks impairment not otherwise recorded.

Concerns about the specific applicability of flashbulb memory findings aside, this model suggests that while integrating emotional content into cognitive rehabilitation for people with MCI or very early AD may be efficacious, it may be less useful for more-impaired populations. Emotional memory decrement may even occur if the rehabilitation protocol in question requires intensive executive control.

We next look to neuroimaging for evidence of the neurological mechanism underlying Borg et al. (2011) and colleagues’ executive resource hypothesis.

TOWARD A NEURAL MECHANISM FOR BORG ET AL. (2011) AND COLLEAGUES’ EXECUTIVE RESOURCE HYPOTHESIS

Neuroimaging of emotional memory enhancement in aging and AD populations remains in its infancy. Most imaging work with patients with AD focuses on memory effects per se in the absence of emotional context. Nevertheless, these memory findings have revealed intriguing effects in emotional brain regions. Patients with AD show abnormal connectivity and activation of emotion-related brain regions even when engaging in putatively non-emotional memory tasks (Rosenbaum et al., 2010). Specifically, increased connectivity between amygdala and pre-frontal cortex for a working memory task was seen in participants with AD relative to controls in spite of no differential indication of arousal correlates during the task. Rosenbaum et al. (2010) and colleagues suggest that this difference represents recruitment of compensatory pre-frontal brain structures and that the amygdala acts as an intermediary through which pre-frontal resources are accessed. During an emotionally neutral delayed match-to-sample task, we found significantly more task-related amygdala activity. Whereas the model might predict that memory of complex events requiring concurrent executive function may suffer given emotional context, flashbulb memory, tending as it does to be associated with emotionally arousing events, shows the opposite trend. One possibility is that recollection per se is what these flashbulb memory paradigms have tested, and that recollection itself does not require extensive executive resources, even though processing the event itself may well have required them. Another possibility is that the patients with AD tested in these protocols have confabulated the details of the various events in ways not accounted for by the various protocols and that this confabulation masks impairment not otherwise recorded.

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activation in participants with MCI than in age-matched older adults (Figure 2) (Broster et al., 2012). Our findings are analogous to those reported by Rosenbaum et al. (2010).

One possible criticism of both these neuroimaging findings is that memory-impaired cohorts may experience anxiety when asked to perform a task at which they are impaired. In other words, the amygdala and frontal responses in these groups could reflect functionally irrelevant anxiety about performance. To address this concern, Rosenbaum et al. (2010) and colleagues point to a lack of arousal difference between groups and argue that similar arousal indicates that anxiety was not responsible for the observed effects.

These findings, if valid, provide a possible neurological mechanism for emotional memory enhancement. They also provide a mechanism for Borg et al. (2011) and colleagues’ claim that executive function resources, implicated as they are in pre-frontal cortical structures, determine whether emotional memory enhancement or emotional memory decrement will occur. Specifically, the pre-occupation of pre-frontal executive function resources with an executive function-intensive task could disrupt pre-frontal recruitment and explain why emotional memory enhancement fails to occur.

The finding that interaction between the amygdala and prefrontal cortices plays a significant role in implementing cognitive-emotional interactions is consistent with the amygdala’s theoretical role as a coordinator of cortical functions regarding affectively significant events (Pessoa and Adolphs, 2010, but see also de Gelder et al., 2011). It is further consistent with anatomical connectivity research in the non-human primate, which purports that the relationship of the amygdala to the prefrontal cortex, and to the orbitofrontal cortex in particular, is likely implicated in emotional processing (Ghashghaei and Barbas, 2002). Non-primate animal research also supports this conclusion (Quirk and Beer, 2006).

Future work should investigate this phenomenon while more aggressively measuring correlates of arousal and anxiety. However, even if the effect turns out not to be differentiable from presumed-functionally irrelevant task anxiety, the mechanism may nonetheless prove interesting. For example, arousal may be an epiphenomenon of the emotional memory enhancement phenomenon.

**SUMMARY AND RELEVANCE FOR COGNITIVE REHABILITATION**

We have reviewed evidence that patients with AD better remember emotional content, but only if that content is not accompanied by complex, mentally taxing phenomena. The neurological mechanism underlying this benefit may lie in the amygdala’s role as an intermediary to the executive resources of the frontal cortex. We assert that the available scientific evidence suggests that patients with MCI or early AD may benefit from integrating emotional memory enhancement into the rehabilitation process. In particular, the evidence implies that coupling positively valenced stimuli to non-complex elements of the cognitive training process may be of particular use in non-depressed patients identified as having amnestic MCI or early AD. Some cohorts, such as patients with history of depression, may have attenuated positivity effects and likewise show reduced benefit from therapy. Another clinical application of Borg et al. (2011) and colleagues’ model might involve using cognitive rehabilitation protocols to target executive function directly. Increased executive resources could increase patients’ ability to benefit from emotional cues in their daily lives (for reviews, see Jean et al., 2010a,b or Martin et al., 2011; for a meta-analysis, see Li et al., 2010).

We have also identified two models, one behavioral (Borg et al., 2011) and one neuroaffective (Rosenbaum et al., 2010; Broster et al., 2012), capable of consolidating at the algorithmic and implementational levels, respectively, seemingly disparate reports of the interactions between emotional memory enhancement, aging, and task difficulty. Stronger tests of these findings, particularly regarding the role of the amygdala in recruiting prefrontal structures in MCI and dementia, would help confirm a promising model that describes how brain systems compensate for early cognitive impairment.

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