Neural correlates of retrieval-based memory enhancement: An fMRI study of the testing effect

Erik A. Wing\textsuperscript{a,b}, Elizabeth J. Marsh\textsuperscript{a}, and Roberto Cabeza\textsuperscript{a,b}
\textsuperscript{a}Department of Psychology & Neuroscience, Duke University, Box 90086, 417 Chapel Drive, Duke University, Durham, NC 27708. United States
\textsuperscript{b}Center for Cognitive Neuroscience, Duke University, Box 90999, B203 Levine Science Research Center, Durham NC, 27708. United States

Abstract

Restudying material is a common method for learning new information, but not necessarily an effective one. Research on the testing effect shows that practice involving retrieval from memory can facilitate later memory in contrast to passive restudy. Despite extensive behavioral work, the brain processes that make retrieval an effective learning strategy remain unclear. In the present experiment, we explored how initially retrieving items affected memory a day later as compared to a condition involving traditional restudy. In contrast to restudy, initial testing that contributed to future memory success was associated with engagement of several regions including the anterior hippocampus, lateral temporal cortices, and medial prefrontal cortex (PFC). Additionally, testing enhanced hippocampal connectivity with ventrolateral PFC and midline regions. These findings indicate that the testing effect may be contingent on processes that are typically thought to support memory success at encoding (e.g. relational binding, selection and elaboration of semantically-related information) in addition to those more often associated with retrieval (e.g. memory search).

Keywords

testing effect; retrieval practice; episodic memory; fMRI; hippocampus; medial PFC

1. Introduction

In the learning literature, there is abundant evidence that practice involving retrieval is more effective than practice involving passive review (Karpicke & Roediger, 2008; Roediger & Butler, 2011). In a typical memory experiment demonstrating this effect, to-be-learned items are first studied under uniform encoding conditions (Study trials) and are then practiced either through additional study (Restudy trials) or through retrieval from memory (Test trials). The reliable finding of this paradigm is that when memory is later assessed on a final memory test, items practiced in Test trials are remembered better than those practiced in Restudy trials (Roediger & Karpicke, 2006; Toppino & Cohen, 2009). The mnemonic advantage of the Test condition over the Restudy condition—known as the testing effect—
illustrates the powerful role that retrieval can play during the course of learning. Research on the testing effect has demonstrated this difference across a wide range of stimuli from foreign-language vocabulary (Carrier & Pashler, 1992) to visuospatial information (Carpenter & Kelly, 2012; Carpenter & Pashler, 2007), and is increasingly directed towards improving learning in real-world settings (Larsen, Butler, & Roediger, 2009; McDaniel, Roediger, & McDermott, 2007).

Behavioral evidence suggests that the testing effect is mediated by the enhancement of cognitive processes typically associated with encoding and/or retrieval. As an example of a process associated with encoding, there is evidence that the testing effect is mediated by semantic elaboration. For instance, in word-pair learning, the testing effect is larger for weakly- rather than strongly-related word pairs, as would be expected if pairs with lower intrinsic relatedness benefited to a greater extent by semantic binding during testing (Carpenter, 2009). Consistent with this idea, testing enhances subsequent memory not only for cue and target words, but also for semantic mediators linking cues and targets (Carpenter, 2011; Pyc & Rawson, 2010). As an example of a process associated with retrieval, there is evidence that the testing effect is enhanced by memory search during Test trials. For instance, there is substantial evidence that the testing effect is larger when Test trials involve recall rather than recognition (Carpenter, Pashler, & Vul, 2006; Glover, 1989; Kang, McDermott, & Roediger, 2007). This advantage holds even when the final test uses a different format (e.g. recognition, Carpenter & DeLosh, 2006; Glover, 1989), suggesting that recall-related processes like memory search may help explain the testing advantage, rather than the mere congruence between initial and final tests (Morris, Bransford, & Franks, 1977).

Despite a wealth of information from behavioral research, little work has been done to directly link the benefits of testing memory to brain function. This is particularly surprising given that functional neuroimaging techniques, such as functional MRI (fMRI), have successfully identified neural mechanisms associated with memory success at encoding and retrieval. A particularly powerful event-related fMRI method for investigating encoding is the subsequent memory paradigm (for a review see Paller & Wagner, 2002), which identifies greater encoding-phase activity for items that were remembered rather than forgotten on a later memory test. This difference in activity is known as the subsequent memory effect (SME) and is assumed to reflect successful encoding processes. A recent meta-analysis described a consistent set of regions that exhibit SMEs, including the medial temporal lobes (MTL), left prefrontal cortex (PFC), and superior parietal cortex (Kim, 2011). Within the MTL, SMEs in the hippocampus have been attributed to the storage of new contextual or semantic associations, which allow later recollection (Davachi, 2006; Eichenbaum, 2004; Norman & O’Reilly, 2003; Prince, Daselaar, & Cabeza, 2005). Within left PFC, SMEs in ventrolateral regions are assumed to reflect processing and evaluation of semantic features (Otten, Henson, & Rugg, 2001; Wagner et al., 1998).

Event-related fMRI can be also used to identify brain regions involved in successful retrieval operations by comparing activity for remembered and forgotten items during the test. Across numerous studies (Spaniol et al., 2009), retrieval success has been associated with activations in the hippocampus, left PFC, ventral parietal cortex, and posterior midline regions (e.g., posterior cingulate). Thus, MTL and left PFC regions are associated with both encoding and retrieval success. A study that directly compared word-pair encoding vs. retrieval (Prince et al., 2005) found that SMEs were stronger in anterior MTL regions, and retrieval success in posterior MTL regions, consistent with other reports in the literature (Lepage, Habib, & Tulving, 1998; Saykin et al., 1999; Strange, Fletcher, Henson, Friston, & Dolan, 1999). Within left PFC, encoding success was associated with ventrolateral regions, and retrieval success with dorsolateral regions (Prince et al., 2005). In contrast to the MTL
and left PFC, ventral parietal cortex and posterior midline cortex are regions very rarely associated with encoding success. In fact, these regions often show “negative SMEs” by displaying greater activity for subsequently forgotten than remembered items (Daselaar et al., 2009; Huijbers et al., 2012; Uncapher & Wagner, 2009).

A few fMRI studies have investigated encoding processes occurring during retrieval by applying the subsequent memory paradigm to new (distractor) items presented during old/new recognition tasks. This method involves testing memory for the distractors through a surprise memory test after scanning, and using performance on this second test to backsort the distractors in the first test as subsequently remembered or forgotten. In these studies, SMEs for the distractors were found in typical encoding regions including MTL (Stark & Okado, 2003) and left ventrolateral PFC (Buckner, Wheeler, & Sheridan, 2001; Huijbers, Pennartz, Cabeza, & Daselaar, 2009). Given that these regions facilitate incidental encoding during retrieval, they may also be important candidate regions involved in strengthening retrieved representations during testing. Nonetheless, these findings have only limited applicability to the testing effect, which necessitates a direct comparison of items from the Test and Restudy conditions. Unlike contrasts using a set of new items during recognition, an ideal testing comparison would examine a single Restudy or Test exposure for a set of items that have been initially exposed under uniform conditions (Study). This format would minimize potential novelty signals elicited by recognition lures and would help ensure that subsequent memory success can more easily be attributed to the single Test or Restudy exposure.

Research on other aspects of memory also offers some insight into questions surrounding the testing effect, including how repeated exposure to an item influences retention. Repeatedly presented items typically produce a reduced neural response in sensory cortex, as compared with their initial presentation (for a review, see Grill-Spector, Henson, & Martin, 2006; Henson & Rugg, 2003), and the extent of reductions has been tied to subsequent memory success or failure in a number of studies (Turk-Browne, Yi, & Chun, 2006; Wagner, Maril, & Schacter, 2000; Xue et al., 2011). This pattern has also been found in the hippocampus (Vannini, Hedden, Sullivan, & Sperling, 2012), which may exhibit memory-related hippocampal reductions that correspond to subsequent gist-based memory rather than recollection (Manelis, Paynter, Wheeler, & Reder, 2013). Related findings have been described in continuous recognition tasks, where activity in the hippocampus decreases across successive presentations but increases during successful recollection (Suzuki, Johnson, & Rugg, 2011a, 2011b), with potential significance for the long-term retention of recollected items.

Neuroimaging studies of retrieval induced forgetting (e.g. Kuhl, Dudukovic, Kahn, & Wagner, 2007) also provide information about repeated processing of a stimulus in differing contexts. In one study, experimental blocks containing selective retrieval of word pair associates were compared against blocks in which items were passively rehearsed (Wimber, Rutschmann, Greenlee, & Bäuml, 2009). The resulting contrast showed increased activity for retrieval blocks in several areas including the hippocampus, ventrolateral PFC, and lateral parietal cortex, even though a final test showed no condition-related memory differences. However, activity in such block-related contrasts may reflect broad differences in task demands between retrieval and restudy conditions that do not relate to the enhancement of later memory—a question better suited to trial-specific subsequent memory analyses. Another recent study directly examined the influence of retrieval on subsequent memory, finding that activity in dorsal anterior cingulate cortex (ACC) increased with the success of previous retrieval attempts, and also predicted across-subject memory performance on a later test (Eriksson, Kalpouzos, & Nyberg, 2011). While this finding suggests the importance of the ACC in test-related memory retention, the absence of a
Restudy comparison complicates the attempt to connect this result with retrieval-specific processes underlying the testing effect. The presence of repeated retrieval practice prior to testing in the scanner also means that memory enhancements cannot be as easily linked to neural activity within the specific test trials that were examined.

The current experiment was designed to address such issues by directly comparing the processes through which retrieval benefits subsequent memory in contrast to simple restudy (see Figure 1). We used pairs of weakly related English words that were both conducive to a cued-recall format and enabled testing over a large number of items after a substantial delay. Each fMRI run consisted of alternating Study and Practice blocks. During Study blocks, participants intentionally encoded a set of word pairs and also rated the relatedness between the left word (cue) and right word (target) in each pair. Study blocks were followed by Practice blocks, in which the previously-shown pairs were evenly split into Restudy and Test trials. In Restudy trials, word pairs were presented intact, and participants intentionally re-encoded the entire pair. In Test trials, only cue words were presented, and participants covertly retrieved the associated targets. Retrieval accuracy was gauged by requiring participants to select the last letter of the recalled target word from several possible options that appeared in the final half of Test trials (Fig 1). In order to equate visual and motor components, this procedure was also included in the final half of Restudy trials, following intentional re-encoding. Unlike testing effect studies in which the same items are practiced several times (e.g. Karpicke & Roediger, 2007), in our study each item was practiced only once (as Restudy or Test) in order to better isolate testing effects within a single learning trial. One day after the fMRI session, participants were given a final cued-recall test outside the scanner. Performance during this final cued-recall phase was used to backsort both Restudy and Test trials as either subsequently remembered or subsequently forgotten. The resulting 2 × 2 factorial design allowed us to compare the size of SMEs (subsequently remembered vs. forgotten) for word pairs presented in Test trials vs. Restudy trials.

In the context of this approach, the mnemonic processes contributing to the testing effect advantage should be reflected in areas showing greater SMEs (subsequently remembered vs. forgotten) for Test than Restudy trials (i.e., a subsequent memory × condition interaction). We investigated this question by measuring not only activation levels but also differences in functional connectivity. Given the that the hippocampus has been associated with successful relational memory during both encoding and retrieval (Davachi & Wagner, 2002; Giovanello, Schnyer, & Verfaellie, 2004; Hannula & Ranganath, 2008; Prince et al., 2005), we reasoned that this region might be particularly involved in strengthening word-pair associations during Test trials. We also expected that Test-related memory enhancement might involve areas of left ventrolateral PFC, particularly if semantic elaboration during Test trials contributes to subsequent memory as suggested by behavioral research. Additionally, we were interested in whether regions more uniquely associated with retrieval than encoding success, such as ventral parietal and posterior midline regions, would contribute to encoding success in the special context of Test trials.

2. Methods

2.1. Participants

Twenty-four right-handed, college-aged participants took part the study. Participants were healthy, native English speakers with no reported history of neurological or psychiatric episodes. All participants gave written informed consent in accordance with a protocol approved by the Duke University Institutional Review Board. One participant was excluded from analysis due to high final cued-recall performance that yielded an insufficient number of miss trials (<10) for the key hit vs. miss contrast, and another participant was excluded...
after failing to follow instructions during the final cued-recall portion of the experiment. All behavioral and fMRI analyses were conducted on the remaining 22 participants.

2.2. Stimuli

Stimuli consisted of 192 weakly related English word pairs drawn from a database of free association norms (Nelson, McEnvoy, & Schreiber, 1998). The mean forward associate strength (i.e., the probability producing the target word given the cue) for the set was 0.043. The pairs were divided into two sets of 96 pairs, which were assigned to Restudy or Test conditions (counterbalanced across participants).

2.3. Scanned paradigm

The scan session consisted of six identically-structured runs (Figure 1), each with four blocks (Study, Practice, Study, Practice). Both types of blocks started with an instruction screen (4 sec) and consisted of 16 trials (4 sec each) separated by jittered fixation periods with a mean of 3 seconds. During each Study block, participants intentionally encoded 16 new word-pairs, while rating the relatedness between the words in each pair (1=moderately related, 4=highly related). In the next Practice block, half of the 16 pairs were practiced in Test trials, and half in Restudy trials. Separate background colors were used for each of the Study, Test, and Restudy trial types. In each Test trial, only cue words were displayed, while a blank space appeared in place of the target. During the first 2 sec of Test trials, participants were instructed to covertly recall the target word, and during the last 2 sec, they had to indicate the last letter of the invisible recalled word from three letter options at the bottom of the screen. In each Restudy trial, word pairs were displayed again in full. During the first 2 sec, participants were instructed to use these trials as an additional opportunity to re-encode the pairs, and during the last 2 sec they had to press a key corresponding to the last letter of the visible target word. Thus, the response component of Test and Restudy trials was identical, and hence, it was subtracted out in direct contrast between these two types of trials. In both Restudy and Test trials, response options appeared during the final half of the trial to ensure participants focused on either restudying or recalling during the first half of the trial. Last letters were used instead of first letters to discourage a strategy in which letter options could be used as retrieval cues, and a post-scan questionnaire confirmed that participants did not rely on such a strategy to successfully recall targets. If participants could not recall the target, or the final letter for the word they retrieved was not among the three letter options provided in the trial, they selected a question mark option. No feedback was given to avoid confounding activity related to recall with activity related to feedback processing. In both Test and Restudy trials, the last letter response options changed across trials. To reduce working memory contributions, pairs presented in the first and second halves of a Study block were re-presented in the first and second halves of the Practice block, respectively (order randomized within each half). Thus, the minimum lag was between the Study and Practice trials for a pair was eight trials.

2.3.2. Post-scan session—Participants returned 24 hours after the scan for a surprise cued-recall test that assessed memory for all 192 word pairs. During this section, which took place in a computer laboratory, each cue word from the scan session was presented for 7 sec on a white background while participants attempted to recall the previously-associated target word. Verbal responses—captured via microphone—were scored as either correct or incorrect, and then used to backsort both Test and Restudy trials from the fMRI session as either subsequently remembered or forgotten. Following the final cued-recall test, participants completed a short reading span task after which they were compensated and debriefed. Even though the correspondence in format between initial and final test is not thought to contribute substantially to the testing effect (Carpenter, 2009; Glover, 1989), we note several elements of the final test that help distinguish it from the initial Test trials,
including response format, trial duration, background color, and physical location/context. Nonetheless, it is difficult to completely rule out the potential influence of cued-recall format similarity across the initial and final tests.

2.4. fMRI methods

All MRI data acquisition was conducted with a 3-T GE scanner. Scanner noise was reduced with ear plugs, and head motion was minimized with foam pads. Behavioral responses were recorded with a 4-key fiber-optic response box (Resonance Technology, Inc.), and when necessary, vision was corrected using MRI-compatible lenses that matched the distance prescription used by the participant. High-resolution structural images were collected using a 3D, T1-weighted FSPGR sequence (256 × 256 matrix, 166 slices, 1mm slice thickness). Functional images were acquired using a SENSE spiral sequence (64 × 64 matrix, TR = 2000ms, TE = 27ms, FOV = 24cm, flip angle = 60). Thirty-four contiguous slices were acquired in an interleaved fashion. Slice thickness was 3.8mm, resulting in 3.75 × 3.75 × 3.8mm voxels.

2.5. fMRI analyses

Preprocessing was performed using SPM8 software implemented in MATLAB (www.fil.ion.ucl.ac.uk/spm/). Segmented tissue probability maps were generated from anatomical volumes, and the VBM8 toolbox was then used to generate deformation fields for each participant based on the DARTEL template brain. After discarding the first 6 volumes of each run, functional images were corrected for slice time acquisition and motion. The corrected images were then coregistered to native space grey-matter tissue maps and normalized to MNI space using the deformations generated during normalization of the anatomical images.

Statistical fMRI analysis at both the subject and group level was performed in SPM5. Data were high-pass filtered using a cutoff of 128s. For each subject, evoked hemodynamic responses to event types were modeled with a delta (stick) function corresponding to the onset of stimulus presentation convolved with a canonical hemodynamic response function and temporal derivative in the context of the general linear model (GLM). Separate trial type regressors were defined for subsequently remembered and forgotten Test and Restudy trials, with additional regressors corresponding to the initial Study trials, instructional screens, and trials with no response. Regressors for session means and motion parameters were also included in the model.

Separate contrasts for subsequently remembered and forgotten Test and Restudy trials were generated for each subject and then submitted to a random effects analysis. For group analyses, a 2 × 2 repeated measures ANOVA was performed, with factors of practice condition (Test vs. Restudy) and subsequent memory (Remembered vs. Forgotten). The main effects of condition and memory were evaluated, along with the testing effect interaction that identified regions associated with memory success for the Test condition in contrast to Restudy (all contrasts were evaluated at p < .001, extent threshold = 5 voxels).

An additional functional connectivity analysis was conducted to explore functional coupling between the hippocampus—identified in the initial memory by condition interaction above—and other regions of the brain. For this analysis, a separate model was constructed in which each trial was entered as a separate regressor, yielding estimates for each individual trial within each participant (for details of this method, see Daselaar, Fleck, & Cabeza, 2006; Daselaar, Fleck, Prince, & Cabeza, 2006; Rissman, Gazzaley, & D’Esposito, 2004). Within each trial type, the mean activity from the hippocampal ROI (suprathreshold voxels from the above interaction contrast within an anatomical mask of the hippocampus) was correlated with that from each other region of interest.
with every other voxel to produce separate correlation volumes. The single subject
correlation volumes for subsequently remembered and forgotten Test and Restudy were then
entered into a 2 × 2 repeated measures ANOVA, allowing for group level comparisons of
hippocampal coupling across conditions. A connectivity interaction contrast was then
generated to detect regions that covaried with the hippocampus during successful Test more
than Restudy trials \( p < .001, k = 5 \), as with the corresponding activity ANOVA, the
interaction effect was further inclusively masked with the Test subsequent memory contrast
\( p < .01 \) to ensure effects were not merely attributable to a reverse memory effect within
Restudy trials).

3. Results

3.1. Behavioral

Confirming that participants were able to successfully recall target words during Test trials,
the last letter option was correctly selected at a rate of 91.3\%, while Restudy trials showed a
predictable ceiling effect of 99.6\%. Final cued-recall demonstrated a clear testing effect.
Memory was higher for words that had been practiced in the Test condition \( M = 0.63, SD = 0.14 \)
than those that had been shown again in the Restudy condition \( M = 0.51, SD = 0.14;\)
\( t(21) = 9.25, p < .0001 \). As illustrated by Figure 2, this testing effect difference \( M = 0.12,\)
\( SD = 0.06 \) was evident in all but one participant, with a range of 0.00 to 0.23.

3.2 Activations

To investigate the brain activity associated with retrieval-based memory enhancement, we
conducted a 2 (subsequent memory: remember vs. forgotten) × 2 (condition: Test vs.
Restudy) ANOVA. As noted before, the critical comparison regarding the testing effect is
the interaction between subsequent memory and condition, which reveals areas where SMEs
are greater for Test than Restudy trials. The main effects of condition are not directly related
to the testing effect because they reflect the particular features of the Test and Restudy
conditions that may not play a role in promoting later memory. Regions showing a main
effect of condition are listed in in Table 1. Compared to Test trials, Restudy trials were
associated with greater activity in inferior parietal, lateral temporal, and dorsal PFC.
Conversely, stronger activations for Test than Restudy trials were found in dorsal ACC,
bilateral ventrolateral PFC extending into anterior insula, and midbrain. A main effect of
subsequent memory was found in a single area in dorsomedial PFC (MNI: 8, 45, 42), which
was more active for subsequently remembered than forgotten trials in both Test and Restudy
conditions. The relative lack of SMEs shared by Test and Restudy trials is consistent with
previous evidence that SMEs are task dependent (Fletcher, Stephenson, Carpenter,
Donovan, & Bullmorel, 2003; Otten & Rugg, 2001a), as well as with substantial SME
differences between Test and Restudy conditions, as revealed by the interaction findings
below.

The main focus of the study was to identify regions showing greater SMEs for Test than
Restudy conditions, which were isolated by memory × condition interactions. As listed in
Table 2, interaction effects were found in the bilateral hippocampus, lateral temporal cortex,
medial prefrontal cortex, and left striatum. The hippocampal interaction occurred in the
anterior hippocampus, with somewhat stronger differences evident in left hemisphere (Fig.
3-B, C). The effects in lateral temporal cortex were also stronger in the left hemisphere,
where they were localized to middle and inferior temporal gyri in contrast to superior
temporal gyrus in the right hemisphere (Fig. 3-A). In addition to showing higher activity for
subsequently remembered vs. forgotten Test trials, some regions from the interaction also
showed an inverted pattern on Restudy trials, with higher activity for subsequently forgotten
than remembered trials. No significant SMEs were found for the reversed contrast (Restudy > Test).

3.3. Functional Connectivity

Motivated by the pattern of findings from the interaction contrast, a functional connectivity analysis was performed to examine the potential for differential coupling between the hippocampus and other brain regions. This analysis addressed the possibility that regions insensitive to condition differences might nonetheless show different patterns in coactivity with a region directly identified as important to later memory. Consequently, a second 2 × 2 repeated measures ANOVA with factors of condition and memory success was constructed using individual subject maps reflecting condition-specific hippocampal correlation instead of activity. The corresponding interaction revealed regions where differential coactivation with the hippocampus predicted subsequent memory success to a greater extent for Test than Restudy trials (Figure 4, Table 3). This pattern was found in the posterior cingulate cortex (Fig. 4-A), medial PFC (Fig. 4-B), and left ventrolateral PFC (Fig. 4-C).

4. Discussion

The current study explored the neural correlates of the testing effect. The capacity of retrieval to promote subsequent memory was examined, as well as the how mnemonic differences during testing differ from simple restudy. On a final memory test one day after the scan, target words that had been practiced through retrieval were remembered at a higher rate than those practiced via restudy, confirming a behavioral testing effect. The critical interaction between practice condition and subsequent memory showed that testing enhanced subsequent memory effects (SMEs) in the hippocampus, left temporal cortex, and medial PFC. Functional connectivity analyses identified regions whose interactions with the hippocampus predicted subsequent memory primarily for the Test condition, including left ventrolateral PFC, medial PFC, and posterior cingulate. Below we discuss these findings in the context of relational encoding processes, which may allow for relevant semantic information to be incorporated into coherent representations through testing. We also discuss how retrieval processes like memory search may contribute to this process, and connect the present findings to related research on how the integration of information during retrieval influences memory consolidation.

4.1. Hippocampus

In contrast to Restudy trials, bilateral anterior hippocampus activity was greater for Test trials that were subsequently remembered than those that were forgotten on the final test. While the hippocampus figures centrally in many mnemonic operations, it is thought to play a particularly important role in binding disparate information into coherent representations at encoding (Eichenbaum, Yonelinas, & Ranganath, 2007; Yassa & Stark, 2011). At retrieval, the hippocampus may coordinate reinstatement of initially-formed associations through pattern completion processes that operate in response to partially-overlapping cues (Norman & O’Reilly, 2003). Differential engagement of relational memory processes suggests one way in which retrieval practice may enhance memory over traditional restudy. In the present study, Test trials might be expected to induce processing of previously formed cue-target associations more often than Restudy trials, wherein such associations are of less strategic importance. The element of memory search during target retrieval may also produce novel associations between word pairs and information (e.g. retrieval candidates). When incorporated into an updated representation, such related information—whether formed during initial encoding or during the process of retrieval—may serve as additional cues during final retrieval, improving retention and facilitating recollection-based memory.
Support for such an account of the present hippocampal findings is evident in both the 
neuroimaging literature on relational memory and in testing effects research. Consistent with 
the localization of observed SME effects, several studies of relational encoding have found 
activity in the anterior hippocampus that tracks subsequent memory success (Jackson & 
Schacter, 2004; Prince et al., 2005; Sperling et al., 2003). Attempts to link brain function 
with recollective experience have also emphasized the critical role of the hippocampus. 
Numerous studies using verbal material have reported hippocampal activity for encoding 
that led to subsequent source or relational memory (Davachi, Mitchell, & Wagner, 2003; 
Ranganath et al., 2004; Uncapher, Otten, & Rugg, 2006), with a similar pattern evident for 
analogous comparisons during the retrieval phase (Giovanello et al., 2004; Yonelinas, Otten, 
Shaw, & Rugg, 2005). Hippocampal activity has also been observed during successful 
encoding (Fernández et al., 1998; Habib & Nyberg, 2008; Staresina & Davachi, 2006) and 
retrieval (Meltzer & Constable, 2005; de Zubicaray et al., 2007) of information during cued 
or free recall tests, which are thought to diminish the opportunity for familiarity-based 
memory responses.

Recent behavioral findings also implicate relational processing in retrieval practice effects. 
In comparison to traditional restudy, testing has been shown to increase memory for 
semantic mediators—words that link cue-target pairs. This is the case both for mediators 
that are explicitly generated to aid retrieval, (Pyc & Rawson, 2010) and for mediators that 
are never explicitly produced but that contain a strong semantic connection between cues 
and targets (Carpenter, 2011). Carpenter (2011) found that when word pairs (e.g. Mother-
Child) were practiced through testing as opposed to restudy, participants were later more 
likely to false alarm to related semantic mediators (e.g. Father). Initial testing also made 
these mediators more effective substitute cues from which to retrieve targets on a modified 
final test. The availability of mediating information to serve as additional memory cues may 
also help explain why retrieval practice also promotes subsequent recollection, as observed 
in studies exploring the testing effect in a dual process framework (Chan & McDermott, 
2007; Verkoeijen, Tabbers, & Verhage, 2011). Although multiple factors likely contribute to 
the testing effect, the pattern of hippocampal findings observed in the present study provides 
further support for the involvement of relational processing, which should be particularly 
beneficial when centered on stimulus-related semantic associations, as detailed in the above 
behavioral studies.

4.2. Lateral Temporal Cortex

The integration of related stimulus associations requires not only a binding mechanism, but 
also access to pre-existing knowledge. Such knowledge is necessary for assessing the 
relatedness of word-pairs during the initial Study trials, and may be tapped to varying 
degrees during Restudy and Test trials. Whereas Restudy trials contain both components of 
the to-be-remembered pair, Test trials require retrieval of previously stored information that 
may be facilitated by initially formed associations. Semantic knowledge would also be 
necessary during the process of memory search, as semantically relevant concepts are 
brought to mind and evaluated in the attempt to produce the specific target.

Findings from the interaction of condition and memory success provide some indication that 
the retrieval practice may benefit from differential processing of semantic information. Most 
notably, activity in left middle temporal cortex was associated with stronger SMEs during 
Test than during Restudy trials. Left lateral temporal cortex has been linked to both semantic 
and episodic memory systems, which are thought to interact closely (Tulving, 1972, 2002). 
Neuroimaging studies of semantic memory retrieval often report activity in left lateral and 
anterior temporal cortices (Simmons & Martin, 2009; Thompson-Schill, 2003), and 
degeneration of the left temporal lobe contributes to memory deficits in semantic dementia 
(Hodges & Patterson, 2007; Snowden, Griffiths, & Neary, 1996). In episodic memory
studies, lateral temporal cortex has been associated with successful memory during both encoding and retrieval (Spaniol et al., 2009). Exploring the relationship between of semantic and episodic memory, Menon et al. (2002) found that activity left lateral temporal cortex (BA 21/22) corresponded to successful retrieval both for the semantic properties of words, and for their mnemonic status in an episodic memory test (Menon, Boyett-Anderson, Schatzberg, & Reiss, 2002). The authors suggest that this finding reflects the use of previously encoded semantic information during episodic retrieval. Such an account fits well with the pattern of current lateral temporal SMEs for the Test condition. Presumably, not all relational information active during Test helps enhance subsequent memory. Associations that contain more specific or detailed connections to target concepts are likely to serve as better subsequent cues. Processes mediating selection and controlled retrieval of unique associations should also be important in producing the type beneficial associations that can be effectively integrated to coherent representations.

4.3. Left ventrolateral PFC

Left ventrolateral PFC effects were observed in an analysis performed to help clarify how hippocampal connectivity during retrieval practice contributes to subsequent memory. Differences in hippocampal connectivity were examined across Test and Restudy trials that varied in final test memory outcome. This comparison produced an expanded and spatially distinct set of regions from those observed in the standard testing effect contrast. Stronger connectivity with ventrolateral PFC was found for Test trials that were effective in producing successful memory on the final recall test. This result indicates that the beneficial aspects of retrieval practice may arise through differential coupling between the hippocampus and other cortical regions that do not differ in overall activity.

The involvement of left ventrolateral PFC in the memory-related cognitive control operations (e.g. Badre & Wagner, 2007) is particularly relevant to test-enhanced learning in the current design. This region has been linked to retrieval of knowledge-based representations in a number of studies comparing semantic vs. non-semantic conditions (e.g. Devlin, Matthews, & Rushworth, 2003; McDermott, Petersen, Watson, & Ojemann, 2003). Damage to ventrolateral PFC has also been shown to impair performance on semantic tasks (Martin & Cheng, 2006; Thompson-Schill, D’Esposito, Aguirre, & Farah, 1997). The nature of current findings in the ventrolateral PFC indicates a more complex relationship when considering the how Restudy or Test trials influence subsequent memory. While this region may operate in a similar capacity to facilitate immediate cued recall during initial Test trials (i.e. a main effect of Test > Restudy), it may not invariably strengthen retrieved items such that they are better remembered in the future. Instead, information selected by the ventrolateral PFC may enhance later memory only when it can be effectively integrated into active representations via the hippocampus. In this scenario, the present connectivity findings would reflect a hippocampally-mediated elaboration effect in which the behavioral advantages of Test draw on the increased likelihood of collaboration between frontal regions, involved in controlled semantic processing, and the hippocampus, which integrates associated content into a more durable representation.

4.4 Medial PFC

The medial PFC was another region where hippocampal connectivity predicted subsequent memory for Test more than Restudy trials. The medial PFC contains considerable anatomical connections with limbic regions including the hippocampal formation (Cavada, Compañy, Tejedor, Cruz-Rizzolo, & Reinoso-Suárez, 2000), and memory studies have also reported functional associations between these regions. Functional connectivity between anterior midline and hippocampus has been found during episodic encoding in the context of working memory (Ranganath, Heller, Cohen, Brozinsky, & Rissman, 2005) and schema
formation (van Kesteren, Fernández, Norris, & Hermans, 2010). Related areas of memory research have explored how retrieval affects pre-existing mnemonic associations. For example, during retrieval-mediated associative learning, newly acquired memories are integrated with (or influenced by) distinct but overlapping representations that have already been encoded (Zeithamova, Dominick, & Preston, 2012). Studies in both humans and rodents have emphasized the importance of the hippocampus and medial PFC during this process, which may enhance consolidation of newly encoded memories (Tse et al., 2007, 2011).

Consolidation processes may also contribute to retrieval practice advantages, as the magnitude of the testing effect typically increases with the temporal separation between retrieval practice and final test (i.e. the retention interval) (Roediger & Butler, 2011; Toppino & Cohen, 2009). Research on the testing effect has only recently begun to consider the role of consolidation, (Eriksson et al., 2011; Finn & Roediger, 2011). While not directly assessing Test/Restudy differences, a recent study found Test-related activity in a more dorsal ACC region that tracked the success of previous retrieval attempts while predicting retention across subjects (Eriksson et al., 2011). In the present study, medial PFC activity was evident in the key testing effect interaction, but this region also demonstrated differential connectivity with the hippocampus during Test and Restudy trials. This behaviorally-selective functional relationship provides a further connection to the research on retrieval-mediated learning for distinct but related material, and supports the notion that testing may strengthen memory through the retrieval and elaboration of initially encoded associative information.

4.5 Posterior cingulate cortex

The functional connectivity difference in posterior midline is interesting because this region in known to play very different role during retrieval than at encoding (Spaniol et al., 2009). Whereas this region is associated with retrieval success, it is rarely associated with encoding success. Examination of encoding effects in posterior midline and other default mode network (DMN) regions including ventral parietal cortex have even shown an inverted success pattern, with higher activity for subsequently forgotten than remembered items (Daselaar, Prince, & Cabeza, 2004; Otten & Rugg, 2001b; Wagner & Davachi, 2001). In posterior midline, reverse SMEs have been shown to overlap with retrieval success activations both within subjects and across a variety of memory studies, underscoring the reliability and pervasiveness of these phase-related differences (Daselaar et al., 2009; Huijbers, Pennartz, Cabeza, & Daselaar, 2011). Furthermore, a study comparing task-related hippocampal activation with resting state default mode activity found greater coupling between the hippocampus and DMN during retrieval, but divergent activity profiles during encoding (Huijbers et al., 2011). These findings illustrate the complex relationship between activity and connectivity in the present results, where the hippocampus appears to act in concert with traditional retrieval regions during Test trials that promote later memory. While many of the present activity findings for Test/Restudy interactions appear to reflect the enhancement of traditional encoding regions (hippocampus, left temporal cortex, left PFC, etc.), patterns of hippocampal connectivity suggest that the testing effect is also associated with recruitment of regions consistently tied to retrieval operations (e.g., posterior cingulate). Further research is necessary to disentangle these two possible mechanisms.

4.6 Conclusion

Following the extensive behavioral research on the testing effect, we used event-related fMRI to investigate the processes through which retrieval promotes subsequent memory in contrast to traditional restudy. Consistent with past findings, final test memory performance was higher for items that had been initially retrieved vs. restudied during scanning. The
critical interaction between practice condition (Test or Restudy) and final memory outcome showed that increased activity in the hippocampus and lateral temporal cortex predicted later memory during testing but not restudy. A connectivity analysis additionally revealed increased coupling between the hippocampus and ventrolateral PFC, medial PFC, and posterior cingulate cortex when retrieval practice was effective at producing subsequent memory. These results suggest that the power of retrieval to promote memory stem in part from relational memory processes that operate on selected semantic associations during testing. The hippocampus may be involved in integrating relevant information into updated representations, supporting memory search, and interfacing with other regions involved in consolidation processes to produce more durable memories.

Acknowledgments
This work was supported by the National Institute on Aging [grant number R56 AG23770]. We thank Henry L. Roediger III for his advice on experimental design and theoretical issues, James E. Kragel and Jared Stokes for their assistance with programming of the paradigm, and Kerry Townsend for her assistance with data collection and analysis.

References


Kang SHK, McDermott KB, Roediger HL. Test format and corrective feedback modify the effect of testing on long-term retention. European Journal of Cognitive Psychology. 2007; 19(4):528–558.10.1080/09541440601056620

Karpicke JD, Roediger HL. Repeated retrieval during learning is the key to long-term retention. Journal of Memory and Language. 2007; 57(2):151–162.


Nelson DL, McEnvoy CL, Schreiber TA. The University of South Florida word association, rhyme, and word fragment norms. 1998

Neuropsychologia. Author manuscript; available in PMC 2014 October 01.


Otten LJ, Rugg MD. Task-dependency of the neural correlates of episodic encoding as measured by fMRI. Cerebral Cortex. 2001a; 11(12):1150–60. [PubMed: 11709486]


Neuropsychologia. Author manuscript; available in PMC 2014 October 01.


Highlights

- We explored the neural correlates of the testing effect
- Subsequent memory effects for retrieval practice vs. passive restudy were compared
- This interaction showed activity in hippocampus/medial PFC/lateral temporal cortex
- VLPFC and midline areas showed task-related hippocampal connectivity differences
- Testing effect may stem from processes benefiting encoding and retrieval
Figure 1.
Schematic of one run from the fMRI portion of experiment. Each Study and Practice block contained 16 trials. Twenty-four hours after the scan session, memory for all initial word pairs was assessed in a final cued-recall test (not shown). Performance on this final test was used to backsort Practice (i.e. Test/Restudy) trials as either subsequently remembered or forgotten.
Figure 2.
Behavioral testing effect. Single subject behavioral performance showing proportion correct recall during final cued-recall test as a function of scan-session practice condition (Test/Restudy).
Figure 3.
Testing effect interaction. Brain regions showing a condition (Test/Restudy) by memory (subsequently remembered/subsequently forgotten) interaction. Interaction effects are evident in left middle/inferior temporal gyri (A) and bilateral anterior hippocampus (B, C). Bars reflect SME (difference in activity between subsequently remembered and forgotten). Error bars denote standard error.
Figure 4.
Hippocampal connectivity interaction. Brain regions showing hippocampal connectivity differences by condition and subsequent memory. Regions in both posterior (posterior cingulate/precuneus, A) and anterior (medial PFC, B) midline showed this effect, along with left ventrolateral PFC (C). Bars reflect difference in hippocampal connectivity between subsequently remembered and forgotten. Error bars denote standard error.
### Table 1

<table>
<thead>
<tr>
<th>Region</th>
<th>Hem</th>
<th>BA</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>Z</th>
<th>vox</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Test&gt;Restudy</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inferior frontal gyrus</td>
<td>R</td>
<td>47</td>
<td>34</td>
<td>19</td>
<td>−8</td>
<td>&gt; 8</td>
<td>1442</td>
</tr>
<tr>
<td></td>
<td>L</td>
<td>47</td>
<td>−30</td>
<td>23</td>
<td>−8</td>
<td>&gt; 8</td>
<td></td>
</tr>
<tr>
<td></td>
<td>L</td>
<td>45</td>
<td>−53</td>
<td>34</td>
<td>4</td>
<td>6.18</td>
<td></td>
</tr>
<tr>
<td>Globus pallidus</td>
<td>R</td>
<td>−</td>
<td>11</td>
<td>8</td>
<td>−8</td>
<td>5.90</td>
<td></td>
</tr>
<tr>
<td>Anterior cingulate gyrus</td>
<td>−</td>
<td>32/24</td>
<td>0</td>
<td>23</td>
<td>34</td>
<td>7.59</td>
<td>756</td>
</tr>
<tr>
<td>Inferior temporal gyrus</td>
<td>L</td>
<td>37</td>
<td>−49</td>
<td>−49</td>
<td>−11</td>
<td>3.76</td>
<td>7</td>
</tr>
<tr>
<td>Parahippocampal gyrus</td>
<td>L</td>
<td>−</td>
<td>−23</td>
<td>−41</td>
<td>0</td>
<td>3.50</td>
<td>6</td>
</tr>
<tr>
<td>Precuneus</td>
<td>L</td>
<td>7</td>
<td>−26</td>
<td>−68</td>
<td>38</td>
<td>4.55</td>
<td>58</td>
</tr>
<tr>
<td>Posterior cingulate gyrus</td>
<td>−</td>
<td>23</td>
<td>0</td>
<td>−26</td>
<td>19</td>
<td>4.19</td>
<td>37</td>
</tr>
<tr>
<td>Middle occipital gyrus</td>
<td>L</td>
<td>19</td>
<td>−34</td>
<td>−83</td>
<td>15</td>
<td>3.44</td>
<td>9</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>−</td>
<td>−</td>
<td>0</td>
<td>−60</td>
<td>−34</td>
<td>4.28</td>
<td>26</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>−</td>
<td>−</td>
<td>15</td>
<td>−41</td>
<td>−46</td>
<td>3.81</td>
<td>5</td>
</tr>
<tr>
<td><strong>Restudy&gt;Test</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Middle/medial frontal gyrus</td>
<td>R</td>
<td>8</td>
<td>23</td>
<td>19</td>
<td>42</td>
<td>6.45</td>
<td>756</td>
</tr>
<tr>
<td>Middle/superior frontal gyrus</td>
<td>L</td>
<td>9</td>
<td>−30</td>
<td>23</td>
<td>38</td>
<td>5.05</td>
<td>125</td>
</tr>
<tr>
<td>Medial frontal gyrus</td>
<td>R</td>
<td>10</td>
<td>11</td>
<td>60</td>
<td>8</td>
<td>4.02</td>
<td>37</td>
</tr>
<tr>
<td>Middle temporal gyrus</td>
<td>R</td>
<td>21</td>
<td>64</td>
<td>−49</td>
<td>−4</td>
<td>4.64</td>
<td>37</td>
</tr>
<tr>
<td></td>
<td>R</td>
<td>21</td>
<td>56</td>
<td>−19</td>
<td>−23</td>
<td>5.49</td>
<td>112</td>
</tr>
<tr>
<td></td>
<td>L</td>
<td>20</td>
<td>−56</td>
<td>−4</td>
<td>−30</td>
<td>3.61</td>
<td>37</td>
</tr>
<tr>
<td></td>
<td>L</td>
<td>21</td>
<td>−45</td>
<td>4</td>
<td>−34</td>
<td>3.55</td>
<td>14</td>
</tr>
<tr>
<td>Superior temporal/Postcentral gyrus</td>
<td>L</td>
<td>22/40</td>
<td>−49</td>
<td>−11</td>
<td>−4</td>
<td>4.94</td>
<td>161</td>
</tr>
<tr>
<td>IPL/Angular gyrus</td>
<td>R</td>
<td>39/40/7</td>
<td>49</td>
<td>−68</td>
<td>30</td>
<td>6.81</td>
<td>329</td>
</tr>
<tr>
<td>Supramarginal gyrus</td>
<td>L</td>
<td>40</td>
<td>−56</td>
<td>−56</td>
<td>27</td>
<td>6.28</td>
<td>172</td>
</tr>
<tr>
<td>Angular gyrus</td>
<td>L</td>
<td>39</td>
<td>−49</td>
<td>−71</td>
<td>34</td>
<td>4.92</td>
<td>172</td>
</tr>
<tr>
<td>Precuneus</td>
<td>R</td>
<td>31</td>
<td>8</td>
<td>−53</td>
<td>34</td>
<td>3.68</td>
<td>15</td>
</tr>
<tr>
<td>Postcentral gyrus</td>
<td>R</td>
<td>40</td>
<td>53</td>
<td>−30</td>
<td>19</td>
<td>3.96</td>
<td>39</td>
</tr>
<tr>
<td></td>
<td>R</td>
<td>40</td>
<td>23</td>
<td>−41</td>
<td>57</td>
<td>3.51</td>
<td>5</td>
</tr>
<tr>
<td>Lingual gyrus</td>
<td>L</td>
<td>18</td>
<td>−23</td>
<td>−83</td>
<td>−8</td>
<td>5.77</td>
<td>264</td>
</tr>
</tbody>
</table>
Region | Hem | BA | x  | y  | z  | Z   | vox |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Cuneus</td>
<td>L/R</td>
<td>18/19</td>
<td>8</td>
<td>−83</td>
<td>19</td>
<td>5.19</td>
<td>96</td>
</tr>
<tr>
<td>Lingual gyrus</td>
<td>R</td>
<td>18</td>
<td>11</td>
<td>−71</td>
<td>−8</td>
<td>5.24</td>
<td>111</td>
</tr>
</tbody>
</table>

Up to 5 local maxima set 16mm apart are reported for each cluster. BA, Brodmann area; Hem, hemisphere. Coordinates are in MNI space.
### Table 2

<table>
<thead>
<tr>
<th>Region</th>
<th>Hem</th>
<th>BA</th>
<th>x</th>
<th>Y</th>
<th>z</th>
<th>Z</th>
<th>vox</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Test &gt; Restudy</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anterior cingulate gyrus</td>
<td>L</td>
<td>32</td>
<td>−15</td>
<td>30</td>
<td>0</td>
<td>3.99</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>L</td>
<td>32</td>
<td>−15</td>
<td>41</td>
<td>−4</td>
<td>3.60</td>
<td>13</td>
</tr>
<tr>
<td>Inferior temporal gyrus</td>
<td>L</td>
<td>20</td>
<td>−49</td>
<td>−8</td>
<td>−23</td>
<td>4.41</td>
<td>18</td>
</tr>
<tr>
<td>Superior temporal gyrus</td>
<td>R</td>
<td>22</td>
<td>53</td>
<td>−8</td>
<td>−11</td>
<td>4.29</td>
<td>23</td>
</tr>
<tr>
<td>Hippocampus</td>
<td>R</td>
<td>−</td>
<td>30</td>
<td>−8</td>
<td>−23</td>
<td>4.1</td>
<td>11</td>
</tr>
<tr>
<td>Hippocampus/Parahippocampal gyrus</td>
<td>L</td>
<td>−</td>
<td>−30</td>
<td>−11</td>
<td>−27</td>
<td>3.67</td>
<td>12</td>
</tr>
<tr>
<td>Insula</td>
<td>R</td>
<td>13</td>
<td>38</td>
<td>−23</td>
<td>15</td>
<td>4.03</td>
<td>23</td>
</tr>
<tr>
<td></td>
<td>R</td>
<td>13</td>
<td>45</td>
<td>−30</td>
<td>19</td>
<td>3.65</td>
<td>9</td>
</tr>
<tr>
<td>Insula/claustrum</td>
<td>L</td>
<td>−</td>
<td>−34</td>
<td>−4</td>
<td>4</td>
<td>4.02</td>
<td>50</td>
</tr>
</tbody>
</table>

Up to 3 local maxima set 8 mm apart are reported for each cluster. BA, Brodmann area; Hem, hemisphere. Coordinates are in MNI space.
### Table 3

**Hippocampal Connectivity Interaction**

<table>
<thead>
<tr>
<th>Region</th>
<th>Hem</th>
<th>BA</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>Z</th>
<th>vox</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Test &gt; Restudy interaction</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inferior frontal gyrus</td>
<td>L</td>
<td>47</td>
<td>-23</td>
<td>19</td>
<td>-19</td>
<td>4.63</td>
<td>20</td>
</tr>
<tr>
<td></td>
<td>L</td>
<td>44</td>
<td>-60</td>
<td>15</td>
<td>8</td>
<td>3.98</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>L</td>
<td>45</td>
<td>-49</td>
<td>26</td>
<td>4</td>
<td>3.49</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>R</td>
<td>47</td>
<td>53</td>
<td>26</td>
<td>-4</td>
<td>3.81</td>
<td>7</td>
</tr>
<tr>
<td>Middle frontal gyrus</td>
<td>R</td>
<td>11</td>
<td>23</td>
<td>38</td>
<td>-19</td>
<td>3.74</td>
<td>5</td>
</tr>
<tr>
<td>Medial frontal gyrus</td>
<td>R</td>
<td>11</td>
<td>11</td>
<td>41</td>
<td>-15</td>
<td>3.69</td>
<td>6</td>
</tr>
<tr>
<td>Superior temporal gyrus</td>
<td>L</td>
<td>22</td>
<td>-64</td>
<td>-38</td>
<td>4</td>
<td>3.52</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>R</td>
<td>21</td>
<td>49</td>
<td>4</td>
<td>-19</td>
<td>4.44</td>
<td>17</td>
</tr>
<tr>
<td>Insula/Superior temporal gyrus</td>
<td></td>
<td></td>
<td>13/22</td>
<td>45</td>
<td>-15</td>
<td>0</td>
<td>3.63</td>
</tr>
<tr>
<td>Middle temporal gyrus</td>
<td>R</td>
<td>37</td>
<td>56</td>
<td>-45</td>
<td>-23</td>
<td>3.76</td>
<td>8</td>
</tr>
<tr>
<td>Inferior temporal gyrus</td>
<td>L</td>
<td>20</td>
<td>-56</td>
<td>-26</td>
<td>-27</td>
<td>4.04</td>
<td>5</td>
</tr>
<tr>
<td>Parahippocampal gyrus</td>
<td>L</td>
<td>36</td>
<td>-34</td>
<td>-23</td>
<td>-23</td>
<td>4.31</td>
<td>5</td>
</tr>
<tr>
<td>Supramarginal gyrus</td>
<td>R</td>
<td>40</td>
<td>60</td>
<td>-53</td>
<td>34</td>
<td>4.25</td>
<td>21</td>
</tr>
<tr>
<td>Postcentral gyrus</td>
<td>L</td>
<td>7</td>
<td>-19</td>
<td>-53</td>
<td>72</td>
<td>4.14</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>R</td>
<td>5/40</td>
<td>26</td>
<td>-49</td>
<td>68</td>
<td>3.66</td>
<td>13</td>
</tr>
<tr>
<td>Precuneus</td>
<td>R</td>
<td>7</td>
<td>15</td>
<td>-49</td>
<td>46</td>
<td>3.93</td>
<td>14</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>R</td>
<td>-</td>
<td>4</td>
<td>-68</td>
<td>-23</td>
<td>3.72</td>
<td>7</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>R</td>
<td>-</td>
<td>23</td>
<td>-86</td>
<td>-42</td>
<td>4.33</td>
<td>11</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>R</td>
<td>-</td>
<td>45</td>
<td>-56</td>
<td>-30</td>
<td>3.81</td>
<td>14</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>L</td>
<td>-</td>
<td>-41</td>
<td>-53</td>
<td>-30</td>
<td>3.59</td>
<td>6</td>
</tr>
</tbody>
</table>

Up to 3 local maxima set 8mm apart are reported for each cluster. BA, Brodmann area; Hem, hemisphere. Coordinates are in MNI space.