Modulation of target recollection and recollection rejection networks due to retrieval facilitation and interference

Caitlin R. Bowman,1,2,3 Shalome L. Sine,1,3 and Nancy A. Dennis1

1The Department of Psychology, The Pennsylvania State University, University Park, Pennsylvania 16802, USA; 2The Department of Psychology, University of Oregon, Eugene, Oregon 97403, USA

To better understand neural recollection processing, we induced interference in target recollection by presenting related lures before their respective targets and facilitated recollection rejection of lures by presenting targets before their related lures. Target recollection following interference recruited visual and prefrontal cortices, showing that these regions support recollection when related information has disrupted target representations. Recollection rejection following target presentation recruited angular gyrus, indicating that this region supports recollection rejection when target representations are strong and highly accessible. Thus, recollection networks are sensitive to the accessibility of target representations that are affected by the presentation of related information during retrieval.

[Supplemental material is available for this article.]
these networks would instead support recollection responses either unhindered by interference or facilitated by target re-presentation.

Data from 21 participants (12 females, mean age = 22.3 yr, SD = 3.04 yr, range = 18–28 yr) were included following six exclusions for incomplete data (1), insufficient trials in a condition of interest (4), and movement >3 mm within a run (1). Participants were right-handed, native English speakers and screened for contraindications for functional magnetic resonance imaging (fMRI). Participants provided informed consent and received financial compensation for participation. Procedures were approved by Penn State’s Institutional Review Board.

Three hundred and sixteen object images were collected from the Bank of Standardized Stimuli database (Brodeur et al. 2010) and an Internet image search. Images were equated for resolution, resized to ∼400 × 400 pixels, and displayed at a screen resolution of 1024 (H) × 768 (V) at 75 Hz. At the viewing distance of 143 cm, the display area was 20° (H) × 16° (V) with experimental stimuli subtending 5° (H) × 4° (V).

Encoding and retrieval were completed in a Siemens Trio 3T scanner with a 12-channel head coil. During encoding, 96 images were each presented for 1500 msec followed by 500 msec additional responding time. Each item was followed by a variable inter-trial fixation (M = 2470 msec, SD = 1760 msec, range: 1000–12,000 msec). Participants made a size judgment about each item (i.e., is this item bigger or smaller than a shoebox?). Next, participants completed a filler task (Raven’s Progressive Matrices) while undergoing a structural scan (MPRAGE: TR: 1650 msec, TE: 2.03 msec, 256-mm field of view (FOV), 160 sagittal slices, 1 mm slice thickness). Approximately 10 min elapsed between encoding and retrieval. Participants completed four runs of retrieval while undergoing echoplanar functional scans (descending acquisition, TR: 2500 msec, TE: 25 msec, 240-mm FOV, 42 axial slices, 3 mm slice thickness). Participants were presented with the target (e.g., a shirt) plus two types of new items from each category: one alternative exemplar of each target (e.g., a different shirt) and one thematically related item (e.g., a sweater). Importantly, only the former (alterative exemplars) were used to define recollection rejection in the present analyses as they maximized these responses (see Fig. 1). Twenty-eight items from novel categories were also included at retrieval but were not the focus of present analyses. Each item was presented for 3000 msec followed by a variable inter-trial interval (M = 2340 msec, SD = 1440 msec, range: 1000–12,000 msec). The target was presented first (T1 condition) for half of the categories while the lure was presented first (L1 condition) for the other half. The distance between a target and its lure ranged between 11 and 285 trials (M = 101.52, SD = 76.15).

Participants made memory responses within an adapted “Remember-Know-New” paradigm (Bowman and Dennis 2016) designed to identify target recollection and recollection rejection. Participants responded “Remember” when they thought an item was old and they remembered specific details of its prior presentation. Participants responded “Familiar” when they thought an item was old but they did not remember specific details of its prior presentation. “Unfamiliar” responses indicated that they believed an item was new because it did not resemble or bring to mind anything from study. “Different” responses indicated that they believed the item was new because they could recall aspects of studied items that provided evidence that the item was not previously presented. Analyses focused on “remember” responses to targets (target recollection) and “different” responses to lures (recollection rejection) in order to focus on recollection-based responding across targets and lures. Following retrieval, participants completed a questionnaire that asked about their strategies and were debriefed.

Functional data were analyzed with SPM8 (http://www.fil.ion.ucl.ac.uk/spm). Scans were checked for movement and artifacts using a diagnostic function, TSDiffAna (Freiburg Brain Imaging), spatially realigned to correct for motion, coregistered with each individual’s high-resolution anatomical image, spatially normalized to a standard stereotaxic space using the Montreal Neurological Institute template, and spatially smoothed using a 6-mm isotropic Gaussian kernel. Trial-related activity was modeled in the General Linear Model with trial onsets convolved with SPM’s canonical hemodynamic response function using trial-by-trial reaction time as the trial duration to account for differences in response times across trials of interest (see below). Statistical parametric maps for each participant were identified by applying linear contrasts to the βs for the events of interest. The current analysis focused on four regressors: T1 Target Recollection

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Figure 1. Behavioral paradigm. During the encoding phase, participants viewed one object image from each of 96 object categories and made a size judgment (bigger or smaller than a shoebox?). During the retrieval phase, participants viewed targets and lures from each of the categories from study. Within each category, the target could be presented before the lure (T1 condition) or the lure could be presented before the target (L1 condition). Participants made recognition judgments using the adapted Remember-Know-New response paradigm.
Table 1. Results of fMRI contrasts

| Region                                              | BA | H | x  | y  | z  | t  | k  |
|-----------------------------------------------------|----|---|----|----|----|----|----|
| **Target Recollection with lure interference (L1 “Remember” hit > T1 “Remember” hit)** |     |   | 36 | 11 | 52 | 5.69 | 38 |
| Middle frontal gyrus                                | 6/8| R |    |    |    |    |    |
| Inferior parietal cortex                            | 40 | M |    |    |    |    |    |
| Lateral occipital cortex                            | 19 | L |    |    |    |    |    |
| Lateral occipital cortex                            | 19 | R |    |    |    |    |    |
| Lingual gyrus                                       | 18 | M |    |    |    |    |    |
| **Target Recollection without lure interference (T1 “Remember” hit > L1 “Remember” hit)** |     |   | 36 | 11 | 52 | 5.69 | 38 |
| Ventromedial prefrontal cortex                      | 25/32| M |  6 | 35 |  8 | 5.29 | 61 |
| **Recollection rejection with target facilitation (T1 “Different” correct rejection > L1 “Different” correct rejection)** |     |   | 36 | 11 | 52 | 5.69 | 38 |
| Angular gyrus                                       | 40 | R |    |    |    |    |    |
| **Recollection rejection without target facilitation (L1 “Different” correct rejection > T1 “Different” correct rejection)** |     |   | 36 | 11 | 52 | 5.69 | 38 |
| Fusiform gyrus                                      | 37 | L |    |    |    |    |    |
| Cingulate gyrus                                     | 24/32| M |  6 |  2 | 40 | 4.35 | 33 |

Regions showing significant differences in activation based on presentation order for target recollection and recollection rejection. (L1) lure presented before target, (T1) target presented before lure, (BA) Brodmann’s Area, (H) hemisphere, (t) t-statistic, (k) cluster extent (number of voxels).

("Remember" hits where the target was presented first), L1 Target Recollection ("Remember" hits where the lure was presented first), T1 Recollection rejection ("Different" correct rejections where the target was presented first), and L1 Recollection rejection ("Different" correct rejections where the lure was presented first). Other responses were modeled separately as regressors of no interest.

To determine the effects of lure presentation on target recollection, we contrasted target recollection responses following lure presentation with those occurring before (target recollection with lure interference) and the reverse (target recollection without interference). To determine effects of target presentation on recollection rejection, we contrasted recollection rejection responses following target presentation with those occurring before (recollection rejection with target facilitation) and the reverse (recollection rejection without target facilitation). All contrasts were computed within a whole-brain gray matter mask defined by the Wake Forest University AAL Pickatlas. To determine a cluster threshold corrected for multiple comparisons at \( P < 0.05 \), 10,000 Monte Carlo simulations were implemented by 3dClustSim in AFNI (Cox and Hyde 1997) with search space (gray matter), average smoothness in millimeters estimated from residuals \((x, y, z = 11.98, 11.73, 10.1)\), no resampling, and the uncorrected \( p \) threshold \((P < 0.001)\) as inputs, resulting in a voxel extent of 30. Full neuroimaging results including peak coordinates are presented in Table 1.

To identify presentation order effects on recollection-based responses, we first computed a 2 (stimulus: target, lure) × 2 (order: T1, L1) repeated-measures ANOVA on “Remember” responses to targets and “Different” to responses to lures. There was no main effect of stimulus \((F_{1,20} = 0.01, P > 0.9, \eta^2 = 0.001)\), but a significant effect of presentation order \((F_{1,20} = 94.62, P < 0.001, \eta^2 = 0.83)\) with lower recollection rates when the lure was presented first \((M = 0.51, SD = 0.14)\) than when the target was presented first \((M = 0.71, SD = 0.13)\). The interaction was not significant \((F_{1,20} = 1.51, P > 0.2, \eta^2 = 0.07)\). We also computed overall d-primes separately for the T1 and L1 conditions and found significantly higher accuracy in the T1 \((M = 2.11, SD = 0.87)\) compared with the L1 condition \((M = 1.28, SD = 0.55; t_{20} = 5.73, P < 0.001)\). Thus, presenting the target before the lure boosted recollection-based responding for both targets and lures compared with presenting the lure first, which also affected overall memory accuracy (see also Supplemental Materials for an analysis of lure responses contingent on the response to the target). These effects are consistent with previous findings showing that priming subjects with the target just before its corresponding lure greatly reduces false recognition (Brainerd et al. 1995), indicating that target presentation facilitates lure rejection. We also found that responses to targets were affected by lure presentation—presenting the lure before the target reduced estimates of target recollection compared with when the target was encountered prior to the lure. Thus, our behavioral results show evidence of both target facilitation and lure interference.

We also tested whether the effects of interference and facilitation were specific to recollection-based responses or instead applied to correct familiarity-based responses as well. We computed a repeated-measures ANOVA on adjusted familiar/unfamiliar rates\(^5\) (i.e., “Familiar” targets and “Unfamiliar” lures). Results revealed a significant main effect of stimulus \((F_{1,20} = 7.35, P < 0.05, \eta^2 = 0.27)\) such that targets were associated with a greater proportion of familiar responses \((M = 0.49, SD = 0.19)\) than lures were with unfamiliar responses \((M = 0.32, SD = 0.19)\). More importantly, the main effect of order was not significant \((F_{1,20} = 0.16, P > 0.6, \eta^2 = 0.008)\) nor was the interaction \((F_{1,20} = 0.36, P > 0.56, \eta^2 = 0.02)\). Thus, the order of target/lure presentation affected recollection-based hits and correct rejections but no other correct responses, indicating that recollection signals are particularly sensitive to reinforcement from targets and interference from lures.

Neuroimaging results revealed that target recollection was associated with greater activation in right middle frontal gyrus, right inferior parietal cortex, and bilateral visual cortices for target recollection following lure interference compared with those occurring prior to lure presentation \((L1 “remember” hit > T1 “remember” hit)\) (see Fig. 2A). Thus, these regions support successful target recollection when interfering information has been presented, altering or disrupting target representations or access to target representations. Such activation may represent, in part, retrieval monitoring that accompanies processing of weaker memories (Henson et al. 1999, 2000). Further, recent work has shown that presenting

\(^5\)Because the rates of “familiar” responses to targets and “unfamiliar” responses to lures are not independent of their respective recollection-based responses, we calculated adjusted rates that correct for this dependence and assess familiarity effects controlling for differences in recollection-based responding. Adjusted familiar hits were calculated as \( p(\text{"familiar" hit})/(1 - p(\text{"remember" hit})) \), and adjusted unfamiliar rejections were calculated as \( p(\text{"unfamiliar" correct rejection})/(1 - p(\text{"different" correct rejection})) \) (Tonelinas and Jacoby 1995; Duarte et al. 2010). These adjusted rates were computed separately for the target first and lure first conditions.
times compared with disrupted target recollection following lure interference, suggesting that it supports strong recollections based on fluently retrieved details from encoding. Taken together, results show that activation in components of the larger target recollection network are differentially modulated by task demands, with some regions showing increased activation to negotiate interference and other regions showing reduced activation following interference.

For recollection rejection, results revealed increased activation in the right angular gyrus following target facilitation compared with recollection rejection occurring prior to target presentation (see Fig. 2B). The angular gyrus is often active during memory retrieval (Hutchinson et al. 2009; Spaniol et al. 2009), with activation that tracks the vividness of recollection (Vilberg and Rugg 2007), and activation patterns that can be used to decode retrieved content (Kuhl and Chun 2014). This region was also part of the recollection rejection network when presentation order was collapsed (Bowman and Dennis 2016), and its sensitivity to target-lure presentation order suggests that activity in right angular gyrus tracks the strength of target representations that are accessed during recollection rejection. Interestingly, previous work has often found left- rather than right-lateralized memory effects in angular gyrus (for review, see Hutchinson et al. 2009). While the effect reached threshold only on the right in the present study, comparison between activation in right and left angular gyrus revealed no significant difference in the overall pattern of activation across hemispheres (see Supplemental Materials).

Finally, a portion of left fusiform gyrus showed recollection rejection activation that was greater prior to target presentation than following target presentation. Activity in higher-order visual regions during retrieval is typically interpreted as reactivation of visual object information from encoding (Garoff et al. 2005). Thus, when target presentation is relatively remote (i.e., target last presented during encoding phase compared with elsewhere in retrieval), behavioral rates of recollection rejection are reduced, responses are slower, and there is more extensive processing of perceptual information due to its reduced accessibility. Taken together, results demonstrate that activation within the recollection rejection network is sensitive to the strength of target representations, which supports the idea that target representations are accessed as part of this recall-to-reject process. Overall, results demonstrate that presenting related information during retrieval modulates activation within both the target recollection and recollection rejection networks, indicating that portions of these networks are sensitive to the strength of target representations that can be either

Table 2. Behavioral response rates and reaction times

|                  | Mean (SD) rates       | Mean (SD) reaction times |
|------------------|-----------------------|--------------------------|
|                  | Target first        | Lure first       | Target first        | Lure first       |
| Targets          |                       |                         |                       |
| “Remember”       | 0.72 (0.14)         | 0.50 (0.17)            | 1165 (175)           | 1261 (220)       |
| “Familiar”       | 0.14 (0.09)         | 0.25 (0.15)            | 1974 (332)           | 1821 (246)       |
| “Different”      | 0.08 (0.06)         | 0.18 (0.09)            | 1837 (313)           | 1675 (311)       |
| “Unfamiliar”     | 0.05 (0.04)         | 0.05 (0.03)            | 1555 (374)           | 1870 (555)       |
| Lures            |                       |                         |                       |
| “Remember”       | 0.06 (0.07)         | 0.14 (0.10)            | 1545 (515)           | 1479 (329)       |
| “Familiar”       | 0.16 (0.15)         | 0.18 (0.12)            | 1882 (333)           | 1969 (409)       |
| “Different”      | 0.71 (0.17)         | 0.53 (0.17)            | 1459 (208)           | 1670 (229)       |
| “Unfamiliar”     | 0.07 (0.05)         | 0.16 (0.08)            | 1869 (662)           | 1603 (282)       |

Results showing proportion of each response for each stimulus type separated by presentation order as well as the corresponding reaction time in milliseconds. (SD) standard deviation.

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strengthened by re-presenting the target at retrieval or weakened by presenting other related information.

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