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Cognitive training-related changes in hippocampal activity associated with 1 recollection in older adults

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ABSTRACT

Impairments in the ability to recollect specific details of personally experienced events are one of the main 22 cognitive changes associated with aging. Cognitive training can improve older adults' recollection. However, 23 little is currently known regarding the neural correlates of these training-related changes in recollection. 24 Prior research suggests that the hippocampus plays a central role in supporting recollection in young and 25 older adults, and that age-related changes in hippocampal function may lead to age-related changes in rec- 26 ollection. The present study investigated whether cognitive training-related increases in older adults' recol- 27 lection are associated with changes in their hippocampal activity during memory retrieval. Older adults' 28 hippocampal activity during retrieval was examined before and after they were trained to use semantic 29 encoding strategies to intentionally encode words. Training-related changes in recollection were positively 30 correlated with training-related changes in activity for old words in the hippocampus bilaterally. Positive cor- 31 relations were also found between training-related changes in activity in prefrontal and left lateral temporal 32 regions associated with self-initiated semantic strategy use during encoding and training-related changes in 33 right hippocampal activity associated with recollection during retrieval. These results suggest that cognitive 34 training-related improvements in older adults' recollection can be supported by changes in their hippocam- 35 pal activity during retrieval. They also suggest that age differences in cognitive processes engaged during 36 encoding are a significant contributor to age differences in recollection during retrieval. 37

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Introduction Q343

The ability to accurately retrieve memories of personally experi-44 enced events is one of the cognitive skills most impaired by aging 45(for reviews see Balota et al., 2000; Jacoby and Rhodes, 2006; 46Kausler, 1994). Dual process models of memory retrieval propose 47 48 that past experiences can be remembered based on cognitivelycontrolled retrieval of an event (recollection) or recognized based 49 on a relatively automatic sense of familiarity in the absence of the 5051ability to retrieve the specific details of an event (familiarity)

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(Jacoby and Dallas, 1981; Mandler, 1980; Yonelinas, 1994). Research 52 comparing memory retrieval in young versus older adults has 53 shown that aging impairs recollection while leaving familiarity rela- 54 tively intact (Hay and Jacoby, 1999; Howard et al., 2006; Jacoby et 55 al., 2001; Perfect and Dasgupta, 1997; for a review see Yonelinas, 56 2002).

Research on the neural correlates of memory retrieval strongly 58 suggests that the hippocampus plays a critical role in supporting rec- 59 ollection. Hippocampal lesions in animals (Fortin et al., 2004; Sauvage 60 et al., 2008) and humans (Aggleton et al., 2005; Cipolotti et al., 2006; 61 Manns et al., 2003; Song et al., 2011; Wais et al., 2006) result in rec- 62 ollection impairments. In healthy young adults, positive correlations 63 have been found between hippocampal volumes and the ability to re- 64 member the spatial locations and temporal order of studied faces 65 (Rajah et al., 2010). Functional neuroimaging studies have also 66 reported hippocampal activity associated with recollection during re- 67 trieval in healthy young adults. For example, the hippocampus is 68 more active when young adults report recognizing words on the 69 basis of recollection as compared to familiarity (Eldridge et al., 70 2000; Wheeler and Buckner, 2004; Yonelinas et al., 2005). Studies 71 Q4 that have modeled recollection memory signals using nonlinear 72

Abbreviations: fMRI, functional Magnetic Resonance Imaging; BOLD, blood oxygen level-dependent; MPRAGE, magnetization prepared rapid gradient echo; ROI, region of interest.

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recognition memory confidence functions have also revealed hippocampal activity associated with recollection (Daselaar et al., 2006a,
2006b). In addition, greater hippocampal activity has been found during correct than incorrect contextual memory retrieval (Cansino et al.,
2002; Kahn et al., 2004; Ross and Slotnick, 2008).

Structural and functional neuroimaging studies suggest that age-78 79related changes in hippocampal function may play an important 80 role in age-related changes in recollection. Healthy aging is associated 81 with reductions in hippocampal volume (Jernigan et al., 2001; Rajah 82 et al., 2010; Raz et al., 2005). Structural equation modeling has 83 suggested that these reductions in hippocampal volume may mediate the relationship between age and recollection (Yonelinas et al., 2007). 84 During episodic memory retrieval, reduced activity in the left sub-85 86 iculum (a subregion of the hippocampal formation) has been found in older relative to young adults when older adults recollect fewer 87 studied words (Cabeza et al., 2004). When recollection memory is 88 modeled using a nonlinear recognition memory confidence function, 89 reduced recollection memory signals have also been reported in the 90 left hippocampus in older adults when they recollect fewer studied 91 words (Daselaar et al., 2006b). Alterations in hippocampal recollec-92tion memory signals as assessed by contextual memory retrieval ac-93 curacy measures have also been found in older adults. Specifically, 94 95Kukolja et al. (2009) reported that older adults had greater activity during incorrect than correct contextual memory retrieval in a left 96 anterior hippocampal region in which young adults had greater activ-97 ity during correct than incorrect contextual memory retrieval. In con-98 trast, Duverne et al. (2008) found that older adults had greater 99 100 activity during correct than incorrect contextual memory retrieval in a left posterior hippocampal region that did not show significant 101 differences in brain activity during correct versus incorrect contextual 102 memory retrieval in young adults. 103

104 Numerous studies have shown that cognitive training can improve 105older adults' memory (for reviews see Lustig et al., 2009; Rebok et al., 2007; Verhaeghen et al., 1992; Zehnder et al., 2009), including their 106 ability to recollect previously experienced events (Belleville et al., 107 2006; Hill et al., 1990; Jennings and Jacoby, 2003; Jennings et al., 108 2005; Kirchhoff et al., 2012; Yesavage et al., 1990). For example, in a 109 study that examined the effects of semantic encoding strategy train-110 ing on older adults' memory performance, self-initiated encoding 111 strategy use, and brain activity patterns during intentional encoding, 112 we found that they were less likely to consciously recollect intention-113 114 ally encoded words than young adults prior to training (Kirchhoff et al., 2012). Teaching older adults to use pleasantness, personal rele-115 vance, and sentence generation strategies to intentionally encode 116 words significantly improved their recollection such that it no longer 117 differed from that of young adults' after training. Semantic strategy 118 119 training also increased older adults' brain activity during encoding in prefrontal and left lateral temporal regions associated with seman-120 tic processing and self-initiated use of verbal encoding strategies 121 (Kirchhoff and Buckner, 2006; Kirchhoff et al., 2005; Petersen et al., 1221989; Poldrack et al., 1999; Vandenberghe et al., 1996). 123

124 To date, the relationship between cognitive training-related in-125creases in recollection and brain activity during memory retrieval in older adults has not been directly explored. Thus, little is currently 126known regarding the role that the hippocampus may play in 127supporting older adults' cognitive training-related improvements in 128129recollection. As noted above, prior lesion, structural neuroimaging, and functional neuroimaging research suggests that the hippocampus 130plays a central role in supporting recollection in both young and older 131 adults. Therefore, the hippocampus may also support cognitive 132training-related improvements in recollection in older adults. Two re-133 cent functional magnetic resonance imaging (fMRI) studies have 134reported that mnemonic training can increase older adults' hippo-135campal activity during memory retrieval (Belleville et al., 2011; 136 Hampstead et al., 2012), which is consistent with the prediction 137 138 that cognitive training-related changes in hippocampal activity during memory retrieval in older adults are associated with improve- 139 ments in recollection. However, the relationship between changes in 140 hippocampal activity and memory performance was not directly ex- 141 amined in these studies. Therefore, it is not clear whether cognitive 142 training-related changes in hippocampal activity supported improve- 143 ments in older adults' memory performance. 144

This paper explores whether cognitive-training related increases 145 in older adults' recollection are associated with changes in their hip-146 pocampal activity during memory retrieval. We investigated the rela- 147 tionship between training-related changes in recollection and 148 hippocampal activity during retrieval in older adults who were 149 trained to use pleasantness, personal relevance, and sentence gener- 150 ation strategies to intentionally encode words. We hypothesized 151 that training-related changes in recollection would be positively cor- 152 related with training-related changes in hippocampal activity during 153 retrieval, which would suggest that the hippocampus can support 154 cognitive training-related changes in recollection in older adults. 155 We also explored the relationship between training-related changes 156 in brain activity associated with self-initiated semantic strategy use 157 during encoding and training-related changes in hippocampal activi- 158 ty during retrieval. We hypothesized that there would be positive 159 correlations between training-related changes in brain activity during 160 encoding in prefrontal and left lateral temporal regions associated 161 with older adults' self-initiated semantic encoding strategy use and 162 training-related changes in hippocampal activity during retrieval. 163

Material and methods

Study design overview

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In this study older adults' brain activity during encoding and retrieval of single words, and their self-initiated encoding strategy 167 use, were assessed before and after two days of semantic encoding 168 strategy training (Fig. 1). The effects of training on memory performance, brain activity during encoding, and self-initiated encoding 170 strategy use have previously been reported for the participants in 171 this study (Kirchhoff et al., 2012). This paper focuses on the relationship between the effects of training on recollection and hippocampal activity during memory retrieval.

Participants

Behavioral and functional magnetic resonance imaging (fMRI) 176 data from fourteen older adults (mean age = 72.0, range 66–81; 177 mean years of education = 14.7, SD = 2.9; 7 Females/7 Males) are 178presented in this paper. Informed consent was provided in accor- 179 dance with Washington University's Human Studies Committee 180 guidelines. Participants were right-handed native English-speakers, 181 had normal or corrected-to-normal vision, reported no significant 182 neurological or psychiatric history, and were not taking psychiatric 183 medications or medications known to influence the blood oxygen 184 level-dependent (BOLD) hemodynamic response. The Short-Blessed 185 (Katzman et al., 1983) was administered to screen for dementia (all 186 participants had <six errors; mean errors = 0.8, SD = 1.1). Partici- 187 pants were also screened for glaucoma, significant heart disease, 188 untreated hypertension, diabetes, kidney disease, thyroid conditions, 189 active cancer, previous chemotherapy treatment, and alcoholism. 190 fMRI data during retrieval were not available for two older adults 191 who were included in our prior paper due to technical difficulties or 192 an inability to complete the retrieval fMRI scans due to fatigue. 193

Pretraining fMRI session

fMRI data acquisition

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A Siemens 3.0 Tesla Allegra scanner (Erlangen, Germany) was 196 used to collect structural and functional magnetic resonance imaging 197

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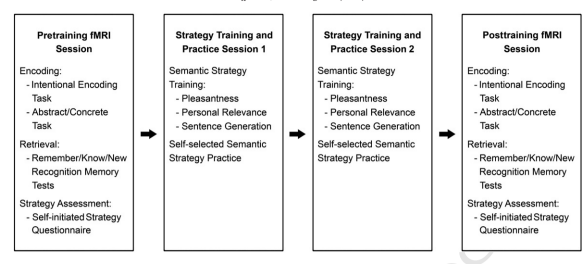


Fig. 1. Study design. During the pre- and posttraining fMRI sessions, older adults' brain activity was measured during encoding (intentional encoding and abstract/concrete tasks) and retrieval (Remember/Know/New task) of single words. Immediately after leaving the MRI scanner, their self-initiated encoding strategy use during intentional encoding was assessed using a strategy questionnaire. Older adults were taught to use three semantic encoding strategies to intentionally encode words (pleasantness, personal relevance, and sentence generation) during two strategy training sessions. They were given extensive practice using each of these strategies individually. In addition, they were given the opportunity to practice using whichever semantic encoding strategy, or combination of semantic encoding strategies, they felt worked best for them at the end of each strategy training session.

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data. An Apple Power Macintosh G4 computer (Apple, Cupertino, CA) 198 199 and PsyScope software (Cohen et al., 1993) controlled the stimulus display and recorded responses from a magnet-compatible fiber-200 optic keypress device interfaced with a PsyScope button box. Stimuli 201 202 were displayed on a screen at the head of the magnet bore with an LCD projector and were viewed using a mirror attached to the head 203 204 coil. Headphones dampened scanner noise and head movement was minimized using padding and tape. High-resolution structural images 205 $(1 \times 1 \times 1.2 \text{ mm})$ were acquired using a sagittal T1-weighted magneti-206 zation prepared rapid gradient echo (MPRAGE) sequence (TR = 2.3 s, 207 208 TE = 2.83 ms, flip angle = 9°, TI = 900 ms). Functional images were acquired using T2^{*}-weighted asymmetric spin-echo echo-planar se-209 quences sensitive to BOLD contrast. Four functional scans of 96 210whole-brain images $(324 \times 4 \times 4 \text{ mm contiguous axial slices acquired})$ 211 parallel to the AC-PC plane, TR = 2.5 s, TE = 25 ms, flip angle = 90°) 212213were collected per participant during encoding. Six functional scans of 88 whole-brain images (32 4×4×4 mm contiguous axial slices ac-214 quired parallel to the AC-PC plane, TR = 3.0 s, TE = 25 ms, flip 215216 angle $= 90^{\circ}$) were collected per participant during retrieval. The first four images in each scan were discarded to allow T1 magnetiza-217218tion to stabilize.

219 Stimuli

Stimuli for the fMRI sessions were four to seven letter English words, which were presented centrally in uppercase letters. Word lists were counterbalanced across tasks and scanning sessions. They were also matched for word frequency, length, and syllable count. Each word list consisted of half abstract (e.g., love, hope) and half concrete (e.g., table, flower) words.

226 Encoding

During the first two scans of the pretraining fMRI session, older 227 adults performed an intentional encoding task in which they were 228instructed to carefully study each presented word in anticipation of 229a later, unspecified memory test. They were also instructed to make 230a right-handed keypress whenever a word appeared to ensure that 231they were attending to the presented words. Older adults performed 232an abstract/concrete incidental encoding task during the third and 233 234 fourth scans of the pretraining fMRI session. They decided whether each word represented an abstract or a concrete entity, and made a 235 right-handed keypress to indicate their decision for each word. Dur-236 ing all encoding scans, three blocks of fixation plus signs (30 s) alter-237 nated with two blocks of words (70 s, 20 words per block) so that a 238 total of 80 words were presented during each encoding task. Ten ad-239 ditional seconds of fixation were acquired at the beginning of every 240 scan to allow T1 magnetization to stabilize. During word trials, a 241 word was presented for 3250 ms and was followed by a fixation 242 plus sign presented for 250 ms.

Retrieval

Immediately following the last encoding scan, older adults' memory 245 for the words studied during intentional encoding was assessed using 246 Remember/Know/New recognition memory decisions (Tulving, 1985) 247 during three fMRI scans (fMRI and behavioral data from three subse- 248 quent abstract/concrete task retrieval scans are not presented here). 249 The retrieval scans consisted of a total of 80 old words, 80 new words, 250 and 80 fixation plus sign trials presented in pseudorandom order so 251 that every trial type was equally likely to be preceded and followed by 252 every other trial type (Buckner et al., 1998). Four additional fixation 253 plus sign trials were presented at the beginning and end of each scan. 254 A plus sign was presented for 3000 ms during fixation trials. Individual 255 words were presented for 2775 ms, and were followed by fixation plus 256 signs presented for 225 ms, during old and new word trials. Participants 257 were instructed to make a Remember response if they recognized that a 258 word had been encountered during the encoding scans and were able to 259 consciously recollect aspects of its prior presentation. They were 260 instructed to make a Know response if they recognized that a word had 261 been encountered during the encoding scans but could not consciously 262 recollect aspects of its prior occurrence. In addition, they were instructed 263 to make a New response if they thought they had not seen the word dur- 264 ing the encoding scans. Although the relationship is imperfect, Remember 265 responses are a measure of conscious recollection while Know responses 266 are a measure of a feeling of familiarity in the absence of recollection 267 (Yonelinas, 2002; Yonelinas and Jacoby, 1995). Participants' responses 268 were recorded using a magnet-compatible fiber-optic keypress device. 269 They used their left index finger to make a Remember response, their 270 right index finger to make a Know response, and their right middle finger 271 to make a New response. 272

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273 Self-initiated encoding strategy questionnaire

274 Immediately after leaving the MRI scanner, older adults completed a self-initiated encoding strategy questionnaire. They rated how often 275276they used twenty-four possible encoding strategies during the intentional encoding scans, including the pleasantness ("Thought about 277whether each word was pleasant or unpleasant"), personal relevance 278("Thought about the personal relevance of each word"), and sentence 279generation ("Constructed phrases, sentences, and/or stories that con-280281 tained one studied word") strategies that they were trained to use in 282 this study, and how often they used no encoding strategy ("Read each 283word but did not use any particular strategy to try to remember the words"). They rated the frequency of their use of these strategies 284using a scale of never, rarely, sometimes, usually, or always. These 285286ratings were converted into numerical values for statistical analyses (1 = never, 2 = rarely, 3 = sometimes, 4 = usually, and 5 = always).287

288 Semantic encoding strategy training

289 Strategies trained

Older adults completed two semantic encoding strategy training 290sessions on separate days after the pretraining neuroimaging session. 291 During the first strategy training session, participants were taught to 292293 use pleasantness, personal relevance, and sentence generation strate-294 gies to intentionally encode lists of words. Specifically, they were taught to decide whether each presented word was pleasant or 295unpleasant and to think about why they felt that way (pleasantness 296strategy), to think about how each word was personally relevant to 297298them (personal relevance strategy), and to form a sentence that contained each presented word (sentence generation strategy). After 299participants were given extensive practice using each of these seman-300 301 tic encoding strategies, they were instructed to study additional word 302 lists using whichever semantic encoding strategy, or combination of 303 semantic encoding strategies, they felt worked best for them. During 304 the second session, they practiced using each of the semantic encoding strategies on multiple word lists and then were again 305 allowed to choose whichever semantic encoding strategy or strate-306 gies they wanted to use to study additional word lists. We allowed 307 308 older adults to choose which semantic encoding strategy or strategies to practice at the end of the cognitive training sessions instead of 309 training them to use just one semantic encoding strategy throughout 310 training because we thought they would be most likely to initiate 311 312 self-selected strategies during the posttraining fMRI session.

313 Stimuli

Stimuli for these training sessions were four to seven letter English words, which were presented centrally in uppercase letters. Word lists were matched for word frequency, length, and syllable count, and consisted of half abstract and half concrete words.

318 Encoding word lists

Older adults practiced using semantic encoding strategies on several lists of words during the strategy training sessions. Each word on these encoding word lists was presented for 3225 ms, and was followed by a 225 ms blank screen interstimulus interval (ISI). Difficulty was gradually increased throughout training by increasing the number of words on these lists (first list: 18, last list: 144).

325 Retrieval word lists and performance feedback

Immediately following each encoding word list, older adults were 326 shown a retrieval word list. Half of the words on these lists were 327 words that had just been studied during the encoding list that pre-328ceded it, and half were new (old/new status was counterbalanced 329across participants). Participants made a Remember/Know/New deci-330 sion for each word, and indicated the outcome of their decisions by 331 making keypresses on a computer keyboard. Each word was pres-332 333 ented for up to 2775 ms, and was followed by a 225 ms blank screen ISI. Immediately after a response was made, or the response window 334 was exhausted, older adults were given visual feedback for 1500 ms 335 on the accuracy of their response ("Correct": Remember or Know re- 336 sponse to an old word or a New response to a new word, "Miss": New 337 response to an old word, "False Alarm": Remember or Know response 338 to a new word, or "Please respond faster": no response to a word). 339 They also received feedback on their performance at the end of 340 every recognition word list. An accuracy summary screen appeared, 341 which informed each older adult of the percentage of the time he/ 342 she was correct when he/she made Remember, Know, and New re- 343 sponses, what percentage of the time he/she did not make a response 344 to a word on time, and what percentage of the time he/she made a 345 correct response for the word list overall. Difficulty was gradually in- 346 creased throughout training by increasing the number of words 347 shown during these retrieval word lists (first list: 36, last list: 288). 348

Posttraining fMRI session

Older adults completed the posttraining fMRI scanning session the 350 day after the second strategy training session. The posttraining scan-351 ning session occurred approximately two weeks after the pretraining 352 scanning session (range 8–18 days). Both scanning sessions used 353 identical structural and functional scanning parameters and encoding 354 and retrieval task designs. Participants were not explicitly told to use 355 the strategies that they learned in the training sessions during the 356 posttraining fMRI session. 357

Behavioral analyses

The effects of training on overall recognition memory perfor- 359 mance, Remember Hits (Remember responses to old words), Know 360 Hits (Know responses to old words), Misses (New responses to old 361 words), and reaction times for keypress responses during retrieval 362 were examined using paired sample *t*-tests. Overall recognition 363 memory performance was measured by subtracting the proportion 364 of Remember and Know False Alarms (Remember and Know re- 365 sponses to new words) from the proportion of Remember and 366 Know Hits. 367

fMRI data analysis

fMRI data preprocessing included adjustment for slice timing dif- 369 ferences using ideal sinc interpolation, correction for odd-even slice 370 intensity differences, mode normalization, and motion-correction 371 using a rigid-body rotation and translation correction. fMRI data 372 were resliced into 3 mm isotropic voxels and transformed into the 373 stereotaxic atlas space of Talairach and Tournoux (1988) using a tem- 374 plate constructed from 16 young and 16 older adult T1-weighted 375 MPRAGE scans acquired on the MRI scanner used in this study 376 (Snyder et al., 2002). The general linear model implemented in an 377 in-house analysis and display package was used to analyze functional 378 data (Miezin et al., 2000). Brain activity during retrieval trials was 379 modeled as an extended gamma function (Boynton et al., 1996) and 380 scaled to percent signal change. Run mean and slope were coded as 381 effects of no interest. Z-transformed reaction times for responses to 382 old words were included as covariates to control for training-related 383 changes in reaction times. Data were smoothed using a two-voxel iso-384 tropic Gaussian filter. 385

Hypothesis-driven analyses of the relationship between training-related 386 changes in recollection and hippocampal activity during memory 387 retrieval 388

The relationship between older adults' training-related changes in 389 recollection and hippocampal activity during memory retrieval was 390 first examined by conducting Pearson Product Moment correlation 391

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analyses between training-related changes (posttraining minus pre-392 393 training) in the proportion of Remember Hits and training-related changes in activity for all 80 old words studied during intentional 394 395 encoding in each voxel of a bilateral hippocampal anatomical mask derived from prior work (Wang et al., 2008). Resulting r statistics 396 were converted to z statistics and plotted over the combined young/ 397 old anatomical image. The activation map was corrected for multiple 398 comparisons using a combined p value/cluster size threshold of 399 400 p < 0.025/26 voxels, which corresponded to a two-tailed false positive rate of *p*<0.05 for the whole anatomical mask. This threshold/cluster-401 402 size requirement provides protection against type I error (Forman et al., 1995; McAvoy et al., 2001) and was chosen based on Monte-403404 Carlo simulations via AlphaSim (Ward, 2000). Activity during all 80 405old word trials was used as the measure of hippocampal activity in fMRI analyses so that training-related changes in the number trials 406 used to estimate hippocampal activity would not confound analyses 407of the relationships between training-related changes in hippocampal 408 activity and memory performance. 409

Voxel-based Pearson Product Moment correlation analyses be-410 tween training-related changes in the proportion of Know Hits and 411 activity for all 80 old words studied during intentional encoding 412 were also conducted within the hypothesis-driven hippocampal ana-413 414 tomical region of interest (ROI) described above using the same p 415 value/cluster size threshold. The goal of this analysis was to examine whether training-related changes in hippocampal activity during re-416 trieval were selectively associated with training-related changes in 417 recollection, or whether they were also driven by training-related 418 419 changes in familiarity.

420 *Exploratory analysis of the relationship between training-related* 421 *changes in recollection and brain activity during memory retrieval*

422 To further examine the relationship between training-related 423 changes in recollection and hippocampal activity, and to investigate whether training-related changes in recollection were associated 424 with training-related changes in activity in regions beyond the hippo-425campus, a whole-brain exploratory analysis of the relationship be-426 427tween older adults' training-related changes in recollection and brain activity during retrieval was conducted. In this analysis, Pearson 428 Product Moment correlations were calculated between training-429related changes in the proportion of Remember Hits and brain activ-430ity in response to old words in each voxel of the brain. Resulting r sta-431 tistics were converted to z statistics and plotted over the combined 432 young/old anatomical image. The statistical significance threshold 433 for the functional activation map was set to p < 0.01 with a minimum 434 voxel size of 5 voxels, uncorrected for multiple comparisons. An auto-435436 mated algorithm identified activation peaks in the functional activation map. ROIs were then created that included all continuous 437 voxels within 12 mm of an activation peak, inclusively masked by 438 the functional activation map. 439

Analyses of the relationship between training-related changes in
 prefrontal and left lateral temporal activity associated with semantic
 strategy use during encoding and hippocampal activity during retrieval

In our prior paper from this dataset that examined the effects of 443 444 semantic strategy training on older adults' brain activity during encoding (Kirchhoff et al., 2012), we identified several prefrontal 445 and left lateral temporal regions previously associated with semantic 446 processing and/or self-initiated verbal encoding strategy use that had 447 strong positive correlations between training-related changes in ac-448 tivity and older adults' memory performance. This suggests that 449 these regions support self-initiated semantic encoding strategy use 450in older adults. However, we did not find strong correlations between 451training-related changes in self-initiated use of the trained semantic 452453 encoding strategies and training-related changes in brain activity in these regions, most likely because participants were allowed to prac- 454 tice whichever encoding strategy or strategies they felt worked best 455 for them at the end of the training sessions. Therefore, to explore 456 whether semantic encoding strategy training altered older adults' 457 hippocampal activity during retrieval by increasing their self- 458 initiated use of semantic strategies during encoding, we examined 459 the relationship between training-related changes in brain activity 460 during encoding in the prefrontal and left lateral temporal regions as- 461 sociated with older adults' self-initiated semantic encoding strategy 462 use (medial superior frontal (BA 6), left middle frontal/precentral 463 (BA 6), left dorsal posterior inferior frontal (6/44/9), left ventral pos- 464 terior inferior frontal (BA 44), left anterior inferior frontal (BA 45), 465 and left middle/superior temporal (BA 21/22)) and training-related 466 changes in brain activity during retrieval in hippocampal ROIs identi- 467 fied from our hypothesis-driven and exploratory analyses using Pear- 468 son Product Moment correlation analyses (p<.05, one-tailed). 469

Results

Semantic encoding strategy training improved older adults' ability to 471

recollect intentionally encoded words 472

Memory retrieval accuracy data for the fourteen older adults in 473 this study who had analyzable brain activity data during retrieval 474 are presented in Table 1 (see Supplementary Table 1 for reaction 475 time data). Training improved older adults' recognition memory 476 overall (t(13) = 5.53, p < 0.001, d = 1.00). It also increased their Re- 477 member Hits (t(13) = 4.43, p < 0.01, d = 1.15), and decreased their 478 Know Hits (t(13) = -2.63, p < 0.05, d = -0.68) and Misses (t(13) = 4.43 suggest that semantic encoding strategy training selectively im- 481 proved recollection of intentionally encoded words in the older adults 482 whose hippocampal activity during retrieval is analyzed in this paper. 483 This result is consistent with the memory retrieval findings in the 484 larger older adult cohort in our prior paper (Kirchhoff et al., 2012).

Training-related changes in recollection were positively correlated with 486 training-related changes in hippocampal activity during memory 487 retrieval 488

Analysis of the relationship between older adults' training-related 489 changes in recollection and hippocampal activity during retrieval in 490 the hypothesis-driven anatomical ROI analysis revealed a positive 491 correlation between training-related changes in Remember Hits and 492 activity for old words in the left hippocampus (-38, -2, -11; 493 Fig. 2). Inspection of the scatterplot from this analysis reveals that 494 older adults with the largest training-related increases in recollection 495 had training-related increases in their activity during retrieval. How- 496 ever, due to the substantial variability in the degree to which older 497 adults benefited from training in this study, training did not signifi- 498 cantly change mean brain activity (t(13) = -0.05, p > 0.1, d = 0.00) 499 in this region. There were no clusters within either hippocampus 500 with a significant negative correlation between training-related 501 changes in Remember Hits and activity for old words.

The relationship between training-related changes in recollection 503 and activity for old words during retrieval throughout the brain was 504

Table 1

Overall recognition memory, Remember Hits, Know Hits, and Misses for older adults before and after semantic encoding strategy training.

	Recognition memory	Remember Hits	Know Hits	Misses	t1. t1.
0	.29 (.15)	.32 (.21)	.27 (.15)	.36 (.19)	t1.
	.49 (.24)****	.61 (.29)***	.15 (.20)**	.23 (.23)**	t1.

Note. Means and standard deviations (in parentheses) for overall recognition memory, Remember Hits, Know Hits, and Misses. Asterisks indicate a significant training effect (**p<0.05, ***p<0.01, ****p<0.001).

t1.6

t1.1

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also examined using a whole-brain exploratory correlation analysis. 505 506 Training-related changes in Remember Hits and activity for old words were positively correlated in the both the left (-22, -21, -21)507508-13) and right (27, -20, -15) hippocampus (Table 2, Fig. 3), further suggesting that the hippocampus supported training-related im-509provements in recollection in this study. However, training did not 510significantly alter mean brain activity in these regions overall (left: 511t(13) = 0.13, p > 0.1, d = 0.04; right: t(13) = -2.04, p < 0.07, d = -2.04512513-0.72), consistent with the results of the hypothesis-driven anatom-514ical ROI analysis. Training-related changes in Remember Hits and brain activity for old words were also positively correlated within 515several additional brain regions, including regions within prefrontal 516cortex (Table 2). There were no significant negative correlations be-517518tween training-related changes in Remember Hits and brain activity for old words. 519

520 Training-related changes in hippocampal activity during memory re-521 trieval were selectively associated with training-related changes in 522 recollection

To explore the specificity of the correlations between training-523related changes in Remember Hits and hippocampal activity, we ex-524525plored the relationships between training-related changes in Know Hits and activity for old words in the hippocampus. First, we con-526 ducted a voxel-based Pearson Product Moment correlation analyses 527within the hypothesis-driven hippocampal anatomical ROI, but did 528not find any significant correlations between training-related changes 529530in Know Hits and activity for old words. We also examined correlations between training-related changes in Know Hits and activity 531for old words in the left hippocampal ROI identified in the 532533hypothesis-driven Remember Hits analysis (Fig. 2), and the left and 534right hippocampal regions identified in the whole-brain exploratory 535Remember Hits analysis (Fig. 3). Training-related changes in Know Hits and brain activity were not significantly correlated in the left 536

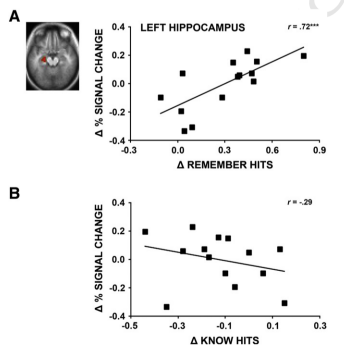


Fig. 2. Training-related changes in recollection and brain activity during memory retrieval were selectively positively correlated in a left hippocampal region identified in a hypothesis-driven anatomical ROI analysis. A) Training-related changes in recollection (as assessed by Remember Hits) and brain activity for old words were positively correlated in the left hippocampus (p<.05, corrected). B) In contrast, there was not a significant correlation between training-related changes in Know Hits and activity in this region. ***p<0.01.

Table 2

Regions with significant positive correlations between training-related changes in recollection and brain activity for old words during memory retrieval identified from a whole-brain exploratory analysis.

Region	BA	x y z (mm)	Ζ	Voxels
L hippocampus	-	-22 - 21 - 13	2.97	10
R hippocampus	-	27 - 20 - 15	3.87	7
L sup frontal	8	- 10 33 51	3.15	12
L mid frontal	46	- 32 46 19	2.81	7
R mid frontal	46	32 39 21	2.85	6
R mid/inf frontal	46	35 41 5	2.88	10
L inf frontal	47	-35 15 -13	3.48	10
R inf frontal	47	29 27 - 13	3.24	10
R precentral/insula	6	37 1 15	3.10	16
Ant cingulate	24/33	-1 22 19	3.04	21
Post cingulate	31	0 - 30 40	2.81	11
R sup temporal	41	34 - 28 16	3.41	13
R lingual	17	13 - 89 4	2.82	8
L cerebellum	-	-45 - 47 - 39	2.72	9

Note. BA = Brodmann's area; x y z (mm) = location of activation peak in Talairach coordinates; z = z score of peak voxel.

hypothesis-driven (r = -.29, p > 0.1) or exploratory (r = -.30, 537 p > 0.1) ROIs, but there was a trend toward a negative correlation in 538 the right hypothesis-driven ROI (r = -.48, p < 0.09). Taken together, 539 the pattern of correlations between training-related changes in Re- 540 member and Know Hits and hippocampal activity for old words sugsts that training-related increases in hippocampal activity reflected 542 increases in recollection but not familiarity. 543

Training-related changes in right hippocampal activity during memory544retrieval were positively correlated with training-related changes in pre-545frontal and left lateral temporal activity during encoding546

Training-related changes in activity in prefrontal and left lateral 547 temporal regions that support older adults' self-initiated semantic 548 strategy use during encoding were not significantly correlated with 549 training-related changes in activity in the left hippocampal region 550 during retrieval identified in the hypothesis-driven anatomical ROI 551 analysis (Table 3). However, there was a trend toward a significant 552 correlation between training-related changes in activity in the left 553 middle frontal/precentral gyrus (BA 6) and training-related changes 554 in activity in the left exploratory hippocampal ROI. Importantly, 555 training-related changes in activity during encoding in all of the re- 556 gions associated with self-initiated semantic encoding strategy use 557 were significantly positively correlated with training-related changes 558 in activity in the right exploratory hippocampal ROI except for the left 559 anterior inferior frontal region (BA 45), which had a trend toward a 560 significant positive correlation. These results suggest that training- 561 related changes in self-initiated semantic encoding strategy use con- 562 tributed to training-related changes in hippocampal activity associat- 563 ed with recollection during memory retrieval. 564

Discussion

This paper explored the effects of a semantic encoding strategy 566 training protocol that enhances older adults' recollection memory 567 on their hippocampal activity patterns during memory retrieval. 568 Training-related changes in Remember Hits were positively correlat- 569 ed with training-related changes in activity for old words in the hip- 570 pocampus bilaterally. Positive correlations were also found between 571 training-related changes in activity in prefrontal and left lateral tem- 572 poral regions associated with self-initiated semantic strategy use dur- 573 ing encoding and training-related changes in right hippocampal 574 activity associated with recollection during retrieval. The implications 575 of these results are discussed below. 576

t2.1

t2.18

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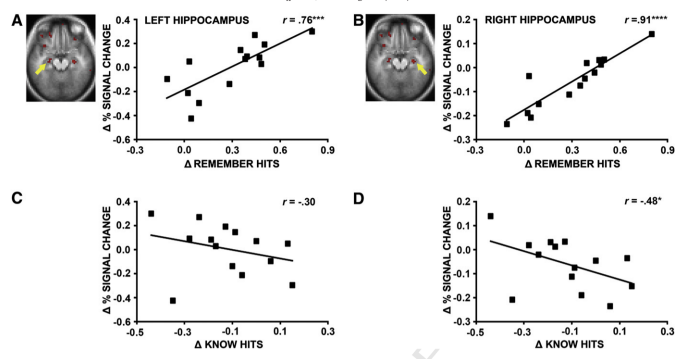


Fig. 3. A whole-brain exploratory analysis revealed selective positive correlations between training-related changes in recollection and brain activity during memory retrieval in the hippocampus bilaterally. Significant positive correlations between training-related changes in recollection (as assessed by Remember Hits) and brain activity for old words were found in the A) left and B) right hippocampus in a whole-brain exploratory correlation analysis (p<0.01, uncorrected). However, training-related changes in Know Hits and brain activity for old words were not significantly correlated in either the C) left or D) right hippocampal regions. *p<0.1, ***p<0.001, ****p<0.001.

577While training-related changes in hippocampal activity during re-578trieval were positively correlated with training-related changes in Re-579member Hits, they were not significantly correlated with changes in Know Hits. This suggests that training-related changes in hippocam-580pal activity associated with training-related changes in Remember 581Hits were not due to differences in scanner signal across scanning ses-582sions, task practice effects, etc., but instead reflected hippocampal 583support of training-related improvements in recollection. The lack 584of significant correlations between training-related changes in Know 585 Hits and hippocampal activity further suggest that training-related 586 changes in hippocampal activity were driven by training-related 587 changes in recollection and not familiarity. 588

The positive correlations between training-related changes in Remember Hits and activity for old words in the left and right hippocampus in this study are consistent with prior research suggesting that the hippocampus plays a central role in supporting recollection in both young and older adults (Aggleton et al., 2005; Cabeza et al., 2004; Cansino et al., 2002; Daselaar et al., 2006b; Dulas and Duarte,

t3.1 Table 3

Correlations between training-related changes in brain activity in prefrontal and left lateral temporal regions that support older adults' self-initiated semantic strategy use during encoding and training-related changes in brain activity in hippocampal regions associated with recollection during memory retrieval.

t3.2 t3.3	Region	Hypothesis-driven left hippocampal ROI	Exploratory left hippocampal ROI	Exploratory right hippocampal ROI
t3.4	Medial BA 6	.31	.30	.54**
t3.5	Left BA 6	.33	.37*	.49**
t3.6	Left BA 6/44/9	.32	.34	.52**
t3.7	Left BA 44	.29	.28	.57**
t3.8	Left BA 45	.27	.27	.46*
t3.9	Left BA 21/22	.24	.24	.55**

Note. ROI = region of interest.

t3.10 * p<0.1.

t3.12 ** p<0.05.

2011; Duverne et al., 2008; Eldridge et al., 2000; Rajah et al., 2010; 595 Yonelinas et al., 2007). It is also consistent with recent studies that 596 demonstrated that cognitive training can increase hippocampal activ-597 ity during memory retrieval (Belleville et al., 2011; Hampstead et al., 598 2012). Importantly, this study extends prior research by beginning to 599 shed light on the relationships between training-related changes in 600 cognitive processing and hippocampal activity and the mechanisms 601 of age-related changes in recollection memory. 602

The positive correlations between training-related changes in Re- 603 member Hits and hippocampal activity during retrieval suggest that 604 even though hippocampal volume is reduced in older adults 605 (Jernigan et al., 2001; Rajah et al., 2010; Raz et al., 2005), the hippo- 606 campus can still support cognitive training-related improvements in 607 their recollection memory. Interestingly, the significant correlations 608 between training-related changes in prefrontal and left lateral tem- 609 poral activity during encoding and right hippocampal activity during 610 retrieval suggest that age differences in cognitive processes engaged 611 during encoding are a significant contributor to age differences in rec- 612 ollection. These age differences in the cognitive processes engaged 613 during encoding may reflect age-related alterations in prefrontal 614 structure and function (for reviews see Gunning-Dixon et al., 2009; 615 Q5 Raz and Rodrigue, 2006). Importantly, the prefrontal and left lateral 616 temporal regions whose activity we found to be correlated with hip- 617 pocampal activity during retrieval are regions associated with self- 618 initiated semantic strategy use during encoding (Kirchhoff et al., 619 2012). Therefore, it is likely that the changes in hippocampal activity 620 in this study resulted from training increasing older adults' use of se- 621 mantic strategies during encoding (and hence increasing activity in 622 these prefrontal and lateral temporal regions), instead of training im- 623 proving the function of the hippocampus per se. Semantic encoding 624 strategy training may improve older adults' recollection by facilitat- 625 ing their ability to form distinctive memory traces during encoding 626 that contain detailed information about the studied words (e.g., what 627 encoding strategy they used to learn the word). During retrieval, these 628 elaborate memory traces could enhance older adults' ability to reinstate 629 contextual information from encoding (e.g., what encoding strategy(ies) 630

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they used) to constrain retrieval and facilitate recollection (i.e., source 631 632 constrained retrieval, Jacoby et al., 2005b; Shimizu and Jacoby, 2005). 633 Prior research has suggested that impairments in source constrained re-634 trieval play an important role in age-related changes in recollection (Jacoby et al., 2005a; Velanova et al., 2007). Multiple prefrontal regions 635 demonstrated positive correlations between training-related changes 636 in activity for old words during memory retrieval and training-related 637 changes in recollection in this study, which is consistent with the possi-638 639 bility that training enhanced older adults' source constrained retrieval.

An important question for future research is whether cognitive 640 641 training can improve older adults' hippocampal function by inducing beneficial changes in its structure and/or improving its processing ef-642 643 ficiency. A recent study demonstrating that aerobic exercise training 644 can increase older adults' hippocampal volumes, and that these volumetric increases are associated with improvements in spatial memo-645 ry, suggests that experiences that enhance older adults' memory can 646 improve older adults' hippocampal function (Erickson et al., 2011). 647

A limitation of this study is that training did not significantly in-648 crease older adults' mean activity in hippocampal regions associated 649 with recollection. This is likely due to the substantial variability in 650 older adults' training-related changes in recollection memory. Al-651 though this variability may have prevented us from finding signifi-652 653 cant training-related changes in mean hippocampal activity, it did allow us to find positive correlations between training-related 654 changes in recollection and hippocampal activity. Another limitation 655 of this study is that it did not include a no treatment control group 656 of older adults who were scanned twice. Therefore, we cannot 657 658 completely rule out the possibility that some of the changes that we observed in hippocampal activity between the pre- and posttraining 659 fMRI scans are due to task practice effects. However, the positive cor-660 relations between training-related changes in activity in prefrontal 661 662 and left lateral temporal regions associated with self-initiated semantic strategy use during encoding and training-related changes in right 663 664 hippocampal activity associated with recollection during retrieval suggest that semantic encoding strategy training was an important 665 contributor to changes in recollection and hippocampal activity dur-666 ing retrieval. Interestingly, hippocampal activity tended to decrease 667 668 following cognitive training in older adults who did not have a substantial improvement in recollection memory from training. Under-669 standing what drives these decreases in hippocampal activity in 670 older adults who do not benefit from semantic encoding strategy 671 672 training is an important topic for future research. In older adults with relatively large increases in recollection memory following 673 training, hippocampal activity tended to increase after training, par-674 ticularly within the left hippocampus. The results of this study sug-675 gest that these increases in hippocampal activity are driven by 676 677 increases in hippocampal dependent recollection.

Conclusions 678

In conclusion, the results of the data analyses conducted in this 679 680 paper suggest that the hippocampus can support cognitive training-681 related improvements in older adults' recollection memory. They also suggest that age differences in cognitive processing during encoding 682 may play an important role in age differences in recollection and hippo-683 campal activity during retrieval. Furthermore, this study demonstrates 684 685 that training-related changes in older adults' cognitive processing during encoding can contribute to training-related changes in recollection 686 and hippocampal activity during retrieval. Therefore, cognitive training 687 protocols that target age differences in cognitive processing during 688 encoding, such as age differences in self-initiated encoding strategy use, 689 may be an effective approach for reducing older adults' impairments in 690 recollection and age-related changes in hippocampal activity during 691 memory retrieval. 692

Supplementary data to this article can be found online at http:// 693 694 dx.doi.org/10.1016/j.neuroimage.2012.06.017.

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