1	Locally distributed abstraction of temporal distance in human parietal
2	cortex
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11	
12	The human brain parsimoniously situates past events by their order in relation
13	to time. Here, we show the posteromedial cortex geometrically abstracts the time
14	intervals separating pairs of event-moments in long-term, episodic memory.
15	Transcranial magnetic stimulation targeted at the precuneus erases these locally
16	distributed multivariate representations, and alters the correlation between
17	precuneal activity patterns and mnemonic judgements, revealing a critical role of
18	the precuneus in abstracting temporal distances during episodic memory
19	retrieval.
20	
21	Time in physics is operationally defined as "what a clock reads". While the
22	passage of time between two moments can be precisely measured by a quartz crystal
23	oscillator or biologically registered by distributed sets of brain regions across intervals

24	of time (Buhusi & Meck, 2005; Meck, Penney, & Pouthas, 2008), the neural
25	mechanisms that abstract such temporal distances separating events in long-term,
26	episodic memory is incompletely understood (Mauk & Buonomano, 2004).
27	Representations of brief elapsed time can be inferred from single neuron
28	activities in the primate brain (Jin, Fujii, & Graybiel, 2009; Leon & Shadlen, 2003).
29	Time-registering neurons are found to code time with high precision in the cortico-
30	basal ganglia circuits (Jin et al., 2009) and inferior parietal cortex (Leon & Shadlen,
31	2003) across short timescales. In contrast, when complex, coherent experiences
32	become consolidated into long-term memories (McGaugh, 2000), the neural circuits
33	that build time representations as an infrastructure for episodic retrieval are theorized
34	to be distinct from those implicated in hippocampal-dependent encoding (Ezzyat &
35	Davachi, 2014; Manns, Howard, & Eichenbaum, 2007) and retrieval (Hsieh, Gruber,
36	Jenkins, & Ranganath, 2014; Nielson, Smith, Sreekumar, Dennis, & Sederberg, 2015),
37	and from those during transient temporal processing (Jin et al., 2009; Leon &
38	Shadlen, 2003). For the recollection of long-term autobiographical memories or
39	episodic events, the posterior medial (PM) memory system plays an instrumental role
40	(Ranganath & Ritchey, 2012). However, the critical issue of how elapsed time
41	between pairs of long-term episodic events – and its interplay with the encoding
42	context – is represented by the PM system has yet to be addressed. Here we
43	investigated the abstraction, at a macro-anatomical level, of temporal distances that
44	were encoded more than 24 hours previously (Kwok, Shallice, & Macaluso, 2012; St
45	Jacques, Rubin, LaBar, & Cabeza, 2008), and determined how several members of

this large cortical system are differentially implicated in this putative mnemonic

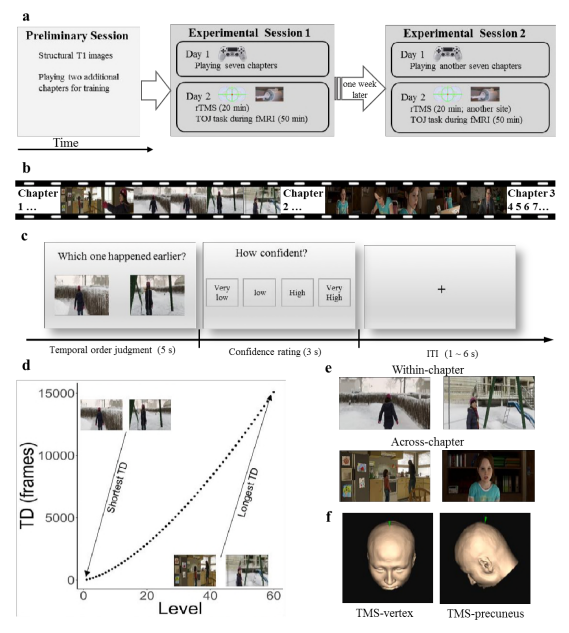
47 function (Richter, Cooper, Bays, & Simons, 2016).

48	Combining functional magnetic resonance imaging (fMRI) with an interactive-
49	video memory paradigm and a temporal order judgement task (TOJ; Fig. 1a)—a
50	validated paradigm to study neural correlates underpinning temporal distances
51	between units of memory traces (Kwok et al., 2012; Manns et al., 2007; St Jacques et
52	al., 2008)-we adopted a two-forked protocol to ascertain how temporal distances
53	separating pairs of past moments-in-time are represented in the human neocortex. We
54	first investigated the neural correlates of temporal distances with a multivariate
55	searchlight representational similarity analysis (RSA) (Nili et al., 2014) tuned to
56	identify a locally distributed neural representation using 9-mm radius spherical
57	searchlight. We parametrized a large set of pairs of event-moments geometrically
58	separated by varying temporal intervals and applied RSA to compare neural
59	representational dissimilarity matrices (RDM) with a number of parametric,
60	condition-rich hypothetical models. Applied across the entire brain, the searchlight
61	approach identifies local multi-voxel patterns driven by structured co-activation at a
62	voxel or sub-voxel level within the size of the searchlight, thereby giving us a
63	snapshot of the locally distributed neural architecture supporting temporal order
64	judgements. To enhance the causal strength of the anatomical associations thereby
65	revealed, we then focally disrupted the identified critical region with repetitive
66	transcranial magnetic stimulation (rTMS), seeking to confirm its functional necessity
67	for mediating the distributed representation of temporal distances. The spatial scale of

rTMS-induced disruption is comparable to that of our chosen searchlight, rendering it
an optimal tool for targeted, reversible disruption of the distributed representation of
interest.

For memory encoding, participants played an interactive video game containing 71 72 seven distinct yet related chapters, each in the range of tens of minutes on day 1 73 (Supplementary Fig. 1, Supplementary Table 1). By the nature of the video game, the within chapter segments contained more coherent narrative strands than those 74 across chapters, yet all chapters were connected by a common plot. After a 24-hour 75 76 retention period (day 2), on each trial, participants judged the temporal order of two images (extracted from their individually-played video game, Fig. 1b), depicting two 77 time-points in their encoded memory, while their blood-oxygen-level-dependent 78 79 (BOLD) activity was measured (TOJ task, Fig. 1c). Assuming a scale-free temporal memory representation(Kwok & Macaluso, 2015), we manipulated the between-80 81 images temporal distances (TD) for all pairs of images so that the TD distribution 82 adhered to a power function permitting scale-invariance across subjects (Gallistel &

#### 83 Gibbon, 2000) (60 levels of TD, **Fig. 1d**).



84 Figure 1. Experiment overview. (a) In experimental sessions 1 and 2, participants played a video game containing seven related chapters with a first-person perspective for encoding, 85 and 24 hours later, received 20 min of repetitive transcranial magnetic stimulation (rTMS) to 86 either one of two cortical sites before performing a temporal order judgement task during 87 fMRI. Order of TMS sites (within-subjects) and choices of video game chapters were 88 89 counterbalanced across subjects (Supplementary Table 1). The two experimental sessions 90 were conducted on different days to minimize rTMS carry-over effects (mean separation = 891 days). Participants underwent structural MRI scans and familiarized themselves with the gameplay using a console prior to experimental sessions proper. (b) Gameplay video: each 92 93 encoding session consisted of seven chapters (Supplementary Fig. 1). (c) Temporal order 94 judgement task. Participants chose the image that happened earlier in the video game and 95 reported their confidence level. (d) 60 levels of temporal distances (TD) were generated for each subject according to their subject-specific video-playing duration. Although the absolute 96

97 TD were different across subjects (Supplementary Fig. 1), we ensured it to be scale-invariant using a power function during image selection. Actual TDs from one subject (subj01) are 98 99 shown. (e) Two pairs of images were extracted from the same chapter (Within-chapter) or two adjacent chapters (Across-chapter). The 60 levels of TD were fully matched within-subjects 100 101 for these two conditions. Note that scenes depicted in Within-chapter tended to be more 102 contextually similar than those depicted in Across-chapter. (f) Stimulation sites, superimposed onto one subject's MRI-reconstructed skull, are marked by a green pointer. The MNI 103 coordinates for precuneus stimulation: x, y, z = 6, -70, 44. 104 105 We first searched for neural representations resembling the matrix of temporal distances using searchlight representational similarity analysis (Nili et al., 2014). 106 Without *a priori* bias for any region of interest, we searched the entirety of the cortex 107 using a RDM consisting of 60 levels of logarithmically-transformed subject-specific 108 109 TD, and identified voxels that contain information of the set of geometrically defined temporal distances in memory (Fig. 2a, Online Methods and Supplementary Fig. 110 2). Within this  $60 \times 60$  matrix of moment-wise distances, we revealed that the neural 111 112 pattern of judging the temporal order of a pair of memories separated with a given temporal distance is more similar to other temporal order judgements which enclosed 113 temporal distances of a comparable scale. These voxels were in the posteromedial 114 parietal areas, bilateral angular gyri, and middle frontal gyri (Fig. 2b, Supplementary 115 Table 2). The scale-invariance of the design allowed us to generalize the resultant 116 mnemonic abstraction of temporal distances across individuals, in line with the scale-117 free representation of time (Gallistel & Gibbon, 2000). 118 The temporal-distance memory representation could be confounded by perceptual 119 similarity in each pair of images. To address this concern, we conducted three 120 separate RSAs, in which we indexed perceptual similarity between the pairs by three 121 different metrics: RGB cross-correlation RDM, RGB-intensity RDM, and RGB-122

123	histogram RDM. No similar representation was observed in the posteromedial parietal
124	cortex using these candidate RDMs, suggesting that our results could not be driven
125	simply by the image properties (rows $3 - 5$ , <b>Supplementary Fig. 3</b> ). To reinforce the
126	point, considering previous work on space-time relationships in memory (Deuker,
127	Bellmund, Navarro Schroder, & Doeller, 2016; Nielson et al., 2015), we also
128	computed the number of locations each participant had virtually traversed in the
129	video, quantified the space displacement embedded between the paired images, and
130	entered them into subject-specific Situational Changes RDM (Supplementary Fig.
131	4). The searchlight RSA results showed the pattern representation observed in Figure
132	<b>2b</b> can be better accounted for by temporal-distance embedded between the images
133	than by spatial-distance that participants had travelled (rows 2, Supplementary Fig.
134	<b>3</b> ). Inferential statistical comparisons of multiple models showed that the neural
135	signals in the selected conjunction ROI (see below) were most attributable to the
136	mnemonic representation of the moment-wise comparison of multiple temporal
137	distances (Supplementary Fig. 5).
138	Having identified a multivariate pattern underlying the temporal memory
139	abstraction in the posterior medial parietal cortex (Fig. 2b), we probed further
140	whether there were voxels whose activities change monotonically as a function of
141	temporal distance, irrespective of any locally distributed representation. Whole-brain
142	parametric modulation analysis (pmod) revealed TD-specific BOLD signals in a
143	cluster within the posteromedial region, including the precuneus (Fig. 2c,
144	Supplementary Table 3). This relationship could not be attributed to difficulty

- 145 (results were same after trial-by-trial reaction times were accounted for,
- 146 Supplementary Fig. 6a). Since the two types of analysis extracted two different
- 147 kinds of neurally-encoded information, their overlap in the precuneus strengthens the
- 148 claim to a critical role for the region. We accordingly created a conjunction map (Fig.
- 149 **2d**), so that both the multivariate and univariate results underlying the temporal
- 150 distance abstraction would be available for the next analysis.

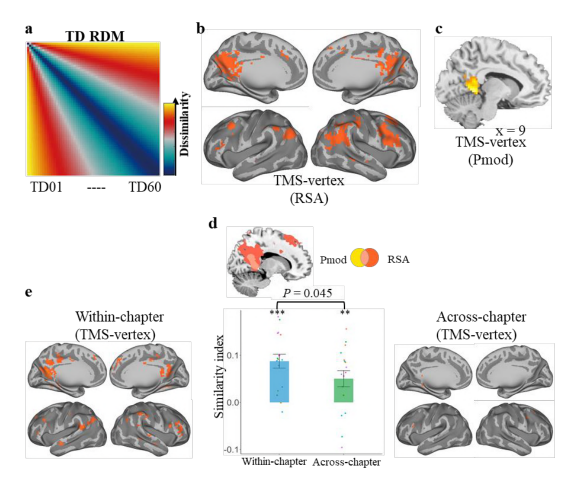




Figure 2. Abstraction of temporal distances in posteromedial cortex. (a) TD representational 152 dissimilarity matrix (RDM) for searchlight RSA. The RDM consisted of 60 subject-specific 153 TD levels. Any two event-moments that are separated by short TD will get increasingly 154 dissimilar with other two event-moments as the TD increases. (b) Using TD RDM for 155 156 searchlight RSA, clusters of voxels that contained TD information were primarily in the 157 posteromedial cortex, bilateral angular gyri and bilateral middle frontal gyri. (c) Activation 158 signal intensity from parametric modulation analysis (pmod). The intensity of these voxels, primarily in the left precuneus, increased as a function of TD. (d) Conjunction mask. The 159 mask was created by intersecting the activation map from pmod and similarity map from RSA 160

161 162 163 164 165	(Supplementary Tables 2 and 3). (e) Stronger multivoxel similarity representing TD variation in Within-chapter condition than in Across-chapter condition in the conjunction region (one-side: $P = 0.045$ , ** $P < 0.01$ ; *** $P < 0.001$ ). Results were consistent when the masks were created using more robust, bias-free leave-one-subject-out method (one-side: $P = 0.040$ ). MRI results are displayed at $P_{\text{uncorrected}} < 0.001$ .
166	Temporal distance or proximity is intertwined with the construct of context. A
167	prominent memory model posits that item representations are linked to a changing
168	"context" at encoding, such that a common retrieved context is triggered during recall
169	for items that were experienced within a similar temporal context (Polyn, Norman, &
170	Kahana, 2009). This theory predicts the TD-neural representation pattern similarity
171	index to be higher when the two event-moment images are extracted from a "similar
172	context" than when they are from two "different contexts". To test this, we
173	manipulated the factor "context" by controlling whether the paired images presented
174	at TOJ task were extracted from the same chapters or two adjacent chapters of the
175	video game while keeping the 60 TDs fully matched between the two conditions
176	(Within-chapter vs. Across-chapter, Fig. 1e). We re-ran the searchlight RSA, now
177	separately for the Within-chapter and Across-chapter trials. The representation of TD
178	was observed only in the Within-chapter condition but not in the Across-chapter
179	condition (Fig. 2e, right). The identified voxels were in the precuneus, retrosplenial
180	cortex, and angular gyri bilaterally (Fig. 2e left). For statistical inference, we
181	extracted the similarity index with the conjunction mask combining the RSA and
182	pmod maps which contained the TD-modulated signals from each subject for
183	comparison (Fig. 2d and Online Methods). In line with our prediction, the voxels in
184	the Within-chapter trials contained higher pattern similarity to the TD RDM than
185	Across-chapter trials ( <b>Fig. 2e</b> , middle panel; one-side: $P = 0.045$ ), confirming the

186	neural pattern similarity related to the TD RDM was indeed stronger in Within-
187	chapter trials. This difference was also found in a voxel-wise univariate analysis. The
188	beta-estimates ( $\beta$ ) from pmod analyses using TD as a regressor were significantly
189	higher in the Within-chapter condition ("Within-chapter > Across-chapter" $P = 0.005$
190	with RT effects regressed out, Supplementary Fig. 6c-d). This confirmed the
191	mnemonic representation of temporal distances was determined by whether the pairs
192	of images were experienced within a similar context, corroborating the interaction
193	between temporally- and semantically-defined factors observed during memory
194	encoding (Ezzyat & Davachi, 2014) and retrieval (Hsieh et al., 2014).
195	To strengthen the claim to a pivotal role of the precuneus in this operation we
196	strategically deployed a disruptive technique, targeting it with repetitive transcranial
197	magnetic stimulation to interrogate changes on both neural and behavioral levels
198	(within-subjects: TMS-precuneus vs. TMS-vertex; Fig. 1f and Supplementary Table
199	1). Strikingly, in the fMRI session of which subjects' precuneus had received TMS
200	stimulation immediately prior to TOJ, the widespread representation of TD was
201	eradicated (rows 1, Supplementary Fig. 3). To the best of our knowledge, no study
202	has previously demonstrated such sharp susceptibility to TMS of such mnemonic
203	representations. The multivariate representations were more vulnerable than the
204	conventional, activation-based analyses: TMS to the precuneus did not induce any
205	discernable changes in the univariate BOLD intensity (Supplementary Fig. 7).
206	Altogether, these findings strengthen our argument that memory traces that are
207	represented during temporal order judgement are indeed conveyed in some localized

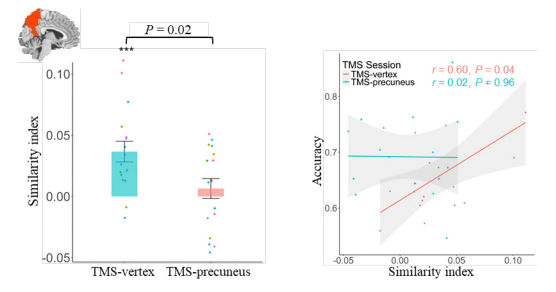
208 multivoxel readouts housed in the PM system cortices, above and beyond the

209 modulated changes in canonical BOLD activation.

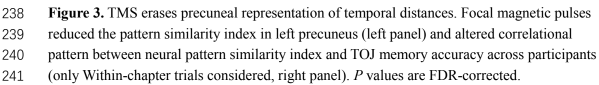
210	In light of the fractionation view for the parietal cortex (Nelson, McDermott, &
211	Petersen, 2012), we further tested the hypothesis that there might be differences in the
212	patterns of neural activity associated with the abstraction of temporal distances in the
213	sub-regions of the PM memory network (Ranganath & Ritchey, 2012). Based on our
214	results (Fig. 2b and Supplementary Table 2) and previous work on the parcellation
215	of the PM memory network(Richter et al., 2016), we chose six anatomical regions-of-
216	interest in the PM memory network (ROIs: bilateral precuneus, bilateral angular gyrus
217	and bilateral hippocampus; see Online Methods), together with the primary visual
218	region (entire occipital cortex) as a control, to test for the disruptive effect caused by
219	the TMS. We extracted the similarity indices from these ROIs and found that the
220	neural-TD pattern similarity was significantly weakened in the left precuneus
221	following TMS to the precuneus (Fig. 3; and to a lesser extent also the left angular
222	gyrus, Supplementary Fig. 8). Such differences were not obtained in the other ROIs
223	(Supplementary Fig. 8). Moreover, we found that changes in individuals' neural-TD
224	pattern similarity in the vertex condition to be associated positively with their TOJ
225	memory performance in this key region (also in the hippocampi, see Supplementary
226	Fig. 8), implying the multivoxel representations are important neurobiological
227	prerequisites for the ability to support temporal order judgement. However, disrupting
228	the precuneus with magnetic field prior to retrieval put this neural-behavioral
229	correlation into disarray (Fig. 3) and slowed response times (Supplementary Fig. 9),

indicating the removal of the TD representations from the precuneal voxels would
causally result in overt memory performance changes. Since the focal perturbation
altered the mnemonic representation in the precuneus, the angular gyri and the
hippocampi, it implied the disruption might have been effective through inducing
alternation in functional connectivity between multiple regions, or more globally
throughout the entire parietal memory network (Nilakantan, Bridge, Gagnon,

236 VanHaerents, & Voss, 2017; Wang et al., 2014).







242 The present parietal representation of temporal distances between pairs of

episodic events, as revealed by both univariate and multivariate pattern analyses,

244 might act in parallel with hippocampal cells that code specific moments in time or

- temporal positions (Eichenbaum, 2014), or act independently as a separate mnemonic
- establishment of episodes over and above the hippocampal memory ensemble (Brodt
- et al., 2016). The current findings align with the temporal context model (Polyn et al.,

248	2009) that the fine-grained TD memory information distributed in this cortex is
249	comprehensively stronger when paired images were associated within the similar
250	context. Building on extant connectivity findings between the hippocampus and
251	neocortical regions (Moscovitch, Cabeza, Winocur, & Nadel, 2016; Ranganath &
252	Ritchey, 2012; Vincent et al., 2006) and the hippocampal role in temporal context
253	memory (Ezzyat & Davachi, 2014; Hsieh et al., 2014), our demonstration of
254	distributed pattern of temporal information in the posteromedial parietal region
255	implied the existence of a higher level parietal mnemonic readout of temporal
256	distances between episodic experiences.
257	In summary, our multivariate searchlight results reveal that the temporal distance
258	representations in the posterior parietal cortex, especially the precuneus, during TOJ
259	retrieval are determined by how distant (and how similar the encoding contexts) two
260	given event-moments the subjects had encountered (Kwok et al., 2012; St Jacques et
261	al., 2008). We also establish that this multivoxel mnemonic abstraction is localized in
262	the precuneal area and perturbation to it alters the neural-behavior relationship
263	across the global parietal memory network, assigning this structure as a locus of
264	flexibly effecting the manipulation of physical time during episodic memory retrieval.
265 266	

267	SUPPLEMENTAL INFORMATION
268	Supplemental Information includes nine figures and three tables.
269	suppremental mormation metudes nine rigures and three tables.
209 270	AUTHOR CONTRIBUTIONS
270 271	Q.Y. designed and conducted the experiments, analyzed data, and wrote the manuscript. Y.H.
272	discussed the results and commented on drafts. Y.K. advised on TMS protocol. K.A. produced
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282	
283	COMPETING FINANCIAL INTERESTS
284	The authors declare no competing financial interests.
285	
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383	

#### 384 METHODS

385 **Participants.** Twenty individuals participated in the study (7 female,  $22.55 \pm 1.54$  years, mean  $\pm$  sd). Data from 3 386 subjects were excluded due to either poor performance (1 subject performed at chance level) or scanner malfunction 387 (projector crashed during scanning for 2 subjects at TMS-vertex session), resulting in a final group of 17 subjects (7 388 female,  $20.65 \pm 1.54$  years, mean  $\pm$  sd). All subjects were unfamiliar with the video game, had normal or correct-to-389 normal vision and did not report neurological or psychiatric disorders or current use of psychoactive drugs. All 390 subjects were eligible for MRI and TMS procedures based on standard MRI safety screening as well as on their 391 answers to a TMS safety-screening questionnaire (Rossi, Hallett, Rossini, Pascual-Leone, & Safety of, 2009). No 392 subjects withdrew due to complication from the TMS or MRI procedures, and no negative treatment responses were 393 observed. All subjects gave written informed consent and were compensated for their participation. All procedures 394 were performed in accordance with the 1964 Helsinki declaration and its later amendments and approved by 395 University Committee on Human Research Protection of East China Normal University (UCHRP-ECNU). The 396 number of participants was determined based on previous studies with similar design (Ezzyat & Davachi, 2014; 397 Wang et al., 2014).

398

399 Experimental design, stimuli, and tasks. Encoding: Interactive video game. The action-adventure video game 400 (Beyond: Two Souls) was created by the French game developer Quantic Dream and played in the PlayStation 4 401 video game console developed by Sony Computer Entertainment. Participants played the game using a first-person 402 perspective. In order to ensure the participants master the operational capability, they were trained to play the game 403 with two additional game chapters (Training chapters: Welcome to the CIA, and The Embassy). The training session 404 varied in duration depending on the dexterity of each participant on using the console (40 - 60 min per chapter). 405 After the training session, participants played 14 chapters in total across two sessions: 7 in Experimental Session 1 406 and then another 7 in Session 2 (Fig. 1). The video game they played were recorded and stored as a single video file 407 in MP4 format (Chapters 1~7: My Imaginary Friend, First Interview, First Night, Alone, The Experiment, Night 408 Session, Hauntings; Chapters 8~14: The Party, Like Other Girls, Separation, Old Friends, Norah, Agreement, 409 Briefing; see Supplementary Fig. 1).

410 Retrieval (scanned): Temporal Order Judgment (TOJ) task. The TOJ retrieval task required participants to 411 choose the image that happened earlier in the video game they had encoded. The task was administrated inside an 412 MRI scanner, where visual stimuli were presented using E-prime software (Psychology Software Tools, Inc., 413 Pittsburgh, PA), as back-projected via a mirror system to the participant. Each trial was presented for 5 s during 414 which participants performed the temporal order judgment. They were then allowed 3 s to report their confidence 415 level following the memory judgement. Participants performed the TOJ task using their index and middle fingers of 416 one of their hands via an MRI compatible five-button response keyboard (Sinorad, Shenzhen, China). Participants 417 reported their confidence level ("Very Low", "Low", "High", or "Very High") regarding their own judgment of the 418 correctness of TOJ with four fingers (thumb was not used) of the other hand. The left/right hand response 419 contingency was counterbalanced across participants. Participants were told they should report their confidence level 420 in a relative way and make use of the whole confidence scale. Following these judgments, a fixation cross with a 421 variable duration (1 - 6 s) was presented. Each participant completed 240 trials in each of the two experimental 422 sessions. Participants were given 15 practice trials using paired images extracted from the two additional chapters 423 they had played in the training session out of the scanner to ensure they understand the task procedure. Participants 424 completed a surprise recognition test after TOJ task outside scanner; data of which are not reported here.

For the TOJ task, we selected static images from the subject-specific recorded videos which the participants had played the day before. Each second in the video consisted of 29.97 static images (frames). For each gameplaying session, 240 pairs of images were extracted from the seven chapters and were paired up for the task based

428 on the following criteria: (1) the two images had to be extracted from either the same chapters or adjacent chapters 429 (Within- vs. Across-chapter condition); (2) the temporal distance (TD) between the two images were matched 430 between Within- and Across-chapter condition; (3) in order to maximize the TD, we first selected the second longest 431 chapter of the video and determined the longest TD according to a power function (power = 1.5), at the same time 432 ensuring the shortest TD to be longer than 30 frames. We generated 60 progressive levels of TD among these pairs 433 (each level repeated twice). In sum, three within-subjects factors regarding the TOJ retrieval task were manipulated: 434 (1) 60 TD levels permitting scale-invariance across subjects between two images (see below); (2) Context (two 435 images extracted from either Within- or Across-chapter); (3) TMS stimulation (TMS-precuneus vs. TMS-vertex, see 436 below).

437 *Selection of 60 levels of temporal distances (TDs).* In order to maximize the range of all TDs, we first selected 438 the second longest chapter of the video game and determined the longest TD (*L*), while ensuring the shortest TD to 439 be longer than 30 frames. The 60 TD levels were selected according to this function,

440

### $TD_n = L * (\frac{n}{60})^{1.5},$

441 where *L* denotes duration of the second longest chapter of the video game in each experimental session, *n* 442 denotes TD level, and value of  $TD_n$  were rounded to the nearest integer using the "round" function in MATLAB. 443 Note that the actual TDs were different across subjects, but since we applied a power function, the scale was thus 444 rendered invariant(Kwok & Macaluso, 2015). Image-pairs extraction from each of the chapters were independently 445 conducted across subjects. The numbers of images-pairs extracted from each of the chapters were approximately 446 equal within-subjects.

447

448 Transcranial magnetic stimulation. TMS procedure and protocol. TMS were applied using a 70 mm Double Air 449 Film Coil connected to a Magstim Rapid2 (The Magstim Company, Ltd., Whitland, UK). In order to localize the 450 target brain regions precisely, we obtained individual anatomical T1-weighted magnetic resonance images and then 451 imported them into BrainSight (Rogue Research Inc., Montreal, Canada) for stereotaxic registration of the TMS coil 452 with the participants' brain. The position of the coil and the subject's head were co-registered with BrainSight, and 453 monitored using a Polaris Optical Tracking System (Northern Digital, Waterloo, Canada) during TMS. Positional 454 data for both rigid bodies were registered in real time to a common frame of reference and were superimposed onto 455 the reconstructed three-dimensional MRI images of the subject using the BrainSight. The center of the coil was 456 continuously monitored to be directly over the site of interest. For all sites (vertex, precuneus, and motor areas for 457 measuring active motor threshold), the TMS coil was held tangential to the surface of the skull and was placed in a 458 rostro-caudal direction. An adjustable frame was used to hold the TMS coil firmly in place, while the participants 459 rested their heads on the chin rest. Head movements were monitored constantly by BrainSight and were negligible. 460 We measured subjects' active motor threshold, defined as the lowest TMS intensity delivered over the motor cortex 461 necessary to elicit visible twitches of the right index finger in at least 5 out of 10 consecutive pulses. The location 462 used to determine the active motor threshold was identified with a single pulse of TMS over the motor cortex at the 463 left hemisphere. The TMS coil was systematically moved until the optimal cortical site was located to induce the 464 largest and most reliable motor response; this stimulus output was then recorded. The TMS intensity was then 465 calibrated at 110% of individual active motor threshold (stimulator output:  $75.2 \pm 6.9\%$ , mean  $\pm$  se, range from 63% 466 to 88%, Supplementary Table 1). In Experimental Session 1 and 2, the TMS was applied at a low-frequency rate 467 of 1 Hz with an uninterrupted duration of 20 min.

468 *TMS stimulation sites.* The target stimulation was delivered to the precuneus(Kwok et al., 2012) (MNI x, y, z = 6, -70, 44), whereas the control stimulation was delivered to the vertex. The vertex was defined individually by the point of the same distance to the left and the right pre-auricular, and of the same distance to the nasion and the inion.
471 Due to the folding of the two cerebral hemispheres, the stimulated vertex site lies at a considerable distance from the

TMS coil, thereby diminishing the effectiveness of the magnetic pulses. Stimulating the vertex is not known to produce any memory task-relevant effects and deemed as a reliable control site. Stimulation magnitude and protocols in the present study were comparable to those used in similar studies that are robust to produce significant memoryrelated changes by targeting at the precuneus (Bonni et al., 2015; Kraft et al., 2015; Mancini et al., 2017) or lateral parietal cortices (Nilakantan et al., 2017; Wang et al., 2014). Immediately after the end of the stimulation, participants performed four runs of Temporal Order Judgment task in the MRI scanner (delay period between the end of TMS and the beginning of MRI:  $M_{\text{precuneus}} = 15.29 \text{ min}, M_{\text{vertex}} = 20.76 \text{ min}, t (16) = -0.87, P = 0.4$ ).

479

480 MRI data acquisition and preprocessing. Data acquisition. All the participants were scanned in a 3-Tesla Siemens 481 Trio magnetic resonance imaging scanner using a 32-channel head coil (Siemens Medical Solutions, Erlangen, 482 Germany) at ECNU. In each of the two experimental sessions, a total of 1,350 fMRI volumes were acquired for each 483 subject across 4 runs. The functional images were acquired with the following sequence: TR = 2000 ms, TE = 30484 ms, field of view (FOV) =  $230 \times 230$  mm, flip angle =  $70^{\circ}$ , voxel size =  $3.6 \times 3.6 \times 4$  mm, 33 slices, scan orientation 485 parallel to AC-PC plane. High-resolution T1-weighted MPRAGE anatomy images were also acquired (TR = 2530 486 ms, TE = 2.34 ms, TI = 1100 ms, flip angle =  $7^{\circ}$ , FOV =  $256 \times 256$  mm, 192 sagittal slices, 0.9 mm thickness, voxel 487 size =  $1 \times 1 \times 1$  mm).

488 Preprocessing. Preprocessing was conducted using SPM12 (http://www.fil.ion.ucl.ac.uk/spm). Scans were 489 realigned to the middle EPI image. The structural image was co-registered to the mean functional image, and the 490 parameters from the segmentation of the structural image were used to normalize the functional images that were 491 resampled to  $3 \times 3 \times 3$  mm. The realigned normalized images were then smoothed with a Gaussian kernel of 8-mm 492 full-width half maximum (FWHM) to conform to the assumptions of random field theory and improve sensitivity 493 for group analyses (Friston et al., 2002). Data were analyzed using general linear models and representational 494 similarity analyses as described below with a high-pass filter cutoff of 256 s and autoregressive AR(1) model 495 correction for auto-correlation.

496

497 Functional MRI data analysis. *Parametric modulation analysis*. First-level models were performed on the fMRI 498 data collected from the TMS-vertex session only (either all the trials altogether or separately for Across-chapter vs. 499 Within-chapter conditions). In all of these models, each of the 240 trials was modeled with a canonical hemodynamic 500 response function as an event-related response with a duration of 5 s.

501 For the TMS-vertex session as a whole (Across-chapter and Within-chapter trials collapsed), we performed 502 three parametric modulation analyses (pmod), each with a different combination of modulatory regressor/regressors 503 (namely, TD; TD + RT; SC). For the TD pmod, we assigned the actual TD values at encoding as the modulatory 504 parameter, and used the polynomial function up to first order. Several regressors of no interest were also included: 505 6 head movement regressors and 1 missing trial regressor (i.e., no-response trials; number of missing trials of Across-506 chapter condition:  $5.65 \pm 6.96$ , of Within-chapter condition:  $5.29 \pm 6.8$ ; n = 17, mean  $\pm$  sd) and the run mean. The 507 purpose of this analysis was to test for any linear TD-dependent modulation of signal intensity in the brain between 508 the TD between the two images at encoding and the brain activity during TOJ retrieval of the same events. For the 509 TD + RT pmod, we aimed to identify the voxels whose activities changed as a function of TD after the removal of the influence of reaction time. Each subjects' RTs corresponding to each TD level were entered as the modulatory 510 511 parameter, together with the regressors of no interest as above. For the Situational Change (SC) pmod, we tested for 512 any linear SC-dependent modulation of signal intensity in the brain between the spatial displacement of two images 513 at encoding and the brain activity during TOJ retrieval of the same pairs of images. We determined the numbers of 514 situational change between the paired of images by analyzing all the subject-specific videos frame by frame to mark 515 out the boundaries at which a situational location had changed (Supplementary Fig. 4), and entered these video-

516 specific and subject-specific SC values as the modulatory parameter.

517 For the Across-chapter vs. Within-chapter comparison, we also performed three pmod analyses with identical 518 sets of regressors as described above (namely, TD; TD + RT; SC). We looked for changes in brain responses as a 519 linear function of the regressor of interest (i.e., TD or SC). Maps were created by multiple regression analyses 520 between the observed signals and regressors. The contrast maps from the first-level model of parametric analyses 521 were taken for second-level group analyses and entered into one-sample t-tests. The group analyses were performed 522 for each contrast using a random effects model (Penny & Holmes, 2004). The statistical threshold was set at p < 0.05523 (FWE corrected) at cluster level and p < 0.001 at an uncorrected peak level according to the SPM12 standard 524 procedure. The activation cluster locations were indicated by the peak voxels on the normalized structural images 525 and labeled using the nomenclature of Talairach and Tournoux (1988) (Talairach & Tournoux, 1988).

526 Searchlight Representational Similarity Analysis (searchlight RSA). RSA were conducted using the RSA 527 toolbox (http://www.mrc-cbu.cam.ac.uk/methods-and-resources/toolboxes/) on the fMRI data following 528 realignment and normalization, but without smoothing. In the Across-chapter vs. Within-chapter comparison, each 529 unique TD level was modeled with a separate regressor and was contrasted to produce a T-statistic map (spmT maps), 530 creating 120 statistical maps in total (Across- vs. Within-chapter conditions; 2 repetitions for each TD level). For 531 the TMS-vertex vs. TMS-precuneus comparison, we collapsed the trials in the Across- and Within-chapter 532 conditions and generated 60 statistical maps in either of the two sessions (4 repetitions for each TD level). Using 533 searchlight RSA, spherical searchlights with a radius of 9 mm (93 voxels, volume  $= 2,511 \text{ mm}^3$ ) were extracted from 534 the brain volume and then the data (i.e., signal intensity) for the 60 TD levels were Person product-moment (1 - r)535 correlated with every other levels to generate a representational dissimilarity matrix (RDM), reflecting the between-536 condition dissimilarity of BOLD signal response. These neural RDMs were then Spearman-rank correlated with a 537 set of candidate RDMs (see Supplementary Fig. 2a), reflecting different predictions of the information carried by 538 similarity structure of neural signal responses and generated correlational maps (r-maps). Finally, these r-maps were 539 converted to z-maps using Fisher transformation. All the z-maps were then submitted to a group-level one-sample t-540 test to identify voxels in which the similarity between the predicted RDM and observed neural RDM was greater 541 than zero. This allowed us to identify voxels in which information of TD at retrieval might be represented (see 542 Supplementary Fig. 2b). The statistical threshold was set as identical to those employed in the univariate analysis, 543 which was at p < 0.05 (FWE corrected) at cluster level and p < 0.001 at an uncorrected peak level.

544 Leave-one-subject-out approach (LOSO), functional and anatomical ROIs. We applied LOSO approach to 545 create functional ROIs to avoid statistical bias (Esterman, Tamber-Rosenau, Chiu, & Yantis, 2010). For instance, in 546 order to identify an ROI (i.e., conjunction mask in Figure 2d) for Subj01, we re-estimated the contrast using a one-547 sample t-test on the whole-brain searchlight z-maps obtained from Subj02 to Subj17. Likewise, we also re-estimated 548 the contrast using a one-sample t-test on the contrast maps obtained from the Pmod analysis of Subj02 to Subj17. 549 We set the same threshold reported above to extract clusters from these two statistical maps. We next overlaid the 550 two resultant maps and extracted a conjunction region (mask01), with which we used to extract the value in the 551 searchlight z-map from Subj01 for further statistical analysis. This procedure was repeated 17 times and generated 552 17 different ROIs, which provided statistically independent regions to extract values for testing differences between 553 conditions (Figure 2e). For the anatomical ROIs (depicted in Figure 3 and Supplementary Figure 8), 7 regions 554 (Hippocampus\_L; Hippocampus\_R; Precuneus\_L; Precuneus\_R; Angular\_L; Angular\_R; Occipital cortex) of AAL 555 template (Tzourio-Mazoyer et al., 2002) were created as masks. We extracted and averaged the similarity value 556 within these masks for each subject for statistical tests.

557

558 **Candidate representational dissimilarity matrices (RDM).** *Model 1* (TD RDM,  $60 \times 60$ ): We ranked the difference across the 60 TD levels, from the shortest to the longest TD, for the Within-chapter condition (or Across-

chapter condition, the two RDMs were identical because of their matched TD). We first log-transformed the subject-

specified TD values for each pair of images and then computed the differences with and among every other TD levels producing  $60 \times 59/2$  values, which were then assigned to the corresponding cells of the RDM.

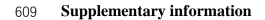
563 *Model 2* (Situational Change RDM,  $60 \times 60$ ): Since the temporal and spatial dimensions were closely inter-564 correlated. We checked whether the situational change might influence the neural patterns in those voxels that 565 represent the TD information. We analyzed the subject-specific videos frame by frame and marked out the 566 boundaries at which a situational location had changed (see illustration in **Supplementary Fig. 4**). Then we 567 computed the numbers of situational changes contained in each of the paired images and then computed the 568 differences with and among every other conditions producing  $60 \times 59/2$  values, which were then assigned to the 569 corresponding cells of the RDM.

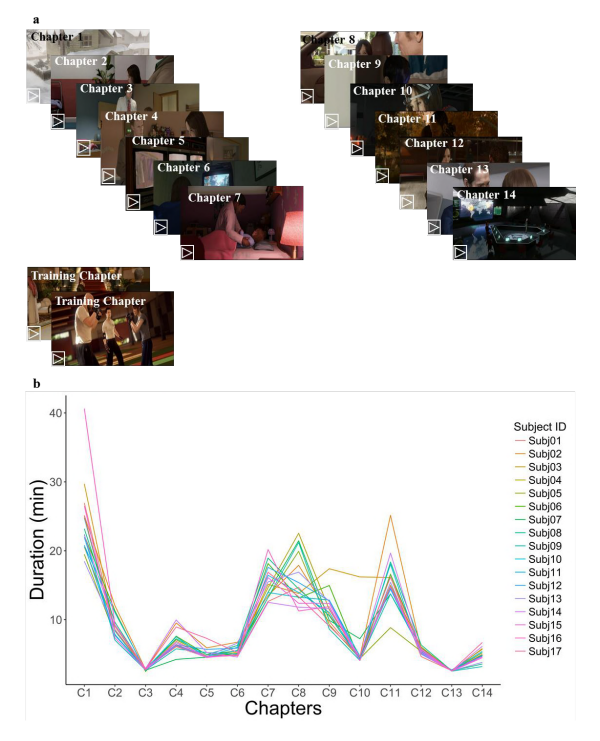
570 Model 3 (RGB-cross-correlation,  $60 \times 60$ ), Model 4 (RGB-intensity RDM,  $60 \times 60$ ) and Model 5 (RGB-571 histogram RDM,  $60 \times 60$ ) considered the perceptual characteristics of the images used in TOJ. For model 3, the 572 similarity measure was based on the cross-correlation value between two images (image of size  $1920 \times 1080$ ) for 573 the three color channels (red, green, and blue; RGB). For every pair of images in each of the three color channels 574 (RGB), we computed the cross-correlation coefficients between the pair. This is a measure of the displacement of 575 one image relative to the other; the larger the cross-correlation coefficient (which ranges between -1 and 1), the more 576 similar the two images was. We then computed the differences with and among every other conditions producing 60 577  $\times$  59/2 values, which were then assigned to the corresponding cells of the RDM. For Model 4, we computed the 578 pixel-wise difference between pair images for the three color channels (RGB). The computed difference is useful 579 when the compared images are taken from a stationary camera with infinitesimal time difference. The output pixel 580 for each color channel is assigned with the value 1 if the absolute difference between the corresponding pixels in the 581 image pair is non-zero, or a value of 0 otherwise. A single value is generated for each of the three color channels by 582 summing all the output pixel values (either 0 or 1). We averaged the sum of difference for all three color-channels 583 for the intensity value of each pair of images and then computed the differences with and among every other 584 conditions producing  $60 \times 59/2$  values, which were then assigned to the corresponding cells of the RDM. For Model 585 5, we constructed color histograms for image pairs and computed the Sum-of-Square-Difference (SSD) error 586 between them for the three color channels (RGB). For each color channel the intensity values range from 0 to 255 587 (i.e., 256 bins), we first computed the total number of pixels at each intensity value and then computed the SSD for 588 all 256 bins for each image pair. The smaller the value of the SSD, the more similar the two images (image pair) 589 was. We then computed the differences with and among every other conditions producing  $60 \times 59/2$  values, which 590 were then assigned to the corresponding cells of the RDM. In contrast to model 4, this approach does not require 591 corresponding pixels in the image pair to be the same, but rather measures the existence of pixel intensity in both 592 images. Overall, the three perceptual-similarity models (3, 4 and 5) look at different similarity measures and they 593 complement each other; thus any difference in the appearance of the two images irrespective of the temporal distance, 594 could be accounted for by at least one of the three models. For any two similar images, the RGB-intensity RDM 595 results in a very small value, thus the corresponding pixels are virtually the same for the entire image. The RGB-596 histogram will also result in a small value as the image pairs will have the same histogram bins. The RGB-cross-597 correlation value will be close to 1, signifying the similarity in the images. When subsections of a scene are visible 598 in both images with varied brightness, the RGB-cross-correlation value will still be closer to 1 but with a very high 599 RGB-intensity RDM value.

600

Behavioral data analysis. To look into the TD-independent effects of precuneal disruption by TMS on memory
 performance, we collapsed the 60 TD levels for each subject and entered their percentage correct (ACC) or response
 times (RT) of the TMS sessions (TMS-vertex vs. TMS-precuneus) and the Context factor (Across-chapter vs.

- 604 Within-chapter) as within-subjects factors into repeated ANOVAs.
- **Data and code availability.** Data and codes are available upon request.



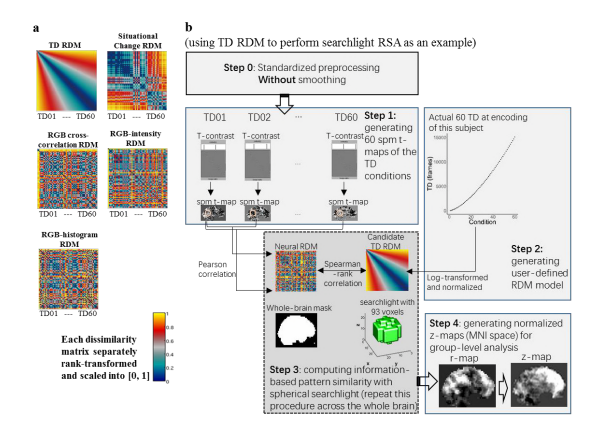




Supplementary Figure 1. Game chapters and playing durations of each chapter for the subjects. (a) 14 different chapters were used: seven in each of two within-subjects TMS sessions (see also Supplementary Table 1). (b) While the playing durations varied across chapters and subjects, the numbers of paired images extracted for the TOJ task were approximately equal across the chapters.

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Supplementary Figure 2. Candidate RDMs and pipeline of RSA analysis. (a) Candidate
RDMs for RSA searchlight analyses, including TD RDM, Situational Change RDM,

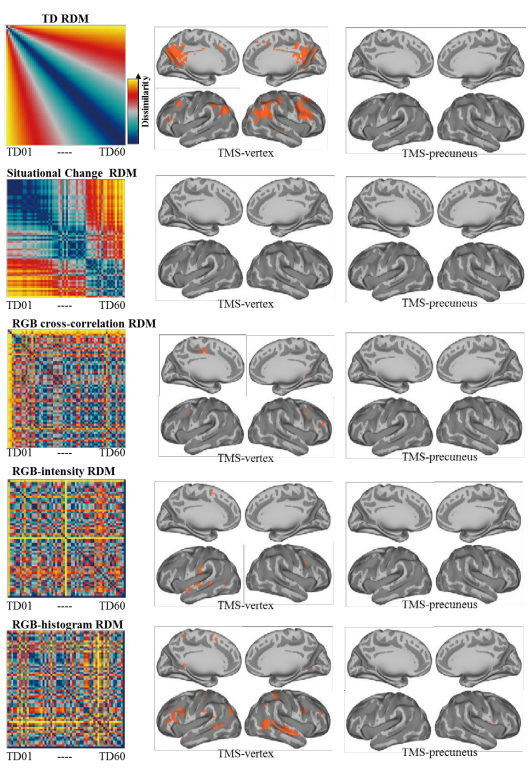
624 RGB cross-correlation RDM, RGB-intensity RDM, and RGB-histogram RDM. Data

depicted in these RDMs were extracted from one subject (Subj01). (b) Using one model

626 (TD RDM) to perform the searchlight procedure as an example in one subject (Subj01).

627 The same principle and procedure applied to all other candidate RDMs and subjects.

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- 629



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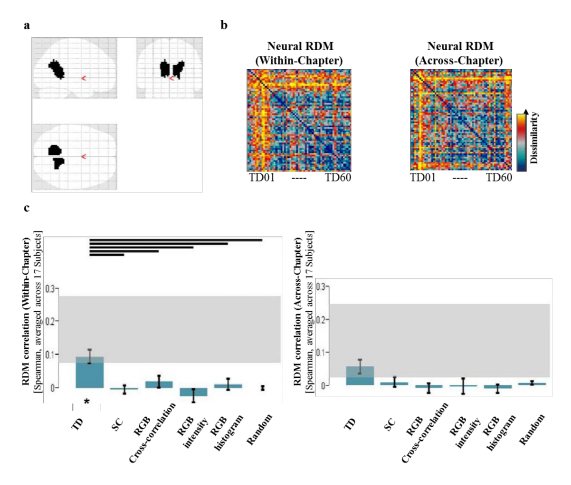
Supplementary Figure 3. RSA searchlight results for TMS-vertex and TMS-precuneus sessions using candidate RDMs (TD RDM; Situational Change RDM; RGB crosscorrelation RDM; RGB-intensity RDM; RGB-histogram RDM). TD representation (top row and cf. **Fig. 2b**) was not found in the other control candidate RDMs (rows 2 – 5). For illustration purpose, we generated these RDMs using one participant's video and TOJ paired-images from his TMS-vertex session (Subj01). MRI results are displayed at  $P_{\text{uncorrected}} < 0.001$ .



638 639

640 Supplementary Figure 4. An excerpt of a subject-specific video illustrates how 641 situational changes were defined frame-by-frame for Situational Change RDMs and 642 pmod analyses. Noting that time and space in our task were partially correlated, we 643 quantified subject-specific situational changes in terms of the number of locations each 644 participant had traversed in the video (i.e., spatial displacement embedded in the paired 645 images presented in TOJ task).

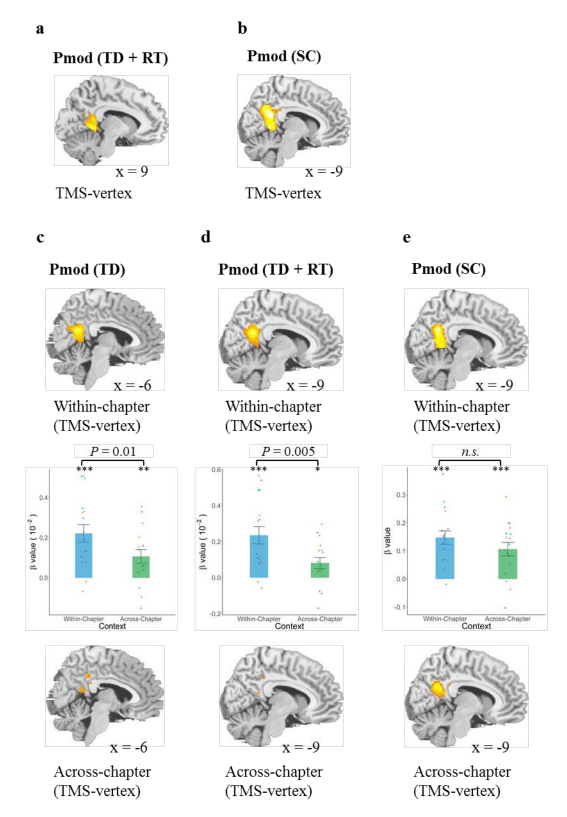
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649 Supplementary Figure 5. (a) Conjunction region for extracting neural signals. (b) 650 Neural RDMs averaged across 17 subjects for Within-chapter and Across-chapter 651 conditions. (c) Candidate RDMs (TD, SC, RGB cross-correlation, RGB intensity, RGB 652 histogram, Random) are tested and compared for their ability to explain the Neural 653 RDM in Within-chapter and Across-chapter conditions. As expected, the TD RDM is 654 the most correlated candidate RDM to the Neural RDM in the Within-chapter condition. 655 \* P < 0.05, FDR corrected.

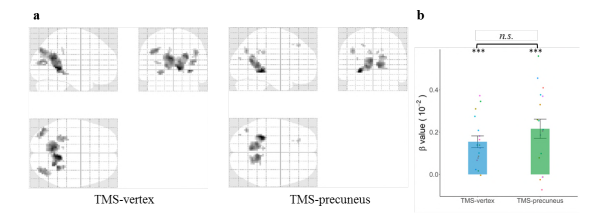
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Supplementary Figure 6. Whole-brain parametric modulation analyses with different regressors (temporal distance (TD), reaction times (RT), and situational change (SC)).
(a) Temporal distance and RT as modulatory regressors for TMS-vertex session (Within-chapter and Across-chapter conditions collapsed). Having removed the influence of RT, neural activity in the precuneus/retrosplenial cortex and posterior

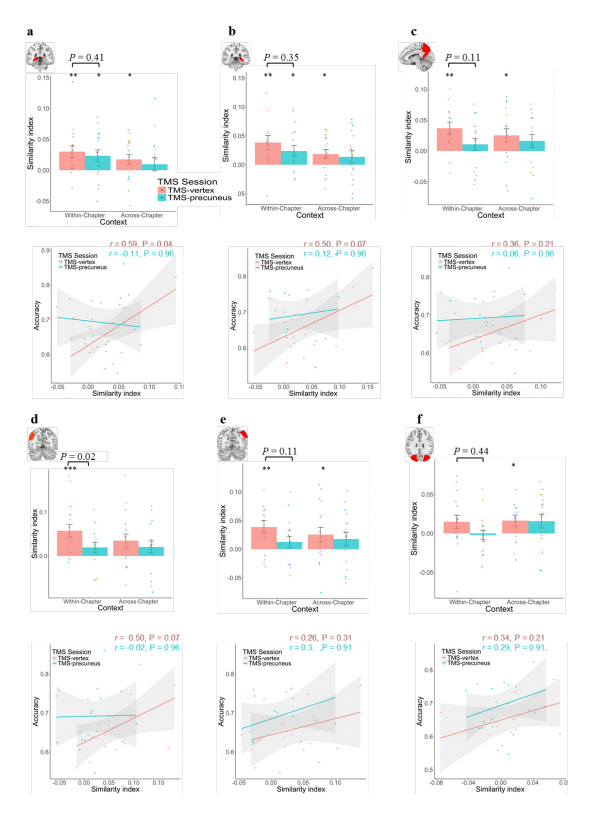
hippocampus increased with TD. (b) Situational changes as a modulatory regressor for 664 TMS-vertex session. Neural activity in the precuneus/retrosplenial cortex and posterior 665 cingulate cortex increased with the number of situational changes. (c) TD as a 666 modulatory regressor separately for Within-chapter (upper) and Across-chapter (bottom) 667 conditions of TMS-vertex session. Neural activity in the precuneus, extending into the 668 retrosplenial cortex and posterior cingulate cortex increased linearly with TD in Within-669 chapter condition. The  $\beta$  values extracted from the cluster was higher in the Within-670 chapter condition than in the Across-chapter condition, P = 0.01. (d) TD and reaction 671 time (RT) as modulatory regressors separately for Within-chapter (upper) and Across-672 chapter (bottom) conditions of TMS-vertex session. After having removed the influence 673 of RT, neural activity in the precuneus, extending into the retrosplenial cortex and 674 675 posterior cingulate cortex, remained linearly associated with TD in Within-chapter condition. The  $\beta$  values extracted from the cluster was also higher in the Within-chapter 676 condition than in the Across-chapter condition, P = 0.005. (e) SC as a modulatory 677 regressor separately for Within-chapter and Across-chapter conditions of TMS-vertex 678 session. Neural activity in the precuneus/retrosplenial cortex and posterior cingulate 679 cortex increased with the number of situational changes in both conditions. No 680 significant difference was found in  $\beta$  values between the two conditions. In the present 681 study the anterior part of the precuneus (including the retrosplenial cortex) was engaged, 682 whereas in previous studies the more dorsal/posterior region of the precuneus was 683 activated (Kwok et al., 2012; St Jacques et al., 2008). This might stem from the fact 684 that the encoding done here during gameplay being different from those involved in 685 watching TV episodes (Kwok et al., 2012) or experiencing a self-guided tour(St Jacques 686 et al., 2008). For visualization purposes, the threshold was set at  $P_{\text{uncorrect}} < 0.001$ . 687 688



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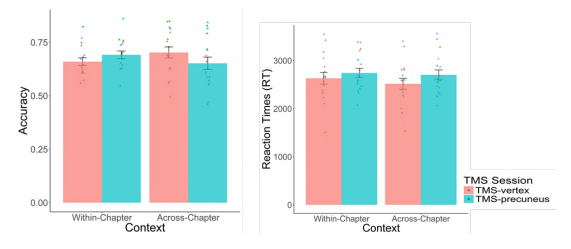
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691 Supplementary Figure 7. (a) In contrast to the removal of the multivoxel representation, 692 TMS to the precuneus did not impact on the activation intensity in the precuneus and 693 retrosplenial cortex. (b) Activation in these clusters (containing the precuneus) 694 increased with temporal distance irrespective of TMS stimulation. Activation maps are 695 displayed at  $P_{\text{uncorrected}} < 0.001$ .



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Supplementary Figure 8. Similarity indices in six ROIs (a-b: bilateral hippocampus, c:
right precuneus, d-e: bilateral angular gyrus, and f: whole occipital cortex) in both TMS
sessions, and their correlation with subjects TOJ accuracy for Within-chapter trials. P
values are FDR-corrected.



Supplementary Figure 9. Memory performance of TOJ task. Behavioral results (Accuracy and RT) of TOJ task by TMS stimulation sites and Context (60 TD levels collapsed). TMS to the precuneus did not result in any retrograde memory amnesia (*Ps* > 0.05), but resulted in a slight slowing in retrieval times (main effect of TMS: *F* (1, 16) = 5.45, *P* = 0.03,  $\eta^2$  = 2.8%; interaction effect: *P* = 0.2) as compared with the TMSvertex condition.

	Motor		Session 1		D I		Session 2	
ID	thresh	Chapters	Duration	TMS Site	Delay (days)	Chapters	Duration	TMS Site
	old	(Playing)	(min)			(Playing)	(min)	
Subj01	66%	Chap 1 ~ 7	89.39	Precuneus	9	Chap 8 ~ 14	35.19	Vertex
Subj02	67%	Chap $1 \sim 7$	102.70	Vertex	6	Chap $8 \sim 14$	44.46	Precuneu
Subj03	80%	Chap 8 ~ 14	35.53	Vertex	9	Chap $1 \sim 7$	114.21	Precuneu
Subj04	58%	Chap $8 \sim 14$	36.41	Vertex	2	Chap $1 \sim 7$	94.70	Precuneu
Subj05	65%	Chap 1 ~ 7	80.50	Precuneus	8	Chap $8 \sim 14$	35.31	Vertex
Subj06	67%	Chap 1 ~ 7	95.05	Vertex	3	Chap $8 \sim 14$	39.63	Precuneu
Subj07	72%	Chap 8 ~ 14	32.41	Vertex	4	Chap $1 \sim 7$	97.06	Precuneu
Subj08	73%	Chap 1 ~ 7	96.08	Precuneus	8	Chap $8 \sim 14$	37.03	Vertex
Subj09	69%	Chap 1 ~ 7	88.81	Vertex	7	Chap $8 \sim 14$	31.46	Precune
Subj10	80%	Chap 8 ~ 14	35.69	Precuneus	7	Chap $1 \sim 7$	89.79	Vertex
Subj11	65%	Chap 8 ~ 14	38.77	Vertex	26	Chap $1 \sim 7$	86.02	Precuner
Subj12	71%	Chap 1 ~ 7	90.56	Precuneus	16	Chap 8 ~ 14	35.89	Vertex
Subj13	68%	Chap 8 ~ 14	35.66	Vertex	7	Chap $1 \sim 7$	87.98	Precuner
Subj14	73%	Chap 1 ~ 7	87.94	Precuneus	7	Chap $8 \sim 14$	40.56	Vertex
Subj15	57%	Chap 8 ~ 14	32.72	Precuneus	5	Chap $1 \sim 7$	91.46	Vertex
Subj16	69%	Chap 8 ~ 14	35.44	Vertex	5	Chap $1 \sim 7$	108.49	Precuner
Subj17	61%	Chap 1 ~ 7	91.10	Precuneus	6	Chap 8 ~ 14	42.13	Vertex

# Supplementary Table 1. Descriptive statistics of TMS motor threshold and full counter-balancing for gameplay chapters by TMS sites.

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Cluster			peak				
k	p-FWE	X	y z		% Cluster	Brain regions	
7554	<.001	30	21	38	5.41	Precuneus_L	
					5.32	Precuneus_R	
					4.38	Frontal_Mid_2_R	
					3.34	Angular_R	
					3.24	Parietal_Inf_L	
					3.15	Frontal_Inf_Tri_R	
					2.99	Parietal_Inf_R	
					2.91	Frontal_Mid_2_L	
					2.83	Frontal_Inf_Tri_L	
					2.74	Angular_L	
					2.73	SupraMarginal_R	
					2.73	Frontal_Inf_Oper_R	
					2.65	Frontal_Sup_2_R	
					2.61	Cuneus_L	
					2.58	Cingulate_Mid_R	
					2.57	Postcentral_R	
					2.55	Temporal_Mid_R	
					2.26	Calcarine_L	
					2.22	Cuneus_R	
					1.87	Precentral_R	
					1.85	Occipital_Mid_R	
					1.62	Calcarine_R	
					1.62	Frontal_Sup_Medial_	
					1.56	Cingulate_Mid_L	
					1.52	Occipital_Sup_R	
					1.47	Frontal_Sup_Medial_	
					1.42	Temporal_Sup_R	
					1.22	Occipital_Mid_L	
					1.02	Cingulate_Post_L	
					1.01	Precentral_L	

## 717 Supplementary Table 2. Brain representation associated with TD RDM in TMS-vertex

719 Note. % Cluster denotes percentage of voxels of the cluster that is contained in each of the anatomical regions

- 720 (Tzourio-Mazoyer et al., 2002).
- 721

Regressor	Clust	er	Voxel				<b>Brain Regions</b>		
	k	р-	Z value	x	Y	Z	-		
		FWE							
TD	555	<.001	4.62	9	-45	-1	Cerebelum_4_5_F		
			4.35	-6	-57	20	Precuneus_L		
			4.27	6	-54	5	Vermis_4_5		
	116	.002	3.87	36	-81	14	Occipital_Mid_L		
			3.78	42	-72	29	Occipital_Mid_L		
			3.75	36	-69	11	Occipital_Mid_R		
	117†	.01	4.34	45	-36	5	Temporal_Sup_R		
			4.17	54	-36	5	Temporal_Sup_R		
			3.81	48	-27	-1	Temporal_Sup_R		

722	Supplementary	Table 3.	Brain	activation	parametrically	modulated	by	temporal
723	distance (TD) in	n TMS-ver	tex sess	sion (cf. Fig	ure 2c).			

724 Note. <sup>†</sup> denotes a negative relationship between TD and brain activity.

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