

Spatial Binding Impairments in Visual Working Memory following **Temporal Lobectomy**

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1	Spatial Binding Impairments in Visual Working Memory following Temporal
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- 32 Arabia.

33	Abstract
34	Disorders of the medial temporal lobe (MTL) adversely affect visual working
35	memory (vWM) performance, including feature binding. It is unclear whether these
36	impairments generalise across visual dimensions or are specifically spatial. To address this
37	issue, we compared performance in two tasks of thirteen epilepsy patients, who had
38	undergone a temporal lobectomy, and fifteen healthy controls. In the vWM task, participants
39	recalled the color of one of two polygons, previously displayed side by side. At recall, a
40	location or shape probe identified the target. In the perceptual task, participants estimated the
41	centroid of three visible disks. Patients recalled the target color less accurately than healthy
42	controls because they frequently swapped the non-target with the target color. Moreover,
43	healthy controls and right temporal lobectomy patients made more swap errors following
44	shape than space probes. Left temporal lobectomy patients, showed the opposite pattern of
45	errors instead. Patients and controls performed similarly in the perceptual task. We conclude
46	that left MTL damage impairs spatial binding in vWM, and that this impairment does not
47	reflect a perceptual or attentional deficit.

49	Significance Statement
50	This study examined color recall in temporal lobectomy patients and healthy controls,
51	to determine whether patients show differential impairments binding color and shape vs color
52	and location of memorised objects. Left temporal lobectomy patients were less accurate
53	recalling color, especially when the target object was identified by the location, rather than
54	the shape it had in the initial display. We found no group difference in a task, which required
55	estimating the centroid of three circles, indicating that the memory impairment was not
56	accounted by perceptual or attentional difficulties. Our findings indicate that lateralised
57	medial temporal circuits are crucial for binding visual features to the location where they had
58	appeared, thus ensuring the primacy of space in organising declarative memories.

The role of the medial temporal lobe (MTL) in episodic memory is well established 61 62 (Squire, 2009). Despite initial reports of preserved immediate memory span in temporal 63 lobectomy patients (Drachman and Arbit, 1966), later studies found that MTL lesions also engender substantial working memory (WM) deficits (Olton et al., 1979; Holdstock et al, 64 65 1995; Hannula et al., 2006). Which WM processes are specifically supported by the MTL is, nevertheless, a matter of ongoing investigations. 66 67 An early, seminal model suggested that visual WM (wWM) contains few discrete 68 "slots", each used to store one and only one object with high fidelity (Luck and Vogel, 1997; 69 Zhang and Luck, 2008). Despite its simplicity, the slot model makes non-trivial predictions. 70 First, the complexity of memorised objects should not affect recall accuracy. Second, 71 recalling feature conjunctions should carry no additional cost over remembering features, 72 since features are stored *ipso facto* as parts of an object into vWM. Wheeler and Treisman 73 (2002) found, instead, that simple objects were recalled more accurately than complex ones, 74 and that recall accuracy was equalised for displays containing the same number of color 75 features rather than the same number of objects. They concluded that memory limitations 76 reflect feature rather than object-based storage mechanisms. Moreover, observers were worse 77 at detecting changes of feature conjunctions than features, indicating that conjunctions are stored or recalled less efficiently than features. Later studies confirmed that changes in 78 79 feature conjunctions are poorly detected (Allen et al., 2006), leading to the suggestion that 80 dimensionally specific registers store features, while an "episodic buffer" is dedicated to 81 holding bound object representations in vWM (Baddeley et al., 2011). The need for binding 82 processes follows logically from the alternative model of vWM which proposes that visual 83 features are stored in dimensionally specific, limited resolution stores (Alvarez and 84 Cavanagh, 2004; Smyrnis et al., 2005; Bays et al., 2009). Clearly, if different feature

Introduction

60

85 dimensions are stored separately, then a binding process is required to ensure that features 86 belonging to the same object, but different feature dimensions, are identified as such 87 (Wheeler and Treisman, 2002; Smyrnis et al., 2005). 88 While the idea that conjunctive binding is required to preserve object identity is not 89 unanimously shared (e.g. Luck and Vogel, 2013), there are several proposals regarding its 90 nature. Treisman and Zhang (2006) concluded that binding is automatic, established initially 91 by the features' shared location, but then becomes location independent. Schneegans and 92 Bays (2017) proposed instead that location information is always required, because features 93 from different visual dimensions are stored in separate retinotopic maps. 94 Investigators examining the neurological underpinnings of declarative memory 95 largely embrace the idea that space plays a crucial role in indexing declarative memories. 96 Animal and patient studies (e.g. Chalfonte et al., 1996; Brown and Aggleton, 2001; Eacott 97 and Gaffan, 2005; Piekema et al., 2006; Ranganath, 2010; Libby et al., 2014) documented a 98 functional parcellation of the MTL with separate structures representing respectively the 99 environmental layout, the objects within it, as well as binding the latter to the former. 100 According to this view, space is crucial for recalling events, but not for binding object 101 features. Olson et al. (2006) for example, reported that patients with post-anoxic or post-102 encephalitic MTL pathology had impaired object-location binding in a WM task. This 103 impairment was unaccounted by either diminished recognition or spatial memory. In animals, 104 MTL lesions are also followed by dissociated impairments in object recognition and recall of 105 object-location conjunctions, suggesting that feature and spatial binding depend on distinct 106 MTL processes (Meunier et al., 1993; Murray and Mishkin, 1998; Malkova and Mishkin, 107 2003). Studies in temporal lobe epilepsy (TLE) patients reported deficits in spatial recall, 108 spatial binding and visual recognition (Abrahams et al., 1997; Bohbot et al., 1998; 109 Stepankova et al., 2004), however it remains unclear whether these impairments should be

attributed to diminished spatial precision (Kolarik *et al.*, 2016) or a binding deficit (Zokaei *et al.*, 2019) and whether binding impairments are dimensionally general (Hannula *et al.*, 2006;
Pertzov *et al.*, 2013) or specific.

113 To examine these issues, we tested TLE patients who had undergone temporal 114 lobectomies and healthy controls with two tasks used in a previous investigation of a stroke 115 patient (Dundon et al., 2017). The first requires recalling the color of one of two polygons 116 identified by either a location or a shape probe, thus directly pitting spatial vs. non-spatial 117 binding. The second probes participants' ability to estimate the average position of three 118 visible disks. In this task healthy participants show a pseudo-neglect pattern of leftward static 119 errors (Baud-Bovy and Soechting, 2001), which suggests that centroid estimation is sensitive 120 to attentional biases (Dundon et al., 2017). The centroid task was therefore used to highlight 121 the presence of unilateral neglect, which can follow lesions of the non-dominant 122 parahippocampal cortex (Mort et a.l, 2003) as well as the integrity of spatial perception and 123 attention (Drew et al., 2010).

125	Material and Methods
126	The aim of the present study was to compare non-spatial and spatial binding
127	performance in TLE patients' with medically refractory epilepsy who had undergone
128	temporal lobectomy and healthy controls. Recruitment and testing took place at the
129	Neuropsychology section of the Department of Neurosciences of the Department of
130	Neurosciences of King Faisal Specialist Hospital & Research Centre in Riyadh, Saudi
131	Arabia. The experimental protocol was approved by the local Institutional Review Board.
132	Participants gave written informed consent prior to engaging in any experimental procedure.
133	
134	Study participants
135	Over a month period, an opportunity sample of patients attending the Neurology and
136	Neurosurgical Clinics were invited to participate in the study. All had been diagnosed with
137	TLE on the basis of clinical presentation and instrumental diagnostic procedures, inclusive of
138	ambulatory EEG and neuroimaging, and after failing medical therapy, had
139	undergone surgical treatment. All patients had normal or corrected to normal visual
140	acuity. Those with an estimated full-scale IQ of less than 75, as assessed with an Arabic
141	version of the Wechsler Abbreviated Scale of Intelligence - Second Edition (WASI-II, Al-
142	Joudi et al., 2019), were excluded, as well as those with a history of traumatic brain
143	injury or major psychiatric disorders. Patients who suffered a seizure in the preceding 48
144	hours had the testing session postponed. Thirteen patients took part in this study. The
145	study's neurosurgeon identified the anatomical structures involved by the resection on the
146	basis of the surgical record and post-surgical MRI scans.
147	Fifteen healthy participants were concurrently recruited from the local community as
148	controls. Potential participants were excluded if they had a history of a major neurological or
149	psychiatric disorder or an uncorrected visual impairment or an estimated IQ less than 75.

151 Testing protocol

152 Testing took place in a quiet, dimly lit room. Participants sat comfortably at a distance of approximately 70cm from a MacBook Pro[®] set at a resolution of 1680x1050 153 154 pixels. Custom-coded Matlab (Mathworks, Natick, Massachusetts) scripts used a set of freely 155 available routines (Brainard, 1997) to control the timing of the displays. Two tasks were 156 conducted using the computerized set-up. 157 158 Cued color recall 159 In each trial, an equilateral triangle and a square, whose side lengths are 1.29° and 160 0.92° respectively, appear side-to-side in the lower half of the screen, as shown in figure 1A. 161 The shapes are centred at an eccentricity of 2.27° along the main screen diagonals and remain visible for 2.0s. The two shapes are of different colors, either red, blue or green. The sample 162 163 display is followed by a 0.2s long pattern mask and a 1.3s blank screen. The recall screen 164 contains three colored rectangles, 0.53° wide and 1.59° high, whose lower edges are aligned 165 1.39° above the screen center and spaced horizontally 4.81° apart. A bright cross (location 166 probe) or the outline of one of the polygons (shape probe) identifies the target. The location 167 probe, which also includes a dark cross, appear at the locations occupied by the two shapes. The shape cue appears 3.0° below the screen center. Participants report the 168 169 target color by placing the cursor over the corresponding colored rectangle and clicking the 170 mouse button. The mouse click prompts the beginning of a new trial after a 1.0s delay during 171 which the screen is blank. Participants practiced the task over 16 trials and then completed 172 two blocks of 96 trials each, including both shape and location cued recalls. Trial order was 173 randomised, minimizing participants' ability to predict whether a shape or location probe

Spatially specific binding impairments

174 would follow the sample display. To ensure that patients had not forgotten the task

175 instructions, at the end of each block they were asked to describe what they had done.

176

177 <u>Centroid estimation</u>

178 This task assesses the accuracy and precision of estimates of the average location of 179 three visible white disks and is illustrated in figure 1B. Each disk's diameter is 0.27°. 180 Participants place a crosshair-shaped cursor at the estimated centroid location and click 181 the mouse. Following a 1.0s interval, a novel set of disks appears and the procedure 182 is repeated. Disks can occupy any three of seven canonical locations, including the screen 183 center and the vertices of a virtual concentric hexagon, with a side length of 3.67°. All 184 permutations of three canonical locations, less any resulting in a collinear configuration, are 185 used as test arrays. Each possible permutation appears twice, for a total of 64 186 trials. Pseudorandom, zero mean, independent circular Gaussian distributions, with a 187 standard deviation of 0.6° , are sampled to jitter each disk's position. Prior to testing, 188 instructions were read to the participants. The centroid was defined as the point where the 189 triangle, whose vertices coincided with the disks' locations, would balance in the horizontal 190 plane (Baud-Bovy and Soechting, 2001). Participants completed 25 practice trials, without 191 feed-back, followed by two blocks of 64 trials each. 192

193 Neuropsychological tests

Three neuropsychological instruments were used to assess participants: 1) Hopkins Verbal
Learning Test – Revised (HVLT-R), 2) Brief Visuospatial Memory Test – Revised (BVMTR), and 3) Color Trails Test (CTT). The Arabic version of these tests were recently validated
(Al-Joudi *et al.*, 2019).

199 Data analysis

In the recall task, participants could either report 1) the color of the target, correct response, 2) the color of the non-target item, that is make a swap error, or 3) the color absent from the sample, that is make a generic error. We approached the group level hypothesis testing as a metanalysis of prevalence data, treating each participant as a separate study. All inferential analysis presented in this study is Bayesian.

205

206 <u>Group differences in recall error rates</u>

207 Group level effects were analysed with mixed Bayesian ANOVAs using the JASP

software (JASP Team, 2021; jasp-stats.org). The between group variable coded whether the

209 participant was 1) a healthy control, 2) a patient following left or 3) right temporal

210 lobectomy, respectively. The within group variables were probe dimension, i.e., whether a

shape or space probe was used to cue recall, and block order, i.e., first or second block. The

212 error rates were normalised with a Friedman-Tukey double arcsine transformation

213 (Barendregt et al., 2013). The transformation stabilizes error rates variances and allows the

214 use of parametric methods to compute group statistics:

215
$$t = \sin^{-1} \sqrt{\frac{c}{n}} + \sin^{-1} \sqrt{\frac{c}{n+1}}$$

where c is the number of either swap or generic errors and n is the total number of trials for each probe dimension and block. The group average proportions were obtained by applying the following inverse transformation to the transformed proportions group means:

$$\hat{p} = \left[1 - sgn(\cos\hat{t}) \cdot \sqrt{1 - \left(\sin\hat{t} - \frac{\sin\hat{t} - 1/\sin\hat{t}}{n}\right)^2}\right] \cdot 0.5$$

219

220 Analysis of centroid task

The analysis of the centroid task was carried out by fitting the following regressionmodel to each participants' reports:

$$\begin{pmatrix} \boldsymbol{r}_{x} \\ \boldsymbol{r}_{y} \end{pmatrix} = \begin{pmatrix} a_{0} \\ b_{0} \end{pmatrix} + \begin{pmatrix} a_{1} & 0 \\ 0 & b_{1} \end{pmatrix} \begin{pmatrix} \boldsymbol{C}_{x} \\ \boldsymbol{C}_{y} \end{pmatrix} + \begin{pmatrix} \boldsymbol{N}_{x} \\ \boldsymbol{N}_{y} \end{pmatrix}$$

where r_x and r_y are the *x-y* screen coordinates of the centroid estimates, C_x , C_y are the centroid horizontal and vertical screen coordinates and N_x and N_y are the normal distributions of the respective residuals. Although incenter biases are also known to affect centroid estimates (Baud-Bovy and Soechting, 2001), we did not include them in the model for sake of clarity and because a preliminary analysis did not reveal appreciable group differences. Group and screen coordinates differences in static offsets, i.e. a_0 and b_0 , accuracy, i.e. a_1 and b_1 , and precision, i.e. the variance of N_x and N_y , were assessed with Bayesian mixed ANOVAs.

230	Results
231	Participants' demographic, clinical and neuropsychometric characteristics
232	Table 1 reports the demographic and clinical characteristics of the left and right
233	temporal lobectomy patients and healthy controls. The groups were matched on age, gender
234	and educational level. Both patient groups showed a lower full-scale IQ that healthy controls,
235	however all of the variables were more likely to reflect a null effect than a group
236	difference. Table 2 details the sex, education level and neuropsychometric performance of
237	the thirteen patients. Both raw scores as well the values normalised on the basis of a reference
238	sample of healthy controls, whose first language is Arabic (Al-Joudi et al., 2019), are shown.
239	We examined whether patients showed a material specific pattern of lateralized deficits (e.g.
240	Saling, 2009) by comparing performance of the left and right temporal lobectomy patients on
241	the HMVT and the BVMT with Bayesian independent samples t-tests (JASP
242	Team, 2021; jasp-stats.org). There was moderate strength evidence for left temporal
243	lobectomy patients having worse delayed verbal recall on the HVLT than controls ($BF_{10} =$
244	7.99), while there was anecdotal evidence for no group difference in the delayed visuo-spatial
245	recall (BF ₁₀ = 0.46). In Table 3 the MTL structures affected by the surgical excisions are
246	listed, patient by patient, while Table 3-1 (see supplementary material) shows representative
247	postsurgical axial, sagittal and coronal multimodal MRI slices for each patient.
248	
249	Cued color recall performance
250	In the cued recall task, participants completed two blocks of 96 trials each. Mixed

effect Bayesian ANOVAs were used to examine the influence of three factors on recall: 1)
group, namely whether participants were controls, patients following left and right temporal
lobectomy, respectively; 2) block and 3) probe dimension. Generic and swap errors were
analysed separately.

Extended data table 4-1 reports the result of the ANOVA model comparison for swap errors. The model with the highest posterior probability included group, probe dimension and block, as well as the interaction of group by probe dimension. Table 4 summarises the evidence for each predictor. There was moderate evidence for an effect of probe dimension and block. There was very strong evidence for an effect of group and strong evidence for an interaction of group by probe dimension.

Extended data table 5-1 summarises the results of the model comparisons for generic errors. The best model was the one which included all three factors, but none of the interactions. Table 5 summarises the analysis of the effects. There was anecdotal evidence for the block and probe dimension affecting the proportion of generic errors. On the other hand,

there was anecdotal evidence for a null effect of group.

Figure 2 shows the mean proportion of swap and generic errors following space and shape probe, respectively. Controls made fewest swap and generic errors. Controls and patients with right temporal lobectomies made more swap errors following shape than space probes. Patients with left temporal lobectomies made most swap errors and three of them made more swap errors following space than shape probes. Participants made more generic errors following space than shape probe. Participants also made more swap and generic errors in the first than second block (data not shown).

273

274 <u>Centroid estimation performance</u>

We examined how group and screen coordinates affected three indices of performance in the centroid estimation task: 1) static offsets, 2) accuracy and 3) precision (see Methods). For static offsets the model with the highest posterior probability included the effect of screen coordinates only. In fact, there was strong evidence, $BF_{inc} = 32.13$, that the horizontal and vertical offsets differed. While there was no appreciable horizontal bias, $m = 0.0^{\circ}$, 95%CI =

280	$[-0.04^{\circ}, 0.05^{\circ}]$, participants showed an upward bias, $m = 0.15^{\circ}, 95\% CI = [0.07^{\circ}, 0.23^{\circ}]$.
281	There was moderate evidence in favour of the null and against both an effect of group, $\mathrm{BF}_{\mathrm{inc}}$
282	= .22, and its interaction with coordinate, BF_{inc} = .28. For accuracy, the model with highest
283	posterior probability included group. However, there was anecdotal evidence for a null effect
284	of group, $BF_{inc} = .76$, with moderate evidence in favour of a null effect of coordinate ($BF_{inc} = .76$)
285	.26), and its interaction with group, $BF_{inc} = .28$. For precision, namely the variance of the
286	variable errors, the null model had the highest posterior probability. There was anecdotal
287	evidence for a null effect of screen coordinate, $BF_{inc} = 0.4$, anecdotal evidence for a null
288	effect of group, $BF_{inc} = 0.53$, and moderate evidence for a null interaction of group and
289	dimension, $BF_{inc} = 0.15$.

290	Discussion
291	In this study we compared performance of healthy controls and temporal lobectomy
292	patients in two tasks, one probing conjunctive binding in vWM, the other perceptual, spatial
293	averaging of disks patterns. The vWM task required the recall of a target's colour, where the
294	target was identified by either a location or a shape probe. The task was thus designed to
295	determine dimensional specificity of working memory conjunctive binding. Controls were
296	more accurate than patients overall. Moreover, they made fewer swap errors following space
297	than shape probes, while left temporal lobectomy patients made more swap errors following
298	space than shape probes. There was no evidence of group differences in static biases,
299	accuracy and precision of perceptual centroid estimates. The implication of these findings for
300	the organization of binding and spatial processes in vWM is discussed in the next paragraph,
301	following a brief overview of prior evidence.
302	
303	MTL lesions specifically disrupt spatial binding in vWM
304	Previous studies addressed whether MTL pathology is associated with impairments in
305	vWM binding. The ability to recall shape-color, shape-location or item-item conjunctions has
306	been found to be diminished in patients with anoxic/ischemic or infectious pathology
307	involving the MTL as well as neurodegenerative disorders, suggesting an impairment in
308	conjunctive and relational binding (Hannula et al., 2006; Olson et al., 2006; Parra et al.,
309	2009). van Geldorp, Bouman, Hendriks and Kessels (2014) compared patients, who had
310	undergone anterior temporal lobectomies for medically refractory epilepsy, and healthy
311	controls' performance in four match-to-sample tasks. The tasks were difficult and required
312	participants to remember three separate frames presented sequentially. Each frame contained
313	the picture of a face and a building, which differed in location and color. A cue, presented
314	before the sample indicated whether participants should only remember the identity of the

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315	items or also their location (spatial binding condition), color (color binding condition) or the
316	item they had been presented with (relational binding condition). Overall, recall was less
317	accurate in patients compared to controls, and particularly so in the relational binding
318	condition. Recall performance in the spatial and color binding conditions were equally
319	affected, suggesting that spatial and non-spatial WM binding were not differentially
320	compromised in these patients. Zokaei et al. (2019) found that patients, who had undergone a
321	temporal lobectomy, made more swap errors when recalling the location of fractal patterns,
322	compared to controls. Since neither fractal recognition nor memory for locations were found
323	to be appreciably impaired in these patients, it was inferred that they suffered a primary
324	binding deficit. Using a similar paradigm, Pertzov et al. (2013) documented both a spatial as
325	well as a non-spatial binding deficit in individuals recovering from autoimmune encephalitis,
326	suggesting that binding impairments due to MTL dysfunction are not dimensionally specific.
327	Braun et al. (2011) concluded instead that TLE patients, who had undergone a right temporal
328	lobectomy, were only impaired when the vWM task required spatial binding, but performed
329	similarly to healthy controls when it required binding of non-spatial features.
330	Our own findings contribute new, crucial evidence for understanding the role of MTL
331	in vWM binding by confirming the association between MTL pathology and spatial binding
332	impairments, unaccounted for by impairments of either spatial vision or feature memory. We
333	found an increase in the proportion of swap errors in the TLE group. Crucially, while healthy
334	participants made significantly more swap errors following the shape probe, left temporal
335	lobectomy patients made more swap errors following the space probe. In healthy participants
336	the results are therefore in keeping with the hypothesis that binding of non-spatial features is
337	mediated by the features' shared location (e.g., Schneegans and Bays, 2017; Treisman and

339 compared to space probes, because in the latter case both target shape and color need to be

Zhang, 2006). In fact, the likelihood of swap errors should be greater following shape

340	bound to the target location before they can be bound to each other (Schneegans and Bays,
341	2017). On the other hand, the observation that some patients made significantly more swap
342	errors following space probes than controls may indicate that patients gained the ability to
343	bind non-spatial features directly, without the mediation of a shared location, allowing them
344	to achieve higher accuracies following shape than space probes. Whether this inference is
345	warranted remains to be established. Regardless, the group level pattern of dimensionally
346	specific binding impairments observed in left TLE patients replicates a previous observation
347	in a stroke patient with bilateral MTL damage, found to be impaired only in vWM tasks
348	requiring spatial binding, but not those requiring non-spatial binding (Dundon et al., 2017).
349	These observations thus allow us to draw the following conclusion: MTL pathology can be
350	associated with WM binding impairments that are spatially specific and reverse the spatial
351	advantage characteristic of healthy controls. If processes underlying spatial binding in vWM
352	are independent from processes devoted to binding of non-spatial features, then the role of
353	space in organising vWM may extend beyond providing a common index for the conjunctive
354	binding of visual features.

356 Hemispheric lateralization and binding

357 Bohbot et al. (1998) found that patients who had undergone thermocoagulation of 358 structures within the right, but not the left MTL were more impaired in a number of spatial 359 WM tasks than those who had not undergone surgery, suggesting that right MTL structures 360 may play an overarching role in spatial memory. Braun et al. (2011) compared patients with 361 right temporal lobectomy and healthy controls' performance on a number of single feature 362 and feature conjunction recall tasks and concluded that these patients are specifically 363 impaired in spatial binding. However, the tasks employed memory samples of different 364 complexity to probe recall of features and conjunctions, respectively, thus introducing a load

365	confound in the comparison. Our own results are in keeping with the idea that left rather than
366	right MTL structures are specifically involved in spatial binding, since patients who had
367	undergone left temporal lobectomies showed greater spatial binding impairments than
368	patients with right temporal lobectomies. While our results need to be interpreted cautiously,
369	given the small sample size, they agree with the conclusion drawn by Kessels et al. (2004),
370	who found that patients who had undergone left, but not right temporal lobectomies were
371	impaired in spatial binding, confirming lateralization effects previously observed by the same
372	group in a sample of patients with cerebrovascular pathology (Kessels et al., 2002). Spiers et
373	al. (2001) found that left temporal lesion specifically affect object location binding while
374	right temporal lesions affect spatial memory more generally. However, the existing literature
375	remains inconclusive to the relation between laterality and working memory spatial binding.
376	A distinct view of lateralization of binding impairments is that the latter reflect attention
377	deficits which follow cortical strokes, especially involving the non-dominant right
378	hemisphere. For example. Cohen-Dallal et al. (2021) reported that patients with unilateral
379	attentional neglect show delay-dependent decrements in spatial binding performance. In light
380	of the fact that parahippocampal damage is associated with unilateral neglect (Mort et al.,
381	2003), Cohen-Dallal et al.'s finding raises the possibility that lateralized attentional deficits
382	may also contribute to the binding impairments observed in our study. However, performance
383	in a centroid estimation task did not show group differences in lateralized static biases,
384	suggesting that lesion in our sample was not associated with unilateral neglect. Moreover,
385	patients showed neither diminished accuracy nor lower precision in the centroid task
386	compared to controls, indicating that spatial perception and attention were not compromised
387	(Drew <i>et al.</i> , 2010).
388	A further concern is that uncontrolled verbal strategies may have confounded the

389 interaction of lesion laterality and probe dimension. We found in pilot studies that healthy

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390 participants maintain a spatial advantage in WM binding under condition of articulatory 391 suppression and therefore concluded that the spatial advantage in WM binding does not 392 depend on verbal strategies (unpublished data). However, we cannot rule out the possibility 393 that left temporal lobectomy patients used a verbal strategy and thus managed to selectively 394 improve binding of shape and color.

395 It is important to note, with regard to the issue of localization and the nature of cognitive 396 impairments encountered in TLE patients, that group level results belie substantial inter-397 individual differences (see Figure 2). Previous electrophysiological recordings from the left 398 MTL indicated that trial by trial changes in the amplitude of low-frequency oscillations, 399 obtained during encoding episodes, predict subsequent recall accuracy of object/place 400 conjunctions in TLE patients (Miller et al., 2018), in keeping with our own conclusion that 401 spatial binding is dependent on left lateralized processes. Interestingly, this was not the case 402 in all suggesting that spatial binding processes are not universally left lateralised in these 403 patients. Unfortunately, the study did not report whether the lateralization of spatial binding 404 processes was affected by the laterality of the seizure focus, precluding firmer conclusion 405 regarding the relation between the two. Other studies have, however, demonstrated 406 anomalous lateralization in high proportion of TLE patients, as memory processes often shift 407 to the contra-lesional hemisphere both pre (Bellgowan et al., 1998; Golby et al., 2002; 408 Janszky et al., 2004) and postoperatively (Sidhu et al., 2016). These findings may provide 409 one possible interpretation of the interindividual differences highlighted above, namely that 410 neural plasticity in some TLE patients modifies the lateralization of memory processes 411 usually encountered in healthy controls. An alternative explanation, initially born out of 412 observations in non-human primates with focal lesions (Browning et al., 2010; Croxson et al., 413 2012), is that effects of MTL functional and structural damage in postsurgical TLE patients 414 may be attenuated by non-lateralized recruitment of neocortical areas (Sidhu et al., 2013). To

415 determine whether either or both of these hypotheses can account for interindividual 416 differences in spatial binding impairments and to what extent other factors, like age of seizure 417 onset, severity and frequency of seizures, neuropsychiatric complications and antiepileptic 418 medications, known to affect neural plasticity and degree of cognitive impairment (Bell et al., 419 2011), also contribute to hemispheric lateralization in TLE patients will require new 420 experimental evidence. 421 Despite the potential confound listed above, the present study indicates that following 422 left temporal lobectomy, vWM binding is diminished in a dimensionally specific manner in

423 the absence of appreciable perceptual, attentional or visual-spatial memory deficits.

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580	Figure 1. Tasks' structure. Panel A shows the event sequence in the color recall task.
581	Participants had to remember the color of a triangle and a square displayed side by side. The
582	sample display was followed by a pattern mask and a blank screen. The recall target was
583	identified either by a space probe, consisting a bright cross displayed at the location.
584	previously occupied by the target, or by a shape probe, consisting of the outline of the target
585	shape. The color was reported by placing the cursor over the corresponding rectangle and
586	clicking the mouse button. Panel B shows the centroid estimation task. The visual display
587	contained 3 bright dots and the participant had to indicate the location of the center of mass
588	of the imaginary triangle whose vertices corresponded to the dots location, by dragging the
589	cursor and clicking.

Legends

590

591 Figure 2. Recall error rates. The bar graph on the left shows the group averaged proportions 592 of swap errors, while the bar graph on the right shows the group averaged proportions of 593 generic errors, following space and shape probes, respectively. Overall patients made more 594 swap errors than controls. Moreover, patients with left temporal lobectomies made more 595 swap errors following space than shape probes, suggesting a specific impairment of spatial 596 binding in this group only. For generic errors, group differences were marginal and were not 597 further affected by probe dimension. For sake of clarity, the data are averaged over the two 598 blocks. Circles are individual participants' error rates. Continuous lines join swap error rates, 599 following space and shape probes respectively, of each left temporal lobectomy patient. 600 Broken lines join the swap error rates of each right temporal lobectomy patient. Error bars 601 are standard errors of the mean.

602

Table 1. Demographic and clinical sample characteristics. Group frequencies were
compared using a Bayesian contingency table. Continuous variables were compared with
Bayesian ANOVAs or Bayesian independent samples t-test. The values in parenthesis are
standard deviations. None of the demographic or clinical variables showed appreciable group
differences since the Bayes factor (BF₁₀) was less than 1.0 for all comparisons.

608

609	Table 2. Demographics a	nd neuro-psychometric	performance of TLE	patients. Raw

610 scores are reported for each test and participant (see methods). The corresponding normalised

611 values are shown in parenthesis. Normalised z-scores values were computed by subtracting

612 the mean score and dividing by the standard deviation of a reference sample (Al-Joudi et al.,

613 2019). To facilitate inspection of the table scores corresponding to better than mean

614 performance are underlined. HVLT-R = Hopkins Learning Test – Revised; BVMT-R = Brief

615 Visuospatial Memory Test – Revised; CTT = Colour Trails Test.

616

617 Table 3. Patients' lesion anatomy. The table lists the pathology and regions affected by the 618 lobectomy for each patient. Table 3-1 of the extended data shows representative MRI slices 619 through the medial temporal lobes. GG, ganglioglioma; MG, meningioma; MTS, medial 620 temporal sclerosis; HIP, hippocampus; ERC, entorhinal cortex; PRC, perirhinal cortex; PHC, 621 parahippocampal cortex; ITC, inferotemporal cortex; MTG, Middle temporal gyrus; ATP, anterior temporal pole; STG, superior temporal gyrus; AMG, amygdala. "0" indicates an 622 623 unaffected subregion, "+" a rostro-caudal lesion extent up to 20 mm, and "++" up to 40 624 mm.

625

Table 3-1 (Extended data). Post-surgical MRI scans for thirteen patients. The images
were obtained with T1 weighted, T2 weighted, Fluid-attenuated Inversion Recovery and

628 Gradient Recalled Echo sequences. Legend: ERC, Entorhinal Cortex; PHC, Parahippocampal 629 Cortex; PRC, Perirhinal Cortex; Hipp, Hippocampus; AMG, Amygdala; Temporal pole, TP; 630 antSTG, anterior Superior Temporal Gyrus; antMTG, anterior Middle Temporal Gyrus. 631 632 Table 4. Swap errors – analysis of effects. The table lists each factor and interaction for 633 swap error rates. Table 4-1 of the extended data lists the models and associated prior and

634 posterior probabilities from which the values in the present table are computed. P(incl) is the

635 prior probability of the effect; P(incl|d) is the posterior probability of the effect; BF_{incl} is

636 Bayes factor. A BF greater than 1.0 favours the effect, a BF less than 1.0 favours a null

instead. Values of the BF greater than 3.0 are in bold, to highlight those effects that have at 637

638 least moderate evidence in their favour. Block, probe dimension, group and the interaction of

639 group by probe dimension all have at least moderately strong evidence in their favour.

640

641 Table 4-1 (Extended data). Best model comparisons for swap errors. The table presents

642 each of the model comparisons from the Bayesian ANOVA. The within factors are block (B)

643 and probe dimension (D). The between factor is group G. P(M) is the a-priori model

644 probability, P(M|d) is the posterior model probability. BF_M is the Bayes factor of the model,

645 BF₁₀ is the Bayes factor of the model relative to the best one. The best model contained all

646 three factors and the interaction of group by probe dimension.

647

648 Table 5. Generic error – analysis of effects. The table lists each factor and interaction for

649 generic error rates. Table 5-1 of the extended data lists the models and associated

650 probabilities, used to compute the effects. P(incl) is the prior probability of the effects;

651 P(incl|d) is the posterior probability of the effect; BF_{incl} is Bayes factor. A BF greater than 1.0

652 favours the effect, a BF less than 1.0 favours a null effect instead. The only predictors with

653 favourable evidence, albeit of very modest entity, are block and probe dimension.

- 655 **Table 5-1 (Extended data). Best model comparisons for generic errors.** The table presents
- 656 the model comparisons obtained from a Bayesian ANOVA. The within factors are block (B)
- and probe dimension (D). The between factor is group G. P(M) is the a-priori model
- probability, P(M|d) is the posterior model probability. BF_M is the Bayes factor of the model,
- BF_{10} is the Bayes factor of the model relative to the best one. The best model included the
- 660 three main factors, namely block (B), probe dimension (D) and group (G).





	left TLE	right TLE	controls	BF_{10}
	(<i>n</i> =6)	(<i>n</i> =7)	(<i>n</i> =15)	
Sex (%males)	83.3	100	93.3	.64
Age (years)	35.2	33.08	33.3	.29
	(±7.9)	(±8.9)	(±7.8)	
Education (highest grade)	15.3	14.3	14.3	.5
	(±1.6)	(±1.9)	(±2.4)	
Full Scale IQ	92.0	87.3	99.5	.9
	(±11.7)	(±9.9)	(±15.5)	
Epilepsy onset age (years)	11.1	22.1		.84
	(±10.2)	(±17.2)		

Table 1. Demographic and clinical sample characteristics.

Frequency group differences were compared using a Bayesian contingency table. Continuous variables were compared with Bayesian ANOVAs or Bayesian independent samples t-test. The values in parenthesis are standard deviations. None of the demographic or clinical variables showed appreciable group differences since the Bayes factor (BF_{10}) was less than 1.0 for all comparisons.

Table 2. Demographics and	l neuro-psychometric	performance of	TLE patients
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		•		•										
Patient	Gender	Education (years)		WA	IS II			HVLT			BVMT		C	T
			Block Design	Vocab	Matrix Reasoning	Similar	Immediate	Delayed	Discrimin	Immediate	Delayed	Discrimin.	CTT1 (sec)	CTT2 (sec)
Pl	М	16	16 (59)	18 (-1.6)	12(41)	24 (77)	19 (-1.43)	6 (-1.47)	9 (-1.9)	12 (-1.01)	4(-1.22)	4 (-1.4)	85 (1.24)	160 (2.0)
P2	М	14	18 (46)	27 (83)	12 (41)	23 (88)	26 (.36)	<u>9 (.11)</u>	<u>12 (1.1)</u>	10 (-1.28)	4 (-1.22)	4 (-1.4)	97 (1.74)	166 (2.22)
P3	F	16	12 (88)	25 (-1.0)	18 (.53)	28 (31)	24 (15)	7 (95)	10 (9)	9 (-1.41)	6 (58)	5 (4)	65 (.41)	130 (.92)
P4	М	14	16 (59)	26 (91)	14 (09)	26 (54)	22 (67)	7 (95)	<u>11 (.1)</u>	14 (75)	5 (9)	4 (-1.4)	66 (.45)	191 (3.12)
P5	М	12	14 (74)	28 (74)	10 (72)	25 (66)	24 (15)	8 (42)	8 (-2.9)	7 (-1.68)	3 (-1.55)	3 (-2.4)	65 (.41)	135 (1.1)
P6	М	16	24 (0)	20 (-1.48)	16 (.22)	22 (-1.0)	24 (15)	7 (95)	<u>11 (.1)</u>	<u>21 (.19)</u>	7 (26)	5 (4)	75 (.82)	150 (1.65)
P7	М	16	14 (74)	28 (74)	8 (-1.03)	25 (66)	19 (-1.43)	5 (-2.0)	7 (-3.9)	12 (-1.01)	5 (9)	4 (-1.4)	87 (1.33)	154 (1.79)
P8	М	16	<u>39 (1.09)</u>	<u>45 (.71)</u>	21 (1.0)	30 (08)	27 (.62)	<u>10 (.63)</u>	<u>12 (1.1)</u>	13 (88)	<u>9 (.39)</u>	5 (4)	32 (97)	85 (7)
P9	М	14	27 (.21)	32 (4)	<u>18 (.53)</u>	26 (54)	21 (92)	8 (42)	<u>11 (.1)</u>	<u>21 (.19)</u>	7 (26)	5 (4)	<u>52 (13)</u>	105 (.02)
P10	М	14	19 (37)	22 (-1.26)	9 (88)	24 (77)	18 (-1.69)	7 (95)	10 (9)	17 (35)	7 (26)	4 (-1.4)	94 (1.62)	165 (2.19)
P11	М	11	26 (.14)	22 (-1.26)	16 (.22)	24 (77)	20 (-1.18)	7 (95)	10 (9)	15 (61)	6 (58)	4 (-1.4)	80 (1.03)	151 (1.68)
P12	М	12	13 (81)	23 (-1.17)	12 (41)	22 (-1.0)	19 (-1.43)	5 (-2.0)	8 (-2.9)	12 (-1.01)	5 (9)	4 (-1.4)	86 (1.28)	153 (1.75)
P13	М	16	17 (52)	28 (74)	11 (56)	28 (31)	21 (92)	8 (42)	10 (9)	14 (75)	6 (58)	4 (-1.4)	83 (1.16)	158 (1.94)

Raw scores are reported for each test and participant (see methods). The corresponding normalised values are shown in parenthesis. Normalised z-scores values were computed by subtracting the mean score and dividing by the standard deviation of a reference sample (Al-Joudi et al., 2019). To facilitate inspection of the table, scores better than mean reference performance are underlined. HVLT-R = Hopkins Learning Test – Revised; BVMT-R = Brief Visuospatial Memory Test – Revised; CTT = Colour Trails Test.

				Temporal lobe structures								
	Age	Pathology	Lesion	HIP	ERC	PRC	PHC	ITG	MTG	ТР	STG	AMG
			laterality									
P1	25	GGs	L	0	+	+	0	0	0	+	0	0
P2	25	MTS	R	++	0	0	0	0	0	+	0	++
Р3	50	MG	L	0	0	+	0	0	+	+	0	0
P4	41	GGs	R	0	+	+	0	0	0	+	0	0
Р5	31	MTS	R	++	+	0	+	0	+	+	+	++
P6	27	MTS	L	++	0	0	0	0	0	0	0	0
P7	40	MG	L	0	0	0	0	0	+	+	0	0
P8	32	MTS	R	++	0	0	0	+	0	++	0	+
P9	33	MTS	R	++	+	0	++	+	0	0	+	+
P10	22	MTS	L	+	+	0	0	++	0	0	+	0
P11	25	MTS	R	++	+	0	0	0	0	0	+	++
P12	47	GGs	L	0	+	0	0	0	++	++	+	0
P13	32	MTS	R	+	+	0	+	+	0	0	+	0

Table 3. Patients' lesion anatomy

The table lists the pathology and regions affected by the lobectomy for each patient. Table 3-1 of the extended data shows representative MRI slices through the medial temporal lobe. GG, ganglioglioma; MG, meningioma; MTS, medial temporal sclerosis; HIP, hippocampus; ERC, entorhinal cortex; PRC, perirhinal cortex; PHC, parahippocampal cortex; ITC, inferotemporal cortex; MTG, Middle temporal gyrus; ATP, anterior temporal pole; STG, superior temporal gyrus; AMG, amygdala. "0" indicates an unaffected subregion, "+" a rostro-caudal lesion extent up to 20 mm, and "++" up to 40 mm.

Table 4. Swap errors - analysis of effects

Effects	P(incl)	P(incl d)	BFincl
В	0.737	0.941	5.673
D	0.737	0.925	4.414
G	0.737	0.997	119.755
B•D	0.316	0.206	0.563
G•B	0.316	0.188	0.500
G•D	0.316	0.891	17.629
G•B•D	0.053	0.017	0.317

The table lists each factor and interaction for swap error rates. In the extended data table 4-1 lists the models and associated prior and posterior probabilities from which the values in the present table are computed. P(incl) is the prior probability of the effect; P(incl/d) is the posterior probability of the effect; BF_{incl} is Bayes factor. A BF greater than 1.0 favours the effect, a BF less than 1.0 favours a null instead. Values of the BF greater than 3.0 are in bold, to highlight those effects that have at least moderate evidence in their favour. Block, probe dimension, group and the interaction of group by probe dimension all have at least moderately strong evidence in their favour.

Effects	P(incl)	P(incl d)	BFincl
В	0.737	0.804	1.463
D	0.737	0.815	1.571
G	0.737	0.641	0.637
B•D	0.316	0.142	0.360
G•B	0.316	0.085	0.202
G•D	0.316	0.087	0.206
G•B•D	0.053	7.157e -4	0.013

Table 5. Generic errors - analysis of effects

The table lists each factor and interaction for generic error rates. Table 5-1 of the extended data lists the models and associated probabilities, used to compute the effects. P(incl) is the prior probability of the predictors; P(incl/d) is the posterior probability; BF_{incl} is Bayes factor. A BF greater than 1.0 favours the predictor, a BF less than 1.0 favours a null effect instead. The only predictors with favourable evidence, albeit of very modest entity, are block and probe dimension.