

Scene-selectivity in CA1/subicular complex: Multivoxel pattern analysis at 7T

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ABSTRACT

Prior univariate functional magnetic resonance imaging (fMRI) studies in humans suggest that the anteromedial subicular complex of the hippocampus is a hub for scene-based cognition. However, it is possible that univariate approaches were not sufficiently sensitive to detect scene-related activity in other subfields that have been implicated in spatial processing (e.g., CA1). Further, as connectivity-based functional gradients in the hippocampus do not respect classical subfield boundary definitions, category selectivity may be distributed across anatomical subfields. Region-of-interest approaches, therefore, may limit our ability to observe category selectivity across discrete subfield boundaries. To address these issues, we applied searchlight multivariate pattern analysis to 7T fMRI data of healthy adults who undertook a simultaneous visual odd-one-out discrimination task for scene and non-scene (including face) visual stimuli, hypothesising that scene classification would be possible in multiple hippocampal regions within, but not constrained to, anteromedial subicular complex and CA1. Indeed, we found that the scene-selective searchlight map overlapped not only with anteromedial subicular complex (distal subiculum, pre/para subiculum), but also inferior CA1, alongside posteromedial (including retrosplenial) and parahippocampal cortices. Probabilistic overlap maps revealed gradients of scene category selectivity, with the strongest overlap located in the medial hippocampus, converging with searchlight findings. This was contrasted with gradients of face category selectivity, which had stronger overlap in more lateral hippocampus, supporting ideas of parallel processing streams for these two categories. Our work helps to map the scene, in contrast to, face processing networks within, and connected to, the human hippocampus.

1. Introduction

There is increasing evidence that the human hippocampus has roles in behaviours beyond purely-mnemonic cognitive functions, including complex visual perception and imagination (Aly et al., 2013; Graham et al., 2010; Hodgetts et al., 2015; Lee et al., 2013; Martin and Barense, 2023). While the precise role played by the hippocampus in supporting such functions is debated (Mayes et al., 2007; Turk-Browne, 2019), there is increasing support for the idea that scenes are central to hippocampal information processing (Gaffan, 1991; Maguire and Mullally, 2013; Murray et al., 2017, 2018; Zeidman et al., 2015b). Indeed,

scene-selective impairments following hippocampal damage are not only seen on memory tasks (Bird et al., 2008; Hartley et al., 2007; Taylor et al., 2007) but also on tasks of complex visual perception (Graham et al., 2010; Lee et al., 2012; Erez et al., 2013; Martin and Barense, 2023). Some of the strongest evidence for this comes from ‘oddy’ simultaneous visual discrimination paradigms (Lee et al., 2005, 2006), modified from the nonhuman animal literature (e.g., see Buckley et al., 2001). In these tasks, participants are presented with an array of scene or non-scene stimuli on each trial (typically 3 or 4 items per trial) and are required to select the odd-one-out as quickly and as accurately as possible (see Fig. 1). Critically, this task has almost no mnemonic

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demand, as stimuli are never repeated during the task, and all compared items are presented concurrently.

Using a four-choice version of this scene-odddity task, Lee and colleagues (Lee et al., 2005, 2006) found that individuals with hippocampal lesions were impaired when having to identify incongruent scenes but not other classes of stimuli (e.g., faces, objects and colour). This scene-selective impairment was observed when scenes were presented from different viewpoints, but not when the scenes were shown from the same viewpoint. In contrast, patients with both hippocampal and perirhinal cortex (PRC) damage demonstrated significant deficits for both scenes and faces shown from different views but performed normally for scenes and faces shown from the same view (Lee et al., 2005; see also Gardette et al., 2023).

Functional magnetic resonance imaging (fMRI) studies in healthy participants have attempted to further elucidate the nature and topography of hippocampal scene processing. Earlier studies, for example, primarily reported involvement of the posterior hippocampus in scene perception, including during oddity discrimination tasks (Barense et al., 2010; Lee et al., 2008). However, more recent studies have since suggested that an anteromedial hippocampal region may be a key ‘hub’ for scene-based cognition, including ‘compositional’ aspects of scene perception that require construction of a global scene ‘model’, such as

across tasks (and indeed individuals) within the hippocampus is unsurprising given that it is not a single homogeneous structure. The hippocampus can be organised into subfields (e.g., CA1, CA2, CA3, dentate gyrus and subiculum) based on cytoarchitectonic features (Schultz and Engelhardt, 2014), and further functional subdivisions may be proposed along both the anterior-posterior and medial-lateral axes based on connectivity patterns (Aggleton and Christiansen, 2015; Christiansen et al., 2017; Dalton et al., 2019; Ezama et al., 2021; Fritch et al., 2021), microstructure (Genon et al., 2021), gene expression patterns (Vogel et al., 2020), and cellular dynamics (e.g., the receptive field size of place cells) (Brunec et al., 2018; Strange et al., 2014). It should be noted that human hippocampal anatomical terms ‘medial’ and ‘lateral’ mostly align with animal hippocampal terms ‘distal’ and ‘proximal’².

Given the methodological challenges associated with characterising such fine-grained functional and/or structural variation within the hippocampus using conventional 3T MRI, further studies have adopted ultra-high field MRI methods to study subfield contributions to scene-related cognition. For example, we previously used high-resolution fMRI (1.2 mm isotropic voxels) to investigate the contribution of different hippocampal subfields (CA1, CA2/3, dentate gyrus, and the subiculum) to scene perception using a 3-choice oddity paradigm (Hodgetts et al., 2017). We found that the subiculum was the only

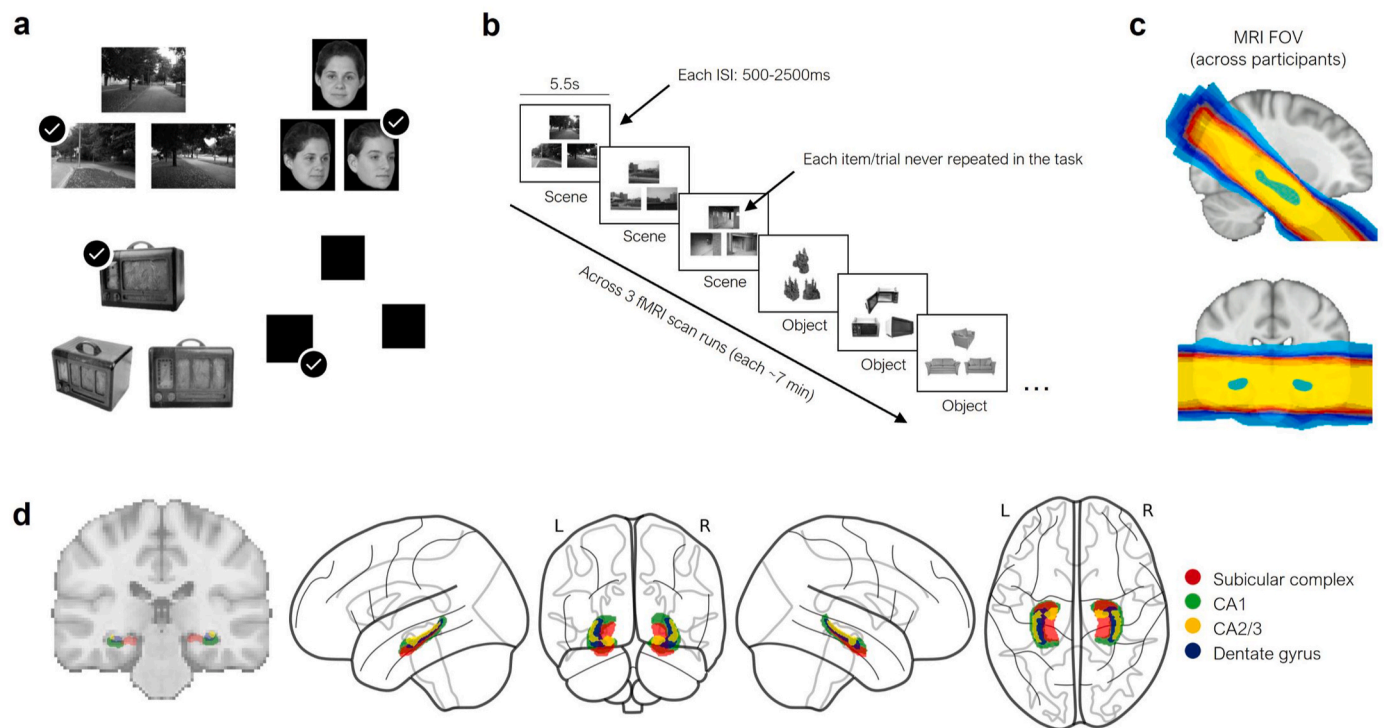


Fig. 1. The oddity behavioral task, the partial FOV, and probabilistic hippocampal subfield ROIs in 1 mm MNI template space. a) Examples of object, face, scene, and size trials (from top left to bottom right). All stimuli were trial unique. b) An illustration of the oddity task procedure. Three trials of the same category were shown sequentially, creating mini-blocks (e.g., here three scene and three face trials are shown). Each trial was presented for 5.5s and trials were separated by a jittered inter-stimulus interval (ISI) of 0.5–2.5s. c) The partial FOV of each participant overlaid in MNI template space. Warmer colours indicate higher overlap between participants. The central yellow area indicates where all participant FOVs overlapped (rendered using FSLeys; McCarthy, 2018). d) The left image shows a coronal slice through the hippocampal subfields on a 1 mm MNI standard brain. The right images show the same ROIs on a glass brain (rendered using Nilearn for Python; see Abraham et al., 2014). The ROIs are colored according to the key on the right. L: Left, R: Right.

matching scenes from different viewpoints (Baldassano et al., 2016; Hodgetts et al., 2016; Zeidman et al., 2015a; Zeidman and Maguire, 2016; Zeidman et al., 2015b). Importantly, this has been shown in larger samples and at different levels of spatial smoothing, addressing prior concerns that posterior hippocampal activations reflected signal bleed from adjacent category-selective regions of posterior parahippocampal cortex (Hodgetts et al., 2016). Such variation in scene activation foci

² The medial-lateral axis can also be described at the distal-proximal axis within the subiculum, the regions which border the presubiculum and CA1, respectively. This is also true to an extent in CA1, where medial or distal CA1 borders the subiculum. However, CA1 curves around to border CA2/3 and this is proximal CA1, which is not consistently its most lateral aspect on the anterior-posterior axis.

hippocampal substructure to show an increased univariate BOLD response during the perceptual discrimination of scenes, but not faces or objects. Further analysis showed that this was solely evident in the anteromedial aspect of the subiculum, a region that we refer to as the anteromedial subicular complex henceforth.³ This finding thus refines the results of previous 3T fMRI studies (Hodgetts et al., 2016; Lee et al., 2013; Zeidman et al., 2015b), by suggesting that the anteromedial hippocampus, and specifically the anteromedial *subicular complex*, may play a unique role in complex scene perception.

While this study and others (e.g. McCormick et al., 2021; Gardette et al., 2022) have provided important insights into subregional functional variation in the hippocampus, as well as the potentially unique importance of the anteromedial hippocampus/subicular complex in scene processing (Zeidman and Maguire, 2016; Zeidman et al., 2015b), these approaches are limited in several respects. First, as noted above, functional gradients in the hippocampus (whether arising via patterns of extrinsic connectivity, gene expression, or cellular dynamics) have been shown to not respect classical subfield boundary definitions (Aggleton and Christiansen, 2015; Dalton et al., 2019, 2022). Electrophysiological studies in animals, for instance, suggest that spatial information in CA1 is not represented uniformly along its transverse axis, but more strongly represented in proximal CA1 (Henriksen et al., 2010; Ng et al., 2018). Notably, this proximal CA1 region (defined as the CA1 region which borders CA2; Amaral et al., 1991) is also preferentially connected to distal subiculum (defined in the animal literature as the subiculum proper region which borders the presubiculum) – a region which likely overlaps with the anteromedial scene region identified in human fMRI studies (Hodgetts et al., 2016; Lee et al., 2013; Zeidman et al., 2015b). This spatially modulated circuit is also seemingly preserved through both entorhinal and parahippocampal cortices (Aggleton, 2012; Aggleton and Christiansen, 2015; Gigg, 2006; Witter and Amaral, 2021).

Extending this work to human data, a recent study from Dalton et al. (2022) applied tract density mapping to diffusion MRI data, and found that areas of high tract endpoint density tended to extend across classical subfield boundaries (e.g., across the distal subiculum-proximal pre-subiculum border, and across proximal subiculum and distal CA1 border). This finding also resonates with functional connectivity studies that show long- and transverse-axis gradients of scene selectivity, not only within entorhinal cortex, but also subicular complex (Grande et al., 2022; Maass et al., 2015; Navarro Schröder et al., 2015; see also Schultz et al., 2015). Overall, therefore, scene selectivity is likely to be distributed across, and vary within, the borders of cytoarchitecturally defined subfields – a pattern that anatomically-defined subfield ROI-based analyses cannot capture without drawing arbitrary sub-divisions (e.g., splitting subicular complex into medial and lateral components as in Hodgetts et al., 2017). Further work is required, therefore, to examine scene-selectivity using methods that enable informational content to be mapped within - and across - ROI boundaries.

Second, most previous studies examining hippocampal scene processing (including at high-resolution) have relied on standard mass-univariate analysis approaches that consider only differences in mean activation magnitude as a marker of category selectivity (e.g., Hodgetts

et al., 2015; Hodgetts et al., 2016; Hodgetts et al., 2017; Zeidman et al., 2015a; Zeidman et al., 2015b; but see Liang et al., 2013; Bainbridge et al., 2021). Critically, it is possible that nonmaximal response patterns carry important (and reliable) category-selective information that cannot be detected using standard methods (Haxby, 2012; Haxby et al., 2014). Spatial/scene information within hippocampal subregions is also likely to be encoded within a distributed neural population code that cannot be captured reliably at the level of individual neurons (Leutgeb et al., 2007; Stefanini et al., 2020) – a principle that may well hold at the level of fMRI voxels (Guest and Love, 2017; Kriegeskorte et al., 2008). Aside from this, there is also considerable evidence (particularly from nonhuman species) that hippocampal subregions outside the subiculum support aspects of spatial and/or scene processing (e.g., see Leutgeb et al., 2007; O'Keefe and Recce, 1993; Oliva et al., 2016 on neurones displaying location-specific firing fields and theta phase precession in CA1, CA2, CA3 and the DG; and Robertson et al., 1998 and Rolls, 1999, 2023 on spatial view cells in CA1, CA3 and the presubiculum).

A key question, therefore, is whether analysis approaches that are sensitive to submaximal patterns of activity, as is Multivariate Pattern Analysis (MVPA) which can identify distinct patterns in fine-grained activity responses to different stimulus categories, would be more sensitive to potential between-category differences outside of the subicular complex (for reviews of this method see Haxby, 2012; Haxby et al., 2014; Norman et al., 2006; Weaverdyck et al., 2020; Yang et al., 2012). However, this approach has not been applied in the context of hippocampal category selective effects in complex perception.

Additionally, although much recent work has placed scenes at the heart of the hippocampal contribution to cognition (Murray et al., 2018; Zeidman and Maguire, 2016), the spatially modulated hippocampal circuit involving anteromedial subicular complex may be complemented by one including lateral hippocampus (corresponding with the location of the prosubiculum/CA1). Unlike the putative spatially modulated circuit, this is proposed to carry face/object information (Dalton and Maguire, 2017; Dalton et al., 2018) due to direct links with the perirhinal cortex (PRC; Insausti and Muñoz, 2001), which is critical to performance of face oddity judgement tasks (Lee et al., 2005; Hodgetts et al., 2015; Behrmann et al., 2016). These parallel scene and face/-object processing streams are also apparent in the entorhinal cortex; human functional connectivity work has shown PRC- and PHC-preferential functional connectivity with anterior-lateral and posterior-medial entorhinal cortex, respectively which, in turn, show different connectivities with lateral subicular complex (and CA1 border) and medial subicular complex (Grande et al., 2022; Maass et al., 2015). It may be possible, therefore, that more sensitive approaches (e.g., MVPA) can identify patterns of activity that are sensitive to face-related information during visual perception, particularly in more lateral parts of the hippocampal formation, thus revealing two parallel processing streams for different information categories, involving different locations of the hippocampus (Ritchey et al., 2015).

Here, then, we addressed two specific questions. First, the extent to which, during perceptual discrimination, hippocampal activity patterns specific to scenes versus other visual categories are focused on a putative anteromedial subicular complex 'hub' (potentially corresponding with the location of distal subiculum and/or pre/para-subiculum) or are distributed more widely throughout the hippocampus. Second, whether, as suggested by Dalton and Maguire (2017), patterns of activity within the lateral hippocampus (corresponding with the location of the prosubiculum/CA1) will carry information relevant to the visual discrimination of faces/objects.

To address these questions, we applied MVPA to the high-resolution fMRI data of our original 7T oddity perceptual discrimination study (Hodgetts et al., 2017). Specifically, we used support vector machine (SVM) searchlights to examine whether (a) scene trials could be distinguished from face, object, and shape-size trials, and (b) face trials could be distinguished from scene, face, and shape-size trials, based on the activity patterns across both the whole hippocampus, as well as the

³ Note that researchers have suggested that this 'anteromedial subiculum' scene-selective region detected in fMRI studies may actually correspond to the pre/para-subiculum subregions (Dalton and Maguire (2017). The pre/para-subiculum: a hippocampal hub for scene-based cognition? *Curr Opin Behav Sci*, 17, 34–40. <https://doi.org/10.1016/j.cobeha.2017.06.001>, Ding, S. L. (2013). Comparative anatomy of the prosubiculum, subiculum, presubiculum, post-subiculum, and parasubiculum in human, monkey, and rodent. *J Comp Neurol*, 521(18), 4145–4162. <https://doi.org/10.1002/cne.23416>, meaning that the term 'subicular complex' may be more appropriate when referring to an ROI that includes multiple subicular subregions (prosubiculum, subiculum proper, the presubiculum, and the parasubiculum). We therefore differentiate between subiculum proper and subiculum complex.

extended fMRI field-of-view (FOV), that encompassed regions of post-eromedial cortex (PMC) and parahippocampal gyrus (PHC) considered to form a 'core' scene processing network (Baldassano et al., 2016; Hodgetts et al., 2016; Epstein and Baker, 2019), and regions of inferior- and superior-temporal regions considered to form a 'core' face network (Haxby et al., 2000; Bernstein and Yovel, 2015; Grill-Spector et al., 2017). Participants' hippocampal subfields were also manually segmented on ultra-high-resolution images and co-registered to a standard template. This generated a novel probabilistic atlas of hippocampal subfields that allowed us to interrogate the location and extent of scene- and object/face-related category information based on classical definitions.

Initially, searchlight classification was carried out within the hippocampus only, to the unsmoothed fMRI data (1.2 mm isotropic voxels) at the individual-level. Uncorrected significance maps (accuracy significantly differing from zero) were then overlaid in MNI space to provide insight into both the spatial distribution and inter-individual variability of scene (and object/face) related information within the hippocampus (building on Hodgetts et al., 2016). Thus, as well as potentially increasing sensitivity to the information contained within hippocampal subfields (Weaverdyck et al., 2020), this more exploratory searchlight approach has the potential to identify the distribution of informational content both within and across classic subfield ROI boundaries. Accordingly, we predicted gradients of scene and face classification overlap would differ across the hippocampus and within the subicular complex and CA1, such that scene classification overlap would be greater more medially, while face classification overlap would be greater more laterally. Second, searchlight classification was applied across the entire FOV at the individual level, but accuracy maps were transformed to MNI space where conservative, multiple comparisons corrected statistical analyses were performed. We predicted that scene selective regions would overlap with the anteromedial subicular complex, CA1 and cortical scene sensitive regions, the PMC and parahippocampal gyrus. Conversely, we predicted that face sensitive regions would overlap with cortical face sensitive regions such as the fusiform cortex and posterior superior temporal sulcus.

2. Methods

This work is a secondary analysis of previously published data (Hodgetts et al., 2017). Despite some aspects being the same, for completeness we describe the methods in full here.

2.1. Subjects

Twenty-five participants (healthy with no history of neurological or psychiatric illness; 16 females; 9 males; age range = 18–35 years; age mean = 25; age SD = 4) were recruited from the University of Oxford and Oxford Brookes University. They were fluent English speakers with normal/corrected-to-normal vision. The research was approved by the University of Oxford Central University Research Ethics Committee and the Medical Sciences Interdisciplinary Research Ethics Committee, and each participant provided written informed consent before the experiment.

2.2. MRI data acquisition

A Siemens 7T Magnetom scanner, with a 32-channel head coil (Nova Medical, MA), was used to acquire MRI data. Whole-head T1-weighted data were produced with a MPRAGE sequence (1 × 1 × 1 mm; TE = 2.82 ms; TR = 2200 ms; flip angle = 7°). Blood-oxygen level-dependent (BOLD) data were acquired using a T2*-weighted echo planar imaging (EPI) sequence. The three, oddity task fMRI runs consisted of 212 vol and took approximately 7 min each (voxel size = 1.2 × 1.2 × 1.2 mm; slices = 30; TE = 25 ms; TR = 2000 ms; flip angle = 90°; partial FOV = 192 mm (Fig. 1C); partial Fourier = 6/8; parallel imaging with GRAPPA

factor = 2; bandwidth = 1562 Hz/Px; echo spacing = 0.72 ms). Slices were oriented parallel to the hippocampal long axis. Slice acquisition occurred in a descending odd-even/interleaved order. To allow for magnetization equilibrium, three volumes were discarded at the beginning of each run. A whole brain T2*-weighted EPI volume was also acquired using identical image parameters, to facilitate co-registration of partial FOV images. To improve registration and reduce image distortion from magnetic-field inhomogeneity, a field map was acquired with the same slice orientation as the functional acquisition (TE 1 = 4.08 ms; TE 2 = 5.1 ms; TR = 620 ms; FOV = 192 mm; flip angle = 39°). Two T2*-weighted ultra-high-resolution structural images were acquired with opposite phase encoding directions (left-to-right; right-to-left; voxel size = 0.6 × 0.6 × 0.6 mm; slices = 44; TE = 25.7 ms; TR = 50 ms; partial Fourier = 6/8; FOV = 192 mm). The experimenters and radiographer aligned slices orthogonal to the hippocampal main axis, by visual inspection.

2.3. MRI pre-processing

FMRIB Software Library (FSL) (Jenkinson et al., 2012) was used to process the fMRI data. The raw data were first converted to NIfTI format. The T1-weighted images were then stripped of non-brain tissue using BET (Smith, 2002). Bias field correction was carried out using the Enhancing Neuro Imaging Genetics through Meta Analysis (ENIGMA) protocol pipeline (Thompson et al., 2020). Prior to analysis, each functional run was motion corrected and co-aligned (registered to the middle volume of the second run) using MCFLIRT (Jenkinson et al., 2002). Additional EPI pre-processing was carried out using the FMRI Expert Analysis Tool (FEAT) Version 6, including high-pass temporal filtering (Gaussian-weighted least-squares straight line fitting, with $\sigma = 50$ s) and field map unwarping using FUGUE (Jenkinson et al., 2002). No spatial smoothing was applied. Time-series statistical analysis was carried out using FMRIB's Improved Linear Model (FILM) with local autocorrelation correction (Woolrich et al., 2001).

2.3.1. MRI data exclusion

Our *a priori* threshold for participant exclusion based on motion was one EPI voxel (1.2 mm): no participant exceeded this threshold. Two participants were removed as one had an incidental finding on their MRI, and another had excessive susceptibility artefacts in the temporal lobe which impeded hippocampal segmentation. Therefore, data from 23 subjects were included.

2.4. Hippocampal subfield segmentation and co-registration

To contextualise the MVPA searchlight results at the group level, we created probabilistic hippocampal subfield ROIs (CA1, CA2/3, dentate gyrus, and subiculum) in MNI space, based on manual segmentations within our participants. Hippocampal subfields were previously manually segmented (by C.J.H.) on participants' ultra-high-resolution T2*-weighted images using a 7T-specific protocol based on Wisse et al. (2012), and then co-registered to each individual's fMRI space (see Hodgetts et al., 2017 for a detailed description of the manual subfield segmentations). For the current study, these individual-subject ROIs were then co-registered to the 1 mm MNI template (using the same warps created during the FSL FEAT analyses of Hodgetts et al., 2017), binarized and summed to create probabilistic subfield ROIs. To aid interpretation and visualization, the ROIs were thresholded so that each ROI covered voxels included in at least 25% of participants (Fig. 1D; see Syversen et al., 2021 for an example of use of this threshold). The full (unthresholded) probabilistic ROIs are freely available at: <https://osf.io/xc4wa/>.

2.5. The oddity judgement task

Participants completed a simultaneous visual discrimination 'oddity'

judgement task (Buckley et al., 2001; Lee et al., 2005). On a given trial, subjects were presented with three stimuli (top centre; bottom left; bottom right) and asked to choose the odd-one-out as quickly and as accurately as possible. In each trial, the triplets of images were presented on a white background (Fig. 2A). The scene stimuli were greyscale photographs of real outdoor locations (in Cardiff City Centre) and were unfamiliar to participants (see Shine et al., 2015). Two scenes depicted a single location from different viewpoints and one scene depicted a different, but highly similar, location. Face stimuli were greyscale photographs of human faces (half male and half female) and were obtained from the Psychological Image Collection at Stirling (PICS, <http://pics.stir.ac.uk/>). Individual faces were overlaid on a black frame of 170×216 pixels. Two faces were the same individual at different viewpoints, or with a different facial expression, and the third (odd) image was a different face, of the same sex, presented from a different viewpoint. Objects were obtained from the Hemera Photo-Objects 50, 000, Volumes 1–3 (Hemera Technologies, Quebec). The trials included two identical objects presented from different viewpoints, and the third object was different but from the same subordinate-level object category. For the ‘size’ trials, three black squares were shown. Two of which

were identical in size and a third square was either slightly larger or smaller; the difference in length between target and non-targets could vary between 9 and 15 pixels. The position of the squares was jittered so that none of the edges lined up along vertical or horizontal axes. All stimuli were trial-unique (i.e., each stimulus was shown only once). Immediately prior to scanning, participants were shown a practice trial for each category (not repeated during the experiment) and indicated to the experimenter their correct response.

The MRI task was projected onto the screen behind the subject using an Eiki LC-XL100 projector system (Resolution: 1024×768 ; Refresh Rate: 60Hz). The paradigm was coded using Presentation (Neurobehavioural Systems, CA), and button presses were recorded with a right-hand MR compatible button box. Trials were presented for 5500 ms with a jittered inter-trial interval (blank screen) of 500–2500 ms (Fig. 2B). The task was carried out over three fMRI runs (each run was ~7 min in duration). Trials were presented in mini-blocks of three consecutive trials of the same condition (either three scenes, faces, objects, or sizes). The order of which was counterbalanced across participants. 15 trials of each stimulus category were presented, per run, totalling to 45 trials per condition. Each condition had the same number

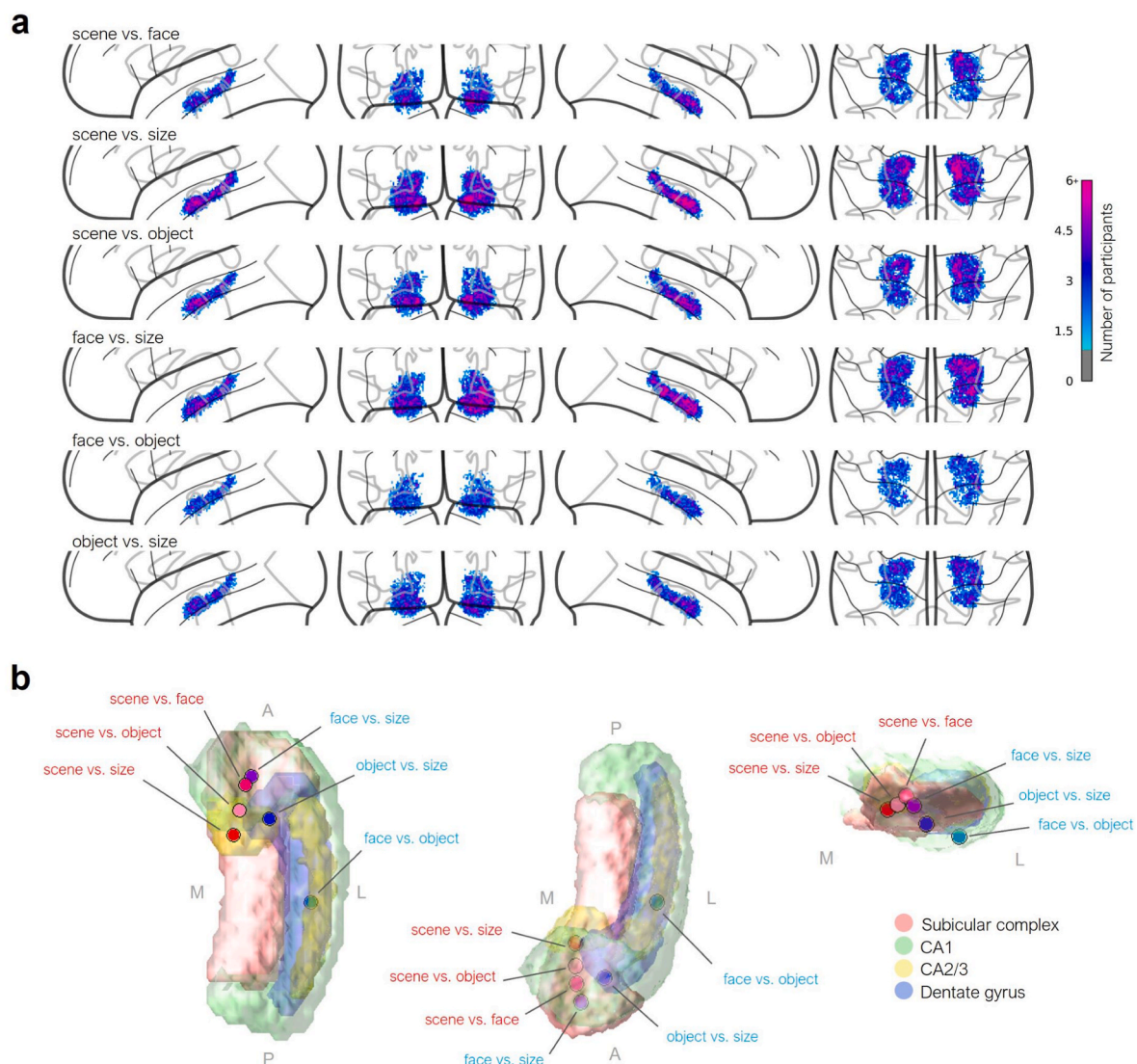


Fig. 2. Overlap statistical hippocampal searchlight maps. A) Overlap maps constructed by summing participant significance maps from individual level Binomial statistical tests against chance level. Maps have a lower threshold of 1, pink colours indicate higher overlap than blue colours. B) Peaks of the overlap maps plotted on three view angles of the right hippocampal subfields (the peaks and ROIs are colored according to the key to the right). Images created using Nilearn (Abraham et al., 2014) for Python and MATLAB.

of targets (odd-one-out) appearing in each triplet position (i.e., top centre; bottom left; bottom right).

2.6. General linear model

Double-gamma hemodynamic response functions were used to model BOLD signals. Each trial, per run, was entered as an explanatory variable (EV) within a General Linear Model (GLM), with event durations of 5.5 s, using FEAT (Woolrich et al., 2001). This resulted in three first-level models (for each of the runs), each containing 60 main EVs (15 per condition: scene, object, face, size) and a motor regressor corresponding to button press onset on each oddity task (duration = 0 s). An additional confound matrix was added to the GLMs to account for volume-wise non-linear motion effects using FSL Motion Outliers. This resulted in 180 t-statistic images for each participant, derived from the parameter estimates. The t-statistic images were then used in subsequent MVPA searchlight analyses.

2.7. Searchlight analysis

Linear support vector machine (SVM) classifier-based searchlight analyses (chosen because of its resilience to overfitting) were conducted using the MVPA-light toolbox (Treder, 2020) in MATLAB R2015A. All trials, correct and incorrect, were used from each category. Neighbours were defined as $3 \times 3 \times 3$ cubes of voxels. Considering the high computational load when searching high resolution data, we used five k-folds (45-9 trials used as the test set), across trials from all runs, with two repeats (a different test set randomly assigned in each repeat, and classifier accuracy values averaged across repeats). Searchlight analyses were carried out in native space and resulting accuracy maps were then warped to the T1-weighted 1 mm MNI standard template (skull stripped) using FNIRT (Jenkinson et al., 2012). We performed two types of searchlight analyses, the first of which was restricted to the hippocampus, and designed to be sensitive to category information, and to allow accommodation of individual differences in intra-hippocampal locations of category selectivity. The second included data from the entire FOV and incorporated traditional conservative group-level statistics, designed to combat the potential bias (predisposition to produce false positives) of the first analysis.

2.7.1. Hippocampal searchlight

First, 2-way classification searchlights of all stimulus category pairs were performed within the hippocampus of each individual subject. As individual variability in the location of category-selective clusters may cause informative clusters to be missed at the group-level (Etzel et al., 2013), we sought to conduct statistical inference at the individual level, prior to creating probabilistic overlap maps on our novel hippocampus MNI template (see Hodgetts et al., 2016; Kaplan and Meyer, 2012; for similar approaches). Statistics were performed at the individual level by carrying out binomial tests on the individual searchlight accuracy result maps (for each 2-way classification) assuming a chance level of 50% (Treder, 2020). Masks for each participant were constructed by selecting voxels where p -values < 0.05 . These individual-level masks were then warped to template space using FNIRT, binarized, and summed across the 23 individuals. Therefore, this resulted in 6 searchlight contrast overlap maps (scene vs. face, scene vs. size, scene vs. object, face vs. size, face vs. object, object vs. size). We also attempted to contrast locations of scene and face information specifically by further combining these overlap maps into scene and face 'hotspot' maps. The scene hotspot map was constructed by summing scene vs. face, scene vs. size, scene vs. object overlap maps, and subtracting face vs. size, face vs. object overlap maps. The face hotspot map was constructed by summing face vs. scene, face vs. size, face vs. object overlap maps and subtracting scene vs. size, scene vs. object overlap maps. Note, the focus on faces in the current investigation (as opposed to faces and objects) is based on previous evidence showing that medial temporal lobe lesions have stronger

influence on face compared to object discrimination performance using similar oddity tasks (Behrmann et al., 2016; Chang et al., 2023).

2.7.2. Whole FOV searchlight

We also sought to examine scene and face selectivity within our extended FOV (e.g., in posteromedial and parahippocampal cortex) using traditional conservative group-level statistics. To do this we applied permutation tests to group-level searchlight data. First, native space accuracy maps of each category pair (2-way classification) were warped to standard MNI space and combined into a 4D file. Second, in order to run a one-sample t -test against zero (as required by FSL Randomise), 0.5 was subtracted from each map so that voxels at chance level were zero (see Ashby and Zeithamova, 2022; Koski et al., 2017 who also adopted a similar technique; simultaneously, a threshold of -0.2 was applied to that regions that were zero outside of the FOV remained zero, rather than becoming -0.5 , and -0.2 was arbitrarily chosen after inspections of image histograms showed a curtailing of voxels with accuracy levels below this value). One-sample permutation tests were carried out using FSL's 'Randomise' (Winkler et al., 2014). We used threshold-free cluster enhancement (TFCE) (Smith and Nichols, 2009) and 5000 permutations, and, as the FOV differed slightly between individuals, a mask that included voxels where all participants had values (Fig. 1C). The resulting t -maps were masked using the corrected p -values maps and thresholded at $p = 0.0083$ (0.05/6 tests). To isolate scene and face information specifically, conjunction analyses were then performed, by combining significance masks from the searchlight contrasts (binary masks multiplied together). Specifically, a scene-selective map was the product of all the significance masks from searchlight contrasts that included scene trials (scene vs. face $\times$ scene vs. size $\times$ scene vs. object) with the product of all the significance masks from the non-scene searchlights (face vs. size $\times$ face vs. object $\times$ object vs. size) removed (and *vice versa* for face selective regions).

2.7.3. Data sharing and open practices

Anonymized output data and the code used within this project are freely available at <https://osf.io/4vgk9/>, so that the figures and values reported in this manuscript are reproducible. However, ethical restrictions, relating to General Data Protection Regulation, do not allow for the public archiving of the raw study data. Access to pseudo-anonymized data could be granted after signing and approval of data-transfer agreements. For this, readers should contact Dr Carl Hodgetts (carl.hodgetts@rhul.ac.uk).

3. Results

3.1. Oddity judgement task behavioural results

Participants achieved high and matched decision accuracies across image categories. The mean proportion correct scores (and SDs) of the scene, face, object, and size conditions were: 0.76 (14), 0.76 (14), 0.77 (16), 0.76 (15), respectively. Detailed analyses on behavioural performance are available in our previous publication (Hodgetts et al., 2017).

3.2. Searchlight MVPA results

3.2.1. Hippocampal searchlight classification

First, we examined the spatial distribution and inter-individual variability of scene and non-scene-selective information within our hippocampal ROI. As noted in the Methods, we conducted a two-category SVM searchlight within each individual's hippocampal ROI for each of the discrimination pairs (scene vs. face, etc), then co-registered the resulting (binarized) maps of significant classification voxels ($> 50\%$) to the standard template before summing the maps across subjects to create overlap maps. The mean number of voxels (and SDs) reaching significance in the binomial tests for each hippocampal searchlight contrast were: scene vs. face: 870 (511); scene vs. size: 1288

(889); scene vs. object: 1047 (402); face vs. size: 1236 (670); face vs. object: 642 (267); object vs. size: 891 (494).

Fig. 2A shows the overlap for each of the SVM hippocampal searchlights (greater values equals larger overlap between subjects). As can be seen, category selectivity appeared to cover most of the hippocampi ROIs, but the highest overlap between participants was generally seen anteriorly (except for the face vs. object overlap map), and in the right hippocampus. When examining the location and distribution of these overlap peaks (voxel locations where most participants had significant voxels) with respect to our probabilistic subfield ROIs (Fig. 2B), we found that the scene-including classification overlap maps had the highest overlaps medially, while the highest overlaps of the face/object-

including classification overlap maps were more lateral. The scene vs. size overlap peak was the most medial, located in the anteromedial subicular complex, close to the superior subicular complex/CA1/CA2/3 ROI borders (maximum overlap between individuals = 10; $x = 18$, $y = -17$, $z = -18$, 43% subicular complex, 20% CA1, 20% CA2/3, 3% DG). The scene vs. object overlap peak was located at the superior subicular complex/CA1 border (7; $x = 19$, $y = -14$, $z = -20$; 52% subicular complex, 36% CA1, 4% CA2/3, 2% DG). The scene vs. face classification overlap map had the highest overlap in anterior CA1 (8; $x = 20$, $y = -11$, $z = -21$; 67% CA1, 34% subicular complex, <1% DG).

With regards to our face/object classification overlap maps, we found that the face vs. object classification overlap peak was the most

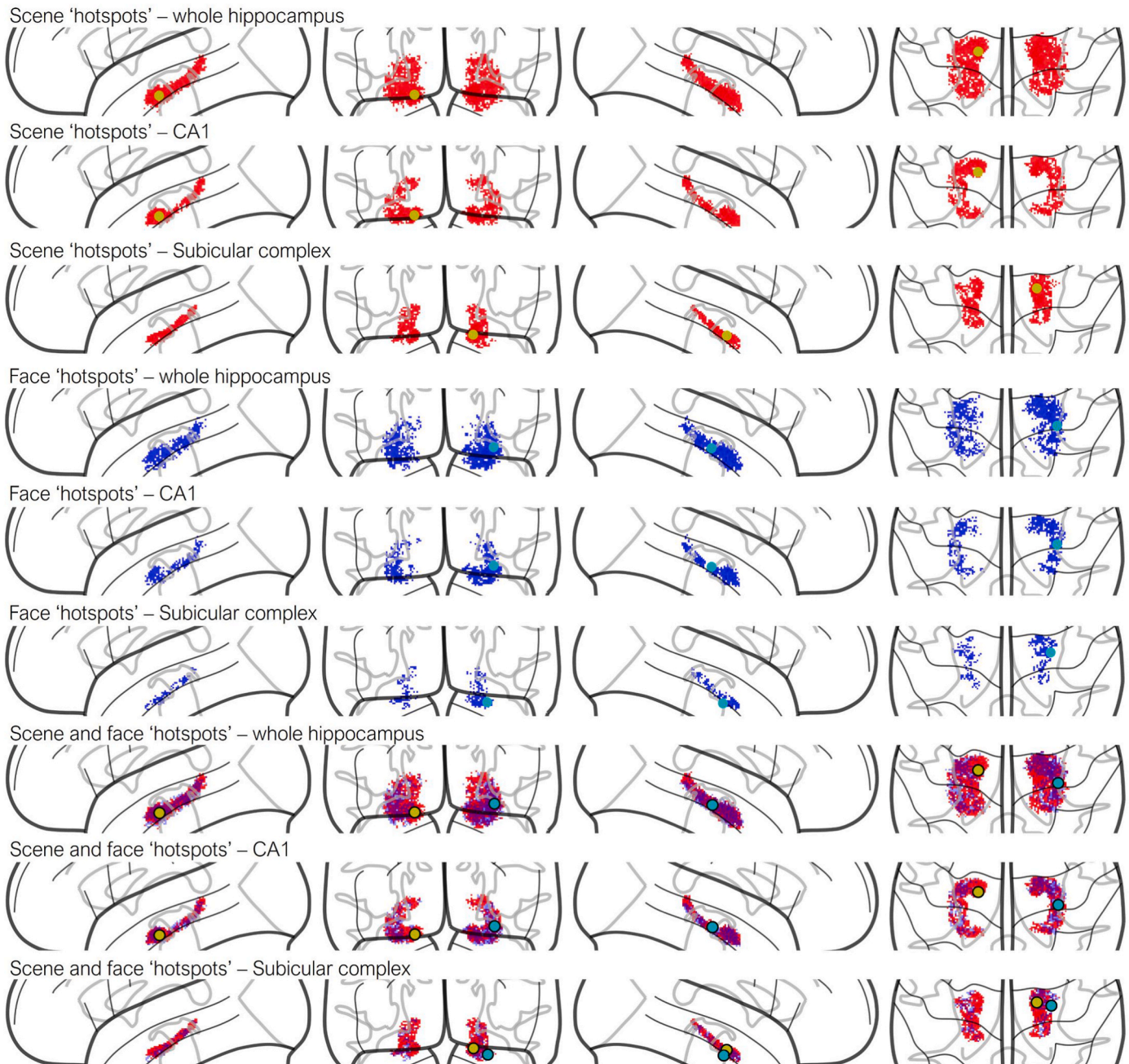


Fig. 3. Overlap statistical hippocampal scene and face 'hotspot' searchlight maps. 'Hotspot' maps constructed by summing together searchlight results from comparisons including scenes, with those results from comparisons including faces subtracted (reds) and constructed by summing together searchlight results from comparisons including faces, with those results from comparisons including scenes subtracted (blues). Overlaid dots represent the locations of maximum overlap for the scene 'hotspot' map (yellow dot) and the face 'hotspot' map (light blue dot). Note that an arbitrary minimum threshold of 3 was applied to increase clarity in the image.

lateral and posterior, located in CA1 (6; $x = 31$, $y = -25$, $z = -15$; 86% CA1, 4% DG). The object vs. size classification overlap map had the highest overlap in the subiculum, but this was more lateral and inferior than the scene classification overlap map peaks, and situated close to the subiculum's lateral border with CA1 (7; $x = 24$, $y = -15$, $z = -23$; 66% subicular complex, 9% CA1, 3% DG). The face vs. size classification overlap map had the highest overlap in the anterior subiculum, close to its superior border with CA1 (8; $x = 21$, $y = -10$, $z = -24$; 63% subicular complex, 28% CA1).

With the aim of comparing where selective scene and face information is most likely to be located, further 'hotspot' overlap maps (Fig. 3) were constructed by 1) summing together hippocampal searchlight overlap maps from comparisons including scenes, and subtracting the sum of overlap maps from comparisons including faces ('scene hotspot'), and 2) summing together hippocampal searchlight results from comparisons including faces, subtracting the sum of overlap maps from comparisons including scenes ('face hotspot'; see Methods). Again, a medial-lateral gradient was apparent, with the scene hotspot map showing higher values medially, and the face hotspot map showing higher values laterally. The maximum value for the scene hotspot maps was in left medial inferior CA1 and superior subiculum boarder ($x = -17$, $y = -15$, $z = -19$; 45% CA1, 39% subicular complex). The maximum value for the face hotspot maps was located in right lateral hippocampus, in the inferior border of the DG with CA1 ($x = 32$, $y = -23$, $z = -14$; 70% DG, 29% CA1; Fig. 3). These peak locations remained when the hotspot maps were restricted to the CA1 ROI (Euclidean distance between scene and face hotspot maps maximum values = 50 mm).

This medial-lateral pattern was still apparent when the hotspot maps were restricted to the subicular complex ROI. Maximum values for the scene and face hotspot maps were located in right anterior medial ($x = 19.0$, $y = -14.0$, $z = -20.0$; 52% subiculum, 36% CA1, 4% CA2/3, 2%

DG) and lateral areas, respectively ($x = 28.0$, $y = -16.0$, $z = -24.0$; 22% subiculum, 21% CA1; Euclidean distance between subiculum scene and face hotspot maps maximum values = 10 mm).

3.2.2. Whole FOV searchlight classification

To examine the whole fMRI FOV, we adopted a more conservative, group statistical approach (group-level nonparametric permutation inference). Specifically, we initially conducted SVM searchlights across each individual's FOV for each discrimination of interest. Accuracy maps were then co-registered to the standard MNI template before Randomise tests (cluster permutation tests against chance level) were run. Each 1 vs. 1 searchlight classification (scene vs. face, scene vs. object, scene vs. size, face vs. size, face vs. object, object vs. size) resulted in significant clusters.

To isolate regions selective to scenes and not faces within the 'scene vs. face' significance mask, significant voxels from 'face vs. size' were removed (Fig. 4A). Then, to isolate regions selective to faces and not scenes within the 'scene vs. face' significance mask, significant voxels from scene vs size' were removed (Fig. 4B). Scene regions were found in right anteromedial subicular complex (maximum ROI probability: 64%, $x = 19$ $y = -18$ $z = -20$; 25 voxels above 25%) and bilateral lateral CA1 (maximum ROI probability: 67%, $x = 31$ $y = -31$ $z = -12$; 229 voxels above 25%). Face regions were also apparent in left anteromedial subicular complex (maximum ROI probability: 29%, $x = -17$ $y = -30$ $z = -13$; 2 voxels above 25%) and lateral CA1 (maximum ROI probability: 68%, $x = 30$ $y = -32$ $z = -11$; 78 voxels above 25%), albeit to a lesser extent.

The same was carried out for 'scene vs. object' (Fig. 4C–D). Again, a scene region was found in right anteromedial subiculum (maximum ROI probability: 75%, $x = 20$ $y = -18$ $z = -19$; 156 voxels above 25%), but CA1 regions were more medial than those in the scene vs. face mask (maximum ROI probability: 78%, $x = 30$ $y = -31$ $z = -11$; 117 voxels

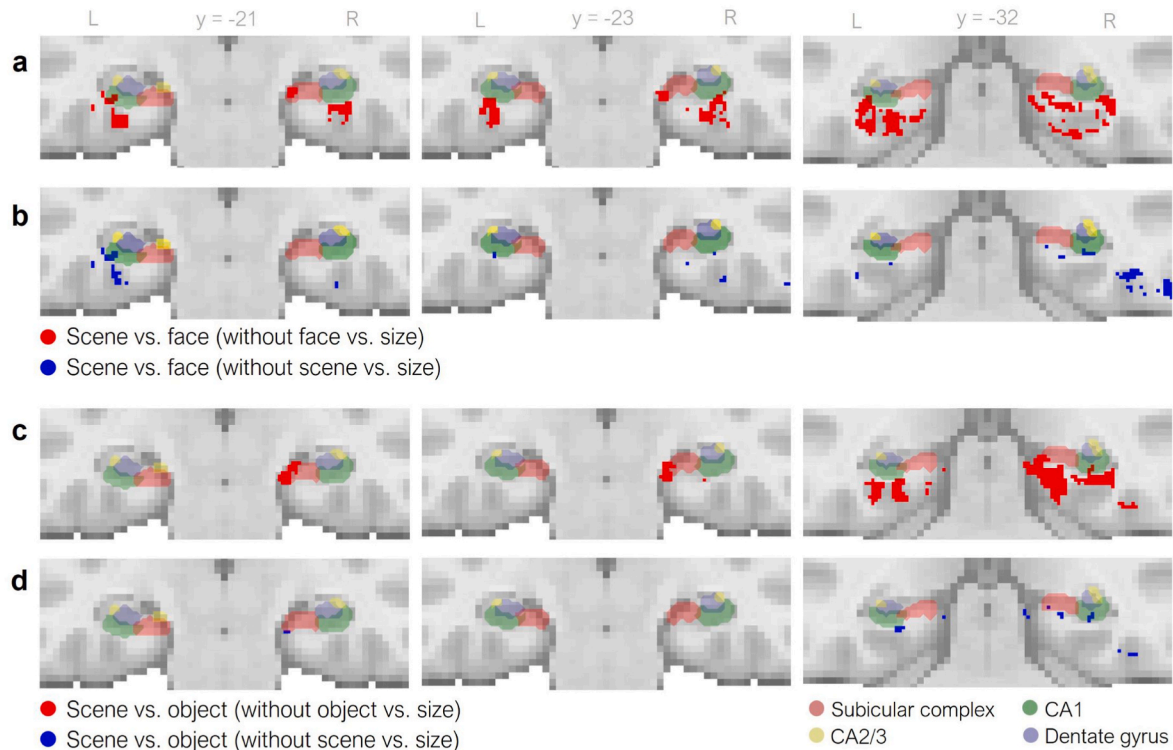


Fig. 4. Scene vs. Face and Scene vs. Object Conjunction maps. A) Scene regions from the scene vs. face searchlight map ('face vs. size' removed). B) Face regions from the scene vs. face searchlight map ('scene vs. size' removed). C) Scene regions from the scene vs. object searchlight map ('object vs. size' removed). D) Object regions from the scene vs. object searchlight map ('scene vs. size' removed). Significant classification regions also extended outside of the hippocampus, into the parahippocampal cortex. The resulting maps are overlaid onto our group subfield ROIs. DG: Dentate Gyrus, L: Left, R: Right. Image created using Nilearn (Abraham et al., 2014) for Python.

above 25%). In addition, the scene region overlapped with medial CA2/3 (maximum ROI probability: 59%, $x = 20$ $y = -18$ $z = -16$; 70 voxels above 25%). Object regions were also apparent in anteromedial subicular complex (maximum ROI probability: 75%, $x = 20$ $y = -18$ $z = -16$; 33 voxels above 25%), CA1 (maximum ROI probability: 78%, $x = 30$ $y = -31$ $z = -11$; 29 voxels above 25%) and CA2/3 (maximum ROI probability: 52%, $x = 19$ $y = -17$ $z = -13$; 14 voxels above 25%), to lesser extents.

Lastly a purely scene-selective map (Figs. 5–6) was defined as the conjunction of significant voxels from all scene-discriminating searchlights ('scene vs. face', 'scene vs. size' and 'scene vs. object'), with the conjunction of significant voxels from all non-scene-discriminating searchlights ('face vs. size', 'face vs. object' and 'object vs. size') removed. Scene-selective regions were found to overlap with the anteromedial portion of the right subicular complex ROI (maximum ROI probability: 55%, $x = 19$ $y = -21$ $z = -18$; 29 voxels above 25%), a region we interpret as corresponding to distal subiculum/presubiculum/parasubiculum (Fig. 5A), and left posterior lateral CA1 (maximum ROI probability: 50%, $x = -33$ $y = -34$ $z = -10$; 31 voxels above 25%), a region we interpret to correspond to posterior proximal CA1 (Fig. 5A). Supporting our interpretations, Fig. 5B shows the locations of these scene-selective map ROI maximum probabilities in relation 1) to the anterior-posterior hippocampus border (defined as the uncus apex at $y = -22$; as defined by Poppenk et al., 2013; Zeidman and Maguire, 2016), 2) a previously defined anteromedial hippocampus location (used by Zeidman et al., 2015b; Zeidman and Maguire, 2016 as a seed region to explore anteromedial hippocampus connectivity, defined through the conjunction of activation during recalling and imagining scenes), and 3) a previously observed peak BOLD location within the anteromedial hippocampus during a scene construction task, interpreted as residing in pre/parasubiculum (Dalton et al., 2018). In addition, the scene-selective map overlapped with known scene selective areas in the posterior PHC

and PMC (note that this PMC area included retrosplenial cortex as defined by Brodman areas 29 and 30; Fig. 6A). The three most overlapping regions within FSL's Harvard-Oxford cortical structural atlas were: precuneus cortex (16% overlap), superior lateral occipital cortex (11% overlap); and lingual gyrus (10%).

In contrast, a face-selective map (Fig. 6B) was defined as the conjunction of significant voxels from all face-discriminating searchlights ('scene vs. face', 'face vs. size' and 'face vs. object'), with the conjunction of significant voxels from all non-face-discriminating searchlights ('scene vs. size', 'scene vs. object' and 'object vs. size') removed. This face-selective map was right lateralised and did not overlap with our hippocampal ROIs. The three most overlapping regions within FSL's Harvard-Oxford cortical structural atlas were: angular gyrus (18% overlap); posterior supramarginal gyrus (12% overlap); and temporooccipital gyrus (13% overlap), and there was also overlap with the temporal fusiform cortex (4% overlap).

4. Discussion

In the current study, we advanced our knowledge of hippocampal contributions to high-level scene perceptual discrimination by: 1) Using MVPA to improve sensitivity to fine-grained category-selectivity patterns of hippocampal BOLD activity, thereby revealing decodable scene information outside of the subiculum (previously identified as a hippocampal 'hub' of scene information based on magnitude of univariate BOLD activity); 2) using a subfield agnostic searchlight approach, thereby revealing precise locations of category selectivity that do not necessarily conform to traditional anatomical subfield boundaries and; 3) using both subject-level and group-based analysis in order to account for the different biases present in each (see Etzel et al., 2013), which helped to reveal a transverse and longitudinal gradient of scene category selectivity in the subiculum and CA1. In sum, this approach allowed us

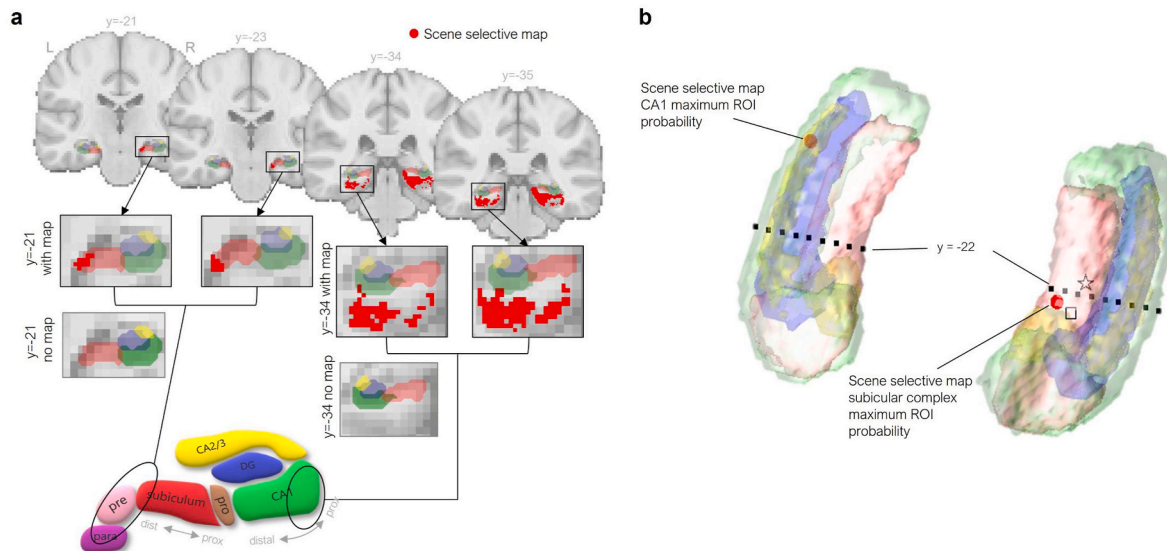


Fig. 5. The scene-selective and face-selective conjunctive maps within the hippocampus. A) The 'scene-selective' map plotted onto multiple coronal template slices. The four coronal 'zoomed-in' images, show inclusion of the right anteromedial subicular complex and inclusion of lateral CA1 regions. MNI slice coordinates are stated by each image. Our subfield ROIs were overlaid onto this map, and their colours correspond to the drawn subfield diagram. For slices $y = -21$ and $y = -34$, additional 'zoomed in' images show the subfields without the scene-selective map to aid visualization of the overlaps of the scene selective map on the subfields. The subfield diagram illustrates our interpretation of the results and includes subfield delineations not possible with our current methods. The circles indicate that we consider the portion of the scene-selective map that overlaps with the medial subicular complex to incorporate distal subiculum/presubiculum/parasubiculum areas. We also consider the portion of the scene-selective map that overlaps with lateral CA1 to incorporate part of the proximal aspect of CA1. B) The locations of these scene-selective map ROI maximum probabilities in relation to the anterior-posterior hippocampus border ($y = -22$; Zeidman and Maguire (2016)). To contextualise these findings, a previously defined anteromedial hippocampus location (Zeidman et al., 2015b; Zeidman and Maguire, 2016) is shown with a star symbol, and a previously found anteromedial hippocampus scene construction peak within the pre/parasubiculum (Dalton et al., 2018) is shown with a star symbol. DG: Dentate Gyrus, L: Left, para: parasubiculum, pre: presubiculum, pro: prosubiculum, R: Right. Images created using Nilearn (Abraham et al., 2014) for Python, MATLAB, and Microsoft products.

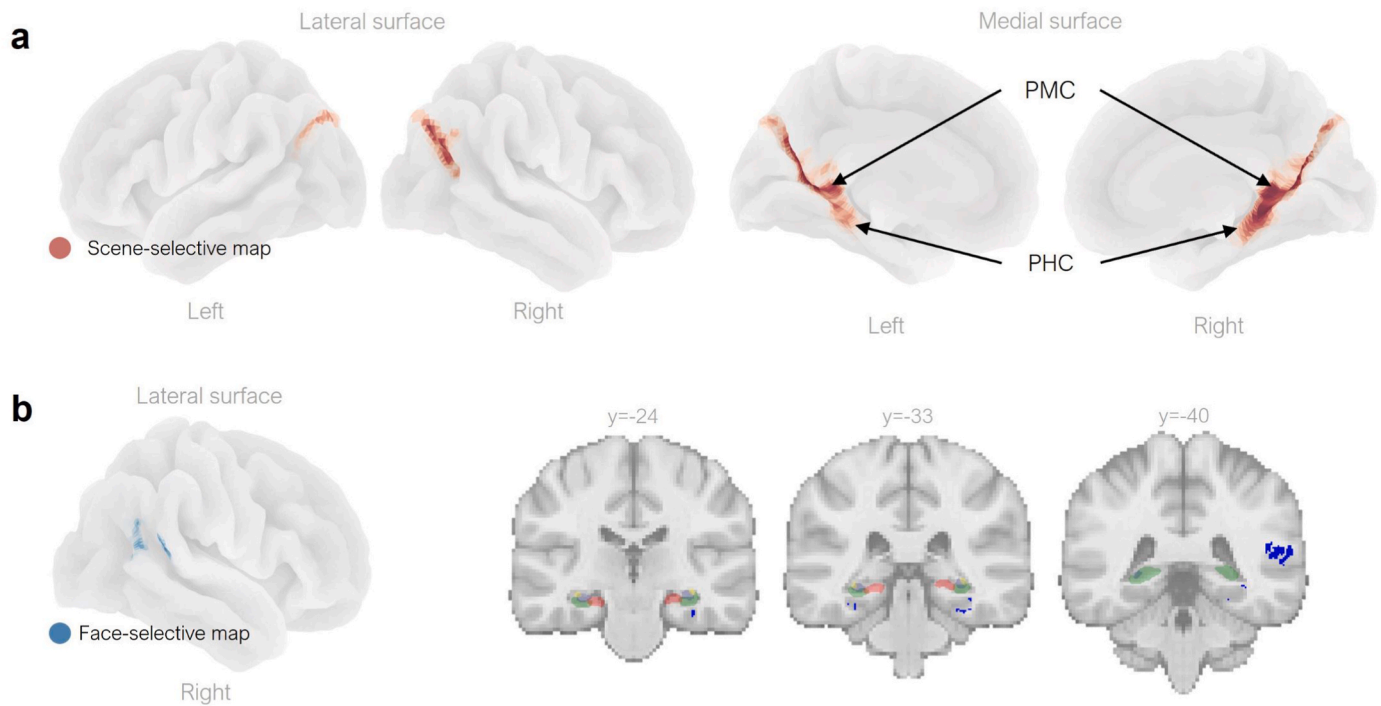


Fig. 6. The scene-selective and face selective conjunctive maps within the whole fMRI FOV. A) The ‘scene-selective’ map plotted onto template brain surfaces, which shows the scene-selective map to overlap with areas of the superior occipital, PHC and PMC regions. D) The ‘face-selective’ map plotted onto a template brain surface (left) and onto multiple coronal slices (right). PHC: Parahippocampal Cortex, PMC: Posteromedial Cortex. Images created using Nilearn (Abraham et al., 2014) for Python.

to map a scene representation pathway in humans, which included regions within the anteromedial (distal) subicular complex and lateral (proximal) CA1, alongside cortical regions that form part of a parieto-medial temporal pathway (Kravitz et al., 2011) implicated elsewhere in spatial navigation and scene-based cognition, namely the posterior PHC and PMC (Boccia et al., 2017; Soto et al., 2012; Margulies et al., 2009; Goodale and Milner, 1992; Epstein and Baker, 2019). In addition, we provide preliminary evidence suggesting that areas within the lateral hippocampus carry information relevant to discriminating faces. We discuss the implications of these findings for accounts of hippocampal contributions to higher-level perception in the following sections.

4.1. Subject-level analysis indicates gradients of hippocampal category sensitivity

Both the ‘overlap’ and ‘hotspot’ maps (see Figs. 2 and 3) demonstrated a gradient-like pattern of scene selectivity, with scene information being preferentially processed anteriorly on the longitudinal (i.e., anterior - posterior) axis and medially along the transverse (i.e., distal - proximal/medial - lateral²) axis. In contrast, face and object information showed a somewhat more diffuse pattern, although there was evidence for increasing face selectivity laterally and anteriorly (see further discussion below). Mapping the data back to individual subfields, we additionally show that the same longitudinal and transverse gradients were present in both the subiculum complex and, notably, CA1. In addition, we report that the subiculum complex carried less face-sensitive information, with most of the face-selective voxels being present along the lateral subiculum border, whereas CA1 contained voxels relevant to the processing of both scenes and faces, with the aforementioned transverse gradient and a less pronounced longitudinal one.

Previous research has demonstrated that both functional and structural connectivity gradients in the hippocampus do not respect classical subfield boundary definitions (Aggleton and Christiansen, 2015; Dalton et al., 2019, 2022). For scene selectivity, this aligns with current

understanding of anterior-posterior hippocampal roles. The scene oddity task required participants to process whole scenes from multiple view-points, and the anterior hippocampus, in general, is proposed to bring together distributed visuospatial, object, and semantic information to construct spatially coherent internal representations of scenes (Sekeres et al., 2018; Zeidman and Maguire, 2016), and to process more global representations of space, relative to the posterior hippocampus (Evansmoen et al., 2013; McCormick et al., 2021; Nadel et al., 2013; Zeidman et al., 2015b; Leferink et al., 2023) potentially as part of a wider role in ‘relational’ (Mayes and Roberts, 2001; Mayes et al., 2007; Olsen et al., 2012; Aly et al., 2013; Turk-Browne, 2019) or ‘compositional’ (Bakermans et al., 2023) processing. For face selectivity, our results demonstrating more lateral processing are in line with recent hippocampal structural connectivity research, which showed that cortical areas important for face processing (such as the temporal pole) preferentially terminate along the lateral borders (Dalton et al., 2022; Barton, 2022).

These results show the importance of studying longitudinal and transverse gradients across the hippocampus and within individual subfields. Further analysis of individual differences in these gradients may be useful in uncovering links to behaviours and pathology. Research on hippocampal connectivity has shown that individual differences in connectivity gradients are associated with phenotypes including mild cognitive impairment, Alzheimer’s disease, and more generalised differences in recollection memory performance (Borne et al., 2023; Przeździk et al., 2019). Larger scale investigations crossed with confirmatory gradient statistics (e.g., Przeździk et al., 2019) will be necessary to confirm the current findings in relation to scene and face processing.

4.2. Group-level analysis demonstrates a scene-processing pathway involving the distal subiculum and proximal CA1

At the group level, we combined significance masks across 2-category classification searchlights to identify scene, face, and object

selective regions within ‘scene vs. face’ and ‘scene vs. object’ searchlight result maps and to create ‘scene-selective’ and ‘face-selective’ maps. Across all scene-discriminating searchlight combinations, regions within the anteromedial subicular complex and some inferior-distal voxels within CA1 were identified as scene-selective. The scene-selective map included the caudal inferior parietal cortex, bilateral anterior PHC, bilateral PMC, and extended into the medial temporal lobes, aligning with well-established scene networks (Baldassano et al., 2016; Epstein, 2008; Epstein and Baker, 2019; Hodgetts et al., 2016; Zeidman et al., 2015b). In the hippocampus, this specifically included inferolateral edges of bilateral CA1, and an anteromedial portion of the right subicular complex (distal subicular complex). The part of the scene-selective map that invaded the subicular complex ROI also extended more medially, providing support for the suggestion, based on univariate fMRI analysis, that the pre/para-subiculum carries information important for scene processing including perceptual discrimination (Dalton and Maguire, 2017). Again, these results mirror recent evidence from structural connectivity analysis, which identified a similar anteromedial hub of hippocampal connectivity to cortical regions associated with scene processing (e.g. medial parietal cortex) (Dalton et al., 2022).

Even with the use of high field (7T) fMRI, it is unclear whether our results show scene selectivity in the distal subiculum proper or pre/para-subiculum, or both (Dalton and Maguire, 2017; Ding, 2013). In terms of both its anatomical connectivity and cellular properties, the subicular complex appears perfectly situated to support scene processing as it is the primary source of hippocampal efferents (Aggleton, 2012; Bubb et al., 2017; Insausti and Muñoz, 2001; Gigg, 2006) and, while many subiculum proper targets overlap with those of neighbouring CA1 (Matsumoto et al., 2019), almost all hippocampal projections to the anterior thalamic nuclei, mamillary bodies, and retrosplenial cortex (RSC) originate in the subiculum proper (Aggleton and Christiansen, 2015; Frost et al., 2021; Gaffan, 1992; O’Mara and Aggleton, 2019). Structural connectivity between the subiculum and CA1 with the entorhinal cortex, along with converging connections to the RSC and PHC, have also been demonstrated (Kloosterman et al., 2003; Witter, 2006; Simonsen et al., 2022). Notably, recent functional connectivity analysis found that medial subiculum – entorhinal connectivity overlapped with entorhinal – PHC connectivity hotspots when participants were engaged in scene processing (Grande et al., 2022). These interconnected structures are key components of an extended hippocampal system thought to support spatial attributes of memory (Aggleton, 2012; see also Murray et al., 2017) as well as contributing to complex scene discrimination (Postans et al., 2014; Hodgetts et al., 2015). The subiculum proper, at least in rodents, also appears to contain several types of spatially-modulated neurons, including cells attuned to spatial position (Sharp, 2006), boundaries (Lever et al., 2009), corners (Sun et al., 2023) and direction of travel (Olson et al., 2017), as well as cells that seemingly fire conjunctively for several such properties needed to re-construct representations of spatial environments (Ledgergerber et al., 2021; Sharma et al., 2022). Additionally, aligning with our results and interpretations, in the rodent there is evidence of a distal-proximal gradient in the amount of environment-specific information conveyed in the subiculum, with neurons in the distal subiculum conveying more information (denser firing rate), and evidence that subiculum neuron firing patterns code more information over a greater proportion of the environment than do CA1 neurons (Kim et al., 2012).

Conversely, while the subiculum proper is the main output region of the hippocampus, the pre/para-subiculum is primarily an input region, receiving input from areas including the subiculum proper, CA1, RSC and inferior parietal cortex (Honda and Shibata, 2017; Honda et al., 2022; Huang et al., 2021; Insausti and Muñoz, 2001; Kravitz et al., 2011). A parieto-medial pathway originates in visuospatial regions of the parietal cortex, and connects to the parahippocampal cortex and RSC (Kravitz et al., 2011), two areas known to contribute to spatial processing (Cukur et al., 2016; Epstein et al., 2017; Hodgetts et al., 2016; Nasr et al., 2013), which in turn connect directly with the

pre/para-subiculum (Kravitz et al., 2011; see also Dalton and Maguire, 2017). Likely related to these inputs, the pre/para-subiculum is known to contain grid, border and head-direction cells (Boccaro et al., 2010; Robertson et al., 1999; Rolls, 2023), and indeed is considered a major component of, and perhaps a site of visual information integration into, the head-direction system (Preston-Ferrer et al., 2016). Future fMRI studies at even higher spatial resolution, combined with directed functional connectivity measures (Reid et al., 2019), will be important in addressing how this network interacts during the processing of scenes (for a proposed account see Bicanski and Burgess, 2018).

The CA1 results are less easy to interpret, only existing at the edges of the ventral distal boundaries. This may simply reflect signal bleed from the PHC which borders CA1, potentially caused by the spatial variability introduced either by the searchlights or from the registrations to standard space (Etzel et al., 2013). Alternatively, given previous literature indicating a role for CA1 in spatial processing, including place cells in rodents (e.g., see Brun et al., 2002; O’Keefe et al., 1998; Park et al., 2011) and humans (Suthana et al., 2009; Ekstrom et al., 2003), as well as allocentric view cells in primates (Rolls, 1999, 2023), it would be perhaps be more surprising if we did not see voxels within CA1 that contained scene-sensitive information. Rodent work has indicated that the CA1 passes spatial information to the subiculum, which carries more spatial location and context information than the sparser CA1 representations, suggesting that the subiculum prepares sparse CA1 spatial codes by condensing them into more dense information, to be communicated to brain regions outside of the hippocampus (Kim et al., 2012).

The results from our two analyses are not necessarily at odds, with the subject-level maps demonstrating that scene information is broadly distributed across CA1, which may be a reason for the lack of ‘scene specific’ significant voxels in the group-level analysis. Indeed, meaningful differences between subject-level and group-level analyses are well documented (Jolly and Chang, 2021; Fedorenko, 2021) and can arise due to individual variation in the spatial distribution of category relevant voxels (Etzel et al., 2013). If, as is suggested by our subject-level maps, category information is not represented in as consistent a topological manner in CA1 compared with the subiculum, then this would result in less power to detect an effect at the group level. This individual variation has also been reported in hippocampal MRI connectivity analyses (Borne et al., 2023; Dalton et al., 2022; Przeźdźik et al., 2019).

Outside of the hippocampus, the face-selective map matched known mapping of face sensitive regions. This was right lateralised (Haxby et al., 2000; Rapcsak, 2019; Rossion, 2014) and also included regions well known to support face processing, such as a portion of the inferior temporal cortex, likely reflecting the fusiform face area, and a portion of the posterior-superior temporal sulcus (Barton, 2022; Kanwisher and Yovel, 2006; Rapcsak, 2019). As mentioned, structural connectivity analysis has shown that these regions connect to the lateral hippocampus (Dalton et al., 2022) (also reported in our individual level analysis). Whilst our group-level results did not reveal face-selective regions within the hippocampus, this may have been due to high individual variation in the location of face-selective voxels (Gao et al., 2022), which we demonstrated were more spatially distributed compared to scene-selective voxels in our subject overlap maps (Figs. 2–3). Such variability might in part account for only very marginal face-oddity impairments (seen in reaction times, rather than accuracy) following hippocampal lesions (Behrmann et al., 2016). In addition, the nature of our analysis was rather conservative as the construction of the face-selective maps necessarily excluded voxels that were sensitive to more than one category. For example, hippocampal regions that may have been sensitive to both faces and objects would not have been included.

4.3. The results align with ideas of parallel processing of category information in networks within, and connected to, the hippocampus

Together considering the differing gradient patterns for scenes and

faces from our probabilistic maps (aligning with Dalton et al., 2022 mapping), as well as the scene selectivity patterns in the distal subicular complex and inferior CA1 (aligning with Aggleton, 2012), our interpretations are encompassed by, and build upon, the PMAT (Posterior Medial and an Anterior-Temporal) framework (Ritchey et al., 2015; Inhoff and Ranganath, 2017). In this framework, two anatomically distinct parallel processing streams (though see Connor and Knierim, 2017; Kinnavane et al., 2017), PM and AT reside in both distributed cortex regions as well as medial temporal lobe regions, the PHC and PrC, respectively, and continue to run in parallel within the distal subicular complex/proximal CA1 and proximal subicular complex/distal CA1, respectively (Brandon et al., 2014; Kinnavane et al., 2017; Grande et al., 2023), and they support behaviours across a range of cognitive domains (e.g., both mnemonic and perceptual) for different modalities (Ritchey et al., 2015). The PM network is proposed to support context representation, encompassing spatial processing required for our scene oddity task, and the AT network supports representation of item information, encompassing face and object processing required for our face and object oddity tasks. Our results add to this framework by providing insight into the topography of the PM within the hippocampus. We also map aspects of the AT within the cortex, however, as mentioned, our methods to reveal face only information excluded regions which process both face and object information, limiting the ability to accurately map AT within the hippocampus.

In addition, our results complement recent findings of two parallel distributed networks within the default mode network, one thought to have spatial functions, which includes a more anterior region of the subiculum, and one thought to have social functions, which includes a more posterior region of the subiculum (Edmonds et al., 2023).

4.4. Limitations

There are some limitations of our study, and our results should be interpreted with these in mind. Our small sample size may mean that comparing classifier performance to theoretical chance level (e.g., 50% for 2 class classification) may have led to inflated results, as actual chance level in small sample sizes can exceed theoretical chance level (Combrisson and Jerbi, 2015). Although our conservative group statistics approach reduced this risk, replication of our results with larger sample sizes would be beneficial. Relatedly, within the k-folds, trials for training and testing were not strictly within different fMRI runs and this lack of independence can produce inflated decoding accuracies. However, our main interpretations are from contrasted SVM results (i.e., results from one SVM pair significance mask are subtracted from another). This means that it is unlikely that the results can be explained by biased classification because systematic false positives would be present in every SVM classification pair.

We opted to use a high-resolution sequence at 7T to optimize SNR in hippocampal regions, but the FOV only had partial brain coverage. Therefore, we have not mapped all the scene- or face-selective regions that could work in conjunction with the subicular complex and CA1, including regions of entorhinal and perirhinal cortex (Grande et al., 2022; Maass et al., 2015). To map all cortical regions processing category information in conjunction with the subicular complex and CA1, future work with whole-brain high-resolution imaging would be ideal but is challenging with current scanning limitations.

It should be noted that this experiment was originally designed to examine univariate activation magnitude differences between categories, and therefore adopted a mini-block design (trials of same category types were shown sequentially in threes) to increase the average BOLD response. This may not be the best approach for MVPA analyses, and future studies may benefit from adopting experimental designs that can improve the power and reliability of MVPA (e.g., see Abdulrahman and Henson, 2016; Aguirre et al., 2011; Weaverdyck et al., 2020; Zeithamova et al., 2017).

Searchlight methods also have limitations in that they are inherently

spatially blurred because the decoding accuracy in each voxel is determined by the information in the surrounding voxels, that is, the 'neighbourhood' (Etzel et al., 2013). This means that a significant finding in one voxel and not its neighbour, does not necessarily translate to category information existing in the first voxel but not the second. We therefore do not interpret the results at this voxel-by-voxel level.

Relatedly, optimal neighbourhood size is unclear (Etzel et al., 2013). Typically, MVPA searchlights use spheres of around 30 voxels (Kriegeskorte et al., 2006; Sunday et al., 2018). In contrast, we chose a $3 \times 3 \times 3$ cube, as it has been used previously with this toolbox (Treder, 2020) and contains only 9 voxels, with the aim of providing enough information to the SVM (to limit false negative results) while maintaining spatial accuracy. It should be noted that, at the level of the maximum subicular complex ROI probability overlapping with the scene-selective conjunction mask ($x = 19, y = -21, z = -18$) no MNI thresholded probabilistic subfield ROI had a height or width of less than 3.6 mm (1.2×3). However, it may be that interpretations of exact hippocampal regions displaying scene information change with variations in voxel and neighbourhood sizes. Moreover, there is unlikely to be optimum voxel/neighbourhood sizes for all types of scene processing information in all brain regions, as the information may be contained at differing spatial frequencies (Coutanche, 2013).

Lastly, it is important to note that although we interpret increased scene classification performance in a region as an indication that this region contributes to scene representations, this does not mean that those representations were available to the SVM. For example, it is very unlikely that the scale of scene representations on the neural level matched our sampling of fMRI information. We can conclude that there is decodable information, but we cannot conclude that the patterns providing the decodable information are category representations (Ritchie et al., 2019; Peelen and Downing, 2023). A future step could be to relate MVPA results of a region (e.g., classification performance, distance from classifier decision boundary or pattern similarity) with a behaviour (such as task performance), as such a relationship would provide stronger evidence that the information available to the decoder was a category-specific neural pattern (Ritchie et al., 2019). Moreover, our analyses included all trials, correct and incorrect, but additionally applying a SVM to classify between correct and incorrect trials, within categories, may also have provided insight into the information available to the decoder (e.g., see Lee et al., 2013). However, it was not feasible in this case to relate the searchlight results to behaviour through, for example, correlating with oddity performance or classifying between correct and incorrect trials, because of the low sample size, high levels of task performance, and low variation in performance across oddity categories.

4.5. Conclusions

In conclusion, using MVPA searchlight analyses, in conjunction with high resolution 7T fMRI data from humans performing complex visual concurrent oddity discrimination tasks, our study provides novel support for the importance of the anteromedial (distal) subicular complex but also the inferolateral (proximal) CA1, alongside traditional cortical scene selective regions in the PHC and PMC, to complex scene discrimination. Therefore, our work has contributed high resolution understanding of hippocampal contributions to the human scene network, demonstrating that analysis across and within hippocampal subfields is crucial for our understanding of hippocampal function and anatomy. In addition, using subject level analysis techniques we demonstrate individual variability of category selective areas within the hippocampus. Finally, contrasting scene selectivity in the medial hippocampus, we provide preliminary evidence for face selectivity along the lateral hippocampus, similar to recent MRI analysis showing lateral hippocampal structural connectivity with face-selective cortical regions, and supporting parallel processing models such as the PMAT framework. Future work could focus on structural and (directed) functional

connectivity between these regions to further our understanding of how category information is communicated within category sensitive hippocampal networks.

CRedit authorship contribution statement

Marie-Lucie Read: Data curation, Formal analysis, Project administration, Visualization, Writing – original draft, Writing – review & editing. **Samuel C. Berry:** Data curation, Formal analysis, Project administration, Visualization, Writing – original draft, Writing – review & editing. **Kim S. Graham:** Conceptualization, Writing – review & editing. **Natalie L. Voets:** Investigation, Writing – review & editing. **Jiaxiang Zhang:** Formal analysis, Writing – review & editing. **John P. Aggleton:** Conceptualization, Writing – review & editing. **Andrew D. Lawrence:** Conceptualization, Funding acquisition, Supervision, Writing – review & editing. **Carl J. Hodgetts:** Conceptualization, Data curation, Formal analysis, Funding acquisition, Investigation, Project administration, Supervision, Visualization, Writing – original draft, Writing – review & editing.

Declaration of competing interest

None.

Data availability

Due to ethical concerns, supporting data cannot be made openly available. Further information about the data and conditions for access are available at <https://osf.io/4vkg9/> DOI: 10.17605/OSF.IO/4VKG9

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