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1	
2	Aberrant phase precession of lateral septal cells in a maternal immune activation model
3	of schizophrenia risk may disrupt the integration of location with reward
4	
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28	

Abstract

30	Spatial memory and reward processing are known to be disrupted in schizophrenia.
31	Since the lateral septum (LS) may play an important role in the integration of location and
32	reward, we examined the effect of maternal immune activation (MIA), a known
33	schizophrenia risk factor, on spatial representation in the rat LS. In support of a previous
34	study, we found that spatial location is represented as a phase code in the rostral LS of adult
35	male rats, so that LS cell spiking shifts systematically against the phase of the hippocampal,
36	theta-frequency, local field potential (LFP) as an animal moves along a track towards a
37	reward (phase precession). Whereas shallow precession slopes were observed in control
38	(CTL) group cells, they were steeper in the MIA animals, such that firing frequently
39	precessed across several theta cycles as the animal moved along the length of the apparatus,
40	with subsequent ambiguity in the phase representation of location. Furthermore, an analysis
41	of the phase trajectories of the CTL group cells revealed that the population tended to
42	converge towards a common firing phase as the animal approached the reward location. This
43	suggested that phase coding in these cells might signal both reward location and the distance
44	to reward. By comparison the degree of phase convergence in the MIA-group cells was weak,
45	and the region of peak convergence was distal to the reward location. These findings suggest
46	that a schizophrenia risk factor disrupts the phase-based encoding of location-reward
47	relationships in the LS, potentially smearing reward representations across space.

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50	
51	Significance statement
52	It is unclear how spatial or contextual information generated by hippocampal cells is
53	converted to a code that can be used to signal reward location in regions such as the ventral
54	tegmental area. Here we provide evidence that the firing phase of cells in the LS, a region
55	that links the two areas, may code reward location in the firing phase of cells. This phase
56	coding is disrupted in a maternal immune activation (MIA) model of schizophrenia risk such
57	that representations of reward may be smeared across space in MIA animals. This could
58	potentially underlie erroneous reward processing and misattribution of salience in
59	schizophrenia.
60	

61	Previous studies suggest that the lateral septum (LS) integrates spatial and locomotor
62	information with reward (Bender et al., 2015; Luo et al., 2011; Wirtshafter & Wilson, 2019,
63	2020, 2021). The primary output of hippocampal CA1 cells that signal location (O'keefe &
64	Nadel, 1978) is to the LS (Risold & Swanson, 1997; Swanson & Cowan, 1977), and the LS in
65	turn has reciprocal connections to several regions involved in reward processing, including
66	the ventral tegmental area (VTA) and the striatum (Groenewegen, Vermeulen-Van der Zee,
67	Te Kortschot, & Witter, 1987; Luo et al., 2011; Zhang, Navarrete, Wu, & Zhou, 2022).
68	Consistent with this connectivity, the integrity of LS transmission is required for the
69	acquisition and flexible maintenance of conditioned place preferences (Cazala, Galey, &
70	Durkin, 1988; Jiang et al., 2018).
71	Recent work has shown that the LS uses both rate and phase coding to represent
72	location (Takamura et al., 2006; Tingley & Buzsáki, 2018; Wirtshafter & Wilson, 2020;
73	Zhou, Tamura, Kuriwaki, & Ono, 1999). For example, rate-coding cells that fire when an
74	animal is in a specific region of space, known as the cell's "place field" have been observed
75	in more dorsal regions of LS, although these place fields are not as robust as those observed
76	in the hippocampus (Takamura et al., 2006; Wirtshafter & Wilson, 2020). However, in more
77	rostral regions of the LS, cells display little evidence of rate coding but appear to encode
78	spatial location via phase precession (Tingley & Buzsáki, 2018).
79	Phase precession refers the observation that, as an animal moves through space, the
80	temporal spiking of principal cells systematically advances relative to the background theta
81	oscillation (O'Keefe & Recce, 1993; Skaggs, McNaughton, Wilson, & Barnes, 1996). When
82	phase precession occurs in an assembly of cells it produces theta sequences (Foster &
83	Wilson, 2007), thereby allowing for the sequential order of experience to be reproduced
84	within a compressed timescale that is suitable for synaptic plasticity (Dan & Poo, 2004). It
85	has been proposed that these phenomena underlie the sequential ordering of information that

86 evolves across time and space (Buzsáki & Tingley, 2018), and that they may play an

87 important role in learning and memory processes (Dragoi & Buzsáki, 2006; Jaramillo &

88 Kempter, 2017). These proposals have been supported by a growing body of work (Feng,

89 Silva, & Foster, 2015; Gupta, Van Der Meer, Touretzky, & Redish, 2012; Terada, Sakurai,

90 Nakahara, & Fujisawa, 2017; Wang et al., 2015; Wikenheiser & Redish, 2015), and recent

91 evidence has described phase precession and theta sequences in humans (Heusser, Poeppel,

92 Ezzyat, & Davachi, 2016; Qasim, Fried, & Jacobs, 2020).

93 Neurodevelopmental abnormalities and dysfunctional activity have been observed in 94 the septum of individuals with schizophrenia, including abnormal spiking activity and LFP 95 oscillations (Heath & Peacock, 2013; Heath & Walker, 1985). Furthermore, changes in LS 96 activity have been observed in both in vivo and in vitro studies following administration of 97 either antipsychotic or dissociative drugs in animal models (Contreras, Dorantes, Mexicano, 98 & Guzmán-Flores, 1986; Sheehan, Chambers, & Russell, 2004; Yu et al., 2002). The spatial 99 and contextual memory deficits (Brébion, David, Pilowsky, & Jones, 2004; Fajnerová et al., 100 2014; Glahn et al., 2003; Hanlon et al., 2006; Park & Holzman, 1992; Rizzo et al., 1996; 101 Salgado-Pineda et al., 2016; Waters, Maybery, Badcock, & Michie, 2004; Weniger & Irle, 102 2008) and abnormal reward processing (Jensen et al., 2008; Strauss, Waltz, & Gold, 2013; 103 Whitton, Treadway, & Pizzagalli, 2015) that have been observed in schizophrenia may 104 therefore be linked to LS dysfunction. 105 In the present study we investigated whether a risk factor for schizophrenia, 106 maternal immune activation (MIA) altered LS activity. The MIA model is based on robust 107 epidemiological evidence that maternal infection during pregnancy increases the risk of 108 schizophrenia in the offspring (Adams, Kendell, Hare, & Munk-Jørgensen, 1993; Brown &

109 Meyer, 2018). When this is modelled in rodents, MIA animals have many schizophrenia-like

behavioral, cognitive and neural deficits (Bitanihirwe et al., 2010; Savanthrapadian et al.,
2013; Wolff, Cheyne, & Bilkey, 2011).

- 112
- 113 Materials and methods

114 ANIMALS AND EXPERIMENTAL DESIGN

115 All subjects were generated using the MIA intervention described previously by 116 Dickerson, Wolff, and Bilkey (2010), Wolff and Bilkey (2015) and Speers et al. (2021). 117 Female Sprague Dawley rats (~3 months old) were time-mated with GD1 considered to be 118 the first day after copulation. On GD 15, pregnant dams were anesthetized with isoflurane 119 (5%; Bayer) and administered either a single injection of polyinosinic:polycytidylic acid 120 (poly I:C; Sigma-Aldrich) 4.0 mg/kg, i.v. dissolved in 0.9% saline (Baxter), or an equivalent 121 saline injection 1 ml/kg. This dosage is the most common induction protocol used for rats 122 (Haddad, Patel, & Schmid, 2020). A number of previous studies have examined the precise 123 timing of injections on developmental phenotypes, with injections performed around GD 15 124 in rats leading to more robust phenotypes associated with schizophrenia than earlier injection 125 protocols, which have been associated more with autism spectrum disorders (Haddad et al., 126 2020). Poly I:C and saline treatments were always performed in pairs. 127 Due to resource limitations, all litters were culled to a maximum of 6 male pups and 128 were housed in open cages prior to weaning. Post-weaning, male offspring were randomly 129 allocated a litter number and then housed in littermate groups of 2-3 in individually ventilated 130 cages (IVC). CTL and MIA animals were housed in a single housing room, which was 131 maintained at a normal 12-h light/dark cycle, and temperature controlled to 20-22°C. Juvenile 132 rats were provided with access to food *ad libitum*, and after 3 months were food deprived to 133 no less than 85% of their free-feeding weight in preparation for the experimental procedure.

Water was available *ad libitum* throughout the entire experimental procedure. All rats
weighed between 400 and 650g at the time of surgery.

136

137 APPARATUS AND TRAINING

138 Animals ran in a rectangular circuit measuring 900 by 800mm (Figure 1a). All arms 139 were 100mm wide with 270mm high side walls and constructed of wood. The entire 140 apparatus was painted in matte black and was devoid of visual cues. A video camera was 141 mounted on the ceiling of the recording room to view the whole apparatus. All experiments 142 were performed in a darkened environment with some ambient light from the recording 143 computer and a small lamp aimed away from the apparatus into one corner of the room. 144 The experimental procedure was identical to the procedure described previously in 145 Speers et al. (2021). Adult male offspring were randomly selected according to their litter 146 number, with a maximum of two animals per litter, and were trained over a period of 5 to 15 147 days. On days 1-5 rats were habituated to the recording room, apparatus and food reward, and 148 were allowed to free-forage for Coco Pops (Kellogg Company) scattered throughout the 149 apparatus. Following successful habituation, whereby rats actively explored the maze and 150 consumed the food reward, the placement of Coco Pops was gradually restricted, first to the 151 top 2 corners of the track and the centre of the reward arm, and then to the reward arm only. 152 During this period, rats were trained to run in a clockwise direction and were turned back to 153 the correct direction with a barrier when necessary. Coco-pops (approx. 6 per reward 154 delivery) were delivered manually by the experimenter. Training was considered completed 155 when rats consistently ran in a clockwise direction for the food reward over a twenty-minute 156 session.

157

158 SURGICAL PROCEDURES

159	All experimental protocols were approved by the Otago University Animal Ethics
160	Committee and conducted in accordance with New Zealand animal welfare legislation.
161	Following successful training, animals were anesthetized with 5% isoflurane (Merial New
162	Zealand) in oxygen and maintained at 1.5 to 2.5% throughout surgery. After animals were
163	anesthetized, they were given a subcutaneous injection of Atropine (1mg/kg) to ease their
164	breathing, as well as the analgesics Carprofen (1mg/kg) and Temgesic (buprenorphine;
165	0.1mL), and a prophylactic antibiotic, Amphoprim (trimethoprim and sulfamethazine,
166	0.2mL). Rats were then mounted on a stereotaxic apparatus (David Kopf Instruments) above
167	a heating pad, and a lubricating eye gel (Visine) was applied. The scalp was shaved and
168	sterilized with Betadine (Povidone-iodine), followed by a subcutaneous injection in the scalp
169	of the local anesthetic Lopaine (lignocaine hydrochloride 20mg mL ⁻¹ ; 0.1mL diluted in
170	0.4mL of saline). After exposing the skull, two openings were drilled above the left
171	hemisphere, one above the dorsal CA1 region of the hippocampus, and one above the septal
172	region. A custom built, 8 channel, adjustable microdrive containing 1 tetrode and 1 tritrode
173	bundle of equal length was implanted at +.5mm AP, -1.5mm ML, and was lowered to \sim 4mm
174	from dura at an angle of \sim 7-8 degrees towards the midline (Figure 1b. Tetrodes consisted of
175	$25\mu m$ nichrome, heavy formvar insulated wire (Stablohm 675 HFV NATRL; California Fine
176	Wire Company), and had been gold electroplated until impedances were reduced to $\sim 200 -$
177	300 k Ω (NanoZ, Neuralynx). A non-movable LFP electrode was separately implanted in CA1
178	at -3.8mm AP from bregma and -2.5mm ML from the midline, and then lowered to 2mm
179	from dura (Figure 1c). Microdrives were secured to the skull with jewellers' screws and
180	dental cement, and a ground wire was secured to an additional screw placed above the right
181	hemisphere. Post-surgery rats received a secondary dose of Amphoprim immediately upon
182	waking, and then an additional dose of Carprofen 24 hours later. Rats were provided with ad
183	libitum food and water post-surgery and were given 8 days to recover.

185

EXPERIMENTAL PROCEDURE AND ELECTROPHYSIOLOGICAL

186 RECORDINGS

187 Following recovery, rats were again food deprived to no less than 85% of their free-188 feeding weight. Post-operative training and test trials were carried out in the recording room. 189 Rats were attached to a multichannel data acquisition system (DacqUSB; Axona Ltd), and 190 single unit data was closely monitored during test trials, which consisted of a 5-10 minute 191 recording session. Extracellular unit activity was first passed through an AC-coupled unity 192 gain amplifier before passing through to the recording system. Single unit data was bandpass 193 filtered between 600 and 6000 Hz, and sampled at a rate of 48 kHz with 24-bit resolution. For 194 each tetrode, one electrode with minimal spiking activity was selected as a reference. Action 195 potential thresholds were set at a minimum of $70 - 80 \mu V$ and recorded for a 1 ms window 196 whenever the spiking amplitude was above this threshold. All spike events were time-197 stamped relative to the beginning of the recording. LFP data was simultaneously recorded 198 from the CA1 region, was bandpass filtered up to 500 Hz (with notch filtering selective for 199 activity at 50 Hz) with a gain of ~500, and sampled at 48 kHz. The animal's location was 200 determined from 3 infrared LEDs mounted on the animal's head-stage and recorded by a 201 camera located above the chamber. Positional data was analysed with a sampling rate of 50 202 Hz and then converted into x and y coordinates by the recording system. 203 During the test period, tetrodes were slowly lowered ($\sim 40 \mu m$ per day) until well-204 isolated single units were identified. Once single unit activity was confirmed, tetrodes were 205 lowered an additional \sim 40µm after every second recording for the remainder of the 206 experimental procedure. Experimental recordings were 20 minutes long, and testing 207 continued for \sim 3-8 weeks, until there was no further evidence of single unit activity, manual

adjustment had reached its limit, or the rat experienced other difficulties that terminated the

209	experiment. Final electrode placements are shown in Figures 1e and 1f. Rats ran no more
210	than one session per day, for $\sim 60 - 80$ laps per session. Single unit, position and LFP data
211	was saved for later analysis. All recordings with at least 1 putative place cell were included in
212	the final dataset.
213	
214	ISOLATION OF SINGLE UNITS
215	For each recording, single units were identified manually offline using purpose
216	designed cluster cutting software (Plexon Offline Sorter, Version 3), primarily via the peak-
217	to-valley distance and principal components analysis of the waveforms. All stable waveforms
218	with clearly observed spike clustering were included in the initial analysis, regardless of
219	spike-width or firing rate. Example waveforms and cluster cutting from both CTL and MIA
220	recordings are presented in Figures 1d. Sorted data was then exported to MATLAB (version
221	R2019a, MathWorks), and analysis of single unit, position and LFP data was carried out in
222	MATLAB with custom-written scripts.
223	
224	SELECTION OF FIRING ONSET LOCATION AND PHASE PRECESSION ANALYSIS
225	Initial inspection of firing properties around the track indicated that, although some
226	cells only fired in a portion of the track, the majority of cells fired indiscriminately across the
227	entire track. Where it did occur, the onset location of firing also appeared to vary from cell to
228	cell, and could appear at any point along the track. Due to this variability and lack of clear

place fields, manual selection of firing starting location and termination was used for phase precession analysis. To these ends all cell recordings were split into groups of ~20 cells and assigned a blinded identifier to ensure experimenter bias was minimized during the manual selection process. These blind groups were then analysed with a custom MATLAB script that first linearized the track, and then allowed the experimenter to select the start and end

234	locations of firing across 2 cycles of the track. For cells that only fired across a portion of the
235	track, firing onset and offset locations were always selected as the locations where robust
236	firing began and ended in a clockwise direction respectively. For cells that fired
237	indiscriminately across the entire track, the start location was selected on the basis of the
238	following criteria, in order: 1) a small pause in the firing, 2) the location where clear phase
239	precession could be observed to begin relative to random noise, and 3, if no clear firing
240	pauses or phase precession relative to noise were observed, then the analysis region was
241	always started just after the reward location, and ended just before the reward location.
242	
243	DATA ANALYSIS
244	LFP activity recorded from electrode located in CA1 was sampled at 4800Hz. To
245	determine theta waveform shape, the LFP was bandpass filtered between 6-10Hz and a phase
246	profile was determined using the Hilbert transform. A sample waveform of 200 ms duration
247	was subsequently captured whenever the phase data indicated a trough had been reached.
248	These samples were then averaged, as were the related phase profiles.
249	Spatial information values, a measure of how informative a spike from a cell is
250	regarding the animal's current location within an environment, were calculated according to
251	the method described in Skaggs, McNaughton, and Gothard (1993). The formula for
252	information content, measured in bits per spike is:
	$Information = \sum_{i=1}^{N} p_i \frac{\lambda_i}{\lambda} \log 2 \frac{\lambda_i}{\lambda}$

where the environment is divided into *N* distinct bins (i = 1, ..., N), p_I denotes the occupancy probability of bin *i*, λ_i is the mean firing rate for bin *i*, and λ is the overall mean firing rate of the cell. Higher information values indicate that cells provide a more reliable prediction of current location than cells with lower information values. Correlations of hippocampal theta frequency and speed were generated for each recording that showed evidence of single unit activity in the LS. This process involved estimating instantaneous values for theta frequency from the Hilbert transform of LFP filtered between 6 and 10 Hz. Estimates of instantaneous speed were determined by monitoring the animals change in position over 500 ms time windows. Speed and theta frequency data were then sampled at one second intervals and correlated. Samples where speed was below 5 cm/s were excluded from the analysis.

264 For all phase precession analyses, the phase reference was always to the LFP signal 265 recorded from the non-movable electrode implanted in CA1, where a phase of zero 266 corresponded to the trough of the oscillation. Phase precession was determined by matching 267 the animal's position to the instantaneous phase of the 6-10 Hz theta rhythm at the CA1 268 reference, as determined from the Hilbert transform. These data were then analysed using 269 procedures described previously (Kempter et al., 2012; Speers et al., 2021). This involves 270 using circular-linear regression to provide a robust estimate of the slope and phase offset of 271 the regression line, and a correlation coefficient for circular-linear data analogous to the 272 Pearson product-moment correlation coefficient for linear-linear data. Phase precession 273 analysis was conducted by pooling spiking data from all passes through the region of interest 274 within a given recording session. The number of phase cycles per track was calculated as the 275 absolute value of the slope (in degrees per mm) multiplied by the length of the full track, and 276 then divided by 360. LS phase precession as described by Tingley and Buzsáki (2018) would 277 generally produce a value of around 1.

Correlations of firing rate and either speed or acceleration were based on the process outlined by Wirtshafter and Wilson (2019). For this analysis, position was sampled every 100 ms to estimate instantaneous speed. These data were then smoothed across a 500 ms window. The animals' occupancy per speed within 2cm/s bins was then established and then

282 spike count as a function of speed was determined. Spike count per speed was then divided 283 by speed occupancy to result in firing rate as a function of speed for each cell of interest. 284 Speeds with less than 2% of total occupancy were excluded from the analysis. The 285 correlation between speed and firing rate was assessed using a linear regression. Correlations 286 with acceleration were determined similarly except that a bin size of 0.5 cm/s^2 was used. 287 To allow between-animal comparison of phase trajectories across the track it was first 288 necessary that any phase shift that might have resulted from variation in the depth of the 289 hippocampal electrode was minimised. To this end the phase relationship between theta 290 activity recorded at the moveable LS electrode was compared to that recorded at the fixed 291 hippocampal electrode using cross correlation of LFP data filtered between 6-10Hz. It was 292 determined that phase shifted systematically as the LS electrode was lowered, but for some 293 animals this phase/depth relationship was offset at equivalent LS electrode depths. This 294 indicated that the depth of the reference hippocampal electrode was different between 295 animals, as theta phase varies depending on electrode position above and below the CA1 cell 296 layer (Brankačk, Stewart, & Fox, 1993; Buzsáki, Rappelsberger, & Kellényi, 1985; Lubenov 297 & Siapas, 2009). With this information theta-frequency LFP phase difference between the 298 two electrodes was normalised to zero when LS electrodes were 4.5 mm deep from the dura 299 (see Figure 6b). All normalised phase trajectories were projected both forward and backwards 300 across the full extent of the linearized track. To determine whether there was any tendency 301 for phase trajectories to intersect at a particular location along the track, the phase angle of 302 each phase trajectory was determined in one cm increments along the track. A population 303 vector was then calculated for each location for all intercepting phase trajectories. The 304 subsequent population vector could vary from zero, indicating no clustering of phase 305 trajectories at this location, to one, which would indicate that all phase trajectories intersected 306 at one phase angle at that location.

307	To determine whether LS cells tended to fire in bursts at near-theta frequencies, an
308	autocorrelation of cell spiking with a +/-500 ms window was conducted across each
309	recording. The dominant frequency between 6 and 10 Hz was determined from the power
310	spectrum of the autocorrelation function.

312 HISTOLOGY

313 Following completion of experiments, animals were anaesthetised with 5% isoflurane 314 in oxygen, and a 2mA direct current was passed through each electrode for approximately 1 315 second to lesion the site of the electrode tip. Rats were then euthanized with an overdose of 316 isoflurane and transcardially perfused, first with 120 ml of 0.9% saline, and then 120 ml of 317 10% formalin in saline. Brains were then carefully extracted from the skull after removal of 318 the Microdrive, and stored in 10 % formalin in saline. One week prior to sectioning, brains 319 were transferred first to 10% formalin in H_2O for 24 hours, and then to a 10% formalin/30% 320 sucrose solution for approximately 3-7 days, until the brain sunk to the bottom of the sucrose 321 solution. Dehydrated brains were then sectioned into 60 µm coronal slices with a cryostat 322 (Leica CM1950). Sections were then mounted on slides and stained with a thionine acetate 323 Nissl stain (Santa Cruz Biotechnology, Inc. After slides were dry (min. 24 hours) electrode 324 placement was imaged with a local power (1.5x) digital microscope (Leica Biosystems, LLC) 325 to verify electrode placement (Figures 1e and 1f).

326

327 STATISTICAL ANALYSES

For all statistical analyses, we performed the following procedure. First, raw data was transformed to a lognormal distribution if appropriate. All data (either in raw form or the log transform) were then checked for assumptions of normality. These checks were performed in GraphPad Prism 8.1.1 (GraphPad Software, Inc., San Diego, CA, USA), using the d'Agostino

332	& Pearson test for normality. If data did not meet the assumptions for normality based on the
333	d'Agostino & Pearson test, visual inspection of histograms and QQ plots was performed, and
334	extreme outliers were removed using the Graphpad function for removal of outliers. All data
335	that failed to meet assumptions of normality based on this procedure were then analysed
336	using the appropriate non-parametric test. Details about the specific tests used are provided in
337	the results section. All t-tests were two-tailed. Data with a normal distribution are presented
338	as mean \pm SEM unless explicitly stated otherwise in the figure legends. For all data that did
339	not meet normality assumptions, the median with 95% confidence intervals is depicted
340	instead. Significance levels were defined as $p < 0.05$. Additional information about
341	significance levels is provided in the figures as: * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$.
342	Additional circular statistics (to compare group differences in the intercept of the circular
343	correlation of phase and position, and to generate the MVL for animal by animal and litter by
344	litter analyses) were performed in Oriana 4 (Kovach Computing Services, Inc., Anglesey,
345	UK). Group differences for angular variance (defined as 1-MVL) were performed using the
346	variance ratio F-test, found at https://www.statskingdom.com/220VarF2.html
347	
348	Results
349	Tetrode verification in the rostral LS
350	Single units were recorded from eight animals (7 litters) in both the CTL and MIA
351	groups respectively. However, one CTL animal was later excluded from further analyses due
352	to electrode misplacement in the medial septum (figure 1e). This decision was made not only
353	based on tetrode placement close to the MS, but also following inspection of other firing
354	properties suggesting that cells recorded from this animal were atypical. For example, cells
355	recorded from this animal had a significantly higher firing rate when compared to the average
356	firing rate of all other cells (figure 1e, lover right corner), and inspection of phase precession

357	plots showed that these cells were tightly coupled to specific phases of the LFP theta rhythm,
358	consistent with the firing properties of GABAergic medial septal neurons (Borhegyi et al.,
359	2004). After excluding this animal, a total of 144 units from seven CTL animals (6 litters)
360	and 362 units from eight MIA (7 litters) were used for all subsequent analyses. All animals
361	were age matched except for one pair, in which the CTL animal was seven months and the
362	corresponding MIA animal was twelve months, and one non-paired MIA animal (12 months).
363	All ages refer to animal age during experimental recordings, and ranged from 4 to 14 months.
364	There was no difference in the mean ages between groups (CTL = 10.43 ± 1.49 , MIA = 11.25
365	\pm 1.13, t(13) = 0.45, p = 0.664).
366	
367	Basic firing properties of LS cells, mean locomotor speed, and comparison of
368	hippocampal theta properties
369	The median firing rate for all LS cells was 0.44 Hz, 95% CI [0.35, 0.51]. The
370	median log transformed firing rate was significantly higher for cells in the MIA group (-0.25,
371	95% CI [-0.34, -0.11] than in the CTL group (-0.71, 95% CI [-0.80, -0.59], Mann Whitney U
372	=16638, p <0.001; Figure 2a). Visual inspection of the distribution of firing rates around the
373	track indicated that activity was distributed relatively diffusely across the track for both
374	groups. In support of this observation, the spatial information content measure for all cells
375	was low (median = 0.69 bits/spike, 95% CI [0.59, 0.77]). It was, however, significantly lower
376	for cells in the MIA group (median = 0.59, 95% CI [0.51, 0.71] compared to cells in the CTL
377	group (median = 0.82, 95% CI [0.71, 0.96], Mann Whitney U = 22555, p = 0.007; Figure 2b).
378	In a further quantification of spatial firing we identified the subset of cells where the
379	contiguous region of firing (as selected by the firing onset and offset locations) was less than
380	half the track length (<150cm), and where the spatial information content was greater than
381	0.8 bits/spike (Wirtshafter & Wilson, 2019). Only 4.7 and 5.8% of cells in the CTL and MIA

groups respectively met these criteria, confirming that spatially selective activity was rare for both groups. There was no significant difference in these proportions between groups ($\chi^2(1) =$ 0.54, p = 0.46).

385 The mean theta-frequency LFP amplitude recorded from electrodes located in the 386 CA1 region were significantly higher in the CTL group (M (log-transformed) = 3.17 ± 0.03 , 387 MIA M = 3.07 ± 0.02 , t(286) = 2.66, p = 0.008; Figure 2d). The mean frequency of theta-388 band CTL LFP recordings (M=7.92 Hz ± 0.02) was also significantly higher than in MIA 389 recordings (M = $7.72 \text{ Hz} \pm 0.02$, t(286) = 7.0, p< 0.001; Figure 2f), although the phase profile 390 was virtually identical for both groups (Figure 2h). 391 Mean running speed was significantly lower in the MIA group (29.61 cm/s \pm 0.50) 392 than for the CTL group (M=31.42 cm/s \pm 0.57, t (279) = 2.35, p = 0.02; Figure 2c). To 393 determine whether this affected the theta-locomotion relationship (Bender et al., 2015) we 394 sampled hippocampal theta frequency and speed values once every second and computed the 395 correlation between these values for each individual recording. A comparison of the resultant 396 r values revealed that the median r value was slightly lower in the CTL group (median = 0.35, 397 95% CI [0.30,0.39]) when compared to the MIA group (median = 0.39, 95% CI [0.34, 0.24], 398 but the difference was not significantly different (Mann Whitney U = 8482, p = 0.059), 399 although only marginally. There were no significant differences for either the slope of these 400 correlations (CTL M = 6.97 ± 0.38 , MIA M = 7.36 ± 0.31 , t(284) = 0.80, p = 0.426), or the 401 intercept (CTL M = -37.12 ± 3.02 , MIA M = -39.86 ± 2.49 , t(284) = 0.7, p = 0.485). 402 403 LS cells in the MIA group continue to demonstrate robust phase precession, but are 404 more likely to precess through multiple theta cycles across the running track 405 In total 34.03% of all cells in the CTL group and 30.12 % of all cells in the MIA 406 group showed evidence of significant phase precession (Figure 4a). This difference was not

407	statistically significant (χ^2 (1) = 0.74. p = 0.391). Examples are provided in Figure 3a (CTL)
408	and 3b (MIA). The circular-linear correlation of phase and position, as calculated across all
409	cells, was significantly higher in the CTL group (median = -0.07, 95% CI [-0.08, -0.05], MIA
410	median = -0.03, 95% CI [-0.04,-0.02], Mann-Whitney U = 21292, p = 0.001), although the p-
411	values of those correlations were not different between groups (CTL median = 0.162 , 95% CI
412	[0.109, 0.269], MIA median = .244, 95% CI [0.191, 0.313], Mann Whitney U = 25017, p =
413	0.481). The same analyses were then performed on the subset of cells showing significant
414	phase precession. The difference between the circular-linear correlations for this subset was
415	non-significant although the result was marginal (CTL mean r = -0.14 \pm 0.02, MIA mean r =
416	-0.10 \pm 0.01, t (156) = 1.94, p = 0.054; Figure 4b). A similar, marginal result was obtained
417	for the p-value of the circular-linear correlation (CTL median = 0.004 , 95% CI [0.001 ,
418	0.012], MIA median = 0.001, 95% CI [<0.001, 0.003], Mann Whitney U = 2155, p = 0.052;
419	Figure 4c).
420	Visual comparison of phase precession trajectories across the track indicated that
421	many cells in the MIA group precessed through several theta cycles over the circuit (Figure
422	3b). By comparison this firing behaviour was observed in few cells from the CTL group.
423	Furthermore, when firing in MIA animals was analysed on a pass by pass basis, there were a
424	number of examples where firing cycled through more than 360 degrees, indicating that the
425	multiple-cycle precession was not simply a cumulative effect produced by variation on
426	individual passes (Schmidt et al., 2009). Analysis of the slope values obtained from the

- 427 circular-linear fit from all cells revealed that MIA group cells had a significantly steeper
- 428 slope when compared to CTL group cells (CTL median = -352 deg/mm, 95% CI [-435.7, -
- 429 188.3], MIA median = -522.2 deg/mm, 95% CI [-623.2, -468.1], Mann Whitney U = 20158,
- 430 p <0.001; Figure 4d, left side). A similar result was obtained for the subset of cells
- 431 demonstrating significant phase precession (CTL median = -173.8 deg/mm, 95% CI [-351, -

432	122], MIA median = -526.4 deg/mm, 95% CI [-700, -424.9], Mann Whitney U = 1536, p
433	<0.001; Figure 4d, right side). The length of track over which these slope values were
434	calculated was not significantly different between groups (CTL median = 2662mm, 95% CI
435	[2523, 2871], MIA median = 2585mm, 95% CI [2468, 2815], Mann Whitney U = 2523, p =
436	0.581). By combining these slope values and the length of the track over which they were
437	calculated, it was possible to determine the number of phase precession cycles that would
438	occur across the entire track length, were precession to continue across the whole region. For
439	cells that demonstrated significant phase precession, the median number of phase precession
440	cycles in the CTL group was 0.79. This was significantly lower than the median number of
441	cycles in the MIA group (1.89 cycles, Mann Whitney U = 1562, p <0.001; Figure 4e).
442	To confirm that these slope differences were not a result of aberrant recordings from
443	a small proportion of MIA animals, mean values were also computed for each individual
444	animal and then compared across groups. For this analysis, only cells that demonstrated
445	significant phase precession were analysed. Comparison of slope values on an animal by
446	animal basis showed that MIA slope values were significantly steeper than CTL slope values
447	(CTL mean = -325.6 deg/mm \pm 50.28, MIA mean = -534.2 deg/mm \pm 69.81, t(13) = 2.36, p =
448	0.035; Figure 4f).
449	Dual oscillator theories of phase precession suggest that a change in the slope of
450	precession in MIA animals might result from an alteration in the theta-frequency/cell-burst-
451	firing-frequency relationship (Kamondi, Acsády, Wang, & Buzsáki, 1998; Magee, 2001;
452	Mehta, Lee, & Wilson, 2002; O'Keefe & Recce, 1993). To test this hypothesis, the burst
453	firing frequency of single cells was calculated. For both MIA and CTL groups mean burst
454	firing frequency was slightly higher than theta frequency, but there was no significant

455 difference in cell-burst frequency between the two groups (CTL median = 8.3 Hz, 95% CI

456 [8.06, 8.55], MIA median = 8.06 Hz, 95% CI [8.06, 8.06], Mann Whitney U 11452, p =

457 0.658; Figure 4h). As described previously, theta frequency was significantly lower in MIA 458 animals. As an approximation of how this difference might affect precession, for controls the 459 theta/burst relationship would result in cell firing precessing through a full 360 degrees in 460 approximately three seconds. In contrast, in MIA animals this precession would occur in 461 around 2.3 seconds. Thus, based on this difference MIA would precess around 30% faster 462 than control animals. This contrasts with the actual difference in precession cycles around the 463 track, which is closer to a 2-fold difference between MIA and controls (figure 4e). 464 465 Starting phase is more variable for MIA cells when compared to CTL cells 466 To determine the phase of cell firing as an animal enters the analysis region, the 467 intercept of the regression line for the circular linear-correlation was examined. Only cells 468 that demonstrated significant phase precession were included in this analysis. LS cells in both 469 the CTL and MIA groups demonstrated significant clustering at a mean phase angle of 470 314.83° (CTL Raleigh Z = 15.78, p < 0.001) and 288.96° (MIA Raleigh Z = 12.36, p < 0.001) 471 0.001), around the starting phase. The circular variance of starting phase for MIA group cells 472 was, however, significantly greater than for CTL group cells (CTL = 0.43, MIA = 0.66, F = 473 0.43, p = 0.001; Figure 4i). The Mardia-Watson Wheeler test, which considers group 474 differences in both the mean and variance for circular data, also returned a significant result 475 (W = 8.99, p = 0.011). As a further test, the circular distance from the mean angle was 476 computed for each cell and then compared across groups. Again, MIA group cells had a 477 significantly higher median distance from the mean angle when compared to CTL group cells 478 (Mann Whitney U = 1996, p = 0.011), indicating that precession starting phase was more 479 variable in the MIA group.

Lead/lag times between the hippocampus and LS vary systematically according to

482	electrode depth	
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483 Previous studies indicate that the firing phase of LS cells changes systematically 484 according to electrode depth (Tingley & Buzsáki, 2018). To examine this possibility, a 485 lead/lag analysis was performed for each recording to determine the phase shift of LS LFPs 486 when referenced to the non-movable hippocampal electrode. These results were then 487 correlated with LS electrode depth on an animal by animal basis (figure 5a). In the majority 488 of animals, the LS was more likely to lead the hippocampus at shallow depths, but as 489 electrode depth increased, the hippocampus was more likely to lead the LS. However, this 490 relationship was not always observed in the MIA group, with 3 animals showing the opposite 491 relationship. A t-test of the regression slopes of the depth/phase relationship indicated that mean slopes were more positive in the CTL group (M = 35.37 ± 6.0 when compared to the 492 493 MIA group ($M = 8.82 \pm 9.72$, t(13) – 2.24, p = 0.043). Inspection of the individual animal 494 phase and LS electrode depth relationship suggested that a number of animals (across both 495 groups) had greater hippocampal lead relative to depth (figure 5a). This was most likely due 496 to variability in hippocampal electrode depth (Brankačk et al., 1993; Buzsáki et al., 1985; 497 Lubenov & Siapas, 2009). To correct for this in further analyses, the phase of all recordings 498 was shifted, on a whole-animal basis, to align with a reference where zero phase shift 499 occurred at an electrode depth of 4500 microns (figure 5b).

500

501 *Relationship between firing phase and reward location*

502 In prior analyses LS cell phase precession had been examined across sections of the 503 track where firing occurred. As a result, data from different cells was often from different 504 (although often overlapping) regions of the track. To examine how the phase trajectories of 505 individual cells (the best fit to the firing phase-position data) would appear if precession was

506	assumed to continue from the start of the linearized track to the end, these phase trajectories
507	were normalized for differences in lead/lag likely resulting from variation in hippocampal
508	electrode position as described above, and then extended in each direction so as to cover the
509	whole track (Figures 5c and f). Inspection of these plots for CTL group cells indicated that a
510	large number of these individual phase trajectories tended to intersect near the location on the
511	track where the reward was delivered. To quantify this effect, each trajectory was allocated a
512	phase angle at each location on the track, based on the phase of firing at this location. Then
513	for each location on the track the average phase angle and mean vector length (MVL) across
514	the whole population of trajectories was determined. This procedure was repeated at one cm
515	increments along the whole track. The MVL thus provided a measure of the degree of
516	intersection of the phase trajectories at each location, which could vary from zero, indicating
517	no clustering of phase trajectories at that location, to one, which would indicate that all phase
518	trajectories intersected at that location. These resultant data indicated that CTL group MVL
519	was highest (0.50) at the reward location (Figure 5d). At the location with the highest MVL,
520	CTL group phase angles were significantly clustered with a mean phase angle of 200.06°
521	(Rayleigh Z = 11.78, $p < 0.001$; Figure 6e). When the same analysis was applied to the MIA
522	data, phase trajectories were overall less convergent (maximum MVL=0.24) with greatest
523	convergence occurring well prior to the reward location (Figure 5g). At the reward location
524	there was little evidence of convergence in MIA trajectories (MVL = 0.08 ; mean angle of
525	236.29°, Raleigh Z = 0.699, p = 0.497; Figure 5h). A Mardia-Watson-Wheeler test revealed
526	significantly greater clustering at the reward location in the CTL compared to the MIA
527	groups (MWW= 12.92, p = 0.002).

Firing rates of a substantial subset of cells located in the rostral LS are significantly

530 correlated with speed for both groups, but positive correlations were significantly more

531 frequent in the MIA group

532 Recent work has demonstrated that the dorsal LS contains a population of cells for 533 which firing rate has a strong linear relationship with either speed or acceleration (Howe & 534 Blair, 2020; Wirtshafter & Wilson, 2019). To examine whether cells located in the rostral LS 535 are also modulated by speed, we computed Pearson correlations of speed and firing rate for 536 each cell (example correlations are provided in Figure 6a. In total, just over 30% of all CTL 537 group cells had firing rates that were significantly correlated with speed, in contrast to almost 45% of all MIA group cells. These proportions were significantly different ($\chi^2(1) = 8.6$, p = 538 539 0.003. From these significant subsets, 45% of all cells in the CTL group and 59% of cells in 540 the MIA group had firing that was positively correlated with speed (Figure 6b). Again, these proportions were significantly different between groups ($\chi^2(2) = 10.61$, p = 0.005). When the 541 542 absolute median r-values generated by these correlations were compared across the entire 543 dataset, they were also significantly higher in the MIA group (CTL median = 0.41, 95% CI 544 [0.31, 0.51], MIA median = 0.56, 95% CI [0.51, 0.60], Mann Whitney U = 20426, p < 0.001: 545 Figure 6c). The median r-values were also compared separately according to the direction of 546 these correlations for those data where significant correlations between firing rate and speed 547 were observed. Results from this reduced data subset showed that there were no group 548 differences for the median r-values for either positive correlations (CTL median = 0.77, 95%549 CI [0.69, 0.88], MIA median = 0.81, 95% CI [0.78, 0.85], Mann Whitney U = 861, p = 0.517) 550 or negative correlations (CTL median = -0.75, 95% CI [-0.80, -0.70], MIA median = -0.79, 551 95% CI [-0.81, -0.74], Mann Whitney U = 674, p = 0.245, Figure 6c). 552 To test the possibility that spatial and locomotor information map onto distinct cell 553 populations, the group of cells demonstrating significant phase precession ("phase coding

554 cells") was compared with the group of cells that had firing rates significantly correlated with 555 speed ("speed modulated cells") to determine if there was any overlap. Of the 49 phase 556 coding cells in the CTL group, 18 were also speed modulated cells, including 12 cells with a 557 positive speed correlation. In total, only 24% of all CTL group cells that had either phase 558 coding or speed modulated properties were involved in both these processes simultaneously. 559 In the MIA group, 59 out of 109 phase coding cells were also classed as speed modulated 560 cells, including 43 cells with positive speed correlations (Figure 6d). In total, this amounted 561 to 28% of cells with overlapping coding properties. These proportions were not significantly different between groups ($\gamma^2(1) = 0.41$, p = 0.52). 562 563 The firing of a relatively small proportion of cells were modulated by the animal's 564 acceleration. In total 8% of CTL group cells and 9% of MIA group cells displaying a 565 significant correlation with either acceleration or deceleration (Figure 6e). These proportions were not significantly different between groups ($\chi^2(2) = 1.57$, p = 0.456). The difference 566 567 between the absolute r-values were also not significant for either acceleration (CTL median = 568 0.25, 95% CI [0.17, 0.29], MIA median = 0.24, 95% CI [0.19, 0.28], Mann Whitney U = 569 14400, p = 0.657) or deceleration (CTL median = 0.23, 95% CI [0.21, 0.27], MIA median = 570 0.22, 95% CI [0.20, 0.26], Mann Whitney U = 24385, p = 0.673; Figure 6e), and similar 571 results were obtained when only those cells with significant correlations were included in the 572 analysis (for acceleration, CTL M = 0.67 ± 0.03 , MIA M = 0.74 ± 0.05 , t(18) = 1.07, p = 573 0.299; for deceleration, CTL M = 0.64 ± 0.05 , MIA M = 0.66 ± 0.04 , t(20) = 0.21, p = 0.834). 574 Discussion 575

We investigated whether MIA altered neuronal coding of location in the rostral LS, a region which is likely to provide an important link between location coding mechanisms and reward systems (Bender et al., 2015; Luo et al., 2011; Wirtshafter & Wilson, 2019, 2020, 579 2021). Most cells fired indiscriminately across the majority of the track, with only a small 580 proportion of cells (~5%) in both the CTL and MIA groups showing evidence of spatially 581 selective firing reminiscent of place cells. This is consistent with previous reports that 582 sampled cells from the rostral LS (Tingley & Buzsáki, 2018) and contrasts with prior studies 583 targeting the dorsal LS, where LS "place fields" (Wirtshafter & Wilson, 2020) (Takamura et 584 al., 2006) have been described. These data therefore provide corroborating evidence that the 585 rate coding of location varies across LS sub-regions.

586 Approximately a third of all LS cells displayed evidence of significant phase coding 587 of location across both groups. This is considerably less than the 89% of cells reported to 588 display phase coding in the Tingley and Buzsáki (2018) study. It should be noted, however, 589 that they used different criteria to determine whether cells showed evidence of phase coding, 590 and our methodology is likely more conservative. During phase coding in CTL cells the 591 phase of firing typically precessed across a single theta cycle as animals navigated the full 592 length of the track (Tingley & Buzsáki, 2018). In contrast, although the MIA manipulation 593 did not compromise the ability of LS cells to precess, there was a significantly steeper slope 594 of precession. In many cells, this resulted in phase precession that circulated through several 595 360-degree cycles as the animal traversed the track. Phase precession in excess of a single 596 theta cycle has not generally been observed in previous studies of the phenomenon (Dragoi & 597 Buzsáki, 2006; Ekstrom, Meltzer, McNaughton, & Barnes, 2001; Geisler et al., 2007; Huxter, 598 Burgess, & O'Keefe, 2003; Kamondi et al., 1998; Kjelstrup et al., 2008; Maurer et al., 2006; 599 O'Keefe & Recce, 1993; Royer, Sirota, Patel, & Buzsáki, 2010; Schmidt et al., 2009; Skaggs 600 et al., 1996; Terrazas et al., 2005; Tingley & Buzsáki, 2018). Instead, phase range has been 601 shown to dynamically shift according to either place field size or route familiarity across a 602 range of experimental paradigms so that that precession remains restricted to 360 degrees 603 across a place field (Ekstrom et al., 2001; Kjelstrup et al., 2008; Royer et al., 2010; Terrazas

et al., 2005). It is possible that the multiple phase precession cycles observed in the MIA
group represent more discrete "place fields" that are entirely decoupled from firing rate, with
discrete "fields" anchored to salient features of the animal's current environment or specific
task demands (Gupta et al., 2012; Maurer et al., 2006). In contrast, CTL animals may be more
likely to collate these discrete environmental "chunks" into a coherent whole as the animal
becomes more familiar with a learned navigational route (see figure 7).

610 When CTL phase trajectories were projected across the full length of the track, as a 611 population they tended to converge so that LS cells were firing at around the peak of theta 612 activity as recorded at the CA1 cell layer. This convergence was maximal at the location of 613 reward delivery. In contrast there was no evidence of phase trajectory convergence at the 614 reward region in MIA animals. The one previous paper that has described and examined 615 phase coding in the LS concluded that the LS phase code was likely agnostic to reward 616 location, however, this analysis was not a major focus of the paper, and some neurons were 617 clearly linked to local cues such as the goal (Tingley & Buzsáki, 2018). Our finding 618 contributes to a growing body of evidence indicating that the LS is involved in the integration 619 of spatial and reward information (Bender et al., 2015; Luo et al., 2011; Wirtshafter & 620 Wilson, 2019, 2020, 2021), and similar to phase precession in the ventral striatum, suggests 621 that firing phase in the LS may contain information about reward proximity or salience (van 622 der Meer & Redish, 2011). It should be noted, however, that we did not systematically 623 manipulate reward location in our study, to isolate it from other local cues. Further study 624 would be required to test this hypothesis. 625 If phase of firing signals reward location and distance to reward then we would 626 predict that in the MIA animals the association between reward and location is 'smeared'

- 627 across the environment such that a far broader range of stimuli and locations become
- 628 associated with the reward. Since the LS has direct connections to the VTA, this effect may

629 model, and potentially provide a mechanism for, some of the changes observed in

630 schizophrenia (Zhang et al., 2022). In particular altered motivational salience (Kapur, 2003),

631 which occurs with dysregulated dopamine signals and a tendency for individuals to mis-

632 assign salience to the elements of experience.

633 One of the most salient landmarks available to animals in this study was the presence 634 of corners and it is interesting to note that the limited phase trajectory convergence that 635 occurred in MIA animals may have been tied to the corners of the apparatus (see figure 5g). 636 Corners not only provide sensory information, but they require bidirectional modifications of 637 locomotor activity to navigate around them. Approximately one third of cells had significant 638 correlations of firing rate and speed, which is about half the figure reported by Wirtshafter 639 and Wilson (2019), but substantially more than that reported by Tingley and Buzsáki (2018). 640 The proportion of CTL group cells with firing rates that were negatively correlated with 641 speed was also just over half of all significantly correlated cells, in contrast to only around 642 one third in the Wirtshafter and Wilson (2019) study, suggesting that cells located in the 643 rostral LS may be particularly important for monitoring speed during tasks that involve 644 bidirectional speed fluctuations. Animals may have been able to move at a more constant 645 speed in the Tingley and Buzsáki (2018) due to the circular running apparatus. 646 A previous study from our lab showed that MIA disrupts theta sequences in the CA1 647 region of the hippocampus (Speers et al., 2021). According to predictions of the dynamic 648 weighting model proposed by Tingley and Buzsáki (2018), disrupted CA1 theta sequences 649 should prevent LS phase precession. However, in the current study, the proportion of cells

- 650 showing evidence of significant phase precession was not statistically different between
- 651 groups. Taken together, these data suggest that the MIA manipulation did not compromise
- 652 the ability of LS to precess relative to the CA1 theta oscillation, suggesting that upstream
- disruptions of phase coding in the hippocampus do not abolish phase coding in the LS. They

654 may, however, interfere with how spatial information is discretized relative to reward 655 locations. Additional studies will be required, however, to test whether the MIA-induced 656 changes in LS phase precession are a direct result of disrupted theta sequences in the 657 hippocampus, or are reflective of some other change. 658 It is unclear what mechanism(s) might account for the steeper phase precession 659 observed in MIA animals. Our results showed that hippocampal theta frequency was 660 significantly slower in MIA animals, while at the same time the intrinsic burst frequency of 661 cells did not differ between groups, consistent with a 'detuned oscillator' (Drieu & Zugaro, 662 2019) explanation of the increase in phase precession slope. The magnitude of change 663 predicted by this model, did not, however, fully explain our observations. Alternatively, 664 according to somato-dendritic interference models (Drieu & Zugaro, 2019), an increase in the 665 rate of excitatory ramping onto LS neurons might produce the same effect, although this 666 ramping would have to be cyclic across the apparatus to produce the effect observed and 667 would fail to produce precession of greater than 180 degrees on a single pass. 668 These results provide further evidence that phase coding may be disturbed following 669 MIA (Speers et al., 2021), and suggest a biophysical mechanism for impaired integration of 670 contextual and reward information, which may explain why MIA animals display memory 671 impairments when multi-sensory integration is required (Ballendine et al., 2015; Howland, 672 Cazakoff, & Zhang, 2012). Impaired spatial-reward integration could also have profound 673 downstream effects on motivation and dopamine signalling, both of which are known to be 674 impaired in schizophrenia (Davis, Kahn, Ko, & Davidson, 1991; Strauss et al., 2013). 675 Consistent with this idea, a recent study has demonstrated that dysregulation of the CA1-LS 676 pathway induces both dopaminergic hyperactivity in the VTA and novelty-induced hyper-677 locomotion that is schizophrenia-like, and that these could be attenuated via inhibition of the 678 LS (Zhang et al., 2022). The apparent smearing of reward representations across space, and

679	potentially t	time, by LS	cells is also	likely to com	promise the develo	opment of conditioned
0.1.2						

- 680 place preferences (Cazala et al., 1988; Jiang et al., 2018; Regier et al., 1990). It may also
- underlie MIA-induced changes in reward and temporal processing (Deane, Millar, Bilkey, &
- 682 Ward, 2017; Millar, Bilkey, & Ward, 2017). The abnormal discretization of spatial
- 683 representations could also contribute to the impaired attentional filtering and aberrant
- 684 salience that have been described in schizophrenia (Kapur, 2003; Luck, Leonard, Hahn, &
- 685 Gold, 2019), and also in autism spectrum disorder (Bodner, Cowan, & Christ, 2019), another
- 686 MIA-associated neurodevelopmental disorder. (Haddad et al., 2020).
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936 Figure 1. Methodological details and verification of electrode placement. (a) Diagram of the 937 rectangular track. Rats were pre-trained to run in a clockwise direction for a food reward delivered at 938 the centre of the bottom arm (marked with the red X). (b) Tetrode placement in the LS (highlighted in 939 vellow). The tetrodes bundles are depicted by the thick black line. Dashed purple lines indicate the 940 initial depth of electrode placement during surgery, while red lines show the range of target location 941 through the rostral LS over the course of the experiment. (c) Diagram of the hippocampus showing 942 the target area for LFP surgical implantation, and an example photograph of histology demonstrating 943 electrode placement in the pyramidal cell layer of CA1. (d) Cluster cutting examples for CTL cells 944 (shown in blue) and MIA (yellow). These examples show some of the different spike widths and 945 inter-spike interval histogram profiles observed in the LS. The top CTL example is similar to a 946 canonical place field, while the second CTL cell has a similar waveform, but the histogram profile 947 shows a more continuous spiking pattern across time. The first MIA example shows a cell with a 948 narrower waveform and a delayed spiking profile, whereas the final example shows a broader 949 waveform, similar to a hippocampal place cell, but the spiking profile is more reminiscent of an 950 interneuron. These spiking profiles were common across both groups, and were chosen to demonstrate 951 the range of different spiking profiles observed rather than systematic group differences. (e) 952 Schematic of final tetrode location in the LS for CTL animals at the termination of the experiment. 953 Modified stereotaxic image taken from Paxinos and Watson (2006). Percentages refer to the 954 proportion of phase coding cells calculated individually for each animal, with phase coding cells 955 defined as cells with a significant (<.05) circular-linear correlation of phase and position. Faded red 956 areas delineate regions where less than 20% of all recorded cells demonstrated evidence of significant 957 phase precession, and faded yellow delineates regions where proportions fell between 21 and 40%. 958 Example images of tetrode placement in the rostral LS is shown below. The second example shows 959 the final electrode placement of the excluded cell in the MS (marked with a cross). Additional 960 inspection of firing rates of the excluded cell compared to all other cells demonstrated that they were 961 atypical (below right). (f) As for e, but for MIA cells.

963	Figure 2. Basic firing properties of LS cells, mean locomotor speed, and comparison of
964	hippocampal theta properties. (a) Median firing rate of LS cells across the entire track. (b) Information
965	content of LS cells, measured in bits/spike. Bar denotes median values. (c) Mean locomotor speed as
966	animals traversed the rectangular track. Bars denote mean and SEM. (d) EEG amplitude of CA1 LFP
967	recordings. Error bars denote mean and SEM. (e) Examples of filtered and raw EEG recordings for
968	both groups. (f) Violin plots of theta frequency in Hz. (g) Average wave form shape of CA1 LFP
969	oscillations in the theta band. Lighter colour denotes standard error. (h) Average phase profile of the
970	theta waveform, from the Hilbert transform. (i) Comparison of median r values for the theta
971	frequency/ speed correlations generated for each individual recording.
972	

975	Figure 3. Comparison of LS phase precession between CTL and MIA animals. (a) Example
976	plots of phase precession in CTL cells. These examples were chosen to demonstrate a range of phase
977	precession (PP) variability, and each example cell is taken from a different animal. For each example
978	cell, the figure on the left displays PP as a function of colour around the track, with dark blue
979	representative of 0° and red representative of 360°. On the right are the corresponding PP plots after
980	linearization of the track for selected segments. Red lines denote the regression slope. In all plots,
981	phase is repeated across 2 cycles for clarity. In the CTL group, example cell 1 demonstrates both
982	spatially selective firing around half of the track in addition to robust PP. Example cell 2 demonstrates
983	robust phase precession across the majority of the track, with PP around 1 full cycle. Example cell 3
984	demonstrates robust PP through the reward area. Example cell 4 demonstrates a shallow PP slope and
985	sparse firing outside the reward area. (b) As for (a), but for example MIA cells. Example cell 1
986	demonstrates robust PP that approaches a full cycle. Example cell 2 demonstrates robust PP across the
987	entire track, in which PP appears to reset at the top left-hand corner. Example cell 3 demonstrates
988	robust PP through multiple cycles across the track. Example cell 4 demonstrates robust PP that
989	precesses through 2 distinct cycles.

991	Figure 4. LS cells in the MIA group are more likely to precess across multiple theta cycles.
992	(a) Proportion of LS cells with statistically significant (p $< .05$) phase precession for each group. (b)
993	Mean r values of the circular correlation of phase and position for both groups. Error bars denote
994	SEM. (c) p values of the circular linear correlation shown in (b). (d) Slope values (deg/mm) for all
995	cells (on right), and for a subset of cells demonstrating significant phase precession (on left). Black
996	lines denote median values. (e) Phase cycles across the full track. Error bars denote median with 95%
997	confidence intervals. (f) Mean slope values of the circular-linear correlation on an animal by animal
998	basis. Error bars denote SEM. (g) Log transformed firing rates of the subset of cells demonstrating
999	significant phase precession. Error bars denote mean and SEM. (h) Theta burst frequency of single
1000	units. (i) Circular histograms of intercept values for both groups, demonstrating greater variability of
1001	phase precession starting phase in the MIA group. Between group differences are based on the
1002	variance ratio F test. Red bars denote the mean angle with 95% confidence intervals.

1005	Figure 5. Relationship between firing phase and reward. (a) Correlations of LS electrode
1006	depth and lead/lag of LFPs recorded simultaneously from CA1 and the LS. Each animal is shown in a
1007	different color. Negative values indicate that the LS is leading CA1, and positive values indicate that
1008	CA1 is leading the LS. (b) Example of phase correction shift. (c) Phase trajectories of each precessing
1009	LS CTL group cell (as in figure 3a). Each trajectory is plotted onto one diagram where x-axis is
1010	linearized track. Vertical lines indicate corners and triangle marks reward location. Note how many
1011	tracks seem to pass through phase/location point at around 180° phase at the reward location. (d)
1012	Clustering of phase trajectories in the CTL group (from data in c), measured as mean vector length
1013	(MVL; y-axis), across the track (x-axis). Mean vector length can vary from $0 - no$ clustering to $1 - no$
1014	tight clustering or focus. Note that control trajectories show greatest clustering (phase focus) at the
1015	reward location (triangle). (e) Circular histogram of corrected phase angle (as shown in shown in c)
1016	when measured at the reward location. (f) Phase trajectory of each MIA group cell (as in figure 3b).
1017	Note that there is no clear clustering of phase trajectories around the reward location. (g) As in d, but
1018	for the MIA group. In contrast to the CTL group, MIA trajectories show much less clustering, with
1019	very little clustering at the reward location and less differentiation across the track. (h) As for e, but
1020	for MIA phase angles.
1021	

1023	Figure 6. Evidence of speed modulated cells in the rostral LS. (a) Example plots
1024	demonstrating robust correlations of firing rate and speed for both groups (first 3 examples in each
1025	row). The final example in each row shows firing rates that were correlated with acceleration rather
1026	than speed. These examples were chosen to demonstrate both positive and negative correlations that
1027	were typically observed across both groups. (b) Proportion of cells with significant correlations of
1028	firing rate and speed. Significance level for the linear correlation was set at 0.05. Sig. + refers to
1029	significant positive correlations and sig refers to negative correlations. (c) Violin plots of the
1030	absolute r value of the firing rate/speed correlation for the entire dataset (left column). Median r
1031	values for the linear correlation of firing rate and speed when only those cells with a significant firing
1032	rate/speed correlation were included are shown on the right. Error bars include 95% confidence
1033	intervals (d) As for (b), but including cells with significant phase precession to demonstrate
1034	overlapping cell properties. (e) As for c, but for acceleration (+) and deceleration (-). Error bars
1035	denote mean and SEM.

1039	Figure 7. Schematic of LS phase precession in CTL and MIA animals. In the CTL diagram,
1040	phase of LS cell firing (PP; coded by colour) occurs across a single cycle as the animal traverses a
1041	familiar route to a reward. In the upper rectangle, the range of PP from the starting location (S) is
1042	determined by a salient external cue marking the end of the route (white square) while other external
1043	cues (grey squares) and the reward itself (grey star) do not affect PP range. Alternatively in the lower
1044	example, PP range is anchored to the reward (white R) itself, while external cues are irrelevant. In
1045	both cases, intact theta sequences(A-D) arriving from CA1 (and potentially CA3) are likely to
1046	contribute to the emergence of single cycle PP in the LS and phase-coded information about the
1047	likelihood of rewards across the trajectory could be transmitted to subcortical regions such as the
1048	VTA. In the MIA example, disordered theta sequences from CA1 (and potentially CA3) may
1049	contribute to PP in the LS that exceeds a single theta cycle as the animal traverses a familiar route.
1050	The range of PP may be anchored to several salient cues across the trajectory (white squares),
1051	providing a more discretized representation of a navigational route in comparison to CTL animals.
1052	Alternatively, multiple cycle PP in MIA animals may reflect erroneous reward expectancies as the
1053	animal traverses the route. As a result, information about incorrect reward expectancies (red arrows)
1054	may be transmitted to the VTA, which could contribute to abnormal dopamine signalling in
1055	subcortical regions.
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4000

5000

5500

-60

electrode depth 4500

CTL lead/lag correlations (by animal)

60

phase shift (°)

0

LS <----> HPC

120

g







e

h











CTL phase angle at reward (corrected)



MIA phase angle at reward (corrected)







