1 Shared and unique properties of place cells in anterior cingulate cortex

2 and hippocampus

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34 Abstract

35 In the brain, spatial information is represented by neurons that fire when an animal is at specific 36 locations, including place cells in hippocampus and grid cells in entorhinal cortex. But how this information is processed in downstream brain regions still remains elusive. Using chronic Ca²⁺ 37 38 imaging, we examined the activity of neurons in anterior cingulate cortex (ACC), a brain region 39 implicated in memory consolidation, and found neurons that fire in a manner consistent with the 40 properties of place cells. While the ACC place cells showed stability, location and context specificity 41 similar to the hippocampal counterparts, they also have unique properties. Unlike hippocampal place 42 cells that immediately formed upon exposure to a novel environment, ACC place cells increased over 43 days. Also, ACC place cells tend to have additional place fields whereas typical hippocampal place 44 cells have only one. Hippocampal activity is required for the formation of ACC place cells, but once 45 they are established, hippocampal inactivation did not have any impact on ACC place cell firing. We 46 thus identified features of ACC place cells that carry spatial information in a unique fashion.

47

48 Introduction

49 Spatial navigation is an essential element of animal behavior that allows animals to forage, 50 return home and avoid dangers. The hippocampus plays a crucial role in this process. It bears neurons 51 called place cells that fire when an animal is located in a particular position of an environment but 52 not in others, providing an allocentric cognitive map of a space [1-4]. Multimodal inputs including 53 visual, tactile, olfaction, auditory, proprioception, and other sensory information are integrated to 54 form the place cells in hippocampus [1-4]. It was also shown that the spatial representation of 55 hippocampal place cell is initially dynamic but gradually stabilized [5-8]. The stabilization selectively 56 occurs at a location with motivational (reward) and environmental (landmark) salience and results in 57 an over-representation in these locations, indicating that the place cells' activity is not merely a 58 representation of the space but also its cognitive value. In this way, hippocampal place cells are well 59 characterized from years of studies [1-4]. However, how spatial information is further processed and 60 chronically represented in downstream brain regions has not been fully elucidated at cellular 61 resolution [5].

We decided to focus on anterior cingulate cortex (ACC), a part of medial prefrontal cortex (mPFC) implicated in memory consolidation [9-11]. Suppression of ACC impaired the recall of the remote but not of the recent memory[11]. This is in contrast to hippocampus, which is required for the recall of only recent but not remote memory [11, 12]. ACC receives a direct input from hippocampus [13] and also from striatum, amygdala and retrosplenial cortex, thereby serving as an 67 integration center of wide variety of sensory and motivational events [14, 15]. Recent work has 68 revealed neurons with spatially specific firing in mPFC including ACC. This spatial firing is regulated 69 by the environmental and task context, consistent with the spatial coding [16-19]. Lesion of the 70 hippocampus abolishes the place code activity in the mPFC indicating that mPFC is situated in 71 downstream of hippocampus for spatial coding [20]. However, how the spatial encoding of ACC 72 neurons develops has not been fully examined.

73 We, therefore, chronically monitored the activity of neurons in ACC while animals navigate 74 through a space. To this end, we used a head-mount miniaturized fluorescence microscope [5] to 75 image Ca²⁺-responses of excitatory neurons in ACC over days from freely moving mice. We found 76 neurons that fire in a manner consistent with the properties of place cells (ACC place cell). ACC 77 place cell showed similar properties with hippocampal place cell such as increase in stability, 78 reliability and context specificity after repeated exposure to the same environment over days. 79 Bayesian decoding verified that those cells indeed carry positional information. On the other hand, 80 we found several properties unique to ACC place cells. The fraction of ACC place cells increased as 81 the task was repeated. ACC neurons tend to have multiple place fields, which might represent 82 association of more than one location in a context. Finally, the formation of the ACC place cells 83 requires hippocampal activity but once established, the cells become independent of hippocampus. 84 Thus, while ACC place cells share properties with hippocampal place cells, they have unique features 85 and mechanism of generation.

86

87 **RESULTS**

88 ACC has place cells similarly to hippocampus

For Ca^{2+} imaging, we used TRE-G-CaMP7-T2A-DsRed2 × CaMKII α -tTA double transgenic 89 90 mice, that coexpress G-CaMP7 and DsRed2 in excitatory neurons [8]. To image ACC, a gradient 91 reflective index (GRIN) lens with a microprism attached at its end was implanted between the two 92 cortical hemispheres so that the center of the tip of the microprism is at 0.5 mm anterior from bregma 93 and 1.4 mm from the cortical surface and the prism faces the left ACC [21] (Fig. 1a and 94 Supplementary Fig. 1a). The dorsal hippocampus was imaged by implanting a GRIN lens above the 95 right dorsal hippocampus after removing a part of overlaying cortex as previously reported [5] (Fig. 1b and Supplementary Fig. 1b). The activity of ACC layer 2/3 neurons or hippocampal CA1 96 97 pyramidal neurons was observed by using a head-mounted fluorescent microscope across multiple 98 days from awake and behaving animals [5] (Fig. 1c).

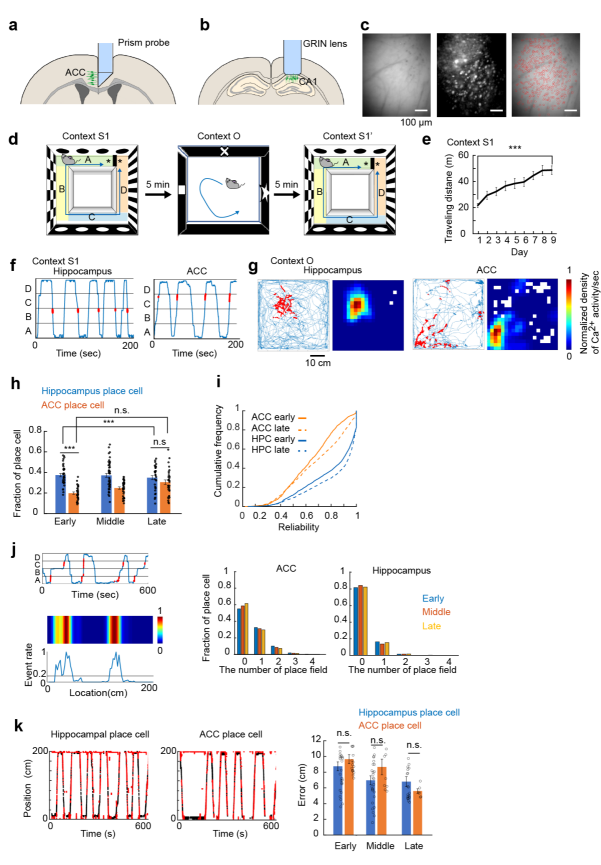
99 The animals were placed first in a square track (one edge 50 cm, context S1) with a wall 100 installed at one of the corners, then in an open arena (50 x 50 cm, context O) with different wall

patterns and scent, and finally in the original square track (context S1') for 10 min each with 5 min intermissions (Fig. 1d). In context S1, the animals ran back-and-forth between the two ends for food rewards, given when the animal reached one end. The animals moved faster and finished more laps over days (Fig. 1e), indicating that the animal became familiar with the context[11]. In the context O, the rewards were randomly thrown into the open arena to motivate animals to navigate around [22].

106 In both hippocampus and ACC, we found neurons that fire in a manner consistent with the 107 properties known for place cells, such as location specificity, reproducibility, and direction selectivity 108 across laps (Fig. 1f and g). The proportion of hippocampal place cells did not change over days, 109 consistent with previous results [5]. The proportion of ACC place cells in early phase (days 1-3) was 110 smaller than in hippocampus (Fig. 1h) but increased as the task was repeated and reached the level 111 comparable to hippocampus in the late phase (days 7-9). We also calculated the reliability index of 112 each cell, an index of how reproducibly a cell fires in a specific location across multiple laps in a 113 single session (Supplementary Fig. 2). In early sessions (days 1-3), the index was lower in ACC than 114 in hippocampus, but gradually increased in the late sessions (days 7-9) though it never reached the 115 level of hippocampus (Fig. 1i). This indicates that the firing reproducibility of ACC place cells is not 116 as high as that of hippocampus place cells. We noticed that nearly a half of ACC neurons had extra 117 place fields (Fig. 1j) whereas hippocampal neurons typically had only one place field. The number of 118 extra place fields remained constant in both ACC and hippocampus during repeated training, 119 indicating that this is an intrinsic difference between these two brain regions.

In order to test if these cells indeed encode spatial information, we used Bayesian decoding (Fig. 1k) by selecting the same number of the cells with the highest activity level from both ACC or hippocampus. The decoder often registered one end of the square track as the other end. This may reflect the fact that the ends are adjacent to each other, separated just by a wall. Nevertheless, the analysis confirmed that the ACC neurons carry positional information comparable to those in hippocampus (hippocampus vs ACC; p = 0.46 (early), p = 0.201 (middle), p = 0.79 (late), One way ANOVA. Fig.1k). Therefore, we called these cells ACC place cells.







129 Figure 1. The basic properties of ACC and hippocampal place cells.

- 130 (a) Imaging of the ACC using a right angle microprism inserted into the fissure.
- 131 (b) Imaging of hippocampus by using a GRIN lens implanted above hippocampal CA1 layer.
- 132 (c) Example images of ACC neurons. Left, max intensity image. Blood vessels appear as shadows.
- 133 Middle, relative fluorescent change ($\Delta F/F$). Right, active cells (red circles) overlaid with the max 134 intensity image.
- 135 (d) Behavior paradigms. The square linear track (context S1 or S1') and the open arena (context O).
- 136 In one session, mice visited two distinct environments (S1 \rightarrow O \rightarrow S1') each for 10 min. Location of
- reward in S1 was indicated by asterisks. In context O, the reward was randomly thrown into the
- arena. One set of experiment was conducted per day for 9 days while monitoring neuronal activityin hippocampus or ACC.
- 140 (e) Behavioral changes induced by repeated training with context S1. Total traveled distance is
- 141 shown (n = 9 mice). Day 1 vs day 9, p = 1.88×10^{-8} , one-way AVOVA. Graphs show means \pm
- 142 SEM.
- (f) Example of place cells in hippocampus in ACC in context S1. Blue lines show the trajectory ofthe mouse and red dots mark calcium events.
- 145 (g) Example of place cells in hippocampus and in ACC in context O. Left, blue lines show the
- mouse's trajectory and red dots mark calcium events. Right, Gaussian-smoothed density maps ofcalcium events, normalized by the mouse's occupancy time per unit area and the cell's maximum
- 148 response.
- 149 (h) The fraction of ACC and hippocampal place cells relative to the number of total identified cells
- 150 in context S1. Hippocampus early vs late p = 0.69, ACC early vs late $p = 2.9 \times 10^{-4}$, hippocampus
- 151 early vs ACC early $p = 5.5 \times 10^{-10}$, hippocampus late vs ACC late p = 0.079; one-way ANOVA. N
- $152 = 36 \text{ data (18 sessions x 2 running directions) in early, 46 \text{ data (23 sessions x 2 running directions)}$
- 153 in middle, 34 data (17 sessions x 2 running directions) in late from 5 mice for hippocampus. N = 32
- 154 data (16 sessions x 2 directions) in early, 38 data (19 sessions x 2 directions) in middle, 32 data (16
- sessions x 2 directions) in late from 4 mice for ACC.
- (i) Reliability of firing of ACC and hippocampus ('HPC') place cells in the early (days 1-3) and thelate sessions (days 7-9). Reliability represents how reproducible is the during multiple laps in the
- 158 same session (Fig. S2). Data were pooled from 3265 cells in the early and 3527 cells in the late
- sessions from 5 mice for hippocampus and 1206 cells in the early and 2494 cells in the late sessions
- from 4 mice for ACC. Hippocampus early vs late $p = 1.61 \times 10^{-12}$; ACC early vs late $p = 7.23 \times 10^{-8}$
- and ACC early vs hippocampus early $p = 1.403 \times 10^{-114}$ and ACC late vs hippocampus late p = 3.68
- 162 x 10^{-171} , one-way ANOVA.
- 163 (j) Example of an ACC place cell with extra place fields in context S1. Left Top, mouse trajectory
- 164 (blue) and calcium events (red). Left middle, linearized heat map of the event rate normalized by
- 165 maximum activity. Left bottom, Normalized event rate in each spatial bin. Vertical line shows the
- 166 threshold for criteria of place field. This cell has 2 place fields. Right, number of place field of
- 167 hippocampus and ACC place cells in the early (days 1-3), middle (days 4-6) and late (days 7-9)

hippocampus group mice. N = 2453 place cells in early, 2812 place cells in middle, 2972 place cells
from 4 ACC group mice.

171 (k) Bayesian decoding of the mouse trajectory (red dots) and actual position (black curves) from 172 hippocampus and ACC place cells. Average median errors of the early (days 1-3), middle (days 4-6) 173 and late (days 7-9) sessions. Not significant by Wilcoxon rank sum test. P = 0.46 (early), p = 0.201174 (middle), p = 0.79 (late). N = 28 (hippocampus) and 16 (ACC) sessions in early sessions; 27 175 (hippocampus) and 9 (ACC) sessions in the middle sessions, 21 (hippocampus) and 6 (ACC) sessions 176 in the late sessions.

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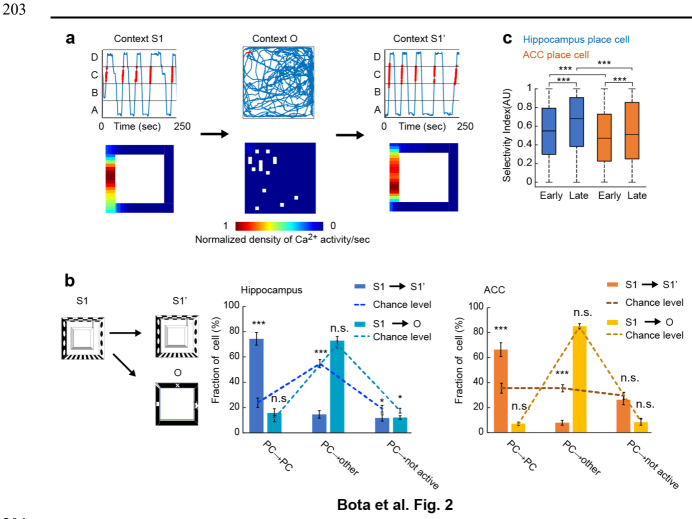
179 Stability of hippocampus and ACC place cells between sessions

We then compared the activity of each cell between two sessions in identical or distinct 180 181 contexts to see whether they switched their encoding mode between sessions (Fig. 2a). When the S1 and S1' sessions interleaved by a session O were compared, $66.1 \pm 5.5\%$ of ACC place cells in context 182 S1 still behaved as place cells in the context S1' (Fig. 2b). In contrast, only $7.0 \pm 1.4\%$ of ACC place 183 cell in context S1 behaved as a place cell in context O. The rest of the S1 place cells were either 184 185 unclassified (84.7 \pm 2.1%) or not active (8.3 \pm 2.4%). This proportion of encoding mode was 186 comparable to the proportion of all cell types in context O (place cells, 6.1 ± 1.1 %; unclassified, 84.8 187 ± 2.2 %; not active 9.0 ± 2.4 %. p = 0.65, 0.96, 0.85, N = 4 mice; one-way ANOVA), indicating 188 random reassignment of the place cells when mice moved to the other context. Likewise, place cells 189 in context O converted their encoding in context S1', in a proportion not different from their 190 proportion among all cells (Supplementary Fig. 3. Place cells, $30.6 \pm 7.0\%$; unclassified cells $25.5 \pm$ 191 5.4%, not active 44.0 \pm 8.1% versus all cells in context S1': place cells, 33.2 \pm 4.6%, unclassified 192 cells, $29.0 \pm 3.6\%$; not active, $37.8 \pm 3.0\%$; p = 0.92, 0.29, 0.50; N = 4 mice; one-way ANOVA). The 193 same tendency was observed in hippocampal place cells. Therefore, this analysis indicates that the 194 place cells in one context are randomly assigned either as place or non-place cells in different contexts 195 and suggests that there is no particular subclass of neurons which preferentially become place cells 196 in different contexts in both brain regions.

Finally, in order to quantitatively assess the change in the activity level of each cell, we defined the selectivity index by calculating the absolute of the difference between firing rate in two contexts normalized by the sum of firing rate in both contexts (Fig. 2c). This analysis revealed that both hippocampal and ACC place cells showed a significant increase in the selectivity over days,

201 indicating that there was a significant differentiation in neuronal activity to one of the contexts during

202 repeated exposure to two different contexts (Fig. 2c).



204

205 Figure 2. Context specificity of ACC and hippocampal place cells.

206 (a) Representative activity of an ACC place cell during one session (context $S1 \rightarrow$ context

207 O→context S1'). Top, heat map for event rate of the cell. Bottom, mouse's trajectory (blue) and

calcium events (red). Firing intensity is normalized by the maximum activity of the cell throughoutthe session.

- 210 (b) Comparison of place cell activity between two identical or different contexts. Encoding mode of
- 211 ACC place cells in context S1 was examined in context S1' or context O. ACC place cells in
- 212 context S1 were classified in 3 groups "place cells (PC)", "other" and "not active" according to its
- 213 activity pattern in context S1' or context O. Figure shows the average fraction of cells in each
- group. Dotted lines show proportion of each class of cells in S1 or O, that serve as chance level.
- 215 Compared with the chance level, hippocampus S1 to S1': place cells to place cells $p = 1.8 \times 10^{-8}$,
- 216 place cells to other cell type $p = 1.2 \times 10^{-9}$, place cells to not active cells p = 0.031. Hippocampus S1
- 217 vs O: in the same order, p = 0.42, 0.58, 0.018. ACC S1 to S1': $p = 0.00027, 1.3 \times 10^{-7}, 0.52$. ACC
- 218 S1 to O: p = 0.65, 0.96, 0.85. One-way ANOVA. N = 4 mice for ACC, n = 5 mice for hippocampus.

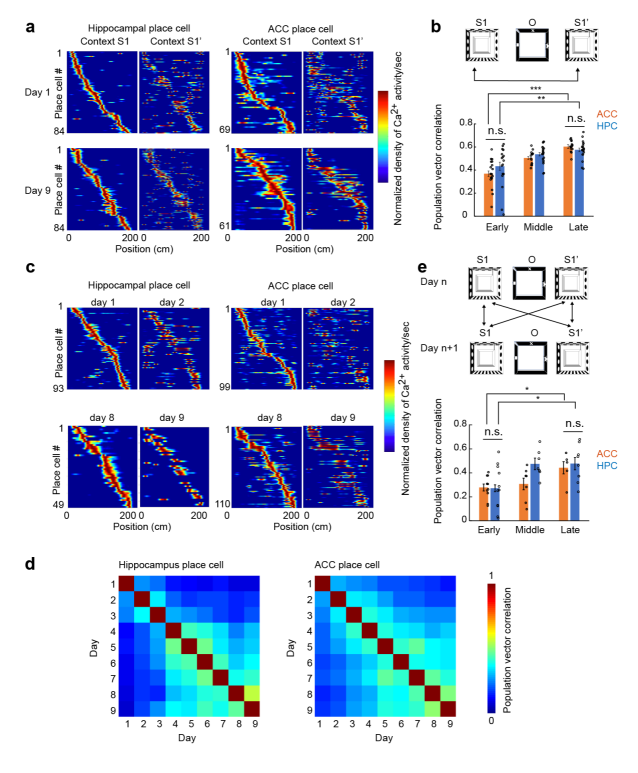
(c) Selectivity index between context S1 versus context O. The index was pooled from 2045
hippocampal place cells (from 5 mice) and 931 ACC place cells (from 4 mice) in early phase (day 13), 1655 hippocampal place cells (from 5 mice) and 1121 ACC place cells (from 4 mice) in late phase
(day 7-9). Whiskers show maximum and minimum value in each data set.

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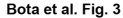
225 Long-term stability of place cells

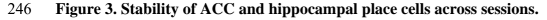
226 We next examined whether these cells stably retain the same positional information on the 227 track across sessions on the same day or different days. We first compared the place field in context 228 S1 and S1' interleaved by a session in context O on the same day by calculating the population vector 229 correlation. In hippocampus the correlation increased during the 9-day sessions (days 1-3 vs 7-9, p = 230 2.5×10^{-3} , one-way ANOVA). This is consistent with the previous results which showed hippocampal 231 place cells become stable after repeating the same task [8, 23]. Also in ACC, we saw an increase of 232 correlation between S1-S1' sessions over days (days 1-3 vs 7-9, $p = 1.6 \times 10^{-7}$, one-way ANOVA). 233 We obtained essentially consistent result by calculating the ratio of cells with stable place fields (less 234 than 12.5 cm shift between S1-S1' sessions) among place cells in both hippocampus ($44.05 \pm 6.1\%$ 235 of cells exhibited stable place field in days 1-3 and $64.7 \pm 5.5\%$ in days 7-9, p = 0.0015, one-way 236 ANOVA. Supplementary Fig.4) and ACC ($28.7 \pm 3.8\%$ of cells exhibited stable place field in days 237 1-3 and 58.1 \pm 3.6% in days 7-9, p = 9.6 x 10⁻⁶, one-way ANOVA).

We then attempted to obtain a more holistic view of stability over days by comparing S1 sessions across days (Fig. 3c-e). We also found that both the hippocampal and ACC place cells showed a gradual stabilization during the sessions repeated over 9 days. These results showed that the ACC place cells have similar properties to the hippocampal place cells with respect to the stability (Hippocampus early vs late p = 0.022; ACC early vs late p = 0.014; ACC early vs hippocampus early p = 0.93; ACC late vs hippocampus late p = 0.69, one-way ANOVA).









- (a) A comparison of place cell map between context S1 and S1' within the same session on day 1
- and day 9. Place-field maps was ordered according to the place field centroid position in context S1.
- 249 The maps were normalized by the maximum activity of each cell.

- (b) Population vector correlation of place cell representation in early, middle, and late sessions.
- Hippocampus early vs late $p = 2.5 \times 10^{-3}$; ACC early vs late $p = 1.6 \times 10^{-7}$; ACC early vs
- hippocampus early p = 0.21; ACC late vs hippocampus late p = 0.24, one-way ANOVA. N = 18
- session pairs in early, 20 session pairs in middle, 20 session pairs in late for hippocampus. N = 18
- session pairs in early, 14 session pairs in middle, 14 session pairs in late for ACC.
- 255 (c) A comparison of place cell map between context S1 and S1' in adjacent sessions.
- (d) Heat map for population vector correlation of place cells in hippocampus and ACC between allpairs of sessions.
- (e) Population vector correlation across early (day 1 vs day2 to day 3 vs day4), and middle (day 4 vs day5 and day 5 vs day 6) and late (day 6 vs day 7 to day 8 vs day 9) sessions for both hippocampus and ACC. Hippocampus early vs late p = 0.022; ACC early vs late p = 0.014; ACC early vs hippocampus early p = 0.93; ACC late vs hippocampus late p = 0.69, one-way ANOVA. N = 11 session pairs in early, 7 session pairs in middle, 8 session pairs in late for hippocampus. N = 10 session pairs in early, 7 session pairs in middle, 5 session pairs in late for ACC.
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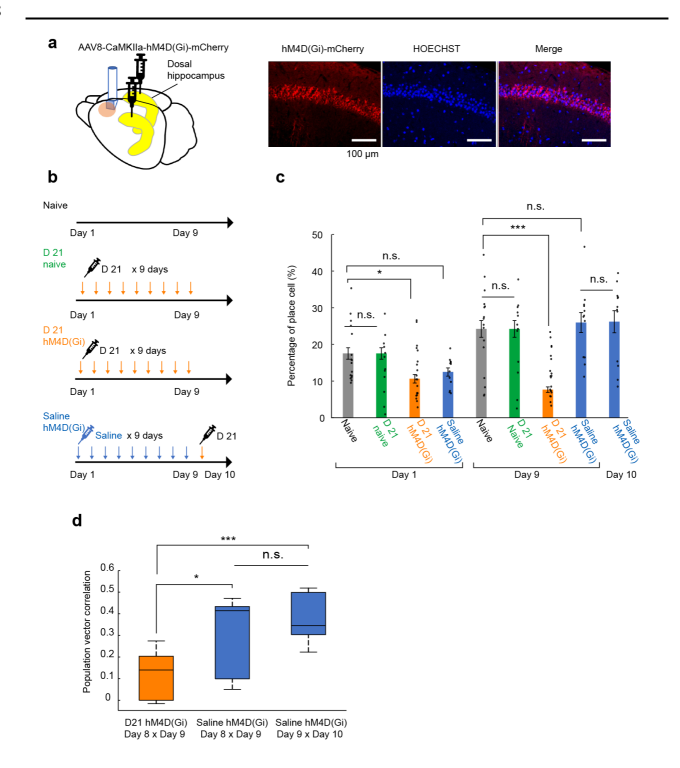
266 ACC place cell activity requires hippocampus for formation but becomes independent after training

Finally, we examined whether the formation of ACC place cells requires hippocampal activity. To this end, we inhibited the hippocampal neuronal activity by administering a Designer Receptor Exclusively Activated by Designer Drugs (DREADD) agonist 21 (D21) to a mouse expressing its cognate inhibitory receptor, hM4Di-mCherry, in dorsal hippocampus (Fig. 4a) [24]. The mice received D21 20 min before each behavior experiments for 9 days. Mice in saline hM4D(Gi) group received saline for 9 days and then received D21 on day 10 (Fig. 4b).

273 In the naïve, D21 naive or saline hM4D(Gi) groups, there was an increase in fraction of ACC 274 place cells during 9-day period (Fig. 4c). However, in the D21 hM4D(Gi) group, the fraction 275 remained low compared with control animals (on day 9. Naive vs. D21 naive, p = 0.83; Naive vs. 276 D21 hM4D(Gi), $p = 1.9 \times 10^{-4}$; Naïve vs saline hM4D(Gi), p = 0.38. One way ANOVA. Fig. 4c). The 277 stability of the map as assessed by the population vector correlation analysis between days 8 and 9 278 was also low stability in D21 hM4D(Gi) group compared saline hM4D(Gi) group (Fig. 4d. D21 279 hM4D(Gi) day 8 and 9 vs Saline hM4D(Gi) day 8 and 9, p = 0.023; D21 hM4D(Gi) day 8 and 9 vs 280 Saline hM4D(Gi) day 9 and 10, $p = 4.7 \times 10^{-6}$. One-way ANOVA). This indicates that the hippocampal 281 activity is required for establishing and maintaining ACC place cell maps.

In contrast, when D21 was administered on day 10 in the saline hM4D(Gi) group after the place cell map was already established, it did not have effect on the proportion of ACC place cells

(Fig. 4c. p = 0.96, one-way ANOVA). Also, the population vector correlation between days 9 and 10 in the presence of D21 was not significantly different with days 8 and 9 when the same animals received saline (Fig. 4d. p = 0.53, one-way ANOVA). This indicates that ACC place cell map, once established, no longer requires hippocampal neuronal activity for maintenance.



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Figure 4. The effect of chemogenetic inhibition of hippocampal excitatory neurons on ACCplace cell.

292 (a) Representative mCherry image in TRE-GaMP7 x CaMKIIα-tTA double transgenic mice

- 293 infected with AAV8-CaMKIIα-hM4D(Gi)-mCherry virus in hippocampi bilaterally. Scale bar =
- 294 100 μm.
- 295 (b) Experimental schedule.

296 (c) Percentage of ACC place cell in each context S1 and context S1'. In the bar graphs, circles 297 represent individual trials from S1 or S1'. (Naive n =16 data (8 sessions x 2 direction) from 4 mice, 298 D21 naive n =12 data (6 sessions x 2 direction) from 3 mice, D21 hM4D (Gi) n = 20 data (10 299 sessions x 2 direction) from 5 mice, Saline hM4D(Gi) n =12 data (6 sessions x 2 direction) from 3 300 mice). Day 1; naïve vs D21 naïve p = 0.23, naïve vs D21 hM4D (Gi) p = 0.033, naïve vs saline 301 hM4D Gi) p = 0.054. Day 9; naïve vs D21 naïve p = 0.83, naïve vs D21 hM4D (Gi) $p = 1.9 \times 10^{-4}$, 302 naïve vs saline hM4D (Gi) p = 0.38, saline hM4D (Gi) day 9 vs day 10 p = 0.96, one-way ANOVA. 303 (d) Stability of place cell representation calculated by population vector correlation between day 8 304 and day 9, day 9 and day 10 in D21 hM4D (Gi) or saline hM4D (Gi) animals. P = 0.023 D21 hM4D(Gi) day 8 and 9 vs Saline hM4D (Gi) day 8 and 9, $p = 4.7 \times 10^{-6} D21 hM4D$ (Gi) day 8 and 9 vs Saline 305 hM4D(Gi) day 9 and 10, p = 0.53 Saline hM4D(Gi) day 8 and 9 vs Saline hM4D(Gi) day 9 and 10, 306 307 one-way ANOVA. N = 10 session pairs for D21 hM4D (Gi) day 8 and day9, 6 session pairs for saline 308 hM4D (Gi) day 8 and day 9, 6 session pairs for saline hM4D (Gi) day 9 and day 10.

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311 Discussion

312 mPFC, in particular ACC, has been implicated in the process of memory consolidation [9, 10]. 313 During this process, coherent theta oscillations coupling between the hippocampus and mPFC 314 facilitates the transfer of memory [25]. However, the exact content of memory carried by the neurons 315 in ACC and whether it changes their properties during consolidation, have not been fully elucidated. In this study, we examined how representation of positional information is processed in the 316 317 downstream of hippocampus. To this end, we recorded calcium activity over days from the ACC of freely moving mice. We found a population of neurons in ACC which shows location specific activity. 318 319 It shares basic properties with hippocampal place cells such as location specific firing and directional 320 selectivity [1-4]. However, unlike hippocampus place cells, the ACC place cells often have extra 321 firing field. This may represent association of information from more than one location on the track. Indeed, mPFC including ACC is involved in the processing contextual information and such 322 323 association of multiple locations is likely to play a role.

324 In addition, we found that ACC place cells gradually increased over days in a manner 325 requiring hippocampal activity for the formation. However, once the ACC place map is formed after 326 9 days, ACC no longer required hippocampal activity to fire and to exhibit a place cell map. This 327 property is reminiscent of systems consolidation process of episodic memory, where hippocampus is 328 required for initial formation of memory but not for recall of remote memory [11, 12, 26]. The 329 memory consolidation theory predicts hippocampal activity after the memory events induces cortical 330 plasticity and consolidates memory in cortical circuit[11]. After memory consolidation, hippocampus 331 is no longer required for recall of remote memory[11]. The same analogy can be applied to the place 332 cells in both hippocampus and ACC. At this point, it is not clear from which brain regions the spatial 333 information is generated under hippocampal inactivation. ACC is bidirectionally connected to other 334 cortical regions such as retrosplenial and entorhinal cortexes, both of which are implicated in 335 processing spatial information [14, 27, 28]. It is therefore possible that spatial information arriving 336 from these cortexes as well other regions can bypass the requirement of hippocampus.

In conclusion, we found that ACC place cells have both shared and unique properties with hippocampal counterparts. Further study is required to elucidate how their formation depends on the hippocampus, but they are maintained without hippocampal activity once they are formed. It will be intriguing to test if these cells indeed overlap with engram cells as defined by *c-fos* expression [6].

341

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- 423

424 Materials and methods

425 Subjects

- 426 All experiments and procedures were approved by the RIKEN and Kyoto University Animal
- 427 Experiments Committees, and conducted according to institutional guidelines. Experiments were
- 428 conducted on 10–24 weeks old TRE-G-CaMP7 x CaMKIIα-tTA double transgenic mice [8]. Mice
- 429 were housed singly in a cage with 12 h-12h light-dark cycle (dark: 6 am-6 pm, light: 6 pm-6 am on
- 430 the next day). All experiments were performed between 6 am and 6 pm.

431 Histology

- 432 Mice were perfused transcardially with phosphate-buffered saline (PBS) followed by 4%
- 433 paraformaldehyde (PFA) in PBS. Brains were extracted and put in 4% PFA for additional fixing.
- 434 After 24h, PFA were transferred to PBS for additional days. Brains were then sliced in 50 µm
- 435 sections using a microslicer (Dosaka). Brain sections were incubated at 4 °C in 0.1 M Tris-HCl,
- 436 0.15 M NaCl, 0.5 % Triton-X and 5 % blocking reagent (Roche) and rabbit anti-GFP antibody
- 437 (A11122, Thermo Fisher Scientific, 1:500) overnight for immunostaining. Brain sections were then
- 438 washed with PBS 3 times for 15 min each and incubated with AlexaFluor 488 (A11008, Thermo
- 439 Fisher Scientific, 1:500) conjugated secondary antibodies. Brain sections were washed again 3
- times for 15 min, mounted, and coverslipped with mounting medium with Hoechst 33258
- 441 (#382061, Calbiochem). Fluorescence images were taken by confocal microscopy (Olympus
- 442 FLUOVIEW FV1500) [29].
- 443

444 Surgery for calcium imaging from Hippocampal CA1

We used isoflurane to anesthetize mice (5% induction, 1.5% during surgery) and mice were fixed in 445 446 a stereotaxic frame. Mice first were implanted a stainless-steel head plate (25 mm length, 4 mm width, 1 mm thickness) with a circular opening (7 mm inner diameter and 10 mm outer diameter, 447 the center is 2.5 mm off relative to the middle of the long side of the plate)[8] to skull with dental 448 449 cement. The skull (excluding the area inside the circle of the head plate) was covered with dental 450 cement including the three anchor screws. Several days later, we removed a circular part of skull 451 (centered 2.0 mm posterior, 2.0 mm lateral from bregma) with a trephine drill and removed the dura 452 and cortex above the CA1 by suction with a 25 or 27 gauge needle washing with sterile cortex buffer (123 mM NaCl, 5 mM KCl, 10 mM glucose, 2 mM CaCl₂, 2 mM MgCl₂, 10 mM HEPES, 453 pH 7.4). We then implanted a metal guide tube with glass window just dorsal to CA1 region and 454

- 455 sealed the space between the skull and guide tube using dental cement[23].
- 456

457 Surgery for calcium imaging from ACC

458 Mice were implanted a stainless-steel head plate to the skull at 1 mm anterior to the bregma 459 centered at midline. Several days later, a part of skull (centered 2.0 mm posterior, 2.0 mm lateral 460 from bregma) and dura was removed to access to the longitudinal fissure. The lens (Inscopix) is 461 comprised of a right angle microprism (1 mm x 1 mm x 1 mm) and GRIN lens (0.85-mm diameter, 462 3.3-mm length). This lens was affixed to stereotaxic frame for implantation. The tip of the prism 463 was positioned at the entrance (midline at the target 1.0-mm anterior to bregma) avoiding the sinus 464 laterally, then we vertically lowed the prism into the longitudinal fissure so that its front faces 465 against the medial surface of the left ACC. The exposed area of skull was covered with dental 466 cement. We then sealed the space between the dental cement and lens using adhesive bond. The exposed lens was sealed with Silicone adhesive (Kwik-Sil, World Precision Instruments) and dental 467 468 cement [21].

469

470 Calcium imaging

For hippocampal imaging, four weeks after the surgery described above, a gradient refractive index 471 472 lens (GRINtech GmbH, 0.44 pitch length, 0.47 NA) was fixed in the guide tube using ultravioletcuring adhesive (Norland, NOA 81). For ACC imaging, the silicone adhesive was removed to 473 474 expose prism lens. The integrated microscope (nVistaHD, Inscopix) with stereotaxic frame was 475 lowed toward the GRIN lens to find G-CaMP7 fluorescence using LED light source (0.12–0.24 476 mW). We then attached the microscope's base plate to the skull using dental cement on suitable 477 imaging focus. The dental cement darkened with black acrylic paint. The base plate was left on the 478 mouse even after the microscope was detached to ensure reproducibility of imaging site. Before 479 each behavior experiment, mice were anesthetized with isoflurane and the microscope was attached 480 to the base plate. The imaging session was started at least after 15 min recovery in the home cage.

In each session of the behavioral experiment, the mice were placed first in a square track (contextS1), then an open arena (context O) with different wall patterns, and then again in the square track

483 (context S1') for 10 min each with 5 min interval. Context S1 consists of a square track of four divisions (each 50 cm long x 5 cm wide) and side walls (8 cm tall). A wall was placed at one of the 484 485 corners to separate the ends. Mice run back-and-forth between the two ends for a sucrose tablet 486 (#1811251 Sucrose Rewards Tablet, Test Diet), given when the animal reached one end. In the 487 context O (50 x 50 cm with wall of 25 cm tall), the rewards were randomly thrown into the open 488 arena. The equipment was wiped by paper towels with different odor (80% ethanol for context S 489 and 0.5% acetic acid for context O) before each behavioral experiment. Before beginning the 490 imaging, they were exposed to context O and context S as pre-training for 3 days. On day 1 of pretraining, mice were let freely move for 10 min in each context without any food reward. After day 1 491 492 pre-training, foods in their home cage are removed to restrict. In day 2 and 3 of pre-training, food 493 reward was given as described above. After each day 2 day3 of pre-training and each daily imaging, 494 mice got 1-3 g food to keep their weight. One S1-O-S1' session per day was performed during 495 calcium imaging and repeated for 9 days. We recorded a total 5 mice from hippocampus and a total 496 4 mice from ACC [5, 23].

497 **DREADD**

498 DREADD agonist 21 (D21; Cayman Chemical Company, #18907), an alternative to CNO is used
499 for DREADD experiments. A stock solution of 5 mg/ml in DMSO was made and then diluted in
500 saline to desired concentration (0.05 mg or 0.1 mg/ml). D 21 was injected intraperitoneally at 1

501 mg/kg 20 min before the behavioral experiment.[30]

502

503 Processing of Calcium imaging data

504 We processed imaging data using Mosaic (Inscopix). First, the original imaging data were down-505 sampled by a factor of four in each dimension to increase processing speed then the down-sampled 506 images were motion corrected. We then created normalized movie by the average (F0) to generate 507 changes over baseline in fluorescence. $\Delta F(t)/F0 = (F(t)-F0)/F0$, where F0 is the value for each pixel 508 averaged over the entire time span of the calcium imaging movie. Finally, the movie was smoothed 509 by applying disk average filter (disk radius: 3x3 pixels). We separately used un-filtered movie for calculation of each neuron's fluorescence intensity, and filtered movie for neuron identification and 510 511 calculating clustering score, respectively, according to the instruction 'Neuron Identification' and 512 'Calcium Event Identification' in the manufacture's users manual.

513 Neuron Identification

- 514 Regions-of-interest (ROIs) was identified using a custom MATLAB routine. First, intensity of each
- 515 pixel is normalized by the mean value of all pixel in that frame of the filtered calcium imaging
- 516 movie. Then spots of fluorescence signal over threshold were detected in each image and
- 517 designated as blobs. The mean intensity of each blob, size and shape were measured by using
- 518 regionprops function of MATLAB. Blobs with suitable size (min = $250 \ \mu m^2$ (30 pixels), max =
- 519 1500 μ m² (180 pixels)) and ratio of long to short axes (max = 2) were employed. Signal traces of
- 520 each employed blob were calculated over all frames of the movie. Finally, spatial correlation among
- all blobs and correlation of signal trace (mean gray value of pixels in the blob) among all blobs

- 522 were calculated respectively. Blob pairs with high spatial correlation (r > 0.4) and high signal
- 523 correlation (r > 0.9) are considered as the same neuron, and smaller one of the two was used as 524 ROI(neuron) for further analysis [29].
- 525

526 Calcium Event Identification

527 We extracted $\Delta F/F$ traces of each ROIs from un-filtered calcium imaging movies. To remove

- 528 crosstalk from neighboring ROIs, clustering score of each ROI is calculated as follows. After ROIs
- 529 (Fig. S5, red) were identified using a custom MATLAB routine, 5-time enlarged ROIs (by area,
- 530 yellow) were made. For a given frame of Ca^{2+} images, the location of pixels with top 20%
- 531 brightness were detected. The proportion of the pixels within the original ROI is defined as the
- 532 clustering score for each frame. Low clustering score indicates high likeliness of crosstalk of
- 533 neighboring ROI. By excluding value with low clustering score from each ROI, crosstalk was
- 534 removed. Second and third steps are performed in each frame of movie. Time points with $\Delta F/F$
- 535 signal >1.5 and clustering score >0.4 were detected as calcium event for each ROI.
- 536

537 Neuron Registration

538 There are two steps in Neuron Registration: session registration and neuron registration.

539 Session registration

- 540 We determined how much imaging field shifted between sessions for each imaging data. We first
- 541 created a median projection of all of imaging frames of each session. All image of each session
- aligned to first session using the 'motion correction' function from the Mosaic and calculated
- 543 transformation object was saved. This process of alignment was applied to each calcium imaging
- 544 movie of all sessions, and saved for further analysis. [29]

545 Neuron Registration

- 546 Neuron Identification step was performed in aligned calcium imaging movie. Each detected neuron
- 547 in a session was mapped to calculate distance of center-of-mass and spatial correlation between
- 548 neurons in other sessions. We designated the neurons with closest center-of-mass (4 pixels (10 μm))
- and high spatial correlation (r > 0.6) as same neuron over sessions. [29]
- 550

551 Place cells and place field

552 We used the following criteria to identify place cells and place fields in the open field and the 553 square track, respectively.

554 Place cells in open field

To identify place cells in the open field, firing rate maps were obtained as follows. First, number of calcium events in each neuron were sorted to 16 cm x 16 cm spatial bins. Second, the number of calcium events in each bin was divided by the number of frames that mouse stayed in the bin.

Then we computed the spatial information using the firing rate maps of each cell, as previously described [31]

560

Spatial information = $\sum_{i} P_{i}(r_{i}/\overline{r}) \log_{2}(r_{i}/\overline{r})$

 r_i is the calcium event rate of the neuron in the *i*th bin; P_i is the number of frames that mouse stayed in *i*th bin divided by total time in the session. \overline{r} is the mean calcium event rate of all bins; *i* is through over all the bins. Then 1000 permutation shuffles were performed, and spatial information for each shuffle were calculated. The probability of higher spatial information was measured from results. Cells with p < 0.05 and mean firing rates higher than 0.1 Hz were classified as place cell.

To detect place fields in the open field, we make firing rate maps with 2.5 cm x 2.5 cm spatial bins, normalized firing rate of each bin by maximum value in the map, and smoothed the normalized map with 1 SD Gaussian kernel. Next, we made binary map based on the firing rate of each bin (spatial bin with firing rate>0.2 is 1, other spatial bins are 0). Connected bins in this binary map are detected by using MATLAB bwlabel function (Pixel connectivity = 8-connected). Each connected bin is considered as individual place field. Place field which contains bin with maximum firing rate is designated as main place field, and other place fields as extra place field. [23] [32]

573

574 Place cell in Square track

To identify place cells in the square track, we employed reliability to measure the coherence of a 575 576 neuron to fire on specific preferred location in the square track (Supplementary Fig. 2) [33]. We 577 calculated reliability following the procedure below. First, we divided track into 72 spatial bins 578 (2.8cm each), and exclude the first and the last 5 bins where food rewards were given. Spatial 579 activity in each lap was transformed into binarized vectors, in which 1 and 0 represent the presence 580 and absence of a calcium events, respectively. Third, we calculated the correlation value between each lap and the mean of all correlation values. Finally, the locations of calcium events in the 581 582 binarized vector are shuffled randomly 1000 times, then reliability of each shuffled data was 583 calculated. Cells with significant reliability (p < 0.05) are considered as place cell in the square track (Supplementary Fig. 2). Number of running lap with calcium event of all cells and population 584 585 vector correlations between a session and all other sessions are calculated to exclude session 586 without spatial activity. The sessions with mean number of running with calcium event lap < 1.5 or 587 mean population vector correlation < 0.005 are excluded from analysis. We separated place cells for 588 forward and backward running directions.

589 To detect place fields in the square track, we normalized the firing rate map by maximum 590 value, smoothed the normalized map with 1 SD Gaussian kernel, and converted the smoothed map 591 to binary map (spatial bin with firing rate > 0.2 is 1, other spatial bins are 0). Connected bins in this 592 binary map are detected, and each connected bin is considered as individual place field. Place field

593 which contains bin with maximum firing rate is designated as main place field, and other place

594 fields as extra place field[23].

595 Selectivity index

- 596 We used the following calculation to determine the selectivity index of each cell: |(activity in
- 597 context S1 activity in context O) / (activity in context S1 + activity in context O)|.

598

599 **Population vector correlation**

We employed population vector correlation to determine the level of similarity between activitypattern of the different sessions. [34] To calculated vector correlation in each spatial bin, matrix of

- 602 each neuron's event rate in each spatial bin was created for each session. We then computed the
- 603 correlation of event rates in one session with that of the matching location in the other session, and
- 604 mean score over all spatial bins. [23]
- 605

606 Statistics

607 Data are expressed as means \pm SEM unless stated otherwise. All statistical tests were performed 608 using MATLAB. One-way ANOVA was used for group/pair-wise comparisons. Where appropriate,

609 Student's paired t-tests or Wilcoxon rank sum tests was conducted. The null hypothesis was

- 610 rejected at the D < 0.05 level
- 610 rejected at the P < 0.05 level.
- 611

612 Bayesian decoder

613 We used a Naive Bayes Classifier [5, 23, 35] to estimate the mouse location based on neuron

614 calcium activity. The computation of the conditional probability for the subject to be at location x is 615 based on Bayes formula:

 $\mathbf{P}(\mathbf{x}|\vec{n}) = \mathbf{P}(\vec{n}|\mathbf{x}) \mathbf{P}(\mathbf{x}) / \mathbf{P}(\vec{n}) .$

617 *n* is the population activity vector of length *N* containing 1 at index *i*, when neuron *i* is 618 considered to be active and 0 when it is not. P(x), the probability for the subject to be at position *x* 619 was obtained from the dwell time distribution at each spatial bin (bin size = 2.8 cm). $P(\vec{n}/x)$, the 620 conditional probability to observe \vec{n} given the subject is at position *x* was computed from the 621 spatial map $P(n_i|x)$ of individual neurons *i* assuming statistical independence between their activity 622 [5].

623
$$\mathbf{P}(\vec{n}|x) = \prod_{i=1}^{N} p(n_i|x)$$

624 The overall probability to observe n is obtained from $P(\vec{n}) = \sum_{j \in X} P(\vec{n}/x_j) P(\vec{n}/x_j) P(x_j)$, with 625 $P(x_j)$ normalized along the spatial dimension *X*. A reconstructed position of the subject is obtained 626 from the peak position of the evaluated $P(x/\vec{n})$, Eq.1 with $\vec{n} = \vec{n}$ (t). This estimate is further refined 627 taking into account the data from dT = 8 frames before the current frame, assuming a negligible 628 change in position of the animal within dT:

629
$$x_{rec} = \operatorname{argmax}_{j} \prod_{\delta t=0}^{dT} P (\vec{n}(t-\delta t)|x_{j}) P(x_{j}) / P(\vec{n}(t-\delta t))$$

630 The estimation error was calculated as the absolute difference between the real and the

631 reconstructed positions. We trained the decoder with the subjects observed positions and activities

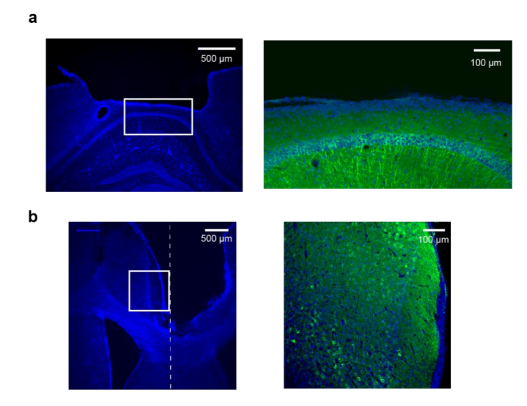
632 of all place cells (place cells with top 6-15% higher firing rate were used) during the first half time

633 of the running period and estimated the trajectory for the following half time. The running time of

the same sessions using the corresponding place cell activities.

636

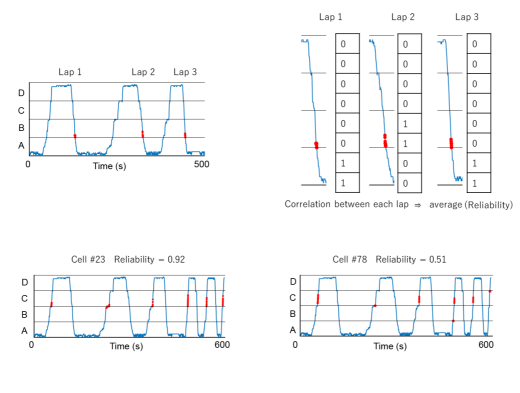
637 SUPPLEMENTARY FIGURES

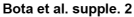


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638 **Figure S1 Histological confirmation of imaging location.**

- 639 (a) Left, a coronal section stained with Hoechst 33258 (blue) and anti-GFP antibody (green) of
- 640 TRE-G-CaMP7 x CaMKIIα-tTA mouse after imaging from ACC. Right, A zoomed image from
- 641 rectangle area. Hoechst image was overlaid with G-CaMP7 immunofluorescence.
- 642 (b) A coronal section stained with Hoechst 33258 and anti-GFP antibody of TRE-G-CaMP7 x
- 643 CaMKIIα-tTA mouse after imaging from hippocampus.
- 644
- 645



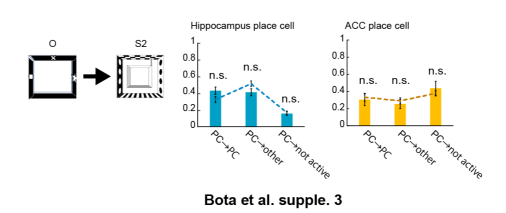


646

647 Figure S2. Calculation of reliability of place cell.

Examples of trajectory of mouse and calcium events of ACC place cells. Spatial activity in each lap
was transformed into binarized vectors, in which 1 and 0 represent the presence and absence of a
calcium events, respectively. We calculated the correlation value between each lap and the mean of
all correlation value. Examples of ACC place cells with high reliability and low reliability are
shown.

653

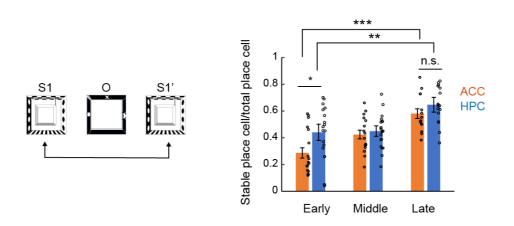


654

656 Figure S3. Conversion of encoding mode of hippocampal and ACC place cells in context O

657 and in context S1.

- 658 Encoding mode of hippocampal or ACC place cells in context O was examined in context S1'.
- 659 Compared with unbiased change, hippocampus: p = 0.082, 0.090, 0.78. ACC: p = 0.92, 0.29, 0.50.
- 660 One-way ANOVA. N = 5 mice for hippocampus, n = 4 mice for ACC.
- 661



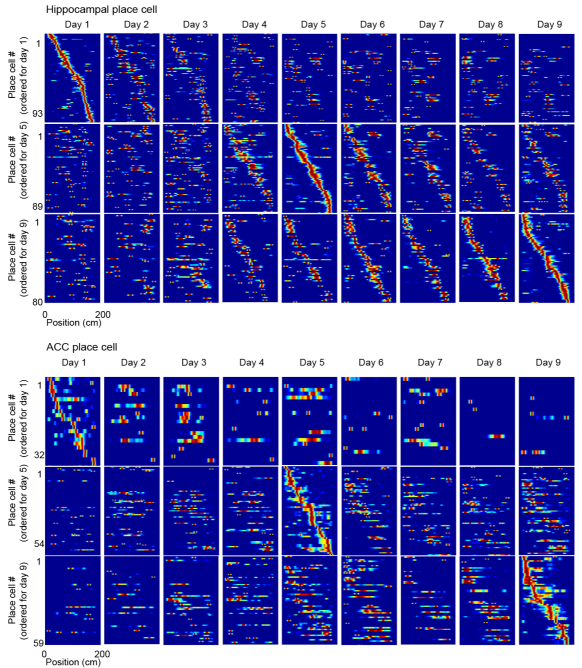
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662

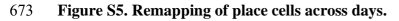
663 Figure S4. The fraction of stable place cells on the same day

664 Place cell stability calculated as the fraction of stable place cells relative to the number of total 665 place cells identified in each session that were compared. Cells in early phase (day 1-3), middle 666 phase (day 4-6) and late phase (day 7-9) were pooled respectively. $p = 9.6 \times 10^{-6}$ (ACC early vs 667 ACC late), p = 0.0015 (hippocampus early vs hippocampus late), p = 0.017 (hippocampus early vs 668 ACC early), p = 0.203 one-way ANOVA. N = 18 session pairs in early, 20 session pairs in middle, 669 16 session pairs in late for hippocampus. N = 18 session pairs in early, 16 session pairs in middle,

- 670 14session pairs in late for ACC.
- 671
- 672

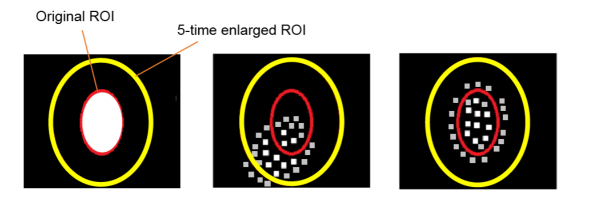


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- 674 Place-field maps of hippocampal and ACC place cells ordered by their centroid positions on day 1
- 675 (top), day 5 (middle) or day 9 (bottom).

677



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678

679 Figure S6. Clustering score.

680 To remove crosstalk from neighboring ROIs, clustering score of each ROI is calculated as follows.

681 After ROIs (red) were identified using a custom MATLAB routine (see method), 5-time enlarged

682 ROIs (by area, yellow) were made. For a given frame of Ca^{2+} images, the location of pixels with top

683 20% brightness were detected. The proportion of the pixels within the original ROI is defined as the

- 684 clustering score for each frame. Low clustering score indicates high likeliness of crosstalk of
- 685 neighboring ROI.

686