Firing rate adaptation affords place cell theta sweeps, phase precession and procession

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- Abstract Hippocampal place cells in freely moving rodents display both theta phase precession
 and procession, which is thought to play important roles in cognition, but the neural mechanism
- for producing theta phase shift remains largely unknown. Here we show that firing rate
- adaptation within a continuous attractor neural network causes the neural activity bump to
- oscillate around the external input, resembling theta sweeps of decoded position during
- ²⁰ locomotion. These forward and backward sweeps naturally account for theta phase precession
- ²¹ and procession of individual neurons, respectively. By tuning the adaptation strength, our model
- ²² explains the difference between "bimodal cells" showing interleaved phase precession and
- ²³ procession, and "unimodal cells" in which phase precession predominates. Our model also
- ²⁴ explains the constant cycling of theta sweeps along different arms in a T-maze environment, the
- speed modulation of place cells' firing frequency, and the continued phase shift after transient
- silencing of the hippocampus. We hope that this study will aid an understanding of the neural
- ²⁷ mechanism supporting theta phase coding in the brain.

²⁹ Introduction

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One of the strongest candidates for temporal coding of a cognitive variable by neural firing is the 30 'theta phase precession' shown by hippocampal place cells. As an animal runs through the firing 31 field of a place cell, the cell fires at progressively earlier phases in successive cycles of the ongoing 32 LFP theta oscillation, so that firing phase correlates with distance traveled (O'Keefe and Recce, 33 1993; Skaggs et al., 1996) (see also (Schmidt et al., 2009)) (Fig. 1a&b). At the population level, 34 phase precession of individual cells gives rise to forward theta sequences once starting phases 35 are aligned across the population (Feng et al., 2015), where neurons representing successive lo-36 cations along the trajectory of the animal display predictable firing sequences within individual 37 theta cycles (Johnson and Redish, 2007). These prospective sequential experiences (looking into 38 the future) are potentially useful for a range of cognitive faculties, e.g., planning, imagination, and 30 decision-making (O'Keefe and Recce, 1993; Skaggs et al., 1996; Hassabis et al., 2007; Wikenheiser 40

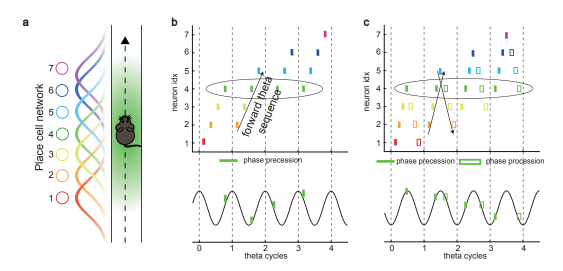


Figure 1. Theta sequence and theta phase shift of place cell firing. a, An illustration of an animal running on a linear track. A group of place cells each represented by a different color are aligned according to their firing fields on the linear track. b, An illustration of the forward theta sequences of the neuron population (upper panel), and the theta phase precession of the 4th place cell (represented by the green color, lower panel). c, An illustration of both forward and reverse theta sequences (upper panel), and the corresponding theta phase precession and procession of the 4th place cell (lower panel). The sinusoidal trace illustrates the theta rhythm of local field potential (LFP), with individual theta cycles separated by vertical dashed lines.

and Redish. 2015: Kav et al., 2020). 41

Besides prospective representation, flexible behaviors also require retrospective representa-42 tion of sequential experiences (looking into the past). For instance, in goal-directed behaviors, it 43 is important to relate the reward information that might only occur at the end of a sequence of 44 events to preceding events in the sequence (Foster et al., 2000; Foster and Wilson, 2006; Diba and 45 Buzsáki, 2007). A recent experimental study (Wang et al., 2020) described retrospective sequences 46 during online behaviors (also indicated by (Skaggs et al., 1996; Yamaguchi et al., 2002)), namely, re-47 verse theta sequences, interleaved with forward theta sequences in individual theta cycles (Fig. 1c). 48 Such retrospective sequences, together with the prospective sequences, may cooperate to estab-49 lish higher-order associations in episodic memory (Diba and Buzsáki, 2007; Jaramillo and Kempter, 50 2017; Pfeiffer, 2020). 51 While a large number of computational models of phase precession and the associated for-52 ward theta sequences have been proposed, e.g., the single cell oscillatory models (O'Keefe and 53 Recce, 1993; Kamondi et al., 1998; Harris et al., 2002; Lengyel et al., 2003; Losonczy et al., 2010) 54 and recurrent activity spreading models (Tsodyks et al., 1996; Romani and Tsodyks, 2015), the 55 underlying neural mechanism for interleaved forward- and reverse-ordered sequences remains 56 largely unclear. Do reverse theta sequences share the same underlying neural mechanism as for-57 ward sequences, or do they reflect different mechanisms? If they do, what kind of neural architec-58 ture can support the emergence of both kinds of theta phase shift? Furthermore, since forward 59 theta sequences are commonly seen, but reverse theta sequences are only seen in some circum-60 stances (Wang et al., 2020), are they commensurate with forward theta sequences? If not, to what 61 degree are forward theta sequences more significant than the reverse ones? 62 To address these questions, we built a continuous attractor neural network (CANN) of the hip-63 pocampal place cell population (Amari, 1977; Tsodyks and Seinowski, 1995; Samsonovich and Mc-64 Naughton, 1997; Tsodyks, 1999). The CANN conveys a map of the environment in its recurrent con-65 nections that affords a single bump of activity on a topographically organized sheet of cells which 66

- can move smoothly so as to represent the location of the animal as it moves in the environment. 67
- Each neuron exhibits firing rate adaptation which destabilizes the bump attractor state. When the 68 adaptation is strong enough, the network bump can travel spontaneously in the attractor space, 60

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- vo which we term as the intrinsic mobility. Intriguingly, we show that, under competition between
- ⁷¹ the intrinsic mobility and the extrinsic mobility caused by location-dependent sensory inputs, the
- network displays an oscillatory tracking state, in which the network bump sweeps back and forth
- ⁷³ around the external sensory input. This phenomenon naturally explains the theta sweeps found
- in the hippocampus (*Skaggs et al., 1996; Burgess et al., 1994; Foster and Wilson, 2007*), where the
- decoded position sweeps around the animal's physical position at theta frequency. More specifi-
- cally, phase precession occurs when the bump propagates forward while phase procession occurs when the network bump propagates backward. Moreover, we find that neurons can exhibit ei-
- ⁷⁷ when the network bump propagates backward. Moreover, we find that neurons can exhibit ei-⁷⁸ ther only predominant phase precession (unimodal cells) when adaptation is relatively strong, or
- ther only predominant phase precession (unimodal cells) when adaptation is relatively strong, or interleaved phase precession and procession (bimodal cells) when adaptation is relatively weak.
- interleaved phase precession and procession (bimodal cells) when adaptation is relatively weak.
 In addition to theta phase shift, our model also successfully explains the constant cycling of
- In addition to theta phase shift, our model also successfully explains the constant cycling of theta sweeps along different upcoming arms in a T-maze environment (*Kay et al., 2020*), and other
- ⁸¹ theta sweeps along difference apcoming arms in a refuze environment (*ray et al., 2020*), and other ⁸² phenomena related to phase precession of place cells (*Geisler et al., 2007*: *Zugaro et al., 2005*).
- ⁸² We hope that this study facilitates our understanding of the neural mechanism underlying the
- rich dynamics of hippocampal neurons and lays the foundation for unveiling their computational
- ⁸⁵ functions.

86 Results

87 A network model of hippocampal place cells

- To study the phase shift of hippocampal place cells, we focus on a one-dimensional (1D) continuous
- attractor neural network (CANN) (mimicking the animal moving on a linear track, see Fig. 2a), but
- ⁹⁰ generalization to the 2D case (mimicking the animal moving in a 2D arena) is straightforward (see
- Discussion for more details). Neurons in the 1D CANN can be viewed as place cells rearranged ac-
- ecording to the locations of their firing fields on the linear track (measured during free exploration).
- ⁹³ The dynamics of the 1D CANN is written as

$$\tau \frac{dU(x,t)}{dt} = -U(x,t) + \rho \int_{-\infty}^{\infty} J(x,x')r(x',t) \, dx' - V(x,t) + I^{ext}(x,t), \tag{1}$$

$$r(x,t) = \frac{gU(x,t)^2}{1 + k\rho \int_{-\infty}^{\infty} U^2(x',t) \, dx'},$$
(2)

- Here U(x, t) represents the presynaptic input to the neuron located at position x on the linear track,
- and r(x,t) represents the corresponding firing rate constrained by global inhibition (*Hao et al.*,
- **2009**). τ is the time constant, ρ the neuron density, k the global inhibition strength, and g is the
- gain factor. The dynamics of U(x,t) is determined by the leaky term -U(x,t), the recurrent input
- ⁹⁸ from other neurons, the firing rate adaptation -V(x,t), and the external input $I^{ext}(x,t)$. The recur-

rent connection strength J(x, x') between two neurons decays with their distance. For simplicity, we set J(x, x') to be the Gaussian form, i.e., $J(x, x') = J_0/(2\pi a) \exp\left[-(x - x')^2/(2a^2)\right]$, with J_0 control-

- ling the connection strength and a the range of neuronal interaction. Such connectivity gives rise
- to a synaptic weight matrix with the property of translation invariance. Together with the global inhibition, the translation invariant weight matrix ensures that the network can hold a continuous
- ¹⁰³ Inhibition, the translation invariant weight matrix ensures that the network can hold a continuous ¹⁰⁴ family of stationary states (attractors) when no external input and adaptation exist (*Tsodyks and*
- ¹⁰⁴ family of stationary states (attractors) when no external input and adaptation exist (*Tsodyks and* ¹⁰⁵ Seinowski, 1995: Samsonovich and McNaughton, 1997: McNaughton et al., 2006: Wu et al., 2008).
 - where each attractor is a localized firing bump representing a single spatial location (Fig. 2b). These
- ¹⁰⁶ where each attractor is a localized firing bump representing a single spatial location (Fig. 2b). These ¹⁰⁷ bump states are expressed as (see Methods. for the parameter settings and SI.2 for the detailed

¹⁰⁸ mathematical derivation):

$$\bar{F}(x,t) = A_r(t) \exp\left\{-\frac{[x-z(t)]^2}{2a^2}\right\},$$
(3)

where $A_r(t)$ denotes the bump height and z(t) the bump center, i.e., the spatial location represented by the network. For convenience, we set the external input to be of the Gaussian form, which

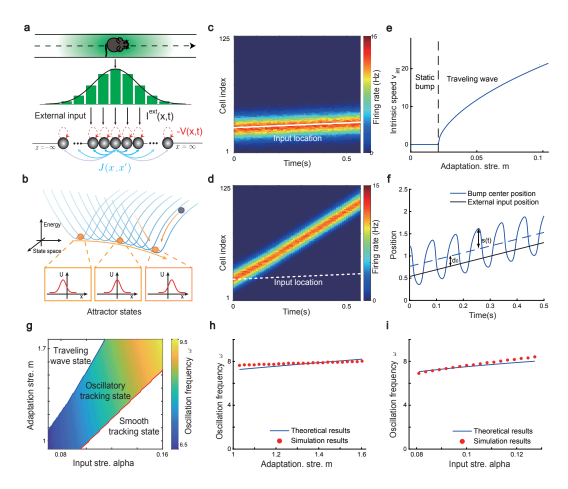


Figure 2. The network architecture and tracking dynamics. a, A 1D continuous attractor neural network formed by place cells. Neurons are aligned according to the locations of their firing fields on the linear track. The recurrent connection strength J(x, x') (blue arrows) between two neurons decays with their distance on the linear track. Each neuron receives an adaptation current -V(x, t) (red dashed arrows). The external input $I^{ext}(x,t)$, represented by a Gaussian-shaped bump, conveys location-dependent sensory inputs to the network. **b**, An illustration of the state space of the CANN. The CANN holds a family of bump attractors which form a continuous valley in the energy space. c, The smooth tracking state. The network bump (hot colors) smoothly tracks the external moving input (the white line). The red (blue) color represents high (low) firing rate. d, The travelling wave state when the CANN has strong firing rate adaptation. The network bump moves spontaneously with a speed much faster than the external moving input. **e**, The intrinsic speed of the travelling wave versus the adaptation strength. **f**, The oscillatory tracking state. The bump position sweeps around the external input (black line) with an offset d_0 . **g**, The phase diagram of the tracking dynamics with respect to the adaptation strength m and the external input strength α . The colored area shows the parameter regime for the oscillatory tracking state. Yellow (blue) color represents fast (slow) oscillation frequency. h-i, Simulated (red points) and theoretical (blue line) oscillation frequency as a function of the adaptation strength (h) or the external input strength (i).

is written as $I^{ext}(x,t) = \alpha \exp\left[-(x - v_{ext}t)^2/(4a^2)\right]$, with v_{ext} representing the moving speed and α

112 controlling the external input strength. Such external moving input represents location-dependent

sensory inputs (i.e., corresponding to the animal's physical location) which might be conveyed via

the entorhinal-hippocampal or subcortical pathways (*Van Strien et al., 2009*). The term -V(x, t)

represents the firing rate adaptation (*Alonso and Klink, 1993*; *Fuhrmann et al., 2002*; *Benda and*

116 Herz, 2003), whose dynamics is written as

$$\tau_v \frac{dV(x,t)}{dt} = -V(x,t) + mU(x,t),\tag{4}$$

where *m* controls the adaptation strength, and τ_v is the time constant. The condition $\tau_v \gg \tau$ holds,

implying that the firing rate adaptation is a much slower process compared to neuronal firing. In ef-

¹¹⁹ fect, the firing rate adaptation increases with the neuronal activity and contributes to destabilizing

the active bump state, which induce rich dynamics of the network (see below).

¹²¹ Oscillatory tracking of the network

Overall, the bump motion in the network is determined by two competing factors, i.e., the external 122 input and the adaptation. The interplay between these two factors leads to the network exhibiting 123 oscillatory tracking in an appropriate parameter regime. To elucidate the underlying mechanism 124 clearly, we explore the effects of the external input and the adaptation on bump motion separately. 125 First, when firing rate adaptation does not exist in the network (m = 0), the bump tracks the external 126 moving input smoothly (see Fig. 2c). We refer to this as the "smooth tracking state", where the 127 internal location represented in the hippocampus (the bump position) is continuously tracking the 128 animal's physical location (the external input location). This smooth tracking property of CANNs 129 has been widely used to model spatial navigation in the hippocampus (Tsodyks and Seinowski, 1995: Samsonovich and McNaughton, 1997: McNaughton et al., 2006: Battaglia and Treves, 1998) 131 Second, when the external drive does not exist in the network ($\alpha = 0$) and the adaptation strength 132 m exceeds a threshold ($m > \tau/\tau_{o}$), the bump moves spontaneously with a speed calculated as 133 $v_{int} = (2a/\tau_v)\sqrt{m\tau_v/\tau} - \sqrt{m\tau_v/\tau}$ (see Fig. 2d&e and Methods. for more details). We refer to this as 134 the **"travelling wave state"**, where the internal representation of location in the hippocampus is 135 sequentially reactivated without external drive, resembling replay-like dynamics during a quiescent 136 state (see Discussion for more details). This intrinsic mobility of the bump dynamics can be intu-137 itively understood as follows. Neurons around the bump center have the highest firing rates and 139 hence receive the strongest adaptation. Such strong adaptation destabilizes the bump stability at 130 the current location, and hence pushes the bump away. After moving to a new location, the bump 140 will be continuously pushed away by the firing rate adaptation at the new location. As a result, the 141 bump keeps moving on the linear track. Similar mechanisms have been applied to explain mental 142 exploration (Hopfield, 2010), preplay during sharp wave-ripple events in the hippocampus (Azizi 143 et al., 2013), and the free memory recall phenomenon in the brain (Dong et al., 2021). 144 When both the external input and adaptation are applied to the CANN, the interplay between 145

the extrinsic mobility (caused by the external input) and the intrinsic mobility (caused by the adap-146 tation) will induce three different dynamical behaviors of the network (see **video 1** for demonstra-147 tion), i.e., 1) when m is small and α is large, the network displays the smooth tracking state: 2) when 148 m is large and α is small, the network displays the travelling wave state: 3) when both m and α have 149 moderate values, the network bump displays an interesting state, called the "oscillatory tracking 150 state", where the bump tracks the external moving input in an oscillatory fashion (Fig. 2f&g). In-151 tuitively, the mechanism for oscillatory tracking can be understood as follows. Due to the intrinsic 152 mobility of the network, the bump tends to move at its own intrinsic speed (which is faster than the 153 external moving input, see Fig. 2d), i.e., the bump tries to escape from the external input. However, 154 due to the strong locking effect of the external input, the bump can not run too far away from the 15 location input, but instead, is attracted back to the location input. Once the bump returns, it will 156 keep moving in the opposite direction of the external input until it is pulled back by the external 157

- input again. Over time, the bump will sweep back and forth around the external moving input,
 displaying the oscillatory tracking behavior.
- Our study shows that during oscillatory tracking, the bump shape is roughly unchanged (see Sec. for the condition of shape variability), and the bump oscillation can be well represented as
- the bump center sweeping around the external input location. The dynamics of the bump center
- can be approximated as a propagating sinusoidal wave (Fig. 2f), i.e.,

$$z(t) = c_0 \sin(\omega t) + d_0 + v_{ext}t = s(t) + v_{ext}t,$$
(5)

- where z(t) is the bump center at time t (see Eq. 3). s(t) denotes the displacement between the bump
- center and the external input, which oscillates at the frequency ω with the amplitude $c_0 > 0$ and a
- constant offset $d_0 > 0$ (see Methods. for the values of these parameters and SI.3 for the detailed
- derivation). When the firing rate adaptation is relatively small, the bump oscillation frequency can
- ¹⁶⁸ be analytically solved to be (see also Fig.S1):

$$\omega = \sqrt{\frac{2\sqrt{\pi\alpha}ak(1+m)}{\tau\tau_{\nu}(J_0 + 2\sqrt{\pi}ak\alpha)}}.$$
(6)

¹⁶⁹ We see that the bump oscillation frequency ω increases sublinearly with the external input strength ¹⁷⁰ α and the adaptation strength *m* (Fig. 2h&i). By setting the parameters appropriately, the bump can ¹⁷¹ oscillate in the theta band (6-10 Hz), thus approximating the experimentally observed theta sweeps ¹⁷² (see below). Notably, LFP theta is not explicitly modelled in the network. However, since theta ¹⁷³ sweeps are bounded by individual LFP theta cycles in experiments, they share the same oscillation ¹⁷⁴ frequency as LFP theta. For convenience, we will frequently use LFP theta below and study firing

175 phase shift in individual oscillation cycles.

Oscillatory tracking accounts for both theta phase precession and procession ofhippocampal place cells

In our model, the bump center and external input represent the decoded and physical positions of
 the animal, respectively, thus the oscillatory tracking of the bump around the external input natu rally gives rise to the forward and backward theta sweeps observed empirically (Fig. 3a&b) (*Wang et al., 2020*). Here we show that oscillatory tracking of the bump accounts for the theta phase
 precession and procession of place cell firing.

Without loss of generality, we select the neuron at location x = 0 as the probe neuron and examine how its firing phase changes as the external input traverses its firing field (Fig. 3c). In the absence of explicitly simulated spike times, the firing phase of a neuron in each theta cycle is measured by the moment when the neuron reaches the peak firing rate (see Methods. for modeling spike times in the CANN). Based on Eqs. 3 & 5, the firing rate of the probe neuron, denoted as $r_0(t)$, is expressed as

$$r_{0}(t) = A_{r}(t) \exp\left[-\frac{[0-z(t)]^{2}}{2a^{2}}\right] = A_{r}(t) \exp\left[-\frac{\left(v_{ext}t + c_{0}\sin\omega t + d_{0}\right)^{2}}{2a^{2}}\right] \equiv A_{r}(t) \exp\left[-\frac{h(t)^{2}}{2a^{2}}\right], \quad (7)$$

where A (t) is the bump height, and h(t) is an oscillatory moving term denoting the displacement 189 between the bump center and the location of the probe neuron. It is composed of a moving signal 190 v_{ext} and an oscillatory signal $c_0 \sin \omega t + d_{0t}$ with c_0 the oscillation amplitude, ω the frequency and d_0 191 an oscillation offset constant. It can be seen that the firing rate of the probe neuron is determined 192 by two factors, $A_{i}(t)$ and h(t). To simplify the analysis below, we assume that the bump height $A_{i}(t)$ 193 remains unchanged during bump oscillations (for the case of time-varying bump height, see Sec.). 194 Thus, the firing rate only depends on h(t), which is further determined by two time-varying terms, 105 the oscillation term $c_0 \sin \omega t$ and the location of the external input v_{ext} . The first term contributes 196

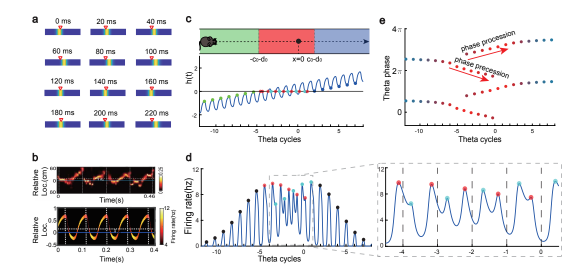


Figure 3. Oscillatory tracking accounts for theta sweeps and theta phase shift. **a**, Snapshots of the bump oscillation along the linear track in one theta cycle (0 ms - 140 ms). Red triangles indicate the location of the external moving input. **b**, Decoded relative positions based on place cell population activities. Upper panel: experimental data, adapted from *Wang et al.* (2020). Lower panel: the relative locations of the bump center (shown by the 10 most active neurons at each timestamp) with respect to the location of the external input (horizontal line) in five theta cycles. **c**, Upper panel: the process of the animal running through the firing field of the probe neuron (large black dot) is divided into three stages: the entry stage (green), the phase shift stage (red) and the departure stage (blue). Lower panel: the displacement between the bump center and the probe neuron as the animal runs through the firing field. The horizontal line represents the location of the probe neuron, which is x = 0. **d**, The firing rates of the probe neuron as the animal runs through the firing field. Colored points indicate firing peaks. The trace of the firing rate in the phase shift stage (the dashed box) is enlarged in the sub-figure on the right hand-side, which exhibits both phase precession (red points) and procession (blue points) in successive theta cycles. **e**, The firing phase shift of the probe neuron in successive theta cycles. **e** form $\pi/2$ to $-\pi/2$ and blues points progress to later phases from $\pi/2$ to $3\pi/2$.

to firing rate oscillations of the probe neuron, and the second term contributes to the envelope of neuronal oscillations exhibiting a waxing-and-waning profile over time, as the external input traverses the firing field (the absolute value $|v_{ext}t|$ first decreases and then increases; see Fig. 3d, also **video 2**). Such a waxing-and-waning profile agrees well with the experimental data (*Skaggs et al., 1996*). In each LFP theta cycle, the peak firing rate of the probe neuron is achieved when |h(t)| reaches a local minima (Fig. 3c&d). We differentiate three stages as the external input passes through the probe neuron (i.e., the animal travels through the place field of the probe neuron), i.e.,

• **the entry stage**. As the external input enters the firing field of the probe neuron (moving from left to right), h(t) < 0 always holds (Fig. 3c). In this case, the peak firing rate of the probe neuron in each oscillatory cycle is achieved when h(t) reaches the maximum (i.e., |h(t)|reaches the minimum). This corresponds to $c_0 \sin \omega t = c_0$, i.e., $\omega t = \pi/2$ (Fig. 3e). This means that the firing phase of the probe neuron at the entry stage is constant, which agrees with experimental observations (*O'Keefe and Recce, 1993; Skaggs et al., 1996*).

• **the phase shift stage.** As the external input moves into the centre of the firing field, h(t) = 0can be achieved in each oscillatory cycle (Fig. 3c). Notably, it is achieved twice in each cycle, once as the bump sweeps over the probe neuron in the forward direction and the other as the bump sweeps over the probe neuron in the backward direction. Therefore, there are two firing peaks in each bump oscillation cycle (Fig. 3d), which are expressed as (by solving $v_{ext}t + c_0 \sin \omega t + d_0 = 0$):

$$\phi_f = -\arcsin\left[\frac{d_0 + v_{ext}t_f}{c_0}\right], \quad \phi_b = \pi + \arcsin\left[\frac{d_0 + v_{ext}t_b}{c_0}\right], \tag{8}$$

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where t_f and t_b denote the moments of peak firing in the forward and backward sweeps,

respectively, and ϕ_f and ϕ_b the corresponding firing phases of the probe neuron. As the external input travels from $(-c_0 - d_0)$ to $(c_0 - d_0)$, the firing phase ϕ_f in the forward sweep decreases from $\pi/2$ to $-\pi/2$, while the firing phase ϕ_r in the backward sweep increases from $\pi/2$ to $3\pi/2$ (Fig. 3e). These give rise to the phase precession and procession phenomena, respectively, agreeing well with experimental observations (*Skaggs et al., 1996; Wang et al.,* 2020; *Yamaguchi et al., 2002*).

• **the departure stage.** As the external input leaves the firing field, h(t) > 0 always holds (Fig. 3c), and the peak firing rate of the probe neuron is achieved when h(t) reaches its minimum in each oscillatory cycle, i.e., $c_0 \sin(\omega t) = -c_0$ with $\omega t = \pi/2$ (Fig.3e). Therefore, the firing phase of the probe neuron is also constant during the departure stage

In summary, oscillatory tracking of the CANN well explains the firing phase shift of place cells when the animal traverses their firing fields. Specifically, when the animal enters the place field. 228 the firing phase of the neuron remains constant, i.e., no phase shift occurs, which agrees with experimental observations (O'Keefe and Recce, 1993; Skaggs et al., 1996). As the animal approaches 230 the centre of the place field, the firing phase of the neuron starts to shift in two streams, one to ear-231 lier phases during the forward sweeps and the other to later phases during the backward sweeps. 232 Finally, when the animal leaves the place field, the firing phase of the neuron stops shifting and 233 remains constant. Over the whole process, the firing phase of a place cell is shifted by 180 degrees. 234 which agrees with experimental observations (O'Keefe and Recce, 1993; Skaggs et al., 1996). 235

²³⁶ Different adaptation strengths account for bimodal and unimodal cells

The results above show that during oscillatory tracking, a place cell exhibits both significant phase precession and procession, which are associated with two firing peaks in a theta cycle. These neurons have been described as bimodal cells (*Wang et al., 2020*) (Fig. 4a). Conversely, previous experiments have primarily focused on the phase precession of place cell firing, while tending to ignore phase procession, which is a relatively weaker phenomenon (*O'Keefe and Recce, 1993*; *Skaggs et al., 1996*). Place cells with negligible phase procession have been described as unimodal cells (Fig. 4b).

Here, we show that by adjusting a single parameter in the model, i.e., the adaptation strength 244 m, neurons in the CANN can exhibit either interleaved phase precession and procession (bimodal 245 cells) or predominant phase precession (unimodal cells). To understand this, we first recall that 246 the firing rate adaptation is a much slower process compared to neural firing and its timescale is 247 in the same order as the LFP theta (i.e., $\tau_{\rm u} = 100$ ms while $\tau = 5$ ms). This implies that when the 248 bump sweeps over a neuron, the delayed adaptation it generates will suppress the bump height 249 as it sweeps back to the same location. Furthermore, since the oscillatory tracking always begins 250 with a forward sweep (as the initial sweep is triggered by the external input moving in the same di-25 rection), the suppression effects are asymmetric, that is, forward sweeps always strongly suppress 252 backward sweeps. On the contrary, the opposite effect is much smaller, since neuronal activities in 253 backward sweeps have already been suppressed, and they can only generate weak adaptation. Be-254 cause of this asymmetric suppression, the bump height in the forward sweep is always higher than 255 that in the backward sweep (see Fig. 4c and Fig. S2a). When the adaptation strength m is small, the 256 suppression effect is not significant, and the attenuation of the bump height during the backward 257 sweep is small (Fig. 4d). In such case, the firing behavior of a place cell is similar to the situation as 258 the bump height remains unchanged as analyzed in Sec., i.e., the neuron can generate two firing 259 peaks in a theta cycle at the phase shift stage, manifesting the property of a bimodal cell of having 260 both significant phase precession and procession (Fig. 4e&g and **video 2**). When the adaptation 261 strength *m* is large, the bump height in the backward sweep attenuates dramatically (see Fig. 4c&d 262 and the video demonstration). As a result, the firing peak of a place cell in the backward sweep 263 becomes nearly invisible at the phase shift stage, and the neuron exhibits only predominant phase 26/ precession, manifesting the property of a unimodal cell (Fig. 4f&h and video 3). 265

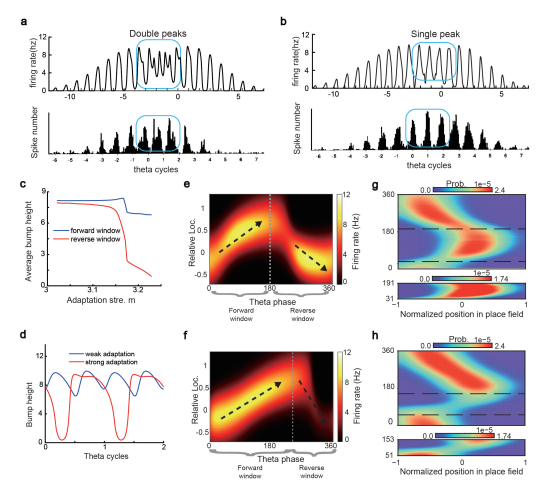


Figure 4. Different adaptation strengths account for the emergence of bimodal and unimodal cells. **a**, The firing rate trace of a typical bimodal cell in our model (upper panel) and the experiment data (lower panel, adapted from (*Skaggs et al., 1996*)). Blue boxes mark the phase shift stage. Note that there are two peaks in each theta cycle. **b**, The firing rate trace of a typical unimodal cell. Note that there is only one firing peak in each theta cycle. **c**, The averaged bump heights in the forward (blue curve) and backward windows (red curve) as a function of the adaptation strength *m*. **d**, Variation of the bump height when the adaptation strength is relatively small (blue line) or large (red line). **e-f**, Relative location of the bump center in a theta cycle when adaptation strength is relatively small (**e**) or large (**f**). Dashed line separate the forward and backward windows. **g-h**, Theta phase as a function of the normalized position of the animal in place field, averaged over all bimodal cells (**g**) or over all unimodal cells (**h**). –1 indicates that the animal just enters the place field, and 1 represents that the animal is about to leave the place field. Dashed lines separate the forward and backward windows. The lower panels in both **g** and **h** present the rescaled colormaps only in the backward window.

In summary, different adaptation strengths explain the emergence of bimodal and unimodal 266 cells. In fact, there is no sharp separation between bimodal and unimodal cells. As the firing rate 267 adaptation gets stronger, the network bump is more attenuated during the backwards sweep, and 268 cells with the bimodal firing property will gradually behave more like those with the unimodal fir-269 ing property (see Fig.S2b). Moreover, our model confirms that even though phase procession is 270 weak, it still exist in unimodal cells (Fig. 4h lower panel), which has been reported in previous stud-271 ies (Wang et al. 2020: Yamaguchi et al. 2002) This implies that phase procession is not a character-272 istic feature of bimodal cells, but instead, is likely a common feature of hippocampal activity, with 273 a strength controlled by adaptation. Furthermore, the experimental data (Fernández-Ruiz et al., 274 2017) has indicated that there is a laminar difference between unimodal cells and bimodal cells. 275 with bimodal cells correlating more with the firing patterns of deep CA1 neurons and unimodal 276 cells with the firing patterns of superficial CA1 neurons. Our model suggests that this difference 277 may come from the different adaptation strengths in the two layers.

278

270

Constant cycling of multiple future scenarios in a T-maze environment We have shown that our model can reproduce the forward and backward theta sweeps of decoded 280 position when the animal runs on a linear track. It is noteworthy that there is only a single hypo-281 thetical future scenario in the linear track environment, i.e. ahead of the animal's position, and 282 hence place cells firing phase can only encode future positions in one direction. However, flexible 283 behaviors requires the animal encoding multiple hypothetical future scenarios in a quick and con-284 stant manner, e.g., during decision-making and planning in complex environments (Iohnson and 285 Redish, 2007: Wikenheiser and Redish, 2015), One recent study (Kay et al., 2020) showed constant 286 cycling of theta sweeps in a T-maze environment (Fig. 5a), that is, as the animal approaches the 287 choice point, the decoded position from hippocampal activity propagates down one of the two 288 arms alternatively in successive LFP theta cycles. To reproduce this phenomenon, we change the 289 structure of the CANN from a linear track shape to a T-maze shape where the neurons are aligned 290 according to the location of their firing fields in the T-maze environment. Neurons are connected 291 with a strength proportional to the Fuclidean distance between their firing fields on the T-maze 292 and the parameters are set such that the network is in the oscillatory tracking state (see details in 293 Methods.). Mimicking the experimental protocol, we let the external input (the artificial animal) move from the end of the center arm to the choice point. At the beginning, when the external 295 input is far away from the choice point, the network bump sweeps back and forth along the center arm, similar to the situation on the linear track. As the external input approaches the choice point. 297 the network bump starts to sweep onto left and right arms alternatively in successive theta cycles (Fig. 5b and **video 4**). The underlying mechanism is straightforward. Suppose that the bump first 200 sweeps to the left arm from the current location, it will sweep back to the current location first due 300 to the attraction of the external input. Then in the next round, the bump will sweep to the right 301 arm, since the neurons on the left arm are suppressed due to adaptation. This cycling process 302 repeats constantly between the two upcoming arms before the external input enters one of the 303 two arms (i.e, before the decision is made). At the single cell level, this bump cycling phenomenon 304 gives rise to the "cycle skipping" effect (Kay et al., 2020: Deshmukh et al., 2010: Brandon et al., 305 2013), where a neuron whose place field is on one of the two arms fires on every other LFP theta 306 cycle before the decision is made (Fig. 5c left panel and Fig. 5d upper panel). For example, a pair of 307 cells with firing fields on each of the two arms will fire in regular alternation on every other theta 308 cycle (Fig. 5c right panel and Fig. 5d lower panel). These cell-level firing patterns agree well the 309 experimental observations (Kav et al., 2020). 310 In summary, our model, extended to a T-maze structure, explains the constant cycling of two 311

possible future scenarios in a T-maze environment. The underlying mechanism relies on delayed 312 adaptation, which alternately causes neurons on one arm to be more suppressed than those on 313 the other arm. Such high-speed cycling may contribute to the quick and continuous sampling 314 among multiple future scenarios in real-world decision-making and planning (see Discussion for 31!

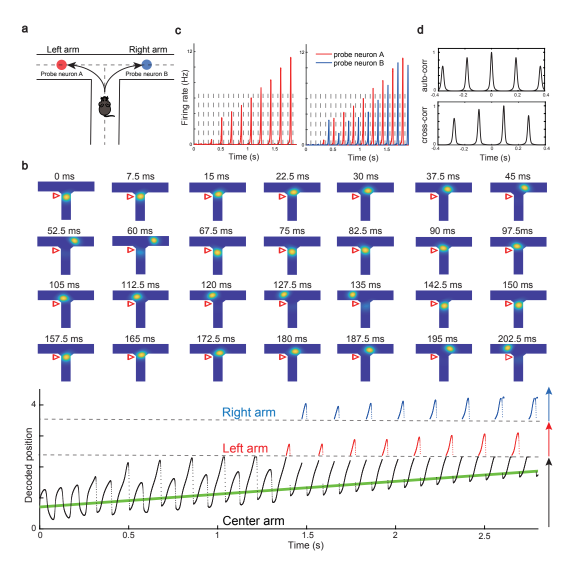
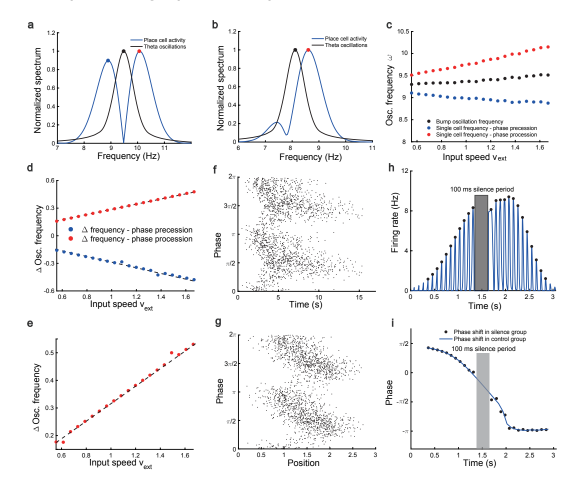


Figure 5. Constant cycling of future positions in a T-maze environment. **a**, An illustration of an animal navigating a T-maze environment with two possible upcoming choices (the left and right arms). **b**, Upper panel: Snapshots of constant cycling of theta sweeps on two arms when the animal is approaching the choice point. Red triangle marks the location of the external input. Note that the red triangle moves slightly towards the choice point in the 200 ms duration. Lower panel: Constant cycling of two possible future locations. The black, red and blue traces represent the bump location on the center, left and right arms, respectively. The green line marks the location of the external moving input. **c**, Left panel: the firing rate trace of a neuron A on the left arm when the animal approaches the choice point. Right panel: the firing rate traces of a pair of neurons when the animal approaches the choice point, with neuron A (red) on the left arm and neuron B (blue) on the right arm. Dashed lines separate theta cycles. **d**, Upper panel: the auto-correlogram of the firing rate trace of neuron A and the firing rate trace of neuron B.

316 more details).



Robust phase coding of position with place cells

Figure 6. Robust phase coding of position. **a**, Normalized spectrum of bump oscillation (black curve) and the oscillation of a bimodal cell (blue curve). The black point marks the peak frequency of the bump oscillation (i.e. LFP theta). The blue and red points mark the peak frequencies of the oscillation of the bimodal cell. **b**, Normalized spectrum of bump oscillation and the oscillation of a unimodal cell. Note that the unimodal cell has only predominant phase precession and hence a single peak frequency (red point), which is higher than that of the bump oscillation. **c**, The peak frequency of bump oscillation versus moving speed (black points), and the oscillation of a bimodal cell versus moving speed (red and blue points). **d**, The difference in peak frequency between the oscillation of a bimodal cell and the bump oscillation versus moving speed. **e**, Same as **d** but for a unimodal cell. **f**-**g**, Theta phase as a function of time (**f**) or position (**g**). Each dot represents a spike generated through a Poisson process (see Method. for more details). **h**, Silencing the network activity for 100 ms (gray shaded area) when the external moving input passes through the center part of the place field of a unimodal cell. **i**, Theta phase shift of the unimodal cell with (black points) or without (blue curve) silencing the network.

As the firing rate shows large variability when the animal runs through the firing field (Fenton 318 and Muller, 1998), it has been suggested that the theta phase shift provides an additional mech-319 anism to improve the localization of animals (O'keefe and Burgess, 2005). Indeed, (Jensen and 320 Lisman, 2000) showed that taking phase into account leads to a significant improvement in the 321 accuracy of localizing the animal. To demonstrate the robustness of phase coding, previous exper-322 iments showed two intriguing findings: a linear relationship between the firing frequency of place 323 cells and the animal moving speed (*Geisler et al., 2007*), and the continued phase shift after inter-324 ruption of hippocampal activity (Zugaro et al., 2005). We show that our model can also reproduce 325 these two phenomena. 326 To investigate the relationship between the single cell's oscillation frequency and the animal's 327

running speed, we consider a unimodal cell with predominant phase precession as studied in the 328 experiment (*Geisler et al., 2007*). Firstly, our model shows that the LFP theta frequency (the bump 329 oscillation frequency ω) is largely independent of the running speed of the animal (the speed of 330 the external input, see Eq. 6) (Fig. 6c). The phase precession implies that the oscillation frequency 331 of a place cell is higher than LFP theta frequency, since firing precesses to earlier phases over theta 332 cycles (Fig. 6a&b). Secondly, we can analytically quantify how the single cell's oscillation frequency is 333 modulated by the external moving speed. As shown in Sec., the distance the animal travels during 334 the phase shift stage is $2c_{0}$, which gives the travelling time $T = 2c_{0}/v_{ext}$ and the number of theta 335 cycles for phase precession $K_f = T\omega$. Since the total amount of phase shift over the whole process 336 is π (i.e, half of the theta cycle, Fig. 3e), it means that K_f firing peaks are generated by a unimodal 337 cell within $T_{f} = (T\omega - 0.5)/\omega$ units of time. Thus, the firing frequency of the cell is calculated to be 338 $\omega_f = K_f/T_f \approx \omega + 0.25 v_{ext}/c_0$ (where the condition $K_f \gg 0.5$ is used), which increases linearly with 339 the animal speed v_{ere} (Fig. 6c-e). This linear relationship ensures that the firing phase of a unimodal 340 cell in each theta cycle is locked with the relative location of the animal in the firing field of that cell, 341 which supports a robust phase-position code. Notably, in our model, the speed modulation of the 342 place cells' firing frequency is not the cause of theta phase shift, but rather a result of oscillatory 343 tracking. This is different from the dual oscillator model (*Lengvel et al.*, 2003), which assumes that 344 phase precession is caused by a speed-dependent increase in the dendritic oscillation frequency 345 (see Discussion for more details). 346 In a different experiment, (Zugaro et al., 2005) found that the firing phase of a place cell contin-347 ues to precess even after hippocampal activity was transiently silenced for up to 250 ms (around 2 348 theta cycles). To reproduce this phenomenon, we also study a unimodal cell by manually turning off 349 the network activity for a few hundred milliseconds (by setting r(x, t) = 0 for all neurons) and then 350 letting the network dynamics evolves again with all parameters unchanged. Based on the theoret-351 ical analysis (Eq. 8), we see that the firing phase of a place cell is determined by the location of the 352 external input (i.e., v_{ert}t), which means that as the external input moves forward on the linear track, 353 the firing phase will precess accordingly in successive oscillatory cycles. Thus, once the network is 354 recovered to the oscillatory tracking state and the external input conveys the new location of the 355 animal to the network, phase precession is resumed from the new location. Therefore, the firing 356 phase in the first bump oscillation cycle after the network perturbation is more advanced than the firing phase in the last bump oscillation cycle right before the perturbation, and the amount of 358 precession is similar to that in the case without perturbation (Fig. 6h&i). This agrees well with the 359 experimental observation, and indicates that the phase-position code is robust to the perturbation 360 of the hippocampal dynamics. 361

Overall, our model reproduces these two experimental findings, and suggests that there exists 362 a one-to-one correspondence between the firing phase of a place cell and the travelled distance in 363 the neuron's place field, which is independent of the animal's running speed or the perturbation 364 duration (Fig.S3). This agrees well with experimental observations (O'Keefe and Recce, 1993) that 365 theta phase correlates better with the animal's location than with time (Fig. 6f&g). In addition to 366 the results for unimodal cells as introduced above, our model predicts new results for bimodal 367 cells. First, in contrast to a unimodal cell, a bimodal cell will have two peaks in its firing frequency. 368 with one slightly higher than the LFP theta baseline (due to phase precession) and the other slightly 360 lower than the LFP theta baseline (due to phase procession). The precession-associated frequency 370 positively correlates with the running speed of the animal, while the procession-associated fre-371 quency negatively correlates with the running speed (Fig. 6d). Second, similar to the preserved 372 phase shift in unimodal cells, both the phase precession and procession of a bimodal cell after 373 transient intrahippocampal perturbation continue from the new location of the animal (see SI), no 374 matter how long the silencing period lasts. The two predictions could be tested by experiments. 375

376 Discussion

377 Model contributions

In this paper, we have proposed a CANN with firing rate adaptation to unveil the underlying mechanism of place cell phase shift during locomotion. We show that the interplay between intrinsic

- mobility (owing to firing rate adaptation) and extrinsic mobility (owing to the location-dependent
- ³⁸¹ sensory inputs) leads to an oscillatory tracking state, which naturally accounts for theta sweeps
- where the decoded position oscillates around the animal's physical location at the theta rhythm.
- At the single neuron level, we show that the forward and backward bump sweeps account for,
- respectively, phase precession and phase procession. Furthermore, we show that the varied adaptation strength explains the emergence of bimodal and unimodal cells, that is, as the adaptation
- tation strength explains the emergence of bimodal and unimodal cells, that is, as the adaptation strength increases, forward sweeps of the bump gradually suppress backward sweeps, and as a
- 386 strength increases, forward sweeps of the bump gradually suppress backward sweeps, and as a 387 result, neurons initially exhibiting both significant phase precession and procession (due to a low
- result, neurons initially exhibiting both significant phase precession and procession (due to a low level adaptation) will gradually exhibit only predominant phase precession (due to a high level adaptation).
- 389 adaptation).

³⁹⁰ Computational models for theta phase shift and theta sweeps

As a subject of network dynamics, oscillatory tracking has been studied previously in an excitatoryinhibitory neural network (*Folias and Bressloff, 2004*), where it was found that decreasing the external input strength can lead to periodic emission of traveling waves in the network (Hopf instability), which is analogous to the oscillatory tracking state in our model. However, their focus was on the mathematical analysis of such dynamical behavior, while our focus is on the biological implications of oscillatory tracking, i.e., how can it be linked to phase precession and procession of hippocampal place cells.

Due to their potential contributions to the temporal sequence learning involved in spatial navigation and episodic memory (Mehta et al., 1997, 2002; Yamaguchi, 2003), theta phase precession 399 and forward theta sweeps have been modelled in the field for decades. These models can be di-400 vided into two main categories, with one relying on the mechanism of single cell oscillation (O'Keefe 401 and Recce, 1993. Kamondi et al., 1998. Lengvel et al., 2003. O'keefe and Burgess, 2005. Mehta et al., 402 2002), and the other relying on the mechanism of recurrent interactions between neurons (Tsodyks 403 et al., 1996; Romani and Tsodyks, 2015; Kang and DeWeese, 2019). A representative example of 404 the former is the oscillatory interference model (O'Keefe and Recce, 1993; Lengvel et al., 2003). 405 which produces phase precession via the superposition of two oscillatory signals, with one from 406 the baseline somatic oscillation at the LFP theta frequency (reflecting the inputs from the medial 407 septal pacemaker (Stewart and Fox, 1990)), and the other from the dendritic oscillation whose 408 frequency is slightly higher. While these models can explain a large variety of experimental phe-409 nomena, it remain unclear how oscillation of individual neurons has a frequency higher than the 410 baseline theta frequency. Here, our model provides a network mechanism for how such higher-411 frequency oscillation emerges. 412

A representative model relying on neuronal recurrent interactions is the activation spreading 413 model (Tsodyks et al., 1996). This model produces phase precession via the propagation of neu-414 ral activity along the movement direction, which relies on asymmetric synaptic connections. A 415 later version of this model considers short-term synaptic plasticity (short-term depression) to im-416 plicitly implement asymmetric connections between place cells (Romani and Tsodyks, 2015), and 417 reproduces many other interesting phenomena, such as phase precession in different environ-418 ments. However, since the asymmetric connections always skew towards the moving direction 419 (along which the connection strength is stronger than that in the opposite direction), the activity 420 bump can only propagate along the moving direction. Therefore, these two models only reproduce 421 theta phase precession. Rather than relying on neuronal asymmetric connections to induce activ-422 ity spreading, our model considers firing rate adaptation at individual neurons, which allows the 423 activity bump to propagate in both directions alternately, and hence generate interleaved phase 424

- ⁴²⁵ precession and procession. Furthermore, to prevent the activity bump from spreading away, their
- model considers an external theta input to reset the bump location at the end of each theta cycle,
- whereas in our model, because of the oscillation of the bump, no external theta input is needed,
- and by choosing the model parameters properly, the bump can oscillate at the theta rhythm. Never-
- theless, experimental studies have suggested that hippocampal neurons receive theta modulation
- from the medial septal pacemaker (Stewart and Fox, 1990; King et al., 1998; Wang, 2002). In our
- model, if we include such an external theta input, the bump oscillation will be locked at the theta rhythm more robustly without the need of fine tuning model parameters. Such a theta input may
- ⁴³² rhythm more robustly without the need of fine tuning model parameters. Such a theta input may
 ⁴³³ also have the role of coordinating theta phase shifts across brain regions. We will investigate this
- 434 issue in future work.

435 Beyond the linear track environment

Besides the linear track environment, the mechanism of generating theta sweeps proposed in our 436 model can also be generalized to more complex environments. For instance, in a T-maze environ-437 ment, our model explains the constant cycling of theta sweeps between left and right arms. Such 438 cycling behavior may be important for high-speed actions such as predating and escaping which 439 require animals to make decision among several future scenarios at the sub-second level. Similar 440 alternative activity sweeps in the T-maze environment has been studied in a previous paper (Ro-441 mani and Tsodyks. 2015), which showed that the frequency of alternation correlates with overtly 442 deliberative behaviors such as head scans (frequency at 1 Hz or less) (Iohnson and Redish, 2007). 443 In contrast to our model, the network activity in their model propagates continuously from the 444 current location on the center arm till the end of the outer arm, which takes a few theta cycles 445 (i.e., 1 second or more). In our model, the network bump alternately sweeps to one of the two 446 outer arms at a much higher frequency (~ 8 Hz), which may be related to fast decision-making 447 or planing in natural environments (Kay et al., 2020). Furthermore, our model can also be easily 448 extended to the multiple-arms (> 2) environment (Gillespie et al., 2021) or the cascade-T envi-449 ronment (Johnson and Redish, 2007) with the underlying mechanism of generating theta cycling 450 remaining unchanged. In addition to the linear and T-maze environments, phase shift has also 451 been reported when an animal navigates in an open field environment. However, due to the lack 452 of recorded neurons, decoding theta sweeps in the 2D environment is not as straightforward as in the 1D case. While theta sweeps in the 1D case have been associated with goal-directed behaviors 454 and spatial planning (Wikenheiser and Redish, 2015), it remains unclear whether such conclusion is applicable to the 2D case. Our preliminary result shows that in the 2D CANN where neurons are 456 arranged homogeneously according to their relative firing locations, the activity bump will sweep along the tangent direction of the movement trajectory, similar to the 1D case (see SI.4 and Fig.S4 458 for details). It will be interesting to explore theta sweeps in the open field environment in detail 450 when more experimental data is available. 460

461 Model implications and future works

In the current study, we have modeled the place cell population in the hippocampus with a CANN 462 and adopted firing rate adaptation to generate theta phase shift. In fact, this model can be easily 463 extended to the grid cell population without changing the underlying mechanism. For instance, 464 we can induce the torus-like connection profile (periodic boundary in the 2D space) (Samsonovich 465 and McNaughton, 1997: McNaughton et al., 2006) or the locally inhibitory connection profile (Burak 466 and Fiete, 2009: Couev et al., 2013) in the CANN structure to construct a grid cell model, and by 467 imposing firing rate adaptation, neurons in the grid cell network will also exhibit phase shift as 469 the animal moves through the grid field, as reported in previous experimental studies (Hafting 460 et al., 2008; Van Der Meer and Redish, 2011). Notably, although for both grid cells and place cells, 470 CANNs can generate theta phase shift, it does not mean that they are independent from each other. 471 Instead, they might be coordinated by the same external input from the environment, as well as by 472 the medial septum which is known to be a pacemaker that synchronises theta oscillations across 47

different brain regions (King et al., 1998; Wang, 2002). We will investigate this issue in future work. 474 Our model suggests that the "online" theta sweep and the "offline" replay may share some 475 common features in their underlying mechanisms (Romani and Tsodyks, 2015; Hopfield, 2010; 476 Kang and DeWeese, 2019: Jahnke et al., 2015). We have shown that the activity bump with strong 477 adaptation can move spontaneously when the external input becomes weak enough (see Sec.). 478 Such non-local spreading of neural activity has a speed much faster than the conventional speed of animals (the external input speed in our model, see Fig. 2d), which resembles the fast spreading of the decoded position during sharp-wave ripple events (Diba and Buzsáki, 2007: Foster and 481 Wilson, 2006: Karlsson and Frank, 2009: Dragoi and Tonegawa, 2011). This indicates that these 482 two phenomena may be generated by the same neural mechanism of firing rate adaptation, with 483 theta sweeps originating from the interplay between the adaptation and the external input, while 484 replay originating from only the adaptation, since the external input is relatively weak during the 186 "offline" state. This hypothesis seems to be supported by the coordinated emergence of theta se-486 quences and replays during the post-natal development period (Muessig et al., 2019), as well as 487 their simultaneous degradation when the animal travelled passively on a model train (*Drieu et al.*, 488 2018). 489 Nevertheless, it is important to note that the CANN we adopt in the current study is an idealized 490 model for the place cell population, where many biological details are missed (Amari, 1977; Tsodyks 491 and Sejnowski, 1995; Samsonovich and McNaughton, 1997; Tsodyks, 1999). For instance, we have 492 assumed that neuronal synaptic connections are translation-invariant in the space. In practice, 403 such a connection pattern may be learned by a synaptic plasticity rule at the behavioral time scale 494 when the animal navigates actively in the environment (*Bittner et al., 2017*). In future work, we 495

will explore the detailed implementation of this connection pattern, as well as other biological

497 correspondences of our idealized model, to establish a comprehensive picture of how theta phase

shift is generated in the brain.

499 Materials and Methods

500 General summary of the model

⁵⁰¹ We consider a one-dimensional continuous attractor neural network (1D CANN), in which neurons

⁵⁰² are uniformly aligned according to their firing fields on a linear track (for the T-maze case, see ⁵⁰³ Methods. below; for the case of the open field (2D CANN), see SI.4). Denote U(x, t) the synaptic

input received by the place cell at location x, and r(x, t) the corresponding firing rate. The dynamics

505 of the network is written as

$$\tau \frac{dU(x,t)}{dt} = -U(x,t) + \rho \int_{-\infty}^{\infty} J(x,x')r(x',t)\,dx' - V(x,t) + I^{ext}(x,t),\tag{9}$$

where τ is the time constant of U(x, t) and ρ the neuron density. The firing rate r(x, t) is given by

$$r(x,t) = \frac{U(x,t)^2}{1 + k\rho \int_{-\infty}^{\infty} U(x',t) \, dx'},\tag{10}$$

where *k* controls the strength of the global inhibition (divisive normalization). J(x, x') denotes the connection weight between place cells at location *x* and *x'*, which is written as:

$$J(x, x') = \frac{J_0}{2\pi a} \exp\left[-\frac{(x - x')^2}{2a^2}\right],$$
(11)

where J_0 controls the strength of the recurrent connection and a the range of neuronal interaction.

Notably, J(x, x') depends on the relative distance between two neurons, rather than the absolute

locations of neurons. Such translation-invariant connection form is crucial for the neutral stability

of the attractor states of CANNs (*Wu et al., 2016*). $I^{ext}(x,t)$ represents the external input which

⁵¹³ conveys the animal location information to the hippocampal network, which is written as:

$$I^{ext}(x,t) = \alpha \exp\left[-\frac{(x - v_{ext}t)^2}{4a^2}\right],$$
 (12)

with v_{ext} denoting the animal's running speed and α controlling the input strength to the hippocam-

⁵¹⁵ pus. V(x,t) denotes the adaptation effect of the place cell at location x, which increases with the ⁵¹⁶ synaptic input (and hence the place cell's firing rate), i..e,

$$\tau_v \frac{dV(x,t)}{dt} = -V(x,t) + mU(x,t),$$
(13)

with τ_v denoting the time constant of V(x,t) and *m* the adaptation strength. Note that $\tau_v \gg \tau$, meaning that adaptation is a much slower process compared to the neural firing.

519 Stability analysis of the bump state

⁵²⁰ We derive the condition under which the bump activity is the stable state of the CANN. For simplic-

ity, we consider the simplest case that there is no external input and adaptation in the network, i.e.,

 $m = \alpha = 0$. In this case, the network state is determined by the strength of the recurrent excitation

and global inhibition. When the global inhibition is strong (k is large), the network is silent, i.e., no

⁵²⁴ bump activity emerges in the CANN. When the global inhibition is small, an activity bump with the

Gaussian-shaped profile emerges, which is written as:

$$\overline{U}(x,t) = A_u \exp\left\{-\frac{[x-z(t)]^2}{4a^2}\right\},$$
(14)

$$\bar{r}(x,t) = A_r \exp\left\{-\frac{[x-z(t)]^2}{2a^2}\right\},$$
(15)

with A_u and A_r representing the amplitudes of the synaptic input bump and the firing rate bump,

respectively. z(t) represents the bump center, and a is the range of neuronal interaction (defined in

Methods.). To solve the network dynamics, we substitute Eqs. 14&15 into Eqs. 9&10, which gives

(see SI.2 for more details of the derivation):

$$\frac{dA_u}{dt} = -A_u + \frac{\rho J_0}{\sqrt{2}}A_r, \qquad (16)$$

$$A_{r} = \frac{A_{u}^{2}}{1 + \sqrt{2\pi k \rho a A_{u}^{2}}},$$
(17)

⁵³⁰ These two equations describes how the bump amplitudes change with time. For instance, if neu-

rons are weakly connected (small J_0) or they are connected sparsely (small ρ), the second term on

the right-hand side of Eq. 16 is small, and A_u will decay to zero, implying that the CANN cannot

sustain a bump activity. By setting $dA_u/dt = 0$, we obtain:

$$A_{u} = \frac{\rho J_{0} \pm \sqrt{\rho^{2} J_{0}^{2} - 8\sqrt{2\pi} 2k\rho a}}{4\sqrt{\pi}k\rho a},$$
(18)

$$A_r = \frac{\sqrt{2}}{\rho J_0} A_u. \tag{19}$$

It is straightforward to check that only when $k < k_c = \rho J_0^2 / 8 \sqrt{2\pi} a$, A_u have two real solutions

(indicated by the ± sign in Eq. 18), i.e., the dynamic system (Eqs. 16&17) has two fixed points. It can be checked that only $A_u = \left(\rho J_0 + \sqrt{\rho^2 J_0^2 - 8\sqrt{2\pi}2k\rho a}\right) / \left(4\sqrt{\pi}k\rho a\right)$ is the stable solution.

537 Analysis of intrinsic mobility of the bump state

⁵³⁸ We derive the condition under which the bump of the CANN moves spontaneously in the attractor

space without relying on external inputs. As the adaptation strength increases, the bump activity

- becomes unstable and has tendency to move away from its location spontaneously. Such intrinsic
- mobility of the CANN has been shown in previous studies (Bressloff, 2011; Wu et al., 2016; Mi et al.,
- **2014**). We set $\alpha = 0$ (no external input), and investigate the effect of adaptation strength *m* on the

bump dynamics. Our simulation result shows that during the spontaneous movement, V(x, t) can

also be represented by a Gaussian-shaped bump, which is written as

$$\overline{V}(x,t) = A_v \exp\left\{-\frac{\left[x - z(t) + d(t)\right]^2}{4a^2}\right\},$$
(20)

where A_r denotes the amplitude of the adaptation bump, and d(t) the displacement between the bump centers of U(x, t) and V(x, t). This displacement originates from the slow dynamics of adaptation, which leads to that the adaptation bump always lags behind the neural activity bump. Similar to Methods. , we substitute the bump profiles Eqs. (14, 15, 20) into the network dynamics Eqs. (9, 10, 13), and obtain:

$$\tau \left[A_{u} \frac{x - z}{2a^{2}} \frac{dz}{dt} + \frac{dA_{u}}{dt} \right] \mathcal{N}(z, 2a) = (-A_{u} + \frac{\rho J_{0}}{\sqrt{2}} A_{r}) \mathcal{N}(z, 2a) - A_{v} \mathcal{N}(z - d, 2a),$$

$$A_{r} = \frac{A_{u}^{2}}{1 + k\rho \sqrt{2\pi} a A_{u}^{2}},$$

$$\tau_{v} \left[A_{v} \frac{x - z + d}{2a^{2}} \frac{d(z - d)}{dt} + \frac{dA_{v}}{dt} \right] \mathcal{N}(z - d, 2a) = -A_{v} \mathcal{N}(z - d, 2a) + mA_{u} \mathcal{N}(z, 2a),$$
(21)

550 where $\mathcal{N}(z, 2a) = \exp\left\{-\left[x-z\right]^2/4a^2\right\}$.

Previous works have shown that the dynamics of a CANN is dominated by very few motion modes (*Fung et al., 2010, 2012*). To solve the CANN dynamics, we can project the network dynamics onto those dominating modes and simplify the analyses significantly. Here, we consider the first two motion modes, corresponding to the changes of the bump height and position, respectively, which are given by,

$$u_0(x,t) = \exp\left\{-\frac{[x-z(t)]^2}{4a^2}\right\},$$
(22)

$$u_1(x,t) = [x - z(t)] \exp\left\{-\frac{[x - z(t)]^2}{4a^2}\right\}.$$
 (23)

⁵⁵⁶ By projecting the network dynamics onto these two motion modes, we obtain:

$$(-A_u + \frac{\rho J_0}{\sqrt{2}}A_r)\sqrt{2\pi}a - A_v \exp(-\frac{d^2}{8a^2})\sqrt{2\pi}a = 0,$$
(24)

$$-A_u + \frac{\rho J_0}{\sqrt{2}} A_r - A_v \exp(-\frac{d^2}{8a^2}) = 0, \qquad (25)$$

$$\tau A_u v_{int} = dA_v \exp(-\frac{d^2}{8a^2}),$$
 (26)

$$\frac{d}{4a^2}\tau_v A_v \exp(-\frac{d^2}{8a^2})v_{int} = -A_v \exp(-\frac{d^2}{8a^2}) + mA_u,$$
(27)

$$\tau_v (1 - \frac{d^2}{4a^2}) v_{int} = d.$$
⁽²⁸⁾

Eqs. 24-28 describes the relationships between bump features A_u, A_r, A_v, v_{int} and d. By solving these

⁵⁵⁸ equations, we obtain,

$$A_{u} = \frac{\rho J_{0} + \sqrt{\rho^{2} J_{0}^{2} - 8\sqrt{2\pi}k\rho a(1 + \sqrt{\frac{m\tau}{\tau_{v}}})^{2}}}{4\sqrt{\pi}k\rho a(1 + \sqrt{\frac{m\tau}{\tau_{v}}})},$$
(29)

$$A_{v} = \frac{\rho J_{0} + \sqrt{\rho^{2} J_{0}^{2} - 8\sqrt{2\pi}k\rho a(1 + \sqrt{\frac{m\tau}{\tau_{v}}})^{2}}}{2\sqrt{2\pi}k\rho^{2}aJ_{0}},$$
(30)

$$A_{r} = \sqrt{\frac{m\tau}{\tau_{v}}} \exp\left[\frac{1-\sqrt{\frac{\tau}{m\tau_{v}}}}{2}\right] \frac{\rho J_{0} + \sqrt{\rho^{2} J_{0}^{2} - 8\sqrt{2\pi}k\rho a(1+\sqrt{\frac{m\tau}{\tau_{v}}})^{2}}}{4\sqrt{\pi}k\rho a(1+\sqrt{\frac{m\tau}{\tau_{v}}})},$$
(31)

$$d = 2a\sqrt{1 - \sqrt{\frac{\tau}{m\tau_v}}},\tag{32}$$

$$v_{int} = \frac{2a}{\tau_v} \sqrt{\frac{m\tau_v}{\tau} - \sqrt{\frac{m\tau_v}{\tau}}}.$$
(33)

Eqs. 29-31 describe the amplitudes of the bumps of synaptic input, firing rate, and adaptation in the CANN, respectively, and Eq. 32 describes the displacement between the neural activity and adaptation bumps. From Eq. 33, we see that for the bump to travel spontaneously, it requires $m > \tau/\tau_v$, i.e., the adaptation strength is larger than a threshold given by the ratio between two time constants τ and τ_v . As the adaptation strength increases (larger *m*), the travelling speed of the bump increases (larger v_{int}).

Analysis of the oscillatory tracking behaviour of the bump state

⁵⁶⁶ When both the external input and the adaptation are applied to the CANN, the bump activity can ⁵⁶⁷ oscillate around the external input if the strengths of the external input and the adaptation are

appropriated. The simulation shows that during the oscillatory tracking, the bump shape is roughly

unchanged, and the oscillation of the bump center can be approximated as a sinusoidal wave expressed as:

$$z(t) = c_0 \sin(\omega t) + d_0 + v_{ext}t,$$
(34)

where c_0 and ω denote, respectively, the oscillation amplitude and frequency, and d_0 denotes a constant offset between the oscillation center and the external input.

Similar to the analysis in Methods. , we substitute the expression of z(t) (Eq. 34) into Eqs. (14, 15, 20), and then simplify the network dynamics by applying the projection method (see SI.3 for more detailed derivation). We obtain,

$$(m+1)A_u - \frac{\rho J_0}{\sqrt{2}} \frac{A_u^2}{1 + \sqrt{2\pi}ak\rho A_u^2} - \alpha = 0,$$
(35)

$$\omega^2 = \frac{\alpha}{\tau \tau_v A_u},$$
 (36)

$$mA_{u} \exp(-\frac{d^{2}}{8a^{2}}) = A_{v}, \qquad (37)$$

$$d_{v} = \tau v. \qquad (38)$$

$$\sqrt{\frac{2(\tau A_u + \alpha \tau_v)}{\alpha \tau_v}} \left[4a^2 (\ln \frac{\tau_v m A_u}{\tau A_u + \alpha \tau_v}) - \tau_v^2 v^2 \right] = c_0,$$
(39)

Eqs. 35-39 describe the relationships among 6 oscillation features A_{ν} , A_{r} , A_{v} , c_{0} , d_{0} and ω . By solving

577 these equations, we obtain:

$$A_u = \frac{J_0 + 2\sqrt{\pi}ak\alpha}{2\sqrt{\pi}ak(1+m)},\tag{40}$$

$$A_r = \frac{A_u^2}{1 + \sqrt{2\pi} a k \rho A_u^2},\tag{41}$$

$$A_v = \sqrt{\left(\frac{\tau A_u + \alpha \tau_v}{\tau_v}\right) m A_u},$$
(42)

$$c_{0} = A_{v} \sqrt{\frac{2}{\alpha m A_{u}}} \left[8a^{2} \ln \frac{m A_{u}}{A_{v}} - \tau_{v}^{2} v^{2} \right],$$
(43)

$$d_0 = \tau_v v, \tag{44}$$

$$\omega = \sqrt{\frac{\alpha}{\tau \tau_v A_u}}.$$
(45)

It can be seen from Eq. 43 that for the bump activity to oscillate around the external input (i.e.,

the oscillation amplitude $c_0 > 0$), it requires that $8a^2 \ln (mA_u/A_v) - \tau_v^2 v^2 > 0$. This condition gives the

boundary (on the parameter values of the input strength α and the adaptation strength m) that

separate two tracking states, i.e., smooth tracking and oscillatory tracking (see Fig. 2g and Fig.S1

⁵⁸² for the comparison between the simulation results and theoretical results).

Note that to get the results in Eqs. 35-39, we have assumed that the amplitudes of neural activity

⁸⁴ bumps and the adaptation bump remain unchanged during the oscillation (i.e., A_u, A_v, A_r are con-

 $_{185}$ stants). However, this assumption is not satisfied when the SFA strength *m* is large (see Sec. and

Fig. 4). In such a case, we carry out simulation to analyze the network dynamics.

⁵⁸⁷ Implementation details of the linear track environment

For the linear track environment, we simulate an 1D CANN with 512 place cells topographically organized on the one-dimensional neuronal track. Since we are interested in how the neuronal 589 firing phase shifts as the animal moves through the firing field of a place cell, we investigate the 590 place cell at location x = 0 and ignore the boundary effect, that is, we treat the linear track with the 591 infinite length. The neural firing time constant is set to be 3 ms, while the time constant of spike 592 frequency adaptation is much longer, which is set to be 144 ms. The density of place cells on the 503 linear track is set to be $256/\pi$. The excitatory interaction range of place cells is set to be 0.4m, while 594 the maximum excitatory connection strength J_0 is set to be 0.2. The gain factor is set to be 5. The 595 global inhibition strength k is set to be 5. The moving speed of the virtual animal v_{ext} is set to be 1.5 506 m/s. For the simulation details, we use the first-order Euler method with the time step δt set to be 597 0.3 the duration of simulation T set to be 10 s. These parameters are commonly used in all plots 598 related to the linear track environment (see Table.1 for a summary). 599 For the two key parameters, i.e., the external input strength α and the adaptation strength m, we 600

vary their values in different plots. Specifically, for illustrating the smooth tracking state in Fig. 2c. 601 we set $\alpha = 0.19$ and m = 0. For illustrating the travelling wave state (intrinsic mobility of the bump 602 state) in Fig. 2d, we set $\alpha = 0$ and m = 0.31. For plotting the relationship between the intrinsic speed 603 v_{im} and the adaptation strength *m* shown in Fig. 2e, we keep $\alpha = 0$, but vary *m* in the range between 604 0 and 0.1 with a step of 0.05. For plotting the overall phase diagram including all three moving states 605 as shown in Fig. 2g, we vary α in the range between 0.05 and 0.16 with a step of 0.001, and m in the 606 range between 0.9 and 1.8 with a step of 0.01. To generate bimodal cell firing patterns in Fig. 3a and 607 Fig. 4a,e&g, we choose $\alpha = 0.19$ and m = 3.02. To generate unimodal firing patterns in Fig. 4b,f&h, we choose $\alpha = 0.19$ but a relatively larger adaptation strength with m = 3.125. The values of these two parameters in different plots are summarized in Table. 2. 610

Parameters	Values
Number of place cells: N	512
Time constant of neural firing: τ	3ms
Time constant of spike frequency adaptation: τ_n	144ms
Neuron density: ρ	$256/\pi$
Recurrent connection range (Gaussian width): a	0.4 m
Recurrent connection strength: J_0	0.2
Gain factor: g	5
Global inhibition strength: k	5
Moving speed of the external input: v_{ext} (m/s)	1.5
Time interval: δt	0.3 s
Simulation duration: T	10 s

Table 1. Commonly used parameter values in the simulation of the linear track environment.

Table 2. Figure specific parameter values for input strength α and adaptation strength *m*.

Figures/parameters	α	m
An example of smooth tracking (Fig. 2c)	0.19	0
An example of traveling wave (Fig. 2d)	0	0.31
Intrinsic speed vs. adaptation strength (Fig. 2e)	0	0:0.05:0.1
Phase diagram (Fig. 2g)	0.05:0.001:0.16	0.9:0.01:1.8
Oscillatory tracking (bimodal) (Fig. 4a,e,g)	0.19	3.02
Oscillatory tracking (unimodal) (Fig. 4b,f,h)	0.19	3.125

611 Implementation details of the T-maze environment

⁶¹² Parameter configurations during simulation

To simulate the T-maze environment, we consider a CANN in which place cells are topographically

organized in a T-shaped area which consists of a vertical central arm and two horizontal left and

right arms (Fig. 5a). The width of the central arm is set to be 0.84 m and the length is set to be 3.14

m. The widths of the two horizontal arms are also set to be 0.84 m, while the lengths of both arms

are set to be 2.36 m. The connection strength between two neurons is determined by the distance

618 between them, which is written as:

$$J(x, x'; y, y') = \frac{J_0}{2\pi a^2} \exp\left[-\frac{(x - x')^2 + (y - y')^2}{2a^2}\right].$$
(46)

Here (x, y) and (x', y') represent the coordinates of two neurons in the T-maze environment, *a* is the recurrent connection range which is set to be 0.3, and J_0 controls the connection strength which is set to be 0.0125. Since we are interested in investigating theta sweeps when the animal is running on the central arm towards the junction point, the external input is restricted on the central arm which is modelled by a Gaussian-like moving bump written as:

$$I^{ext}(x,y) = \alpha \exp\left[-\frac{(x-x_0)^2 + (y-y_0)^2}{2a^2}\right],$$
(47)

where $x_0 = 0$ and $y_0 = v_{ext}t$ represent the center location of the external input with a moving speed $v_{ext} = 1.5$ m/s. In the simulation, we used the first-order Euler method with the time step $\delta t = 0.3$ s and the duration of simulation T = 4.2s. The parameters used are summarized in Table.3.

²²⁷ Calculating auto-correlogram and cross-correlogram

To show the "cycle skipping" effect of a single place cell in the T-maze environment, we calculate

²⁹ the auto-correlogram of the firing rate trace of a place cell whose firing field encodes a location on

the left arm (the upper panel in Fig. 5d). Assume the firing trace of the place cell is f(t) (showed in

left panel in Fig. 5c), the auto-correlogram is calculated as:

Parameters	Values
Number of cells central/left/right: N_1, N_2, N_3	3000/1500/1500
Time constant of neural firing: τ	3 ms
Time constant of spike frequency adaptation: τ_v	144 ms
Neuron density: <i>ρ</i>	$(128/\pi)^2$
Recurrent connection range (Gaussian width): a	0.3
Recurrent connection strength: J_0	$1.25 * 10^{-2}$
Gain factor: g	20
Global inhibition strength: k	1.25
Moving speed of the external input: v_{ext} (m/s)	1.5
Input strength: α	2
Adaptation strength: m	3.96
Time interval: δt	0.3 s
Simulation duration: T	3.3 s

Table 3. Parameters values in the simulation of the T-maze environment

$$(f * f)(\tau) \stackrel{\Delta}{=} \int_{-\infty}^{\infty} f(t)f(t+\tau)dt,$$
(48)

 $_{\mathbf{632}}$ where τ represents the time offset.

To show the "alternative cycling" effect of a pair of place cells with each of them encoding a location on each of the two outward arms, we calculate the cross-correlogram between their firing traces (the lower panel in Fig. 5d). It measures the similarity of the two firing traces as a function of the temporal offset of one relative to the other. Assume the firing traces of the two place cells are f(t) and g(t), respectively, the cross-correlogram is calculated as:

$$(f * g)(\tau) \stackrel{\Delta}{=} \int_{-\infty}^{\infty} f(t)g(t+\tau)dt,$$
(49)

 $_{638}$ where τ represents the time offset.

⁶³⁹ Details of generating the probability heatmap of theta phase shift

In Fig. 4g&h we described the smoothed probability heatmaps of theta phase versus normalized position in the place field of both bimodal and unimodal cells. Generally, these two plots are similar to the traditional spike plot of phase and position traveled in the place field (*O'Keefe and Recce,* **1993; Skaggs et al., 1996**). However, in our rate-based model, the phase of neuronal spike is not directly modelled, rather we use the phase of firing rate peak to represent the phase shift in neuronal firing. Here we describe the implementation details of generating the heatmaps.

The x-axis denotes the normalized position in the place field, with -1 representing the position 646 where the animal just enters the place field, and 1 representing the position where the animal just 647 leaves the place field. In our simulation, the firing field of a place cell with preferred location at 648 x_0 is defined as $x \in (x_0 - 2.5 * a, x_0 + 2.5a)$, with a roughly the half size of the firing field. Consider 649 the animal is at x_t at time t (note that $x_t = v_{ext}t$), then its normalized position \tilde{x}_t is calculated as 650 $\tilde{x}_{t} = (x_{t} - x_{0})/(5a)$. The y-axis represents the phase of neuronal activity, which is in the range of 651 $(0^{\circ}, 720^{\circ})$. To calculate the phase at every time step, we divide the duration of the animal traversing 652 the linear track into multiple theta cycles according to the bump's oscillation. We can calculate the phase by $\theta_{t} = (t - t_{0})/T$, with t_{0} referring to the beginning of the present theta cycle and T referring 654 to the theta period. Denote the firing rate of the *i*-th neuron at time t as $r_i(\tilde{x}_i, \theta_i)$, the probability 655 heatmap is calculated by, 656

$$p(\tilde{x}, \theta_i) = C \sum_{i=1}^{N_c} \theta_i r_i(\tilde{x}, \theta_i),$$
(50)

where $C = 1 / \sum_{t} \sum_{i=1}^{N_c} \theta_t r_i(\tilde{x}, \theta_t)$ is the normalization factor.

Spike generation from the firing rate

- ⁶⁵⁹ To understand phase shift based on spiking time rather than the peak firing rate, we convert the
- ⁶⁶⁰ firing rate into spike trains according to the Poisson statistics (note that our analysis is rate-based,
- ⁶⁶¹ but converting to spike-based does not change the underlying mechanism). For the *i*th place cell
- which encodes position x_i on the linear track, the number of spikes n_i it generates within a time
- interval Δt satisfies a Poisson distribution, which is expressed as,

$$P(n_i|z) = \frac{\left[f_i(z)\Delta t\right]^{n_i}}{n_i!} e^{-f_i(z)\Delta t},$$
(51)

where z is the animal's location, and $f_i(z)$ is the tuning function of cell *i*, which is given by

$$f_i(z) = A_r \exp\left[-\frac{(x_i - z)^2}{2a^2}\right],$$
 (52)

where A_r denotes the amplitude of the neural activity bump and a the range of recurrent interaction.

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- **Competing interests:**Authors declare that they have no competing interests.
- **Data and materials availability:** All code for reproducing the figures in the main text are available
- ⁸²⁶ in the supplementary materials.
- List of material contained in the Supplementary Material
- Supplementary text (pdf file)
- Figures S1-S4
- Video 1-4
- Code for reproducing all the figures in the main text