Short-Term Plasticity Based Network Model of Place Cells Dynamics

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ABSTRACT: Rodent hippocampus exhibits strikingly different regimes of population activity in different behavioral states. During locomotion, hippocampal activity oscillates at theta frequency (5–12 Hz) and cells fire at specific locations in the environment, the place fields. As the animal runs through a place field, spikes are emitted at progressively earlier phases of the theta cycles. During immobility, hippocampus exhibits sharp irregular bursts of activity, with occasional rapid orderly activation of place cells expressing a possible trajectory of the animal. The mechanisms underlying this rich repertoire of dynamics are still unclear. We developed a novel recurrent network model that accounts for the observed phenomena. We assume that the network stores a map of the environment in its recurrent connections, which are endowed with short-term synaptic depression. We show that the network dynamics exhibits two different regimes that are similar to the experimentally observed population activity states in the hippocampus. The operating regime can be solely controlled by external inputs. Our results suggest that short-term synaptic plasticity is a potential mechanism contributing to shape the population activity in hippocampus.

KEY WORDS: neural network; phase precession; hippocampus; place cells; sharp waves

INTRODUCTION

Neuronal population activity recorded in the hippocampus shows strikingly different profiles during different behavioral states of the animal. During exploration, the hippocampus exhibits a prominent oscillation in the 5–12 Hz range (theta frequency) (Vanderwolf, 1969; Buzsaki, 2002). During rest, this regular oscillatory profile switches to global irregular bursting activity (sharp-waves, SW) (Buzsaki, 1986). It is believed that SWs originate in the CA3 sub-field of the hippocampus (Buzsaki, 1989; Csicsvari et al., 2000), a region rich in recurrent collaterals. Artificial disruption of SWs impairs the acquisition of memory tasks, providing a causal link to memory consolidation (Ego-Stengel and Wilson, 2009; Girardeau et al., 2009; Jadhav et al., 2012).

Spiking activity observed in these two regimes also reveals profound differences. During exploratory behavior, cells preferentially fire at specific locations, the place fields (O’Keefe and Dostrovsky, 1971; O’Keefe, 1976). When the animal moves across a neuron’s field, spikes are emitted at progressively earlier phases of the underlying theta rhythm (phase precession) (O’Keefe and Recce, 1993; Skaggs et al., 1996; Cei et al., 2014). During SWs, when the animal rests in a linear track or in a 2D environment, bursting activity occasionally spans sequences which cover—an expanded compressed time scale—a possible trajectory of the animal (Foster and Wilson, 2006; Diba and Buzsaki, 2007; Davidson et al., 2009; Karlsson and Frank, 2009; Gupta et al., 2010; Pfeiffer and Foster, 2013). We term these “virtual” runs non-local events (NLEs), since neural activity is not confined to represent the current location of the animal.

Is it possible to account for the regimes of neural dynamics described above with a single network? In this contribution we propose a neural network model of hippocampus that can exhibit both activity regimes. Our model rests on the observation that place selectivity as well as NLEs appears as spatially coherent activity patterns on the map defined by the place fields of the neurons. We therefore construct a recurrent network model of CA3 that can maintain spatially localized activity “bumps” (Tsodyks and Sejnowski, 1995). We assume that the map of the environment is stored in the strengths of synaptic connections, which depend on the distance between place fields of the corresponding neurons. Recurrent connections in the network are endowed with short-term synaptic depression (STD) that generates a movement of the bump accounting for NLEs (York and van Rossum, 2009). Depending on external inputs, networks with STD can also produce a burst-like behavior that is similar to SWs (Tsodyks et al., 2000; Loebel and Tsodyks, 2002).

MATERIALS AND METHODS

A basic unit of the network represents a population of neurons with highly overlapping place fields. Unit $i$ is described by the firing rate at time $t$, $m_i(t)$, which follows the dynamics

$$\tau m_i(t) = -m_i(t) + f(I_{\theta}^i(t) + I_F^i(t)).$$

$f(I) = a\log(1 + e^{I/a})$ is an F-I curve with an exponential sub-threshold tail and a linear supra-threshold
SHORT-TERM PLASTICITY BASED NETWORK MODEL OF PLACE CELLS DYNAMICS

The input current to unit $i$, is a sum of a recurrent $I_{ii}(t)$ and an external contribution $I_{i}^{E}(t)$. The recurrent contribution to the current is

$$I_{i}^{R}(t) = \frac{1}{N} \sum_{j=1}^{N} W_{ij} m_{j}(t) x_{i}(t),$$

where the sum extends over the $N$ units in the network. The synaptic strength $W_{ij}$ depends on the geometry of the environment and will be specified in each case below.

The network parameters used are: $\tau=10\ ms$, $f_{0}=30\ Hz$, $f_{1}=15\ Hz$, $r_{th}=0.8$, $U=0.8$, $f_{0}=10\ Hz$, $\alpha=1\ Hz$, $N=100$. Parameters for the external currents can be found in Table 1.

Multiple Circular Environments

Each unit $i$ in the network is characterized by a binary vector of selectivity for the $K$ circular environments, $\xi_{i}^{\mu}$.
where \( i = 1 \ldots N \), \( \mu = 1 \ldots K \). If the unit is selective for environment \( \mu \) (i.e., the unit participates to the encoding of that environment) then \( z_i^\mu = 1 \), or 0 otherwise (e.g., Solstad et al., 2014). Selectivity is assigned randomly, with the constraint that \( \sum_i z_i^\mu = fN \), where \( f \) determines the fraction of units selective for any given environment. Each unit is also characterized by the vector of place field locations in the \( K \) circular environments, \( \theta_i^\mu \) [see e.g., (Battaglia and Treves, 1998; Romani and Tsodyks, 2010)].

The synaptic weights are defined as

\[
W_j = \frac{1}{f} \sum_{\mu=1}^{K} \sum_{\nu=1}^{N} \delta_{\nu,j} \cos \left( \theta_i^\mu - \theta_j^\nu \right) - J_0.
\]

The place field locations of the units selective for an environment are a random permutation of equally spaced place field locations in a circular environment. The place specific input is absent in this case, as we only consider the intrinsic dynamics of the network.

The network parameters used are: \( j_1 = 35 \), \( j_0 = 25 \), \( N = 300 \), \( f = 0.3 \), \( K = 3 \). The remaining network parameters are the same of the previous section. Parameters for the external currents can be found in Table 1.

The T-Maze Environment

The T-maze is described by two 1D segments, representing the stem and branches, in a 2D plane. Setting the intersection point at the origin of the plane, a point \( p_i \) on the maze is described by \( p_i = (x_i, 0) \) where \( x_i \in [-\pi, 0) \) (stem, horizontal segment) or \( p_i = (0, y_i) \), \( y_i \in [-\frac{\pi}{2}, \frac{\pi}{2}) \) for the branches (vertical segment, see Fig. 7A). Similarly to the circular map case, units are equally spaced on the maze. The synaptic weight between a pair of units decays exponentially with the distance between their place field locations (Tsodyks et al., 1996)

\[
W_j = j_1 e^{-\frac{d(p_i,p_j)}{\lambda_I}} - J_0.
\]

The function \( d() \) is the Euclidean distance in 2D, \( d(p,p') = \sqrt{(p_x - p'_x)^2 + (p_y - p'_y)^2} \), and \( \lambda_I \) sets the spatial scale of the interaction (see Fig. 7B).

The place specific input is described by

\[
g_i(t) = e^{-\frac{d(p_i,p(t))}{\lambda_p}}.
\]

This input decays exponentially around the location of the virtual animal \( p_i(t) \), with a spatial scale dictated by \( \lambda_p \). For the simulations shown in Figures 7C,D, the peak of the spatial input \( p_i(t) \) moves with a constant speed on the stem of the T-maze for 5 s, and then remains constant for 3 s.

The network parameters used are: \( j_1 = 34 \), \( j_0 = 30 \), \( U = 0.6 \), \( f_0 = 8 \) Hz, \( \lambda_I = 0.3 \), \( \lambda_P = 0.4 \), \( N = 1000 \). The remaining network parameters are as in the section describing the circular environment. Parameters for the external currents can be found in Table 1.

The Toroidal Environment

For the case of a network storing a map of a toroidal environment, unit \( i \) is characterized by two angles, \((\theta_i^1, \theta_i^2)\) (Fig. 1B). The synaptic interaction has the form (Romani and Tsodyks, 2010)

\[
W_{ij} = j_1 [\cos(\theta_i^1 - \theta_j^1) + \cos(\theta_i^2 - \theta_j^2)] - j_0
\]

and the spatially tuned external input is

\[
g_i(t) = [\cos(\theta_i^1 - \theta_{t1}(t)) + \cos(\theta_i^2 - \theta_{t2}(t))]\]

Parameters: \( j_1 = 35 \), \( j_0 = 25 \), \( N = 1024 \), the rest of parameters as in 1D case. Parameters for the external currents can be found in Table 1.

Note that for a given environment, all the network parameters are kept constant and only the external input parameters are varied to produce the different dynamical regimes described in Results. For a given regime, both the network and the external input parameters are varied depending on the environment. We did not attempt to search for network parameters that would allow us to keep the external input parameters fixed for a given regime in all environments.

## RESULTS

### Spatially Coherent Large Irregular Activity

Simulating the network storing a map of a circular environment with a constant uniform input (see Materials and Methods), we observed the emergence of a novel regime of intrinsic activity. For a range of inputs, the network produces elevated activity that is quickly terminated due to fast depression of synaptic transmission (Fig. 2A). A new period of elevated activity is generated after recovery of the depressed synapses. Because of lateral inhibition and local excitation, this activity involves only units with neighboring place field locations (Fig. 2B). Intermittently, a spatially coherent activity bump moves along the map, producing NLEs. The activity is
highly irregular, even though the equations representing the network dynamics are deterministic. This is due to the spatial localization of the neural activity: a bump emerging during a period of elevated activity locally alters the spatial profile of synaptic depression, which in turn influences the timing and localization of subsequent activity. In order to better characterize the network dynamics we performed a longer simulation (1,000 s). Following (Davidson et al., 2009), burst events were defined as contiguous epochs of average population activity above a certain threshold (taken as activity averaged across neurons and the whole simulation period). A total of 2,275 burst events were identified, ranging from \( \sim 100 \) ms to \( \sim 500 \) ms duration (Fig. 2C). The average population activity is multi-peaked; in our network the fraction of burst events with 1 to 4 peaks is 78%, 12%, 8%, and 2%, respectively. The number of peaks in an event is growing with the duration of the event. One circle for each event (black line: linear regression, 7.9 peaks/s). E: Length of the traveled path during an event vs. event duration. Each dot corresponds to a burst event (black line: linear regression, 16.4 rad/s). Longer paths are observed for longer lasting events. The path length saturates at full circle length. F: Distribution of average bump speed during an event, computed from events with more than one peak (average 12 rad/s). See Materials and Methods for the parameters used in the simulation. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

FIGURE 2. Network activity in the spatially coherent bursting regime. A: Average population activity from 10 s of simulated network dynamics in the case of constant uniform input. Periods of elevated activity are interspersed within periods of near silence, resembling multi-unit activity recordings from the rat hippocampus during immobility. B: Corresponding raster plot of single unit activity in the network. Units are orderly arranged in rows according to their place field position in the circular environment. The color code represents unit firing rates. Elevated activity is characterized by the transient activation of localized groups. C: Distribution of event duration. Events are extracted from the average population activity in a 1,000 s simulation (see text). D: The number of peaks in an event correlates with the duration of the event. Ecoenzym: one circle for each event (black line: linear regression, 7.9 peaks/s). E: Length of the traveled path during an event vs. event duration. Each dot corresponds to a burst event (black line: linear regression, 16.4 rad/s). Longer paths are observed for longer lasting events. The path length saturates at full circle length. F: Distribution of average bump speed during an event, computed from events with more than one peak (average 12 rad/s). See Materials and Methods for the parameters used in the simulation. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]
event duration (Fig. 2D). Note that the duration of a single peaked burst event, \(~100\) ms, is comparable with the experimentally observed duration of a single SW event (Foster and Wilson, 2006). Each peak in a burst event contributes \(~100\) ms to the total duration of the event, accounting for the structure observed in Figure 2C. These results are comparable to the experimental finding of chains of SWs in rodents exposed to long linear tracks (Davidson et al., 2009, Gupta et al., 2010), with an experimentally estimated number of \(~10\) SWs/s, compared with the model outcome of 7.9 peaks/s (Fig. 2D). The path length traveled by the bump increases with the duration of the event (Fig. 2E). Some of the burst events, but not all them, contain NLEs (path length > 0). NLEs occur typically, but not only, during burst events with two or more peaks. The distribution of average propagation velocity during events with more than one peak is unimodal, with a characteristic speed of \(~12\) rad/s (Fig. 2F), which can account for the close to linear relationship between path length and duration seen in Figure 2E.

Spatially Coherent Large Irregular Activity in Multiple Environments

One of the hallmarks of place cells activity in the hippocampus is the ability to code for spatial locations in multiple environments. For instance, when exposed to two distinct environments of similar shape, most place cells are active in only one environment, while the minority active in both environments exhibits place fields at different spatial locations, a phenomenon called global remapping (see e.g., Leutgeb et al., 2005). In this scenario, NLEs can be observed in both maps (Karlsson and Frank, 2009). To test whether the network model can qualitatively account for these observations, we stored the maps of three circular environments in the strength

FIGURE 3. Bursting regime in a network storing multiple maps. A: Raster plot of single unit activity during 4 s of simulated dynamics with constant uniform input. Units are orderly arranged in rows according to their place field position in the first circular environment. All the units selective for environment 1 (30% of the total number of units) are included in the raster plot. Color code, unit firing rates. Periods of elevated activity are characterized by the activation of either localized groups or spatially incoherent subsets of units (B,C) As in (A), for environments 2 and 3. Note that spatially incoherent activity in one of the maps occurs whenever localized groups of neurons are active in another map. See Materials and Methods for the parameters used in the simulation. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

FIGURE 4. Activity of the network with spatially tuned input. The same network of Figure 2 but receiving a spatially tuned input around location \(p\) in the circular environment, representing the current location of the animal. A: Dynamics of network activity (color coded) shows moving spatially coherent activity bumps. Red line, location of the peak of the external input. B: Position of the maximally active unit in the network at every time-step. The color code denotes the population averaged activity. Red line as in (A). C: Circular histogram of starting locations for burst events containing more than one peak in the average population activity. NLEs preferentially originate near the location biased by the spatially selective input (red line). D: Histogram of stopping locations for the events considered in (C). [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]
of synaptic connections (see Materials and Methods). We observed the emergence of an irregular regime of activity in a network receiving a constant uniform input. Periods of elevated activity are sometimes characterized by activation of units with nearby place field locations in one of the maps (see e.g., the first 500 ms in Fig. 3A). In contrast with the dynamics that results from a network storing a single map, some periods of elevated activity are characterized by spatially unstructured patterns of units’ activation (e.g., Fig. 3A, from 500 ms to 1 s). The unstructured activity results from spatially coherent activity occurring in other maps; A fraction of the units whose activity is spatially localized in one map, also participates to the encoding of other maps, where their activity appears unstructured (e.g., Figs. 3A,B, from 500 ms to 1 s). Within 4 s of network dynamics, spatially coherent activity and NLEs emerge in all three maps (Fig. 3). Note that there are no instances of spatially localized activity in more than one map simultaneously, due to global feedback inhibition.

Nonlocal Activity With Selective Spatial Input

Based on the similarity between the network behavior and the hippocampal activity during rest, we propose that the novel regime of intrinsic network activity described above corresponds to the experimentally observed NLEs. To establish a tighter link with the experimental observations, we add a place-specific input to the network pointing to a certain location on the map. This input provides information about the current location of the animal. The increased current to the units with place fields close to the position of the animal, biases the starting location of the moving bump on the map, as shown in Figures 4A–C. NLEs typically terminate at a short distance from the current location of the simulated animal (Fig. 4D), sometimes after travelling almost one full circle.

Place Fields and Phase Precession

When the animal is just entering a place field, the first spikes are emitted at the late phase of the ongoing theta rhythm, which is approximately the same for all cells. Subsequent spikes precess toward earlier phases of the theta cycle as the animal traverses a place field. One immediate implication of phase precession is the formation of “theta-sequences” in a population of place cells (Skaggs et al., 1996). Indeed, consider two place cells with overlapping place fields: when the animal moves from the first to the second field and its current location is in the overlapping region of the fields, the first cell will fire earlier than the second one within each theta-cycle, resulting in a wave of place cells activity which propagates towards the direction of motion of the animal. This effect has been recently observed experimentally with multi-unit recordings (Foster and Wilson, 2007; Davidson et al., 2009).

In our network, theta sequences emerge due to a combination of a moving place-specific input and spatially uniform oscillatory input at theta frequency (10 Hz), which could represent the input from the medial septum or entorhinal cortex (Buzsáki, 2002) (Fig. 5). The time course of network activity in Figure 5A,B shows that during each theta cycle, an activity bump emerges, peaked at the neurons with the place fields centered near the current position of the animal (see Zugaro et al., 2004 for experimental evidence of extrahippocampal location reset at the beginning of a theta cycle). The bump subsequently propagates towards the direction of motion of the animal and then disappears at the phase of the theta cycle when inhibition is at its maximum. The packet of activity does not propagate backward, since the recent passage through the just visited locations excited the corresponding place units, thus temporarily depressing the connections between them. Hence, the packet of activity propagates towards unvisited locations.

This mechanism of phase precession is similar to an earlier network model based on asymmetric recurrent connections (Tsodyks et al., 1996), but in the current model theta sequences are generated by STD, hence asymmetry is not required. The absence of hard-wired asymmetry allows the model to produce theta sequences propagating along the direction of motion of the place-specific input, irrespective of its forward or backward direction. Hence the model results account for the experimental evidence of bi-directional theta sequences in one-dimensional environments (Cei et al., 2014).

Single unit activity shows a shift of peak firing from the end of the cycle at the entrance of the field, to the beginning of the cycle while the simulated animal is leaving the place field (Fig. 5C). Phase precession is evident also from the firing rate map in the phase-position space (Fig. 5D). As in (Tsodyks et al., 1996), the intrinsic speed of propagation of the bump during a theta cycle is independent of the speed of the moving animal, thus ensuring a tighter correlation between phase and position, rather than phase and time spent in the place field (Fig. 6), in agreement with experimental results (O’Keefe and Recce, 1993). The circular-linear correlation between theta phase and position estimated from the data shown in Figure 6B is $-0.14$, comparable with experimental results (Huxter et al., 2008). However, the range of precession, defined as the difference between the average firing phases on field entry (10% of the entire field) and field exit, is $65\degree$. The observed range of precession is closer to $180\degree$ (Harris et al., 2002; Huxter et al., 2008). Note that a theta-modulated input is not necessary in order to obtain place specific firing. A strong selective spatial input alone is sufficient to stabilize the place field, accounting for the observation of place cell firing without theta-modulation in bats (Ulanovsky and Moss, 2007; Yartsev and Ulanovsky, 2013).

Activity Sweeps During Ongoing Theta Modulation

Another occurrence of NLEs was observed during alert immobility accompanied by theta oscillations. When an animal temporarily stops at the turning point of a T-maze, the reconstructed position showed alternating sweeps along the unvisited branches of the T-maze (Johnson and Redish, 2007).

We observed a similar behavior in a network encoding a map of a T-maze composed of two segments representing the stem and the branches (Fig. 7A). The maze is stored in the

Hippocampus
synaptic weights between pairs of units, which depend on 2D Euclidean distance between the corresponding fields on the map (Fig. 7B; see Materials and Methods for details). We simulated the network dynamics with a moving spatially tuned input in the presence of theta modulation to mimic a 5-s run along the stem of the maze (Figs. 7C,D). We assume that during the subsequent 3-s period of alert immobility, both spatially tuned input and theta-modulated inputs have reduced amplitude. Under these conditions, the network shows alternating sweeps of activity “exploring” the two branches (Figs. 7C,D). The network does not exhibit backward activity sweeps on the stem of the T-maze, because synapses connecting corresponding units were depressed when the animal moved along the stem.

Dynamics in Two Dimensional Environments

So far we considered the network representing a 1D environment, where most of the cells only fire in one direction (McNaughton et al., 1983). We now extend our model to 2D environments, such as those used in random-foraging experiments, where place fields are nondirectional (Muller and Kubie, 1987; Skaggs et al., 1996; Huxter et al., 2008; Pfeiffer and Foster, 2013; Jeewajee et al., 2014).

For technical reasons, in order to avoid effects of the boundary, we considered an environment with opposite sides “glued” together, i.e., a torus (Romani and Tsodyks, 2010) (see Materials and Methods for details). We observed that the novel regime of irregular activity characterized by spatially coherent activity moving along the map is maintained also in this case. By numerically solving the network dynamics over a long period of time, we observed NLEs preferentially starting closer to the current position of the animal, but ending in random locations (Fig. 8A). To explore phase precession on the torus, we started with a straight run along one of the circles defining the torus. The activity bump grew and disappeared within a theta cycle, while moving toward the direction of motion of the animal, similar to the behavior observed in the circular environment (not shown). As in the 1D case, this movement results in phase precession. The correlation between phase and
NLEs should preferentially avoid the recent path traveled by the animal, at least during SW occurring shortly after the animal stops (Fig. 9). This prediction could be tested with a careful analysis of the experimental results in (Pfeiffer and Foster, 2013). A second prediction is related to the velocity of the animal and its influence on the level of depression on synapse. In the T-maze, when the animal is approaching the decision point at elevated speed, the backward sweeps should be more probable since the synapses had less time to reach a highly depressed state. The few backward sweeps observed in the experiment of (Johnson and Redish, 2007) could be analyzed in order to test this prediction.

**DISCUSSION**

We proposed a neural network model that accounts for the diverse range of hippocampal activity states. The model has two major ingredients: the recurrent connections encode a map of the environment, and synaptic transmission exhibits short-term depression. We observed a novel regime of network activity in the form of temporally irregular bursts of spatially coherent subsets of neurons, occasionally moving along the neural map. Due to the assumed pattern of recurrent connections, the intrinsic activity of the network is shaped by the stored map of the environment, resembling the spontaneous emergence of the orientation maps observed in the visual cortex (Kenet et al., 2003). This novel form of network activity also accounts for the activity replays observed during slow-wave sleep (Wilson and McNaughton, 1994; Lee and Wilson, 2002) and, in line with experimental findings, supports the hypothesis that the intrinsic mode of operation of CA3 is bursting activity (Buzsáki, 1986). In the absence of oscillatory modulation the model also accounts for place selectivity without phase precession (Ulanovsky and Moss, 2007; Yartsev and Ulanovsky, 2013). An additional oscillatory modulation controls the movement of the activity packet during each cycle, resulting in phase precession.

Manipulation of the input when the simulated animal is steady produces escaping activity that resembles the observed sweeps at decision points. In 2D environments, the model exhibits phase precession and predicts NLEs that preferentially avoid the recent trajectory of the animal during the first SWs following immobility. The emergence of NLEs in the model could potentially account for activity “preplays” (Dragoi and Tonegawa, 2011; 2013), where sequences of place cells recorded during exploration of a novel environment were pre-played during SWs preceding the experience. NLEs generated by a network storing multiple environments could be primed for use in subsequent exploratory behavior, when a map of the novel environment can be stored via Hebbian plasticity mechanisms.

It is widely believed that cholinergic modulation plays a major role in the switching between dynamic regimes, but the observation of theta activity interspersed with SWs (O’Neill...
et al., 2006) supports the assumption that switching can be a fast process occurring without the influence of neuromodulators. Since external currents control the different regimes of the network, opto-genetics tools could provide an effective way to probe the model dynamical regimes by depolarizing/hyperpolarizing CA3 neurons or directly manipulating its afferents.

The movement of the activity packet on the neural map caused by STD is an extension of previous recurrent network models of phase precession, where an asymmetry in the synaptic structure generates the movement of the packet (Tsodyks et al., 1996; Jensen and Lisman, 1996; Wallenstein and Hasselmo, 1997). This model has been shown to be consistent with the

FIGURE 7. Activity sweeps in a T-Maze. A: Geometry of the environment and the animal's trajectory. Two segments of equal length are used as the map of a T-maze. Units have place fields arranged on a regular grid on the map. During the simulation, the virtual animal runs for 5 s from the left-most point on the maze stem to the stopping point at the crossing, followed by additional 3 s at the crossing. B: Synaptic weights (color coded) of recurrent connections from the unit at the stopping point to the rest of the network. C: Network activity on the stem of the T-maze. Units are sorted according to their place field position on the horizontal segment of the map. Firing rate is color coded. Red line: movement of the spatial input during the first 5 s. At rest, no sweeps of activity are observed on the stem. D: Network activity on the unvisited branches of the maze. Units sorted according to their place field position on the vertical segment. Units close to the crossing activate when the virtual animal approaches the stopping point at 5 s. For the remaining 3 s the average external input to the network is increased, while the amplitudes of the theta-modulated and spatially selective inputs are decreased. Sweeps of activity cover the unvisited regions of the segment. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

FIGURE 8. Local and nonlocal activity in a 2D environment. A: NLEs in a toroidal map, with a place specific input around the \((\pi, \pi)\) location. Red and black dots denote respectively the location of origin and ending of the NLEs during identified bursts events from a simulation of 50 s (same method as for the 1D case). NLEs typically start close to the location imposed by the selective external input. NLEs terminate at locations covering the torus more uniformly. B: Phase precession in 2D. The curve represents the simulated trajectory of the animal (correlated random walk with a constant speed of \(\frac{2\pi}{5}\) rad/s, and 0.05 angular s.d.) in a run of 300 s. The firing rate (thresholded at 0.5 Hz) of the peaks of the unit with place field at \((\pi, \pi)\) is size-coded, bigger dots correspond to higher firing rates. The color denotes the phase of the external theta-input at the time of the peaks in the firing rate. For illustration purposes, the phases are binarized (red and blue). [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]
experimental findings about phase precession, the robustness to transient perturbation of activity (Zugaro et al., 2004), and the intracellular signatures of place coding (Harvey et al., 2009; Romani et al., 2011). However, the model relies on the directionality of the neurons, and cannot account for phase precession in 2D, where place cells are not directional, or reverse phase precession in 1D during backward travel (Cei et al., 2014). In our model, the movement of the activity packet is caused by STD, hence connections can be symmetric. Strong STD has been indeed observed in CA3 synapses in vitro (Selig et al., 1999), but other studies report a mixture of short-term synaptic facilitation and depression (Miles and Wong, 1986). The behavior of the network with such connections, in the conditions we considered, is qualitatively similar to the one presented in the paper (not shown). Networks with mixed depressing/facilitating synapses have a richer repertoire of behaviors; a full description of the dynamical regimes of the network with depressing, facilitating, or mixed synapses, and the dependence of network behavior on internal and external parameters is outside the scope of the present contribution and will be presented in future studies.

This study shows that attractor networks with short-term synaptic plasticity in recurrent connections can account for a wide range of experimental observations on hippocampal dynamics. We cannot exclude that additional or alternative dynamics at the single neuron or single synapse level may contribute to the observed hippocampal circuit dynamics. For instance, spike frequency adaptation in attractor networks has been shown to produce spontaneous movement of the activity packet (Hansel and Sompolinsky, 1996; Itskov et al., 2011; Azizi et al., 2013), and bursting activity in unstructured spiking networks (Gigante et al., 2007). Whether spike frequency adaptation alone could be sufficient to reproduce the results of our model, and how it would be possible to experimentally discriminate between the two alternatives, will be matter for future studies.

Several studies used facilitation mechanisms to generate firing sequences in models of hippocampal or entorhinal circuits (Leibold et al., 2008; Thurley et al. 2008; Navratilova et al., 2012). In (Leibold et al. 2008; Thurley et al., 2008) a facilitating excitatory input interacted with oscillating excitability in CA3, resulting in temporal compression of sequential activity, as in phase precession. In this class of facilitation models, phase precession results from feed-forward mechanisms, in contrast with the primary role of recurrent connections in our model.
In (Navratilova et al. 2012) path integration network model, a velocity specific input was used to generate an asymmetry in the network, responsible for the movement of the bump; neuronal facilitation (after-depolarization) contributed to resetting the bump at the beginning of a theta cycle, resulting in phase precession. In contrast, in our model, the asymmetry is generated with STD while the reset is achieved with a place-specific input.

It has been suggested that nonlocal hippocampal activity observed during sharp-waves in wakefulness and sleep contributes to memory consolidation. The model we propose exposes a potential mechanism for the generation of such activity, thus constituting an important first step towards the mechanistic understanding of memory encoding.

Our study shows that short-term synaptic plasticity allows the network to be effectively controlled by afferent inputs to produce vastly different dynamic activity regimes. This could be a way for the network to adapt its properties to specific computational demands associated with different behavioral states of the animal. We suggest that this flexibility could be a general feature of cortical networks.

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