Remapping in a recurrent neural network model of navigation and context inference

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Abstract

Neurons in navigational brain regions provide information about position, orientation, and speed relative to environmental landmarks. These cells also change their firing patterns (“remap”) in response to changing contextual factors such as environmental cues, task conditions, and behavioral state, which influence neural activity throughout the brain. How can navigational circuits preserve their local computations while responding to global context changes? To investigate this question, we trained recurrent neural network models to track position in simple environments while at the same time reporting transiently-cued context changes. We show that these combined task constraints (navigation and context inference) produce activity patterns that are qualitatively similar to population-wide remapping in the entorhinal cortex, a navigational brain region. Furthermore, the models identify a solution that generalizes to more complex navigation and inference tasks. We thus provide a simple, general, and experimentally-grounded model of remapping as one neural circuit performing both navigation and context inference.

Keywords: Recurrent neural network models, dynamic coding, latent state, attractor manifolds, population coding, medial entorhinal cortex, navigation.
Introduction

Neural circuit computations throughout the brain, from primary sensory cortex\(^1\)–\(^4\) to higher cognitive areas,\(^5\)–\(^9\) are shaped by combinations of internal and external factors. Internal state changes, such as shifts in attention,\(^6\)–\(^11\) thirst,\(^12\) arousal,\(^13\) and impulsivity,\(^14\) can profoundly alter neural activity across multiple brain areas. This raises a question: how can individual brain regions with specialized functions integrate global state changes without compromising their local processing dynamics?

For example, neurons in the medial entorhinal cortex typically represent one or more features such as spatial position, heading direction, and environmental landmarks and are therefore thought to support navigation.\(^15\)–\(^22\) At the same time, these neurons change their firing rates and shift their spatial firing positions—or “remap”—under a variety of circumstances, even when navigational cues remain stable.\(^6\)–\(^8,23\)–\(^26\) It is difficult to pinpoint the reason for these spontaneous remapping events—i.e., remapping not driven by changes in navigational features—in biological experiments. Theoretical models of this phenomenon propose that remapping occurs because these cells are responding to global contextual cues (like arousal or attention) in order to decorrelate related experiences with distinct contextual relevance.\(^27\),\(^28\) This process could enable animals to form distinct memories or choose appropriate actions for a given set of circumstances. However, these normative models\(^29\) (i.e., theories for why remapping occurs) do not address how a biological system might implement this strategy.

To bridge the gap between existing theoretical models and biological observations of remapping in the entorhinal cortex, we sought to establish a minimal set of task constraints that could reproduce the essential dynamics of remapping in a computational model. Specifically, we tested the normative hypothesis that remapping occurs when a population of neurons must maintain its local navigational processing, while at the same time responding to global latent state changes (e.g., changes in behavioral state, task conditions, etc.; see Sanders et al.\(^28\)). We trained recurrent neural network models (RNNs) to maintain an estimate of position in a simple environment, while at the same time reporting a changing, transiently-cued latent state variable. In isolation, neither of these tasks is novel to the RNN literature—e.g., Cueva and Wei\(^30\) trained RNNs to path integrate in complex environments while Sussillo and Barak\(^31\) trained RNNs on a “1-bit flip-flop” memory task akin to our latent state inference task. Here we combine these two tasks to ask how a network would solve them simultaneously and to probe how this combination of tasks relates to remapping in navigational circuits.

We found that RNNs trained to navigate while inferring latent state changes exhibited network-wide activity patterns that were strikingly similar to those found in the brain,\(^26\) suggesting a possible function for spontaneous remapping in the entorhinal cortex and other navigational brain areas. These activity patterns comprise a geometrically simple solution to the task of combining navigation with latent state inference. Importantly, we found that RNNs identified a modular solution:\(^32\),\(^33\) the RNN dynamics converged to a particular geometry in a simple task, and the geometry and algorithmic principles from this basic task readily generalized to more complex settings. Furthermore, we performed a new analysis of experimental data published in Low et al.\(^26\) and found a similar modular structure in neural activity from a subset of sessions with more than two stable spatial maps. Overall, these results provide an interpretable
and experimentally grounded account of how a single neural population might flexibly represent global brain state changes (corresponding here to remapping) and localized circuit computations (corresponding here to navigation) in orthogonal subspaces.\textsuperscript{34,35}

Results

A recurrent neural network model of 1D navigation and context inference remaps between aligned ring manifolds.

To investigate a putative functional role for spontaneous remapping in an unchanging environment, we developed a task that requires simultaneous latent state inference and navigation in a single neural circuit. To ground our model in experimental data, we designed our task to reflect the basic structure of a recent study\textsuperscript{26} (Figure 1A–D). In this study, Low et al. demonstrated that remapping in the medial entorhinal cortex simultaneously recruited large populations of neurons across the entorhinal cortical circuit. Remapping comprised discrete transitions between aligned neural activity manifolds, which each represented a distinct map of an unchanging, virtual reality 1D environment (Figure 1C–D). Remapping was correlated with transient decreases in running speed (Figure 1B), which could correspond to discrete changes in latent state (such as shifts in arousal, task engagement, or other behavioral states). Thus, we developed a task that requires a single neural circuit to navigate a 1D circular environment while inferring transiently cued, discrete latent state changes. We hypothesized that these task constraints would produce a manifold structure similar to that observed in Low et al. (Figure 1D).

We trained recurrent neural network models (RNNs) with $N = 248$ units to integrate a 1D velocity input along a circular environment (equivalent to a 1D virtual track with seamless teleportation, as in Low et al.\textsuperscript{26}) (Figure 1E, top) and to concurrently remember a binary latent state signal (Figure 1E, bottom). Trained RNNs achieve high performance in both tasks, indicating that they can correctly form a persistent representation of each latent state while maintaining a stable position estimate across states (100% correct state estimation; average angular position error after 300 steps (mean \pm standard deviation): $8.13^\circ \pm 0.51^\circ$) (Figure 1F). To visualize trial-by-trial RNN activity, we provided the trained model with nonnegative velocity inputs and divided the resulting session into track traversals, labeling each traversal by the predominant latent state (Figure 1G, red, context 1; black, context 2). As in biological data (Figure 1C), single units and network-wide activity alternated between distinct maps of the same environment across the two latent states (Figure 1G). Similar to biological neurons, single RNN units remapped heterogeneously (Supplemental Figure 1). Units changed their spatial firing field locations to a similar extent as biological neurons, but changes in firing rate were more common in the model units.
Figure 1: RNN models and biological neural circuits remap between aligned spatial maps of a single 1D environment

(A - D) are modified from Low et al. 26

(A) Schematized task. Mice navigated virtual 1D circular-linear tracks with unchanging environmental cues and task conditions. Neuropixels recording probes were inserted during navigation.

(B) Schematic: slower running speeds correlated with remapping of neural firing patterns.

(C, left) An example medial entorhinal cortex neuron switches between two maps of the same track (top, raster; bottom, average firing rate by position; red, map 1; black, map 2).

(C, right/top) Network-wide trial-by-trial correlations for the spatial firing pattern of all co-recorded neurons in the same example session (colorbar indicates correlation). (C, right/bottom) k-means map assignment.

(D) PCA projection of the manifolds associated with the two maps (colorbar indicates track position).

(E) RNN models were trained on a simultaneous 1D navigation (velocity signal, top) and latent state inference (transient, binary latent state signal, bottom) task.

(F) Example showing high prediction performance for position (top) and latent state (bottom).

(G) As in (C), but for RNN units and network activity. Map is the predominant latent state on each trial.

(H) Example PCA projection of the moment-to-moment RNN activity (colormap indicates track position).

(I) Total variance explained by the principal components for network-wide activity across maps (top 3 principal components, red points).

(J) Normalized manifold misalignment scores across models (0, perfectly aligned; 1, p = 0.25 of shuffle).

(K) Cosine similarity between the latent state and position input and output weights onto the remap dimension (left) and the position subspace (right).
When we projected the hidden layer activity into the subspace defined by the first three principle components, the activity occupied two distinct rings, where position along each ring corresponded to position on the linear track (Figure 1H, red to blue color map). Together, these top 3 components explained ~50% of the variance (Figure 1I, red points). As in Low et al., we used Procrustes shape analysis to demonstrate that these rings were more aligned in high-dimensional activity space than expected by chance for all trained models, such that the position along one ring matched the position on the second ring (Figure 1J). Thus, these task constraints are sufficient to organize randomly initialized synaptic weights into a network model that qualitatively reproduces the remapping dynamics and representational geometry that Low et al. observed in the entorhinal cortex.

RNN geometry is interpretable

We next considered the network geometry in more detail, asking how much of the geometry arises necessarily from the task design. Some basic features of the network structure follow intuitively from components of the task. First, the RNN must maintain an estimate of 1D circular position, which is best achieved through approximate ring attractors. Thus we expect the model to form two ring attractors, one for each of the two distinct latent state conditions, which would explain the two ring-shaped manifolds that we observe. Second, the network must track two statistically independent information streams and should therefore develop separate orthogonal subspaces for each stream. One subspace, the “position subspace,” should contain the position tuning curves for all neurons, as well as the RNN readout weights for position. The other subspace, the “remapping dimension,” should be tuned to changes in latent state and contain the readout weights for state. We confirmed that these dimensions were orthogonal to one another (Figure 1K, Supplemental Figure 2), such that changes in latent state do not interfere with changes in position and vice versa.

However, as we show in detail below, the task does not require the two ring manifolds to be strictly aligned in high dimensional activity space. The RNN maintains a stable position estimate in spite of switches in latent state—which we call “remapping events”—that may occur anywhere along the circular track. To do so, the RNN must implement an invertible remapping function to match track locations across the two ring manifolds. This remapping function could be complex and high dimensional (Figure 2A), resulting in misaligned manifolds, or it could take the form of a simple translation in firing rate space (Figure 2B), resulting in aligned manifolds. The RNN could implement a complex remapping function using its recurrent dynamics, but this implementation could lead to a delay between the latent state signal and remapping. We therefore reasoned that the RNN might converge to the simpler configuration, allowing the linear input layer to implement remapping and thereby enabling rapid remapping in a single timestep.

It is useful to mathematically formalize these ideas to show that the alignment of the two ring manifolds is not strictly imposed by the task. Consider an RNN with $N$ neurons and, for simplicity, consider a discrete grid of $P$ position bins indexed by $p \in \{1, \ldots, P\}$. Let $x_p^{(1)}$ denote an $N$-dimensional vector corresponding to the neural firing rates in spatial bin $p$ along the first ring attractor (i.e. position $p$ in state 1). Likewise, let $x_p^{(2)}$ denote the corresponding firing rate
vector in the second ring attractor (i.e. position \( p \) in state 2). We can compute \( \mathbf{x}_p^{(1)} \) and \( \mathbf{x}_p^{(2)} \) by averaging the RNN activations across many simulated trials, similar to how spatial tuning curves are estimated in biological data.

Let \( W \) denote the \( 2 \times N \) matrix holding the readout layer weights used to decode angular position \( \theta \) by predicting \( \cos \theta \) and \( \sin \theta \) (see STAR Methods). The linearity of this decoder imposes a constraint that \( W\mathbf{x}_p^{(1)} = W\mathbf{x}_p^{(2)} \) for all \( p \); otherwise, the decoded position will erroneously depend on which latent state is active. Importantly, this constraint does not imply that the two rings must have the same shape nor that they must be aligned. To see this let \( \xi_1, \ldots, \xi_p \) denote any arbitrary set of \( N \)-dimensional vectors in the nullspace of \( W \) (i.e., we have \( W\xi_p = 0 \)) and define \( \mathbf{x}_p^{(2)} = \mathbf{x}_p^{(1)} + \xi_p \). Then it is easy to see that the constraint is satisfied,

\[
W\mathbf{x}_p^{(1)} = W\mathbf{x}_p^{(1)} + W\xi_p = W(\mathbf{x}_p^{(1)} + \xi_p) = W\mathbf{x}_p^{(2)}.
\]

Because each \( \xi_p \) was chosen arbitrarily from the nullspace of \( W \), an \( (N-2) \)-dimensional subspace, there are many configurations of the two rings that are compatible with linear decoding of position (Figure 2A). An alternative, lower-dimensional network geometry would instead remap along a constant \( N \)-dimensional translation vector \( \mathbf{v} \), such that we have \( \mathbf{x}_p^{(2)} = \mathbf{x}_p^{(1)} + \mathbf{v} \) (approximately) for all positions \( p \) (Figure 2B).

We now explore each of these intuitions in our data to see how well the trained RNN matches our expectations. First, how closely do the manifolds match the best alignment wherein \( \mathbf{x}_p^{(2)} = \mathbf{x}_p^{(1)} + \mathbf{v} \) ? We computed the empirical remapping vectors \( \xi_p = \mathbf{x}_p^{(2)} - \mathbf{x}_p^{(1)} \) for each position bin and verified that \( W\xi_p \approx 0 \) for all positions (mean \( W\xi_p \pm \text{sem}: 9 \times 10^{-6} \pm 10^{-5} \)). We then defined \( \mathbf{v} \) to be the average of these remapping vectors, \( \mathbf{v} = \langle \xi_p \rangle \). If the manifolds were perfectly aligned then we would observe \( \xi_p = \mathbf{v} \) for all positions \( p \).

We instead find that there is some variability in the remapping vectors, such that \( \xi_1, \ldots, \xi_p \) are not exactly equal to one another (Figure 2C). Indeed, when we perform PCA on the \( p \times N \) matrix formed by concatenating the vectors \( \xi_1, \ldots, \xi_p \), we find that remapping dimensions lie within a 2-dimensional subspace (Figure 2D, E), in contrast to our original conjecture that remapping vectors would be effectively zero-dimensional (i.e. \( \xi_p = \mathbf{v} \) for all positions).

Nonetheless, the idealized model in which each \( \xi_p \approx \mathbf{v} \) is a much better fit to the observed RNN dynamics than would be expected by chance. When we randomly rotate the orientation of the two rings in the nullspace of \( W \), we find that this approximation is much worse (Figure 2C, dashed line).
Figure 2: The ring attractors are more aligned than strictly required by the task.
(A) Schematic illustrating how the two attractor rings could be misaligned in the nullspace of the position output weights (left) while still allowing linear position decoding (right) (colormap, track position; solid arrows, remapping vectors; dashed arrow, projection onto the position subspace).
(B) Schematic illustrating perfect manifold alignment (colors as in A).
(C) Normalized difference between the true remapping vectors for all position bins and all models and the ideal remapping vector (dashed line, p = 0.025 of shuffle; n = 50 position bins, 15 models).
(D) Dimensionality of the remapping vectors for an example model. (Right) Total variance explained by the principal components for the remapping vectors (red points, top 2 PCs). (Left) Relative variance explained by the remapping vectors (red) and the position rings (blue) (1 = total network variance).
(E) The remapping vectors vary smoothly over position. (Right) Projection of the remapping vectors onto the first 2 PCs. (Left) Normalized covariance of the remapping vectors for each position bin (blue, min; yellow, max).

Altogether, these findings suggest that RNN models trained on a simultaneous latent state inference and navigation task converge to a geometrically simple solution out of the space of all possible, high-dimensional solutions. This simpler solution recapitulates the geometry of entorhinal cortical dynamics during remapping in virtual reality environments. Notably, neither the RNN nor the biological data are consistent with the simplest 3-dimensional solution, as evidenced by the imperfect ring alignment (Low et al. and Figure 1J), the variable remapping vectors (Fig. 2C-D), and the dimensionality of the network (which is > 3; Figure 1I).

RNN dynamics follow two ring attractor manifolds

While neural manifold geometry can provide clues about the computational mechanisms at play in the system, one advantage of RNN models is that we can precisely characterize the structure and logic of their dynamics using tools from nonlinear systems analysis. As we describe
below, these tools reveal several insights into the underlying network computations that are not easy to experimentally demonstrate in biological networks.

Each RNN defines a nonlinear, discrete-time dynamical system, $x_{t+1} = f(x_t, u_t)$, where $f(\cdot, \cdot)$ is a nonlinear function parameterized by synaptic weight matrices and $u_1, \ldots, u_T$ defines a sequence of latent state cues and velocity inputs to the network. Using methods pioneered in Sussillo & Barak,\textsuperscript{31} we used numerical optimization to identify approximate fixed points, which are $N$-dimensional vectors $x_*$ that satisfy $f(x_*, u) \approx x_*$ for a specified input $u$. In particular, we studied the case where $u = 0$, corresponding to a situation where no velocity or context input is provided to the network. Intuitively, the network should approach a fixed point when no velocity or context input is provided because the position and latent state is unchanging.

The fixed points of the RNN provide a backbone for understanding its dynamics. While the global RNN dynamics are complex and nonlinear, the dynamics near any fixed point $x_*$ can be approximated as a linear dynamical system governed by the $N \times N$ Jacobian matrix of partial derivatives $\partial f / \partial x_j$ evaluated at $x = x_*$ and $u = 0$ (see STAR Methods).

We computed these Jacobian matrices across 988 fixed points in the trained RNN shown in Figure 1. Roughly 60\% of these fixed points were located on one of the two previously described ring manifolds and largely had marginally stable linear dynamics (largest Jacobian eigenvalue \( \approx 1 \); Figure 3, color-coded green). The remaining fixed points were located between the two ring manifolds and had unstable dynamics (largest Jacobian eigenvalue $> 1$; Figure 3, color-coded gold). In essence, this analysis confirms that the RNN dynamics indeed implemented a pair of ring attractors. Further, a collection of unstable fixed points form a boundary between the two stable ring attractor basins. In line with observations by Susillo & Barak\textsuperscript{31} on a discrete flip-flop task, these intermediate fixed points are unstable along a small number of dimensions (i.e. saddle fixed points; Figure 3D, gold points) which “funnel” neural activity to the appropriate location during a remapping event.

This interpretation is supported by examining the principal eigenvector—i.e. the eigenvector associated with the largest magnitude eigenvalue—for each fixed point. For the fixed points along the two ring attractors, this eigenvector corresponds to a slow dimension along which $x$ does not grow or decay (i.e., its associated eigenvalue $\lambda \approx 1$)(Figure 3D, green points). Consistent with a mechanism for integrating positional information, these eigenvectors were nearly orthogonal to the remapping dimension and aligned with the position subspace (Figure 3E-G, green). Conversely, for the unstable fixed points, the principal eigenvector corresponds to a dimension along which $x$ moves rapidly away from the fixed point (i.e., its associated eigenvalue $\lambda > 1$)(Figure 3D, gold points). Consistent with a mechanism for “funneling” activity during remapping events, these eigenvectors were aligned with the remapping dimension and nearly orthogonal to the position subspace (Figure 3E-F, yellow).
Figure 3: RNN dynamics follow stable ring attractor manifolds mediated by a ring of saddle points

(A) Fixed points (gray) reside on the two ring manifolds and in a ring between them (gray, all fixed points; colors, example fixed points for B and D).

(B) The maximum real component of each fixed point (shading, marginally stable or unstable points; colored points, examples from A). Dashed red lines indicate cut-off values for fixed points to be considered marginally stable ($\lambda \approx 1$).

(C) Projection of fixed points onto the remapping dimension (-1, map 1 centroid; 1, map 2 centroid; 0, midpoint between manifolds).

(D) Distribution of eigenvalues (colored points) in the complex plane for each example fixed point from (A) (black line, unit circle).

(E) Cosine similarity between the eigenvectors associated with the largest magnitude eigenvalue of each fixed point and the remap dimension (top) or the position subspace (bottom).

(F) Eigenvector directions for 48 example fixed points (black lines, eigenvectors).

(G) Projection of the example fixed points closest to each manifold onto the respective position subspace (top) and zoom on an example fixed point for each manifold (bottom, red box)(purple, estimated activity manifold; dashed line, approximate position estimate).

Green indicates marginally stable points and gold indicates unstable points throughout.
Aligned toroidal manifolds emerge in a 2D generalization of the task.

Virtual 1D tracks are an ideal setting to experimentally study spontaneous remapping: the environmental cues can be tightly controlled and it is possible to sample an identical spatial trajectory hundreds of times, such that remapping events can be easily identified from the neural activity alone. But navigation is often studied in 2D environments, in which it is more difficult to control the animal’s experience and the animal can pursue an essentially infinite number of trajectories through the environment. Thus, while it is of interest to understand what remapping in the entorhinal cortex might look like in 2D spaces, it remains challenging to identify spontaneous remapping in biological data. In contrast, the RNN modeling framework that we have developed here can be readily generalized to 2D spatial environments. Are the computational solutions identified by the RNNs fundamentally different in this case? Or do RNNs use similar geometric structures and algorithmic principles across these related tasks?

To investigate this question, we again trained models to simultaneously integrate velocity inputs and estimate latent state from transient state cues, but this time we provided two velocity inputs and asked the models to estimate 2D position (Figure 4A, right). As before, the models performed well on both components of the task (mean loss ± sem: position estimate, 0.036 ± 1.1x10^-3; latent state estimate, 0.002 ± 1.9x10^-5)(Figure 4A) and single unit activity was modulated by both spatial position and latent state (Figure 4B). When we projected the activity into a subspace defined by the first three principal components, the activity occupied two distinct toroidal manifolds with position on each torus corresponding to the position in the 2D space (Figure 4C). In keeping with these qualitative observations, four principal components explained ~50% of the variance in network activity (Figure 4D) and the manifolds were again highly, though not perfectly, aligned in the full-dimensional activity space (Figure 4E). By holding either the horizontal (X) or vertical (Y) position variable constant during the RNN simulation, we recover a pair of 1D aligned ring manifolds (Figure 4F). That is, we can recover the geometry of the original 1D task (see Figure 1) by taking “slices” through the toroidal manifolds. The remapping and position dimensions were again orthogonalized in these models (Figure 4G). Thus models trained on a 2D navigation task with latent state inference identified a geometrically similar solution to those trained on a 1D task. These findings demonstrate that spontaneous remapping is possible in 2D and may operate under similar mechanisms as in 1D.
Figure 4: An RNN model of 2D navigation and context inference remaps between aligned toroidal manifolds

(A) (Left) Schematic illustrating the 2D navigation with simultaneous latent state inference task. (Right) As in Fig. 1E, but RNN models were trained to integrate two velocity inputs (X, Y) and output a 2D position estimate, in addition to simultaneous latent state inference.

(B) Position-binned activity for three example units in latent state 1 (top) and latent state 2 (bottom) (colormap indicates normalized firing rate; blue, minimum; yellow, maximum).

(C) Example PCA projection of the moment-to-moment RNN activity from a single session into three dimensions (colormap indicates position; left, X position; right, Y position).

(D) Average cumulative variance explained by the principal components for network-wide activity across maps (top 4 principal components, red points).

(E) Normalized manifold misalignment scores for all models (0, perfectly aligned; 1, p = 0.25 of shuffle).

(F) Example PCA projection of slices from the toroidal manifold where Y (left) or X (right) position is held constant, illustrating the substructure of RNN activity.

(G) Cosine similarity between the latent state and position input and output weights onto the remap dimension (left) and the position subspace (right), defined for each pair of maps.
Manifold alignment generalizes to three or more maps

It is simplest to consider remapping as switches between two maps, but neural activity can conceivably switch between any number of maps. Indeed, while Low et al. most commonly observed remapping between two maps of the same virtual track, they occasionally found transitions between more than two maps. Because these “multi-map” sessions were rare, Low et al. predominantly limited their analysis to the more common “2-map” sessions. Nonetheless, there were notable similarities between the 2-map and multi-map sessions. In particular, remapping was correlated with changes in running speed and position was preserved across remapping events. We reasoned that we could study the geometry of “multi-map” sessions using RNN models to gain insight into what we might expect to see in biological data. In particular, do multiple ring manifolds corresponding to multiple spatial maps emerge and are these ring manifolds still geometrically aligned with each other?

We trained models to integrate a 1D velocity input, while tracking three (instead of two) binary state switch cues (Figure 5A). Models performed well on both task components (mean loss ± sem: position estimate, 0.013 ± 3.6x10⁻⁴; latent state estimate, 0.0039 ± 4.9x10⁻⁵) and single unit activity was modulated by both spatial position and latent state (Figure 5B). Importantly, we found that the same essential geometry of the original task was preserved. When we visualized each pair of maps using PCA, the ring manifolds were again qualitatively aligned (Figure 5C, right). Projecting the activity from all three maps into the same subspace revealed that they were further organized as vertices of an equilateral triangle (Figure 5C, left)—i.e. the acute angle between any two remapping dimensions was 60° (Figure 5D). Again, the network geometry was relatively low-dimensional (4 principal components explained ~60% of the variance; 14 principal components, ~90% of the variance)(Figure 5E). Procrustes analysis revealed that all pairs of manifolds were highly aligned relative to chance, with a similar degree of alignment across ring manifold pairs and across RNNs (Figure 5F). Finally, positional and latent state information were orthogonalized, as before (Figure 5G).

In Supplemental Figure 3, we show that RNNs are capable of solving this task with larger numbers of latent states (more than three). Further, the RNN dynamics and geometry generalizes accordingly: each latent state is associated with a different ring attractor and every pair of ring attractors is highly aligned. Motivated by these observations, we revisited a subset of experimental sessions from Low et al. (N = 4 sessions from 2 mice) that exhibited remapping between 3 – 4 stable maps of the same virtual track (Figure 6A-B; Supplemental Figure 4) for a pilot comparison with the RNN models, which we hope will inspire future experimental analysis. As in Low et al. we first confirmed that these remapping events did not reflect recording probe movement by comparing the waveforms from different maps across the session, which were highly stable (Figure 6B, right; Supplemental Figure 4).
Figure 5: An RNN model of 1D navigation and identification of three latent states remaps between three aligned ring manifolds.

(A) (Left) RNN models were trained to navigate in 1D (velocity signal, top) and discriminate between three distinct contexts (transient, binary latent state signal, bottom). (Right) Example showing high prediction performance for position (top) and context (bottom).

(B) Position-binned activity for six example single RNN units, split by context (colors as in (A)).

(C) Example PCA projection of the moment-to-moment RNN activity into three dimensions (colormap indicates track position) for all three contexts (left) and for each pair of contexts (right).

(D) Schematic: (Top) The hypothesized orthogonalization of the position and context input and output weights. (Bottom) Across maps, corresponding locations on the 1D track occupy a 2D remapping subspace in which the remapping dimensions between each pair are maximally separated (60°).

(E) Total variance explained by the principal components for network-wide activity across maps (top 4 principal components, red points).

(F) Normalized manifold misalignment scores between each pair of maps across all models (0, perfectly aligned; 1, p = 0.25 of shuffle).

(G) Cosine similarity between the latent state and position input and output weights onto the remap dimension (left) and the position subspace (right), defined for each pair of maps.
To examine the manifold structure of the neural data, we projected the neural tuning curves associated with each pair of maps into the subspace defined by the first three principle components. In many sessions, population-wide activity across the two maps occupied distinct, qualitatively aligned rings, where position along each ring corresponded to position along the virtual track (Figure 6C). Procrustes analysis revealed that these pairs of maps were largely more aligned than expected by chance (Figure 6D; 13/18 map pairs more aligned than shuffle). Notably, 5 map pairs from one mouse (3 in session A, 2 in session D) were not aligned (Figure 6D, black and teal points), suggesting that manifold alignment does not always emerge in biological data. Finally, we asked whether the remapping dimensions from the biological sessions were organized symmetrically, as in the model (Figure 6E, left). We found that there were a range of acute angles between pairs of remapping dimensions (Figure 6E, right), suggesting that there was more asymmetry in the biological network geometry than in the model.

Figure 6: Biological recordings with more than two maps recapitulate many geometric features of the recurrent neural network models. (A - B) are modified from 26
(A) Schematic: Mice navigated virtual 1D circular-linear tracks with unchanging environmental cues and task conditions. Neuropixels recording probes were inserted during navigation.
(B) Examples from the 3-map Session A. (Left/top) Network-wide trial-by-trial correlations for the spatial firing pattern of all co-recorded neurons in the same example session (colorbar indicates correlation). (Left/bottom) k-means map assignments. (Middle) An example medial entorhinal cortex neuron switches between three maps of the same track (top, raster; bottom, average firing rate by position; teal, map 1; red, map 2; black, map 3). (Right) Overlay of average waveforms sampled from each of the three maps.
(C) PCA projection of the manifolds associated with each pair of maps from the 4-map Session D (colorbar indicates virtual track position).
(D) Normalized manifold misalignment scores between each pair of maps across all sessions (0, perfectly aligned; 1, p = 0.25 of shuffle).
(E)(Left) Schematic: maximal separation between all remapping dimensions for 3 and 4 maps. (Right) Angle between adjacent pairs of remapping dimensions for all sessions (dashes, ideal angle).
Discussion

Previous experimental studies have found that neurons in the medial entorhinal cortex change their firing patterns in response to changes in task conditions, behavioral state, or visual and motor cues.6–8,23–25 In virtual reality environments, these remapping events can recruit neurons across the entorhinal cortical circuit to rapidly switch between distinct maps of the same track.25,26 Here, we used RNN models to explore a normative hypothesis that these remapping dynamics reflect hidden state inference.27,28 We showed that RNNs initialized from random synaptic weights recapitulate the essential features of biological data—aligned ring manifolds26—when trained to remember a binary latent state variable and to simultaneously integrate a velocity signal in a circular environment. RNNs learn to represent positional and state change information in orthogonal subspaces such that navigation and latent state inference co-occur without interference. Further, we demonstrated that the geometry and algorithmic principles of this solution readily generalize to more complex tasks including navigation in 2D environments and tasks involving three or more latent states. These findings provide a jumping off point for new analyses of remapping in neural data, which we demonstrated in a pilot analysis of neural data from Low et al.26

These results complement an existing body of theoretical and experimental work on the neural basis of navigation. Fenton et al.11 proposed that the hippocampus constructs multiple spatial maps that are anchored to different landmarks; when the animal’s attention switches between these reference points, hippocampal cells remap (see also ref.38). This proposal is consistent with the idea that the hippocampal circuit groups navigational episodes into discrete categories by combining internal context with external landmarks.27,28,39 Related experimental work demonstrates that an animal’s prior experience with an environment can shape how the hippocampus delineates these categories.40 Each of these hypotheses can be seen as layering a discrete latent variable (e.g. changes in reference landmarks, task context, or prior experience) on top of a distributed neural code of position, which are the essential ingredients of our RNN task. While we draw explicit comparisons with spontaneous remapping in the entorhinal cortex,26 Sheintuch et al.41 reported similar experimental findings in the hippocampus, highlighting the broad relevance of this remapping phenomenon and our modeling efforts. We explored these topics in a general modeling framework applicable to any circuit that supports navigation through physical space and even navigation of abstract cognitive spaces.42–44

In our task, discrete switches in latent state are signaled by brief impulses to the RNN, such that the navigational circuit must maintain a persistent representation of the latent state based on these transient cues. This simple task design allowed us to clearly identify the minimal set of constraints that produces aligned ring attractors. In particular, our results suggest that aligned ring attractors could emerge even if upstream circuits trigger latent state changes and signal these changes to downstream navigational circuits. Indeed, Low et al. found that remapping was correlated with brief decreases in running speed,26 suggesting that this temporary behavioral state change—which is known to have widespread impacts on global brain activity13—may serve as an external trigger of remapping in the entorhinal cortex. Extensions to our task could build on this basic framework by asking the network to infer state changes given
a noisy input or using a more complex interaction with the environment (e.g., through reinforcement learning paradigms). The mechanisms of remapping in biological circuits are still poorly understood, but have been modeled using multistable attractor dynamics for several decades. Classically, these models were engineered and hand-tuned to produce the desired attractor dynamics. In contrast, RNN models are indirectly engineered by specifying task constraints and a learning algorithm. Thus, our observation that trained RNNs produce multistable attractor manifolds is nontrivial, because different solutions might have, in principle, emerged. Despite this key similarity, there are notable differences between our models and classical multistable attractor models. Classical models typically store completely decorrelated spatial maps, while our RNNs produce distinct maps that are, by construction, perfectly correlated in the position readout dimensions. Romani & Tsodyks studied the effects of adding correlation to spatial maps in forward-engineered multistable attractor networks, as did Low et al. Notably, our model captured aspects of the data that these previous forward-engineered models did not explore—namely, that the ring manifolds corresponding to the correlated spatial maps were much more aligned than expected by chance and than strictly required by the task. Most fundamentally, forward-engineered models provide insights into how neural circuits may remap, but do not answer why they do so. We investigated the latter question in this work by identifying a minimal set of task constraints that provide a putative explanation for why the entorhinal cortex spontaneously remaps.

Other work has studied remapping in trained artificial networks performing navigation. Unlike our results, these papers typically consider remapping across different physical environments. Whittengton et al. propose a normative model and a neural circuit that supports non-spatial remapping, which is perhaps most similar to the task constraints we studied. However, our investigation focused on a simpler and more targeted computational task to draw a tighter link to a specific biological finding and to perform a deeper examination of the resulting population geometry and dynamical structure.

While we were motivated to study remapping in the specific context of navigational circuits, our results have broader implications for understanding how RNNs perform complex, context-dependent computations. This topic has attracted significant interest. For example, RNNs trained in many computational tasks develop modular neural populations and dynamical motifs that are re-used across tasks. When the RNN architecture is explicitly modular in its design, these subpopulations can improve model performance on particular types of tasks. Thus, there is an emerging conclusion that RNNs use simple dynamical motifs as building blocks for more general and complex computations, which our results support. In particular, aligned ring attractors are a recurring, modular motif in our results, appearing first in a simple task setting (2 maps of a 1D environment) and subsequently as a component of RNN dynamics in more complex settings (e.g., as sub-manifolds of toroidal attractors in a 2D environment, see Figure 4). Intriguingly, this modular motif also emerged in our novel analysis of neural data from Low et al., suggesting that biological systems may also leverage this strategy.

Ultimately, our model provides a strong foundation for future experimental investigations of the functional role of remapping in navigational circuits. Our findings suggest that latent state
changes can drive remapping; an experimental task that explicitly requires animals to report a latent internal state would provide substantial insight into this hypothesis. We also identify concrete predictions for how the representational geometry of neural populations generalizes from the dynamics found in 1D virtual reality environments\textsuperscript{25,26} to more complex settings. We found direct support for one of these predictions by re-analyzing an existing experimental dataset. Our work therefore provides a parsimonious, plausible, and testable model for the neural population geometry of remapping navigational circuits under a variety of task conditions.
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Author contributions

I.I.C.L., conceptualization, methodology, formal analysis, investigation, writing – original draft, writing – review & editing, and visualization; L.M.G., conceptualization, writing – review & editing, supervision, and funding; A.H.W., conceptualization, methodology, formal analysis, investigation, writing – original draft, writing – review & editing, visualization, supervision, and funding.

Declaration of interests

The authors declare no competing interests.
Methods

Resource Availability

Lead Contact:
Further information and requests for resources and reagents should be directed to and will be
fulfilled by the Lead Contact, Alex H. Williams (alex.h.williams@nyu.edu).

Materials Availability:
This study did not generate new unique reagents.

Data and Code Availability
Code to train RNN models and reproduce the figures of this paper are provided in a GitHub
repository (https://github.com/ahwillia/rnn_remapping_paper).

Any additional information required to reanalyze the data reported in this paper is available from
the lead contact upon request.

Experimental Model and Subject Details

RNN Model and Training Procedure
We examined Elman RNNs ("vanilla" RNNs), which are perhaps the simplest RNN architecture
capable of theoretically representing any nonlinear dynamical system,\textsuperscript{52} and which can be
viewed as an approximation to continuous time firing rate models of neural circuits.\textsuperscript{53} At each
time index \( t \in \{1, \ldots, T\} \) the activation vector of \( N \) hidden units is denoted by \( x_t \in \mathbb{R}^N \). Loosely,
we can think of \( x_t \) as the firing rates of \( N \) neurons in a biological circuit at time \( t \). The activation
vector is updated according to:
\[
x_{t+1} = \text{ReLU}(Ax_t + Bu_t + \beta)
\]
where \( \text{ReLU}(x) \) denotes a rectifying linear unit function (i.e., an element-wise maximum
between the vector \( x \) and a vector of zeros), \( A \in \mathbb{R}^{N \times N} \) is a matrix holding the recurrent synaptic
connection weights, \( u_t \in \mathbb{R}^M \) is a vector of input signals at time \( t \), \( B \in \mathbb{R}^{N \times M} \) is a matrix holding
the input connection weights, and \( \beta \in \mathbb{R}^N \) is a vector holding bias terms for each hidden unit.
The output of the network at time \( t \) is defined by:
\[
y_t = Cx_t + \alpha
\]
where $\mathbf{y}_t \in \mathbb{R}^L$ is a vector of $L$ output units, $\mathbf{C} \in \mathbb{R}^{L \times N}$ is a matrix holding output connection weights, and $\mathbf{\alpha} \in \mathbb{R}^L$ is a vector holding bias terms for each output unit. Finally, the initial condition $x_0 \in \mathbb{R}^N$ for each dynamical sequence was set by:

$$x_0 = \mathbf{D} \mathbf{z} + \mathbf{y}$$

where $\mathbf{z} \in \mathbb{R}^{M_0}$ is a vector of $M_0$ inputs used to define the initial condition, $\mathbf{D} \in \mathbb{R}^{N \times M_0}$ is a matrix holding connection weights, and $\mathbf{y} \in \mathbb{R}^{M_0}$ is a vector holding bias terms. As described below, the vector $\mathbf{z}$ is used to define the initial position on the circular track, which is randomized on each trial. Altogether, these equations define an RNN model with trainable parameters $\{\mathbf{A}, \mathbf{B}, \mathbf{\beta}, \mathbf{C}, \mathbf{\alpha}, \mathbf{D}, \mathbf{y}\}$.

The number of inputs, $M$, and outputs, $L$, varied depending on the computational task the RNN was trained to perform. Specifically, $M$ is given by the number of latent states (“contexts”) plus the number of spatial dimensions. Thus, for the 1D navigation task with binary state cues diagrammed in Figure 1E, the number of network inputs was $M = 3$ (whereas the tasks diagrammed in Figure 4A and Figure 5A each have $M = 4$ inputs). The number of network outputs, $L$, is given by the number of latent states plus two times the number of spatial dimensions. Thus, for the task diagrammed in Figure 1E, $L = 4$ (for Figure 4A, $L = 6$; for Figure 5A, $L = 5$). The additional spatial output dimensions can be understood as follows: Due to the periodic boundary conditions, the network must output a predicted spatial position $\theta_t \in [0, 2\pi)$ for each spatial dimension. Predicting this raw angular position would require the network to implement something akin to an $\arctan(\cdot)$ function. Because this function is highly nonlinear and discontinuous, the linear readout layer of the RNN will struggle to predict $\theta_t$ directly in a fashion that is numerically stable. We therefore trained the networks to predict $\sin \theta_t$ and $\cos \theta_t$ for each spatial dimension, which requires an extra factor of two spatial output dimensions. Similarly, for the initial condition $x_0$, the number of input variables $M_0$ is given by two times the number of spatial dimensions, and the input vector $\mathbf{z}$ is formed by concatenating $\sin \theta_0$ and $\cos \theta_0$ for each spatial dimension.

The network was trained by randomly generated input sequences with a ground truth target output. The input vector at each time step, $\mathbf{u}_t$, contained the angular velocity along each spatial dimension as well as state change cues (see schematic in Figure 1E). The output vector at each time step, $\mathbf{y}_t$, contains disjoint dimensions that predict spatial position and the latent state or “context” (see schematic in Figure 1F). For each sequence, the overall loss function is a sum of two terms: (i) the mean-squared-error between the ground truth sine and cosine of angular position, $\sin \theta_t$ and $\cos \theta_t$, and the network’s prediction of these terms, and (ii) the cross-entropy of the true latent state and the network’s prediction. This second term is the usual loss function...
for multi-class classification tasks; it coincides with a negative log-likelihood criterion on a categorical variable (see `torch.nn.CrossEntropyLoss` class in the PyTorch library\textsuperscript{54}).

We trained networks with $N = 248$ hidden units using stochastic gradient descent with a batch size of 124 sequences and gradient clipping (gradient norm clipped to be less than or equal to 2). At the beginning of training, we trained RNNs on sequence lengths of $T = 1$ and increased the sequence length by one every 50 parameter updates. We performed 30,000 parameter updates, so that by the end of training the RNNs were training on sequence lengths of $T = 600$. We found that this gradual increase in task complexity along with gradient clipping were necessary to achieve good performance. Intuitively, training on short sequences at the beginning helps the network learn suitable parameter values for $\{C, \alpha, D, \gamma\}$ without worrying about the typical challenges (e.g. exploding and vanishing gradients) associated with RNN training. Then, the remaining parameters $\{A, B, \beta\}$ can be fine-tuned with gradually increasing sequence lengths.

Each sequence was randomly generated. For each spatial dimension, the initial angular position, $\theta_0$, was sampled uniformly between $[0, 2\pi)$. The angular velocity at each time step was given by $\Delta \theta_t = \bar{\theta} + \epsilon_t$ where $\bar{\theta}$ denotes the mean velocity and $\epsilon_t$ was sampled randomly from a normal distribution with mean zero and standard deviation of 0.3 radians. For each sequence the mean velocity, $\bar{\theta}$, was sampled from a normal distribution with mean zero and standard deviation of 0.1 radians. The initial latent state was chosen randomly from the available states. State change cues occurred randomly according to a homogeneous Poisson process with an expected rate of one state change per 50 time steps. In Figure 5, we trained networks to switch between three or more states—for each state change one of the inactive states was chosen uniformly at random to be the new active state. State changes were cued by a pulse lasting two time steps.

For Figure 1G, the trained model was provided velocity inputs with an initial position of $\theta_0 = 0$ and non-negative angular velocity at each time step, $\Delta \theta_t = abs(\bar{\theta} + \epsilon_t)$, so that the RNN output would follow a trial structure comparable to the biological data. Similarly, state changes occurred less frequently (at an expected rate of once per 500 time steps) to better match the biologically observed remapping rate. For comparison with the biological data, we truncated each sequence to remove incomplete track traversals and concatenated 50 sequences into a single session. For visualization purposes, we computed the smoothed, position-binned (n bins = 50) firing rates for 5 example units and labeled each track traversal according to the most commonly reported latent state for that traversal.

**Mice**

All experimental data reported here were collected for a previous publication, Low et al.\textsuperscript{26} and were approved by the Institutional Animal Care and Use Committee at Stanford University School of Medicine. More information on data collection and analyses can be found in Method Details, below, and in the Methods section of Low et al.\textsuperscript{26}
Method Details

Manifold Geometry Analyses

We used Procrustes shape analysis\textsuperscript{55} according to the methods described in Low et al.\textsuperscript{26} to determine the degree to which manifolds from different maps were aligned in the high-dimensional activity space. Briefly, we divided the track into 50 position bins and computed the average activity for all units within each position bin for each latent state to obtain an estimate of the manifold associated with each map. We then mean-centered these manifolds and rescaled them to have unit norm. We compute the root-mean-squared error (RMSE) between these two manifolds (the “observed” RMSE). We then find the rotation matrix that optimally aligns the two manifolds and calculate the RMSE between the optimally aligned manifolds. We report the observed RMSE relative to the RMSE after optimal (misalignment = 0) and random (misalignment = 1) rotation. For Figure 5 and Supplemental Figure 3, which had more than two latent states, we computed this score for all pairs of manifolds.

In Figures 1, 3, 4, and 5 we consider the network in terms of subspaces tuned to the two task components—position and latent state. We define the “position subspace” as a two-dimensional subspace containing the position-binned average firing rates of all units. We divided the track into 250 position bins and computed the average activity for all units within each position bin for each latent state. To find the position subspace across maps—as in Figure 1K, Figure 3E, Figure 4G, and Figure 5G—we performed 2-factor Principal Components Analysis (PCA) on the position-binned activity across both latent states. To find the position subspace for a single map—as in Figure 3G—we performed 2-factor PCA on the average activity from just one latent state. We define the “remapping dimension” as the dimension separating the manifold centroids, which we find by computing the average activity for each unit within each map and taking the difference across the two maps.

Fixed Point Analysis

We numerically identified fixed points according to the methods described in Sussillo & Barak.\textsuperscript{31} Briefly, we used stochastic gradient descent to minimize $||x -ReLU(\mathbf{A}x + \mathbf{b})||_2$ over hidden layer activation vectors $x$. Values of $x$ that minimize the expression close to zero, correspond to approximate fixed points of the recurrent RNN dynamics when the input is held constant $u_t = \mathbf{0}$. At each numerical fixed point $x_*$, we can use standard autodifferentiation tools in PyTorch to compute the $N \times N$ Jacobian matrix $\frac{\partial x_{t+1}}{\partial x_t}$ evaluated at $x = x_*$. The eigenvalues and eigenvectors of this matrix then provide a local linear dynamical approximation to the full system as explained in Sussillo & Barak\textsuperscript{31} and in the main text.

Single Unit Analysis

To characterize single unit remapping properties for Supplemental Figure 1, we performed the rate remapping versus global remapping analysis described in Low et al.\textsuperscript{26} For each model, we computed the average firing rate for all units in each map, smoothing with a Gaussian filter
(standard deviation, 2 position bins). We then calculated the percent change in peak firing rate (i.e., rate remapping). To compute a spatial dissimilarity score (i.e., global remapping), we subtracted 1 from the cosine similarity between firing rate vectors (dissimilarity score of 0 indicates identical spatial firing, 1 indicates orthogonal spatial representations).

**Experimental Data**

The experimental data included in Figure 1, Figure 6, and Supplemental Figure 4 were collected for a previous publication. Briefly, mice were trained to navigate a 1D virtual reality track with tower landmarks and floor cues to provide optic flow. The landmarks repeated seamlessly every 400 cm such that the track was circular-linear. The mice received randomly distributed, visually cued rewards within the middle 300 cm of the track. During behavior, neural activity was recorded using Neuropixels 1.0 silicon recording probes, which were acutely inserted into the medial entorhinal cortex. Behavioral and neural data were processed as described in Low et al. Figure 1A–D are modified from Low et al.

The pilot analyses in Figure 6 and Supplemental Figure 4 are performed on a subset of the data from Low et al. (n = 684 cells from 4 sessions in 2 mice). As described in that publication, we used k-means clustering to divide these sessions into 3 or 4 maps. We then assessed the trial-by-trial spatial stability of the population-wide neural activity within each map in order to restrict our analysis to stable trials. We divided the session according to the k-means map labels and computed the Pearson correlation between the position-binned firing rates (n bins = 80) of all neurons for each pair of trials within each map. We excluded trials that were spatially unstable from our analysis (average correlation with all other trials < 0.25).

To assess the geometry of the neural population activity, we used the k-means cluster centroids as an estimate for the neural activity manifold associated with each map. We then performed Procrustes shape analysis to assess manifold alignment and identified the remapping dimensions, as described above.

To ensure that remapping was not an artifact of probe movement or multi-unit activity, we compared the spike waveforms for all cells across remapping events, as described in Low et al. Briefly, we identified the longest epoch of trials for each map and extracted waveforms for 100 spikes for each cell from each epoch. We then computed the average waveforms within each epoch. To determine waveform similarity, we computed the Pearson correlation between the vectorized average waveforms for each pair of maps and then calculated the average correlation across pairs. For all waveform analyses, we used waveforms from the 20 channels closest to the Kilosort2-identified depth for each cell.

**Quantification and Statistical Analysis**

**Statistics**

All data were analyzed in Python, using the scipy stats library to compute statistics. Unless otherwise noted, all tests are two-sided, correlation coefficients represent Pearson’s correlation, and values are presented as mean ± standard error of the mean (SEM). Statistical tests are
listed following the relevant result given in the Results, figure legend, or Method Details. Unless otherwise stated, $p < 0.05$ was taken as the criterion for significance.
Supplementary Information for: Remapping in a recurrent neural network model of navigation and context inference

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Supplemental Figure 1: single RNN units remap heterogeneously.

(A) Absolute fold change in peak firing rate versus spatial dissimilarity across latent state changes for all units from 2-map RNN models (points, single units; histograms, density distributions for each variable). Median change in peak firing rate (horizontal red dashes) = 1.9-fold; 95th percentile (horizontal gold dashes) = 3-fold. Median spatial dissimilarity (vertical red dashes) = 0.025; 95th percentile (vertical gold dashes) = 0.21.

(B) Modified from Low et al.26 As in (A) but for biological neurons from 2-map sessions. Median change in peak firing rate (horizontal red dashes) = 1.28-fold. Median spatial dissimilarity (vertical red dashes) = 0.031; 95th percentile (vertical gold dashes) = 0.22.
Supplemental Figure 2: RNN geometry is interpretable.

(A) Schematic showing the orthogonalization of the position and context input and output weights.
(B) Reproduced from Figure 1K.
(C-D) Schematic: How a single velocity input (blue arrows) updates the position estimate (yellow to red points) from the starting position (blue points).
(C) Velocity input lies in the position tuning subspace (gray plane) (hypothetical). Note that the same velocity input results in different final positions.
(D) Velocity input is orthogonal to the position tuning subspace (observed).
(E) Schematic of possible flow fields in each of the three planes (numbers correspond to planes in C and D), which would result in the correct positional estimate given orthogonal velocity inputs at different positions (D).
Supplemental Figure 3: Manifold geometry generalizes for up to 10 latent states.
(A) (Left) RNN models were trained to navigate in 1D (velocity signal, top) and discriminate between many (2–10) distinct contexts (transient, binary latent state signal, bottom). (Right) Example showing high prediction performance for position (top) and context (bottom) for 5 latent states.
(B) Position-binned activity for six example single RNN units, split by latent state (colors as in (A)).
(C) Models accurately estimated position (top) and latent state (bottom) for different numbers of latent states or "maps."
(D) Angle between remapping dimensions for different numbers of maps. The remapping dimensions were always maximally separated.
(E) Manifold misalignment score for all pairs of maps across all models (0, perfectly aligned; 1, $p = 0.25$ of shuffle). All pairs of manifolds were more aligned than expected by chance.
Supplemental Figure 4:

**A1** network similarity

**A2** correlations all cells

**A3** neural activity manifolds

**A1** example cell 1

**A2** waveform overlays cell 1

**A3** maps 1 & 2

mean misalignment = 1.0

**A1** example cell 2

**A2** waveform overlays cell 2

**A3** maps 1 & 3

**B1** network similarity

**B2** correlations all cells

**B3** neural activity manifolds

**B1** example cell 1

**B2** waveform overlays cell 1

**B3** maps 1 & 2

mean misalignment = 0.55

**B1** example cell 2

**B2** waveform overlays cell 2

**B3** maps 1 & 3

**B1** example cell 3

**B2** waveform overlays cell 3

**B3** maps 2 & 3


**C1** network similarity

**C2** correlations all cells

**C3** neural activity manifolds

**C1** example cell 1

**C2** waveform overlays cell 1

**C3** maps 1 & 2

mean misalignment = 0.78

**C1** example cell 2

**C2** waveform overlays cell 2

**C3** maps 1 & 3

**C1** example cell 3

**C2** waveform overlays cell 3

**C3** maps 2 & 3


**D1** network similarity

**D2** correlations all cells

**D3** neural activity manifolds

**D1** example cell 1

**D2** waveform overlays cell 1

**D3** maps 1 & 2

mean misalignment = 0.51

**D1** example cell 2

**D2** waveform overlays cell 2

**D3** maps 1 & 3

**D1** example cell 3

**D2** waveform overlays cell 3

**D3** maps 2 & 4

**D1** example cell 4

**D2** waveform overlays cell 4

**D3** maps 3 & 4
Supplemental Figure 4: Medial entorhinal cortex can remap between 3 or 4 maps and many geometric features are preserved.

Experimental data for 4 example multi-map sessions from Low et al.\textsuperscript{26}

(A) Examples from the 3-map Session A ($n = 142$ cells).

(A1)(Left) Network-wide trial-by-trial correlations for the spatial firing patterns of all co-recorded neurons in the same example session (colorbar indicates correlation). (Left/bottom) k-means map assignments. (Right) Three example medial entorhinal cortex neurons switch between three maps of the same track (top, raster; bottom, average firing rate by position; teal, map 1; red, map 2; black, map 3).

(A2) Comparisons of average spike waveforms sampled from each of the three maps (approximate sampling epochs indicated by black lines, A1 far right). (Left) Across map correlations for average spike waveforms from all cells (horizontal bar, median = 0.978; vertical line, 5\textsuperscript{th} – 95\textsuperscript{th} percentile, 5\textsuperscript{th} percentile = 0.935). (Right) Overlay of average waveforms sampled from each map for the three example cells from (A2).

(A3) PCA projection of the manifolds associated with each pair of maps (colorbar indicates virtual track position). Text: average normalized manifold misalignment score for all pairs of maps (0, perfectly aligned; 1, $p = 0.25$ of shuffle).

(B–D) As in (A), but for the 3-map Session B ($n = 184$ cells; median waveform correlation = 0.988), 4-map Session C ($n = 196$ cells; median waveform correlation = 0.995), and 4-map Session D ($n = 162$ cells; median waveform correlation = 0.99).
References


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