
Topological Neural Data Analysis with Behavioral Constraint

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Abstract

1 Recently, Topological Data Analysis (TDA) has revealed insights into the topo-
2 logical structure of neural population activity. However, existing TDA methods
3 for neural population activity are computationally demanding, noise-sensitive,
4 and sometimes difficult to interpret. We develop a simple and more interpretable
5 analysis approach to infer the topological structure of behaviorally relevant neural
6 response variability. Our approach first maps the neural activity onto firing rate
7 maps of behavioral variables, and then performs analysis based on these rate maps.
8 Application of our method to grid cell recordings demonstrates its effectiveness
9 without sophisticated preprocessing as required in prior methods. Further test of
10 the methods based on synthetic data suggests that our method is more informative
11 of the deviations from standard topological shapes. Our results also point to the
12 importance of joint analysis of the geometry and topology of neural manifolds.

13 1 Introduction

14 While classic work in neuroscience emphasized individual neurons, recently there is surge of interest
15 to understand the population-level structure of neural activity [Vyas et al., 2020] by conceptualizing
16 it as a “neural manifold” [Seung and Lee, 2000, Kriegeskorte and Wei, 2021, Perich et al., 2025,
17 Chung and Abbott, 2021]. At each moment, the neural population activity represents a point on the
18 manifold, which is determined by stimulus, internal states, and noise. The geometrical [Kriegeskorte
19 and Wei, 2021] and topological [Giusti et al., 2015] structure of neural manifolds can be informative
20 of the underlying neural computation.

21 Advances in modern recording techniques for simultaneously recording large populations of neurons
22 [Grienberger and Konnerth, 2012, Jun et al., 2017] provide unprecedented opportunity to investi-
23 gate the structures of neural manifolds. Recent studies applied Topological Data Analysis (TDA;
24 [Wasserman, 2017]) to analyze neural data in various neural systems, e.g., V1 [Singh et al., 2008],
25 the hippocampus [Giusti et al., 2015], head direction cells [Chaudhuri et al., 2019], and grid cells
26 [Gardner et al., 2022]. In particular, [Gardner et al., 2022] provided evidence for toroidal structures
27 in the grid cells by evaluating the persistent homology [Zomorodian and Carlsson, 2004]. However,
28 existing methods based on neural population activity are prone to noise and rely on the pre-processing
29 steps, thus complicating the interpretation and the reliability of the result.

30 Here, we introduce a simple method that extracts the topological structures of neural population
31 activity that are relevant for encoding certain behavioral variables. Our approach achieves comparable
32 topology characterization with simpler data curation steps compared to previous methods, while
33 enabling robust interpretation of topological features. We also demonstrate a challenge for Neural
34 TDA: when neural manifolds deviate from standard shapes (e.g., circle/torus), it can be difficult to
35 detect these deviations with TDA. We show that our method better captures these deviations. These

36 findings suggest that future research can benefit from the joint analysis of the geometry and topology
 37 of neural manifolds [Kriegeskorte and Wei, 2021, Ye and Wessel, 2025].

38 2 Methods

39 2.1 Analysis pipeline

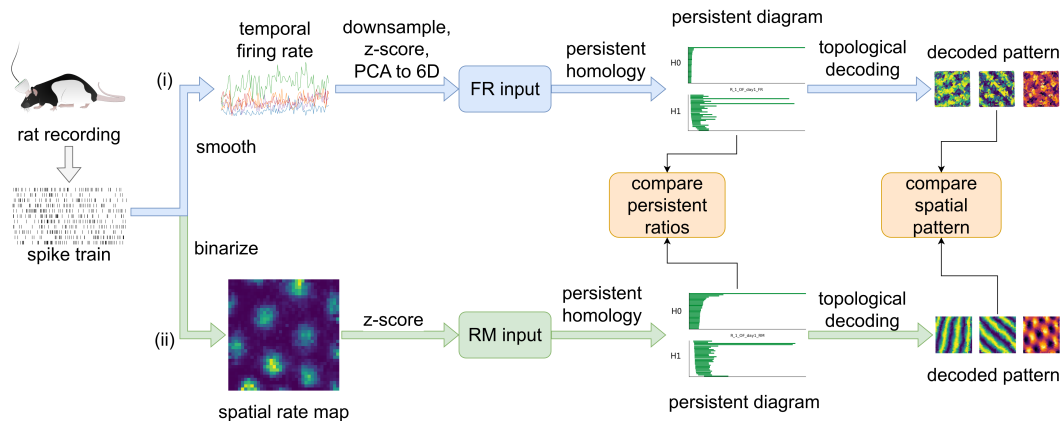


Figure 1: A comparison of the two TDA pipelines.

40 Fig. 1 illustrates the comparison of the previous analysis approach and ours. For the previous
 41 approach (see Fig. 1 (i)) developed by Gardner et al. [2022], first the firing rate of individual neurons
 42 were computed based on the spike trains. Substantially downsampling were needed to select the
 43 most active time bins. After further downsampling, z-scoring, and projecting the data onto the first 6
 44 principal components, the *persistent diagrams* were computed. The *persistent barcodes* in a persistent
 45 diagram indicate topological features of a certain dimension. When applying to grid cell data, two
 46 outstanding long barcodes in the H1 diagram indicated two independent circular features, jointly
 47 composing the torus manifold. Each circular feature was used to assign a circular parameter to all
 48 data points. The dual circular parameters were summed up across neurons to compute two arrays
 49 of spatial firing strengths over time bins, visualizing two circular firing patterns with a 60° angle,
 50 together yielding a hexagonal grid pattern.

51 In our analysis pipeline (Fig. 1 (ii)), we first project the neural activity onto the spatial locations
 52 to yield the firing rate maps of individual neurons. We then compute the persistent homology
 53 using the z-scored firing rate maps. No additional preprocessing steps are needed. By projecting
 54 the neural activity onto behavioral variables, our method enables analyzing topological features of
 55 behavior-relevant neural variability.

56 2.2 Persistence ratio

57 After applying topological data analysis (TDA) to the neural activity, we obtain two persistent
 58 diagrams denoted as H0 and H1 diagrams, each indicating the 0-th and first dimensional topological
 59 features. Intuitively, H0 features indicate disconnected clusters within the data, while H1 topological
 60 features correspond to circles in the data manifold.

61 Since our focus is on neural manifolds with hypothesized toroidal (grid cells) or circular (“circular
 62 cells”) topology, we analyze the longest bars in the H1 diagram to determine whether the data supports
 63 the presence of one or two prominent circles. To quantify this, we first rank the lengths of all the
 64 barcodes in the bottom H1 diagram (see Fig. 1), and then compute the ratio of the i -th and $(i + 1)$ -th
 65 longest barcodes. We will refer to it as the i -th *persistence ratio*, denoted as $PR(i)$.

66 A larger $PR(i)$ indicates stronger significance of the first i dominant circular features relative to other
 67 features, thus indicating the whole manifold closer to a product of i circles in its shape. For instance,
 68 the ratio of lengths between the second and third longest barcodes in the H1 persistent diagram is the
 69 second persistence ratio $PR(2)$, and a manifold with a larger $PR(2)$ has a shape closer to a torus.

70 **2.3 Topological decoding**

71 Once circular features are identified, the next step is to interpret what they represent. The decoding
 72 procedure follows [Gardner et al., 2022].

73 Each circular feature assigns a *cocycle* value to the data points, with which we solve a least-square
 74 problem ($Ax = b$) to obtain an angular parameter for each data point, so that the angular parameters
 75 can span the edges to obtain the distance matrix among cocycles.

76 To connect these circular parameters with spatial behavior, we compute weighted activity maps.
 77 Specifically, we compute a scaled activity value for each spatial bin across all neurons by multiplying
 78 the RM value with the angular parameter. Finally, by summing up across all neurons, we can obtain
 79 spatial coordinates associated with each spatial bin, and visualize them as a circular spatial firing
 80 pattern.

81 In [Gardner et al., 2022], however, the FR input was first downsampled to 1,200 temporal bins. To
 82 extend decoding to the full dataset, they introduced an interpolation step: each neuron was assigned
 83 a weighted factor by summing up the scaled values across downsampled temporal bins, then the
 84 original firing rates were scaled with the weighted factors, where each factor is shared for all original
 85 activity across one neuron. Then temporal coordinates were computed by summing up across all
 86 neurons, and scattered on a 2D open-field map in correspondence to their x-y positions to visualize a
 87 circular spatial firing pattern.

88 **3 Results**

89 **3.1 Rat grid cell recordings**

90 We analyzed the Neuropixels [Jun et al., 2017] recordings of rat MEC provided in [Gardner et al.,
 91 2022]. We performed TDA analyses on all the open-field (OF) foraging task recordings. See Appendix
 92 B.2 for more information about the dataset.

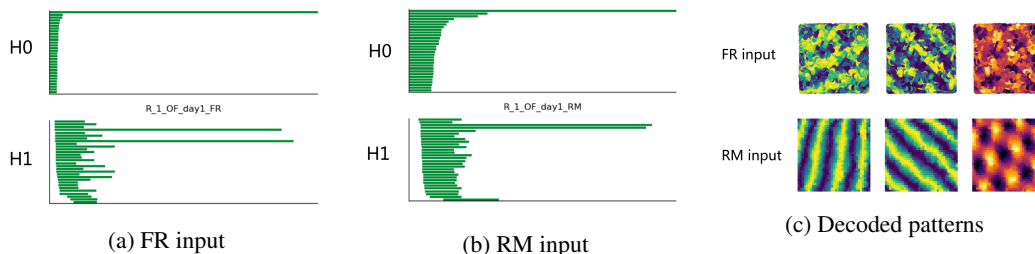


Figure 2: A comparison of two TDA methods with different types of input.

93 We first replicated the results reported in [Gardner et al., 2022] by following their exact analysis
 94 procedure. The firing rate (FR) inputs to TDA computation are arrays of $[1200, N]$ where N is the
 95 number of recorded neurons in one session. For fair comparisons, in our method (RM), we chose a
 96 spatial bin size so the dimensionality of the RM inputs are comparable to theirs.

97 Fig. 2 reports the TDA results from the OF session of rat R, module 1, recording day 1.

98 Fig. 2a shows the persistent diagrams computed with firing rate (FR) input. Two significantly longer
 99 barcodes in the bottom (H1) diagram indicates two dominant circular feature that are irrelevant to
 100 each other, hence a toroidal topological feature.

101 Fig. 2b displays persistent diagrams computed with our rate-map (RM) method.

102 Fig. 2c depicts decoded circular features for FR (top) and RM (bottom) inputs. Two circular patterns
 103 tend to have a 60 degree angle, and form a hexagonal grid together.

104 The results with all OF recording sessions are shown in Fig. 4. We find the mean $PR(2)$ for FR input
 105 is slightly higher than that for the RM input (3.43 v.s. 3.13), suggesting that method based on FR has
 106 more evidence for toroidal topological shapes. For decoded circular patterns, the results from our
 107 proposed RM method are significantly cleaner.

108 We further analyzed the robustness of two methods when removing certain data curation processes.
 109 Results in Fig. 5 suggest that our method, while simple, can robustly recover clear topological
 110 features. In contrast, firing-rate-based method may fail to recover clear features when missing certain
 111 preprocessing steps.

112 3.2 Synthetic heterogeneous 1D “grid cells”

113 While applying TDA on grid cells reveals a toroidal feature, in reality the underlying topology often
 114 deviates from a perfect torus due to the heterogeneity within firing rate tuning. To examine how
 115 well the TDA methods can detect deviations from simple shapes like circle/torus, we conducted an
 116 analysis on simulated data, focusing on populations of 1-dimensional “circular cells” due to their
 117 simplicity and high interpretability.

118 In this simulation (for details see Appendix B.3), we introduce heterogeneity in the response gain of
 119 individual firing fields for each neuron. Specifically, we construct the tuning curves by multiplying a
 120 periodic tuning with a linear function over space. The slope of the linear function (linear decay factor)
 121 controls the heterogeneity of firing. Importantly, when the decay factor is larger than 0, the manifold
 122 is topologically equivalent to a line, but not a circle. As the decay factor increases, the ground-truth
 123 manifold becomes like a spring being stretched by force.

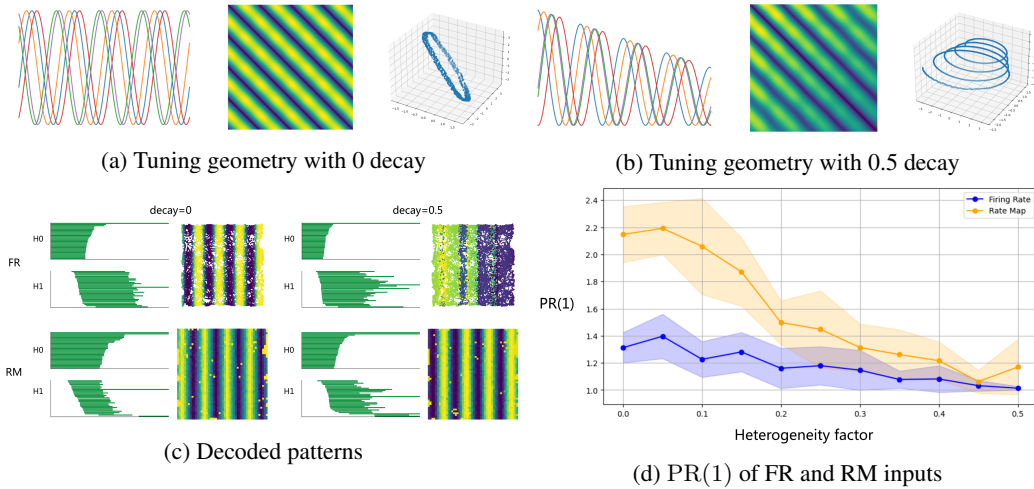


Figure 3: Visualization of heterogeneous tuning amplitude.

124 Fig. 3a and 3b show the representational distance matrix and a 3D visualization using Multi-
 125 dimensional Scaling [Kruskal and Wish, 1978] when the decay factor is 0 and 0.5, respectively.
 126 It is clear that the ground-truth tuning geometry of the perfect simulation looks like a ring, while
 127 noticeably deviates from a ring under heterogeneity.

128 Fig. 3c shows the representative persistent diagrams and decoding results with heterogeneity. The
 129 results suggest that the H1 (bottom) diagram becomes messy with an induced decay factor, and our
 130 RM-based method can better decode the circular pattern. We then systematically analyzed models
 131 with a range of decaying factors from 0 to 0.5 with a 0.05 increment, repeating the computation of
 132 persistent diagrams 10 times for each setting.

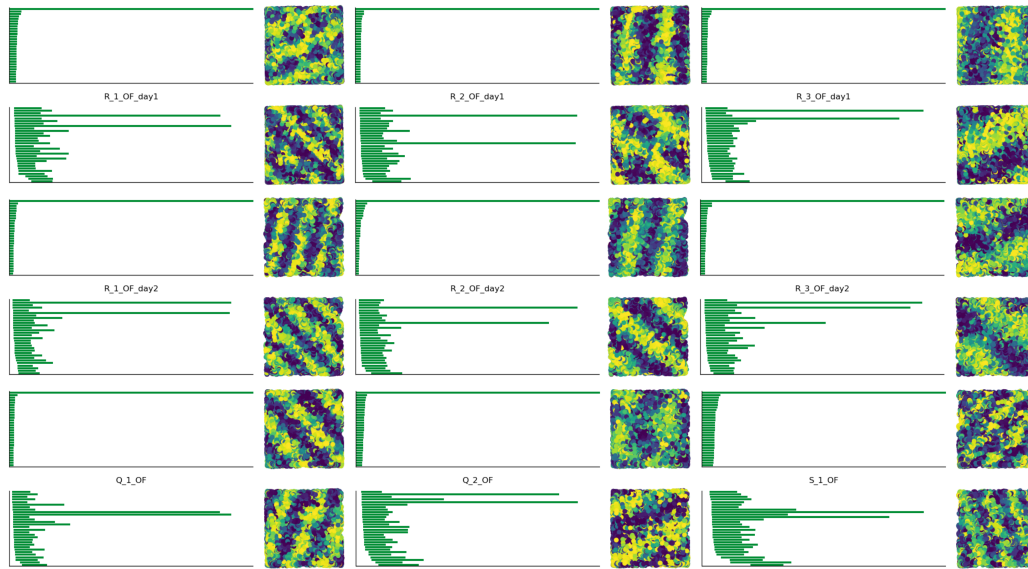
133 Fig. 3d shows the inferred persistence ratios for different decaying factors. We find that the circular
 134 feature becomes less dominant with the decay, in accordance with tuning geometries shown above.
 135 While the persistence ratio inferred from both methods are affected by increasing heterogeneity, our
 136 method is substantially more informative of the change of the decaying factor and thus the deviations
 137 in tuning amplitudes.

138 Putting together, these results suggest that practically it may be challenging for the existing TDA
 139 method to detect mild deviations from standard shapes. Our RM-based method is more informative
 140 for revealing these deviations, at least for the settings we have tested. Integrating TDA with the
 141 analysis of geometry may lead to more powerful approaches.

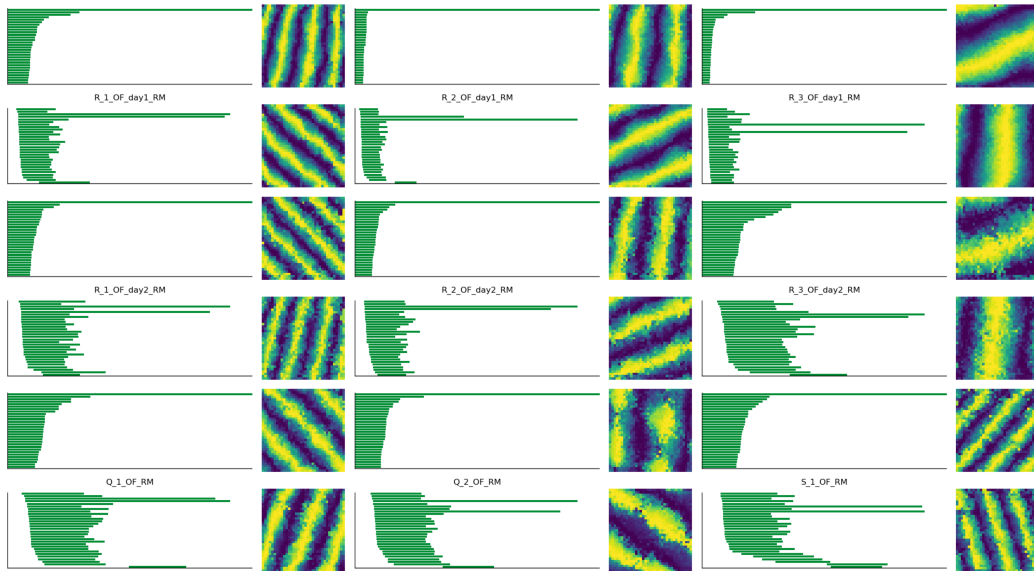
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(a) TDA results with FR input



(b) TDA results with RM input

Figure 4: Computed persistent diagrams of all the open-field (OF) sessions, together with two circular patterns decoded from the two longest barcodes in the bottom (H1) diagram.

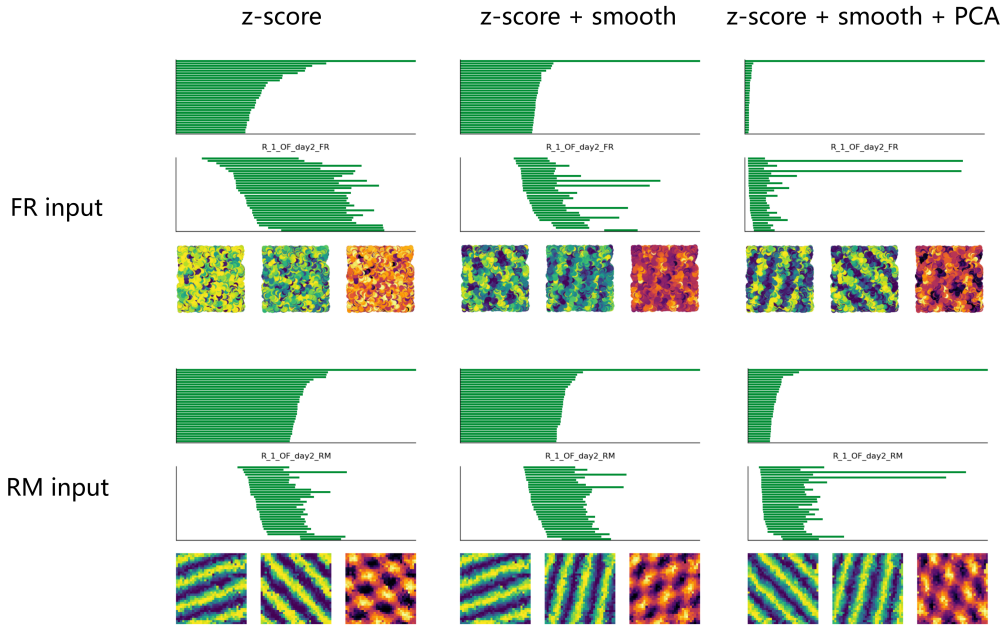


Figure 5: Computed persistent diagrams of rat R, grid cell module 1, open field session in recording day 2; together with two circular patterns decoded from the two longest barcodes in the bottom (H1) diagram, and their combined grid structure. FR inputs are downsampled to 1,200 time points in advance, while RM inputs are set to 1,225 (35×35) spatial bins. Each row: results with firing rate and rate map inputs. Each column: results with different data curation steps.

203 **B Experimental details**

204 **B.1 Code base and computing devices**

205 We use NumPy [Harris et al., 2020] to store, formulate, and compute our data as arrays, and Ripser
206 [Bauer, 2021] to perform all the topological data analysis and topological decoding steps. All the
207 code runs on a CPU with 32GB RAM.

208 **B.2 Grid cell recording dataset**

209 The grid cell recording dataset is provided in [Gardner et al., 2022], and can be avail-
210 able at [https://figshare.com/articles/dataset/Toroidal_topology_of_population_](https://figshare.com/articles/dataset/Toroidal_topology_of_population_activity_in_grid_cells/16764508)
211 [activity_in_grid_cells/16764508](https://figshare.com/articles/dataset/Toroidal_topology_of_population_activity_in_grid_cells/16764508).

212 The dataset consists of recordings of rat Medial Entorhinal Cortex (MEC) with Neuropixels silicon
213 probes [Jun et al., 2017]. In the dataset, there are recordings of 3 animals named "rat Q", "rat R", and
214 "rat S". The grid cells in rat Q are classified as 2 modules, and the grid cells in rat R are classified as
215 3 modules. The recording sessions cross 2 days, and are classified as open-field foraging sessions
216 (OF), wagon-wheel foraging sessions (WW), rapid-eye-movement sleeping sessions (REM), and
217 slow-wave sleeping sessions (SWS). For a detailed introduction of the experimental techniques and
218 procedures, please refer to "Methods" of [Gardner et al., 2022].

219 In [Gardner et al., 2022], the authors retracted toroidal topological features from all of the recording
220 sessions with firing rate based inputs. While our rate map based method requires an input of animal
221 movement positions, so we only performed our method with OF sessions in comparison with their
222 OF results.

223 **B.3 Simulation settings**

224 We performed simulations of circularly tuned "1-dimensional grid cells", or "circular cells".

225 We first constructed circular tuning curves for 100 artificial "circular cells", while each cell has a
226 random shift in its tuning phase, sampled from a uniform distribution of $[0, 2\pi]$. After that, we apply
227 a linear decay on the tuning curves, with a heterogeneity factor set as the lowest peak amplitude.

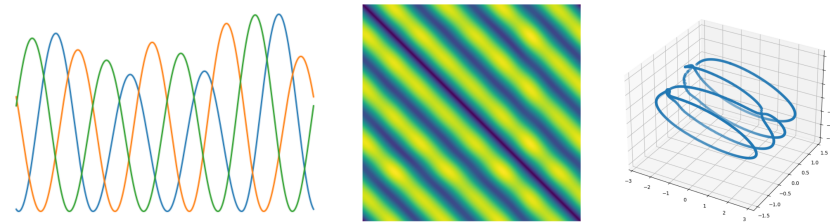
228 We then apply the spatial tuning curves on the real movement trajectory of rat R, OF session, day 1 to
229 obtain a ground-truth firing rate. After that, we apply a Poisson firing model to get spike trains of 100
230 circular cells. We then smooth the spike trains into a $[15000, 100]$ empirical firing rate tensor.

231 The firing rate tensor was z-scored, projected to 6 principal components, and downsampled to shape
232 $[2500, 6]$ as the FR input; and the firing rate tensor was binned against 50×50 positional bins to
233 form a firing rate map, z-scored, projected to 6 principal components, and sent in as a shape $[2500, 6]$
234 RM tensor.

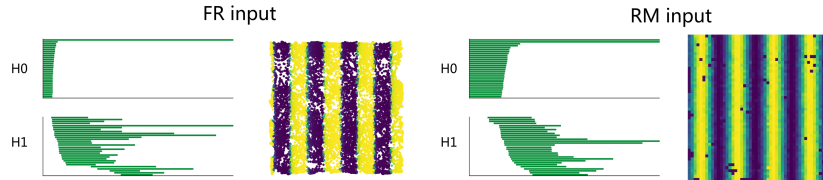
235 **C Alternative simulation experiments**

236 In addition to the “linear decay” simulation scheme described in the main text, we also implemented
 237 an alternative simulation setting. Each artificial “circular cell” retains an intrinsic phase shift sampled
 238 from a uniform distribution over $[0, 2\pi]$. To introduce heterogeneity in tuning amplitude, we assign a
 239 “fluctuation factor” within $[0, 1]$ and sample a scaler for each waveform (between two local minima)
 240 of a circular cell, hereby allowing each waveform to exhibit a random scaling of tuning amplitude.
 241 We argue that this mechanism of tuning heterogeneity better approximates realistic neural variability
 242 compared to the linear decay scheme, although its effects on the topology of tuning curves, firing
 243 rates, and open-field rate maps are less straightforward and need further investigation.

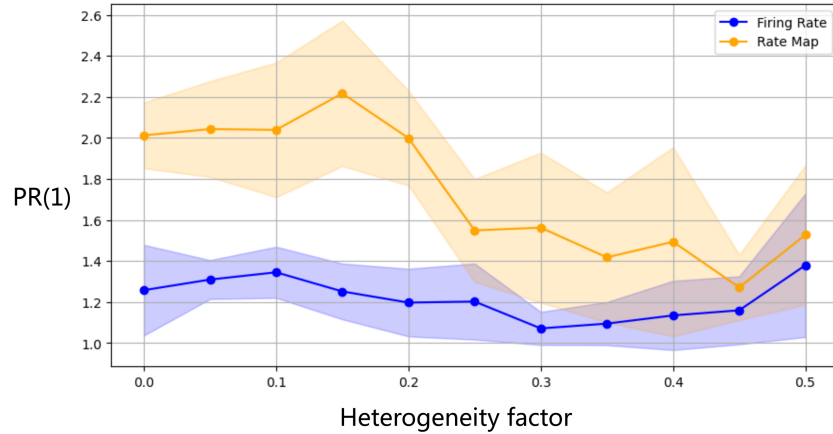
244 Following the simulation, we derived empirical firing rate tensors and converted them into FR and
 245 RM inputs using the same preprocessing steps described in Appendix B.3. We then applied the same
 246 TDA analysis and topological decoding procedure in the main text.



(a) Tuning geometry with fluctuation factor 0.5



(b) Decoded patterns with fluctuation



(c) PR(1) of FR and RM inputs

Figure 6: A demonstration of “fluctuating” tuning curve simulation.

247 As shown in Fig. 6a, with a fluctuation factor of 0.5, the tuning geometry deviates noticeably from a
 248 perfect ring topology. The tuning curves of three representative circular cells illustrate variability
 249 in both phases and amplitudes across neurons. Correspondingly, the distance matrix and MDS
 250 visualization of the tuning curves also deviate from the idealized setting. In Fig. 6b, we present the
 251 persistent diagrams computed with FR and RM inputs. Notably, the RM-based method recovers a one-
 252 dimensional circular spatial pattern with discernible color gradient, whereas the FR-based decoding
 253 result presents a step-like switching pattern. Finally, Fig. 6c plots the average PR(1) values across

254 11 fluctuation factors with 10 repeated runs each. The results indicate that our RM-based method
255 captures the influence of tuning heterogeneity more effectively, demonstrating greater sensitivity to
256 perturbations under behavioral constraints in neural data.