Discovering Precise Temporal Patterns in Large-Scale Neural Recordings through Robust and Interpretable Time Warping

Highlights

- Trial-to-trial variability in spike timing is correlated across co-recorded neurons
- Unsupervised time warping reveals precise spike patterns from neural data alone
- Simple and interpretable warping functions (piecewise linear) are often sufficient
- Spike time precision may be systematically underestimated without warping

Authors
Alex H. Williams, Ben Poole, Niru Maheswaranathan, ..., Bence P. Ólveczky, Krishna V. Shenoy, Surya Ganguli

Correspondence
ahwillia@stanford.edu (A.H.W.), sganguli@stanford.edu (S.G.)

In Brief
The timing of neural dynamics can be highly variable across trials due to uncontrolled behavioral variability or unobserved cognitive states. Williams et al. describe an interpretable statistical model to control for these misalignments and use this approach to uncover fine-scale temporal structure that is imperceptible in raw data.
Discovering Precise Temporal Patterns in Large-Scale Neural Recordings through Robust and Interpretable Time Warping

Alex H. Williams,1,16* Ben Poole,9 Niru Maheswaranathan,8 Ashesh K. Dhawale,10,11 Tucker Fisher,1 Christopher D. Wilson,14 David H. Brann,10,11 Eric M. Trautmann,1,8 Stephen Ryu,4,12 Roman Shusterman,13 Dmitry Rinberg,14,15 Bence P. Olveczky,10,11 Krishna V. Shenoy,3,4,6,7,8 and Surya Ganguli2,3,4,6,7,9,*

1Neuroscience Program, Stanford University, Stanford, CA 94305, USA
2Applied Physics Department, Stanford University, Stanford, CA 94305, USA
3Neurobiology Department, Stanford University, Stanford, CA 94305, USA
4Electrical Engineering Department, Stanford University, Stanford, CA 94305, USA
5Bioengineering Department, Stanford University, Stanford, CA 94305, USA
6Bio-X Program, Stanford University, Stanford, CA 94305, USA
7Wu Tsai Stanford Neurosciences Institute, Stanford University, Stanford, CA 94305, USA
8Howard Hughes Medical Institute, Stanford University, Stanford, CA 94305, USA
9Google Brain, Google Inc., Mountain View, CA 94043, USA
10Department of Organismic and Evolutionary Biology, Harvard University, Cambridge, MA 02138, USA
11Center for Brain Science, Harvard University, Cambridge, MA 02138, USA
12Department of Neurosurgery, Palo Alto Medical Foundation, Palo Alto, CA 94301, USA
13Institute of Neuroscience, University of Oregon, Eugene, OR 97403, USA
14Neuroscience Institute, New York University School of Medicine, New York, NY 10016, USA
15Center for Neural Science, New York University, New York, NY 10016, USA
16Lead Contact
*Correspondence: ahwillia@stanford.edu (A.H.W.), sganguli@stanford.edu (S.G.)
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SUMMARY

Though the temporal precision of neural computation has been studied intensively, a data-driven determination of this precision remains a fundamental challenge. Reproducible spike patterns may be obscured on single trials by uncontrolled temporal variability in behavior and cognition and may not be time locked to measurable signatures in behavior or local field potentials (LFP). To overcome these challenges, we describe a general-purpose time warping framework that reveals precise spike-time patterns in an unsupervised manner, even when these patterns are decoupled from behavior or are temporally stretched across single trials. We demonstrate this method across diverse systems: cued reaching in nonhuman primates, motor sequence production in rats, and olfaction in mice. This approach flexibly uncovers diverse dynamical firing patterns, including pulsatile responses to behavioral events, LFP-aligned oscillatory spiking, and even unanticipated patterns, such as 7 Hz oscillations in rat motor cortex that are not time locked to measured behaviors or LFP.

INTRODUCTION

The role of spike time precision in neural computation has been widely examined from both experimental and theoretical perspectives, engendering intense debates in systems neuroscience over the last several decades (Softky and Koch, 1993; London et al., 2010; Bruno, 2011; Brette, 2015; Denève and Machens, 2016). Assessing the degree of temporal precision in multi-neuronal spike trains is challenging, since variations in behavioral and cognitive variables can mask precise spike time patterns. For example, the temporal precision of olfactory coding may be underestimated by factors of two to four when spike times are aligned to stimulus delivery instead of inhalation onset (Shusterman et al., 2011; Cury and Uchida, 2010; Shusterman et al., 2018).

Thus, experimental estimates of spike time precision hinge on the choice of an alignment point, which defines the origin of the time axis on each trial. This choice can often be challenging and subjective. Even relatively simple tasks often involve a sequence of stimuli, actions, and rewards that occur with varying latencies, thus presenting multiple choices for temporal alignment. Moreover, in addition to choosing an origin of time, we must also choose its units. Should spike times be measured in absolute clock time relative to some measured event or in units of fractional time between two events? Could any one of these choices unmask spike-timing precision that is otherwise invisible? These questions are most challenging to answer in systems far from the sensory or motor periphery, where neural responses may not be
locked to any measurable behavior and instead reflect internal decisions or changes of mind.

Past studies have addressed these challenges in a number of ways: grouping trials together with similar durations before averaging spike counts (Murakami et al., 2014; Starkweather et al., 2017; Wang et al., 2018), manually stretching or compressing time units between measured task events (Leonardo and Fee, 2005; Shusterman et al., 2011; Kobak et al., 2016; Aronov et al., 2017), or repeating statistical analyses around different choices of alignment points (Feierstein et al., 2006; Harvey et al., 2012; Jazayeri and Shadlen, 2015; Shushruth et al., 2018).

These complications, and the diversity of heuristic approaches used to address them, underscore a need for statistical frameworks to assess the temporal precision of neural computation. Of particular interest are unsupervised statistical methods that reveal precise patterns in multi-neuronal spike trains without reference to behavioral measurements. Such methods would be broadly applicable, as they make few restrictive assumptions about the experimental design. Furthermore, by moving beyond pre-conceived alignments designed by human experts, these data-driven methods may discover novel and unexpected results.

While time series and image alignment methods are a well-studied topic in signal processing (Marron et al., 2015; Pnevmatikakis and Giovannucci, 2017), these techniques have rarely been applied to large-scale neural recordings (but see recent work by Poole et al., 2017; Lawlor et al., 2018; Duncker and Sahani, 2018). Neuroscientists have historically utilized simple alignment operations, namely, translating (Baker and Gerstein, 2001; Ventura, 2004) and potentially stretching and/or compressing activity traces between pairs of behavioral events (Shusterman et al., 2011; Leonardo and Fee, 2005; Perez et al., 2013; Kobak et al., 2016). In contrast, popular statistical methods, such as Dynamic Time Warping (DTW) (Berndt and Clifford, 1994), allow signals to be non-uniformly compressed and dilated on each trial. While such nonlinear warping models can be useful, we demonstrate that they can be difficult to interpret and sensitive to the high level of noise that is typical of neural data.

To identify interpretable alignments for high-dimensional spike trains, we developed a framework for linear and piecewise linear time warping that encompasses existing human-annotated procedures (Leonardo and Fee, 2005; Kobak et al., 2016). We applied these methods to multielectrode recordings collected from three experiments spanning different animal models (rodents and primates), brain regions (olfactory and motor cortex), and behavioral tasks (sensation and motor production). In each case, time warping revealed precise spike patterns that were imperceptible in the raw data. Moreover, some of these results were not easily captured by any common behavioral alignment. For example, in rodents performing a motor timing task, we uncovered ~7 Hz oscillations in spike times that were not aligned to the LFP or any measured behavior.

RESULTS

Time Warping Framework

Our ultimate goal is to identify firing patterns that reliably occur on a trial-by-trial basis. If these activity patterns are tightly time locked to a sensory or behavioral event, then we can characterize the neural response by averaging over trials. This is illustrated in Figure 1A (left), which shows 100 trials of simulated neural activity with additive Gaussian noise. The average activity trace (red; bottom) extracts the neural response from noisy single-trial instantiations. This simulated example loosely resembles calcium fluorescence traces, but the methods we describe can be flexibly applied to any multi-dimensional time series, including spike trains, fMRI data, or LFP traces.

More formally, if N neurons are measured at T time points over K trials, the trial average is given by:

\[ \mathbf{X} = \frac{1}{K} \sum_{k=1}^{K} \mathbf{X}_k. \]  

(Equation 1)

where \( \mathbf{X}_k \) is a \( T \times N \) matrix denoting the activity of all neurons on trial \( k \). In the context of spiking data, each column of \( \mathbf{X} \) corresponds to a peri-stimulus time histogram (PSTH) of a recorded neuron. Trial averaging is also a common step in population-level statistical analyses (Kobak et al., 2016).

Despite its widespread use, trial averaging can produce inaccurate and misleading results when neural activity is misaligned across trials. For example, introducing a random shift to each simulated trial produces a less informative result (Figure 1A, right). Such jitter commonly arises in practice, leading many research groups to develop custom-built alignment procedures. For example, in songbirds it is common to manually segment and cluster song syllables and warp intervening spike times on a per-syllable basis (Leonardo and Fee, 2005). In olfaction, detailed measurements and fluid dynamics modeling of the sniff cycle have been pursued to understand the accuracy of sensory responses (Shusterman et al., 2011, 2018).

Time warping methods address these challenges through a data-driven, statistical approach. The key idea is to fit a response template time series that is shifted and stretched—i.e., warped—on a trial-by-trial basis to match the data. The response template, denoted \( \mathbf{X} \), is an \( T \times N \) matrix of activity traces that captures the average activity across trials after correcting for variability in timing.

The time axis of the response template is transformed by a warping function on each trial. Formally, we denote the warping function for trial \( k \) as \( \omega_k(t) \); this function maps each time bin \( t \) (clock time) to a new index \( \omega_k(t) \) (template time). If \( \omega_k(t) \) is an integer between 1 and \( T \), then the warping transformation for every neuron \( n \) on trial \( k \) amounts to the transformation \( X_{t,n} \rightarrow X_{\omega_k(t),n} \). If \( \omega_k(t) \) is not an integer, then time warping is implemented by linear interpolation (see STAR Methods). Note that this model assumes that all recorded neurons share the same warping function on a trial-by-trial basis, though this assumption could be relaxed by future work.

Figure 1B illustrates how different classes of warping functions account for single-trial variability in timing. We focus on three main model classes: shift-only warping, linear warping, and piecewise linear warping (Figure 1B, top three models). Shift-only warping is the simplest model: the warping functions are constrained to be linear with slope equal to one, and only a
single parameter (the y intercept of $u_k(t)$) is fit on each trial. As its name suggests, the shift-only model can only account for trial-to-trial differences in response latency. In contrast, a linear warping model, which fits the slope in addition to the intercept of $u_k(t)$, can account for variable latencies as well as uniform stretches and compressions of the response template. A piecewise linear warping model adds further complexity by adding one or more knots (points where the slope of $u_k(t)$ can change). Most generally, nonlinear warping functions may be used, which non-uniformly stretch and compress portions of the template on each trial (Figure 1B, bottom). In all cases, we constrain the time warping functions to be monotonically increasing. Loosely speaking, this ensures that the model cannot go backward in time while making a prediction. This constraint also implies that the warping functions are invertible, which we later exploit to align spike times across trials.

The model parameters are fit to minimize reconstruction error over all neurons, trials, and time points. For simplicity, we chose the mean squared error to quantify model performance. Ignoring

Figure 1. Illustration of Time Warping Models
(A) Single-neuron synthetic data from 100 trials. When data are aligned (top left), the trial average provides a good description of activity (bottom left). When the data are misaligned by introducing jitter (right), the trial average does not produce a useful description (bottom right).
(B) Time warping models estimate a template (black line, left) that is transformed on a trial-by-trial basis by warping functions (middle column) producing single-trial estimates of neural activity (right column). Colored lines represent warping functions and estimated firing rates on three example trials. Top row illustrates shifting the template activity (shift-only warping), second row illustrates stretching and compressing the template in addition to shifting (linear warping), third row illustrates piecewise linear warping with two-line segments, and the bottom row illustrates a fully nonlinear warping.
(C) Results of shift-only warping (top), linear warping (middle), and nonlinear warping (bottom) (Petitjean et al., 2011) fit to the synthetic data from (A), right. For each model, we show the warping templates (left, black lines) and warping functions (right, lines colored by ground truth shift; red indicates rightward shift, blue indicates leftward shift).
(D) The predicted firing rates of shift-only warping provide a denoised trial-by-trial estimate of neural activity.
(E) The raw data sorted by the learned per-trial shift parameter.
(F) Data aligned by the shift-only model; the per-trial shift is applied in the opposite direction to the raw data.
Figure 2. Recovery of Ground Truth Warping Functions in Synthetic Data

(A) Synthetic spiking data from N = 5 units, T = 150 time bins, and K = 75 trials. Data were simulated from a ground truth model with piecewise linear warping functions with 1 knot.

(B) Data re-aligned by a shift-only warping model.

(C) Data re-aligned by a linear warping model.

(D) Data re-aligned by a piecewise linear (1-knot) warping model.

(E) Data re-aligned by a piecewise linear (2-knots) warping model.

(F) Data re-aligned by the ground truth model. Note similarity with (D) and (E).

(G) Ground truth neural response templates (black) and estimated response templates (red) from the piecewise linear (1-knot) model. y axis denotes the probability of spiking in each time bin.

(H) Ground truth warping functions (black) on six representative trials and estimated warping functions (red) from the piecewise linear (1-knot) model.

(I) Model performance ($R^2$) in training, validation, and test partitions of various warping models. Thin lines represent maximum and minimum values; thick lines represent mean ± standard error. Results were computed over 40 randomized cross-validation runs. The horizontal dashed blue line reflects the performance ceiling on held-out data achieved by the ground-truth model.
the interpolation step of time warping for the sake of clarity (see STAR Methods), the objective function is:

\[
\frac{1}{NTK} \sum_{n=1}^{N} \sum_{t=1}^{T} \sum_{k=1}^{K} \left( \hat{X}_{nk}(t) - X_{nk}(t) \right)^2 \quad \text{(Equation 2)}
\]

This expression is minimized with respect to the warping functions on each trial, \( \omega_k(t) \), and the response template, \( X \). Other loss functions may be substituted for the mean squared error. In particular, a loss function based on Poisson noise is popular in neural modeling (Paninski, 2004), and our accompanying Python package supports this option.

For illustration, we fit shift-only, linear, and nonlinear warping models to the misaligned synthetic data shown in Figure 1A. By design, the shift-only model is sufficient to capture the ground truth variability. As expected, this model identifies a highly accurate template firing pattern (Figure 1C, top left), along with warping functions that tightly correlate with the ground truth delay (Figure 1C, top right). The linear warping model is a gentle extension of the shift-only warping model, which only introduces one additional parameter on each trial: the slope of each warping function. Yet even this minor extension produces a slightly worse estimate due to overfitting (Figure 1C, middle).

To demonstrate a more severe case of overfitting, we fit a nonlinear warping model using Dynamic Time Warping (DTW) (Berndt and Clifford, 1994) combined with a standard barycenter averaging procedure (Petitjean et al., 2011). This method can be highly effective on datasets with low levels of noise and complex temporal deformations. However, as we will soon see, neural datasets often exhibit the opposite: high levels of noise and simple temporal deformations. In this regime, DTW barycenter averaging identifies a noisy template (Figure 1C, bottom left) and unnecessarily complex warping paths (Figure 1C, bottom right).

These results demonstrate that time warping models, particularly with complex warping functions, are susceptible to overfitting. We adopted four strategies to mitigate this possibility. First, as mentioned above, we assume that each trial’s warping function is shared across neurons, thus mitigating the possibility of overfitting to noise in any single neuron. Second, as illustrated by the progression of models in Figure 1C, we always compare the estimates of complex warping models (e.g., with piecewise linear warping functions) to the performance of simpler models (e.g., shift-only warping). Third, we include a roughness penalty on the response template, which directly encourages the model to identify smooth firing rates. Fourth, we place a penalty on the area between each warping function and the identity line, which penalizes the magnitude of warping on each trial. We include these roughness and warp-magnitude penalties in subsequent results, but we show the results of unregularized time warping in Figure 1C for the sake of illustration.

These models enable several strategies for visualizing and understanding neural data. First, one can directly inspect the model parameters (Figure 1C): the response template for each neuron captures the shape of the neural response, while the warping functions capture trial-to-trial variability in timing. Second, one can view the model prediction as a denoised estimate of firing rates on a single-trial basis (Figure 1D). Third, one can re-sort the trials by the slope or the intercept of the warping function, producing a multi-trial raster plot that is easier to visually digest (Figure 1E). Finally, one can invert the warping functions on each trial to transform the raw data into an aligned time domain (Figure 1F). This alignment procedure simply entails plotting each activity trace as a function of \( \omega_k^{-1}(t) \) instead of raw clock time \( t \). Intuitively, this amounts to reversing the flow diagram shown in Figure 1B, which is possible as long as the warping functions are monotonically increasing and thus invertible.

### Extraction of Precise, Ground Truth Spike Patterns on Synthetic Data

Before proceeding to biological data, we examined a more challenging simulated dataset consisting of spike trains from \( N = 5 \) neurons, \( T = 150 \) timebins, and \( K = 75 \) trials. On each trial, the neural firing rates were time warped by randomized piecewise linear functions with one knot (the “ground truth” model). This resulted in spike trains that appear highly variable in their raw form (Figure 2A).

Time warping successfully reveals the spike patterns corresponding to the ground truth process. Figures 2B–2E show model-aligned spike trains (as in Figure 1F) across warping models of increasing complexity. The patterns evident in the ground truth data (Figure 2F) are partially revealed by shift-only and linear time warping (Figures 2B and 2C), but these models are too simple to capture the fine-scale temporal structure in the data. A piecewise linear warping model with one knot (piecewise-1 model; Figure 2D) accurately captures these details and represents a parsimonious and “correct” model, since it matches the data generation process. Furthermore, the parameters of this model closely matched the ground truth response template (Figure 2G) and warping functions (Figure 2H). Using a piecewise linear model with 2 knots did not result in substantial overfitting and indeed closely matched the result of the piecewise-1 model (Figure 2E).

Identifying a parsimonious warping model is challenging in real-world applications where there is no observable ground truth. To select the appropriate model and regularization strengths, we developed a nested cross-validation scheme, which correctly identified the piecewise-1 model as having optimal performance on a held-out test set (Figure 2I). This approach fits the neural response templates to a subset of trials (Figure 2J, shaded blue regions) and fits the warping functions to a subset of the neurons (Figure 2J, shaded red regions). We refer to the intersection of these two sets as the training set (Figure 2J, shaded purple regions), while the intersection of the complement sets is used for validation and testing (Figure 2J, unshaded...
regions). This procedure is then repeated many times with different randomized partitions of the data.

We used the proportion of variance explained (\(R^2\)) to quantify model performance. Unlike in trial-averaged analyses, \(R^2\) values will typically be very low because spike trains exhibit substantial single-trial variability. In the case of this synthetic dataset in Figure 2, the ground truth model achieves an \(R^2\) of approximately 0.131 (blue dashed line, Figure 2I). This \(R^2\) value, while low due to unexplainable trial-to-trial spiking variability, nevertheless constitutes an upper bound on the \(R^2\) value achievable by any model on a held-out test set. The piecewise-1 model closely matches this performance, achieving an average \(R^2\) of 0.113 on the test set (Figure 2I). However, in real experimental data, it is not possible to know how close we are to this performance ceiling. Thus, while \(R^2\) can be useful as a relative measure when comparing two different models, the absolute magnitude should be interpreted cautiously.

After using cross-validation to select the number of warping function knots and the regularization parameters, we are often interested in using time warping to visualize single neuron raster plots (as in Figures 2A–2F). To further validate these raster plots, we can hold out single neurons, fit the model to the remaining data, and apply the inverse warping functions to align the held-out neuron (Figure 2K). All model-aligned raster plots shown in subsequent results are generated using this procedure.

**Alignment of Olfactory Responses to Sniff Cycle**

Mitral/tufted cells in the mouse olfactory bulb exhibit variable firing patterns across trials when naively aligned to odor delivery (Figure 3A). If spike times are instead aligned to the onset of inhalation, neural responses are drastically more reliable (Shusterman et al., 2011).

We reasoned that simple time warping models could be used to align mitral/tufted cell activity using purely neural activity, bypassing the need to measure inhalation directly. We tested this hypothesis on a multielectrode recording from \(N = 30\) neurons over \(K = 45\) trials of odor presentation at a fixed concentration (\(x\)-pinene, \(10^{-2}\) dilution from standard vapor pressure). We experimentally measured intra-nasal pressure to detect sniff onset and offset, but these measurements were not provided to the model and spike times were instead aligned to odor presentation. As expected, this initial alignment strategy produced highly disordered spike rasters (Figure 3B).

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**Figure 3. Time Warping of Mitral/Tufted Cell Activity Recovers Sniff-Locked Activity Patterns**

(A) A head-fixed mouse sampled odorized air (\(x\)-pinene). Variability in inhalation onset from trial to trial caused jitter in the olfactory response. (B) Spike raster plots for six representative cells over all \(K = 45\) trials with spike times aligned to odor delivery. Black dots denote spike times, and blue dots denote sniff onsets. Blue histogram indicates the distribution of sniff onsets. (C) Same as (B) but with trials re-ordered by the magnitude of shift warping. (D) Same as (B) but with the time axis aligned by shift-only warping. (E) Same as (B) but with the time axis aligned to sniff onset. (F) Trial-to-trial reliability (\(R^2\)) for all \(N = 30\) cells before (x axis) and after (y axis) alignment by the shift-only model. Dashed gray lines indicate a 2-fold increase or decrease in \(R^2\). The p value is computed using a Wilcoxon signed rank test. (G) Same as (F) but comparing the alignment of linear warping to shift-only warping. (H) Same as (F) but comparing the alignment of shift-only warping to sniff onset alignment.
We found that a shift-only time warping model captured precise sensory responses from these raw data, as revealed by re-sorting the trials based on the model’s shifts (Figure 3C) or by applying these shifts to align the raw spike times (Figure 3D). Here, as well as in all subsequent results, we use the leave-one-out validation procedure shown in Figure 2K; thus, any temporal structure seen in Figure 3D is unlikely to arise as an artifact of overfitting.

As expected, aligning spike times to inhalation onset revealed similar patterns in these data (Figure 3E). Indeed, the shifts learned by the model correlated very tightly with the onset of sniffing (see blue dots and histograms, Figures 3B–3E). These results are nonetheless a useful demonstration, since the model inferred these precise responses from the neural data directly. Furthermore, closer examination suggested that the unsupervised, shift-only model may enjoy slight performance advantages relative to the simple supervised alignment method. For example, when aligned to sniff onset, cells 4 and 5 in Figure 3 exhibit subtle, but perceptible, jitter in their responses (Figure 3E), and this variability is visibly corrected by time warping (compare to Figure 3D).

To explore these effects on a neuron-by-neuron basis, we quantified trial-to-trial variability before and after time warping by computing the coefficient of determination ($R^2$) of the neuron’s PSTH. Larger $R^2$ values imply that the trial-averaged PSTH better approximates single-trial firing rates. We fit time warping models while holding out neurons one at a time and computed $R^2$ on the held-out neuron before and after warping. Relative to the raw spike times, shift-only time warping improved $R^2$ in nearly all neurons, with many increasing over 2-fold (Figure 3F; average 107% increase in $R^2$, geometric mean; Wilcoxon signed rank test, $p < 10^{-4}$, $n = 30$). Linear warping did not produce consistent improvements at the level of single neurons (Figure 3G; average 6.6% decrease in $R^2$, geometric mean; Wilcoxon signed rank test, $p = 0.24$, $n = 30$). Relative to sniff onset alignment, shift-only time warping improved the $R^2$ criterion mildly (Figure 3H; average 11% increase in $R^2$, geometric mean; Wilcoxon signed rank test, $p = 0.15$, $n = 30$). Overall, we conclude that timing variability in these olfactory responses is well described by a per-trial shift. Further, the optimal shift correlates tightly with, but may not coincide exactly with, sniff onset (Shusterman et al., 2018).

**Figure 4. Time Warping of Reach Dynamics in a Nonhuman Primate**

(A) Reaches toward a 90° target were analyzed.

(B) Spike data from four example multiunits over all trials. Units 1 and 4 were from primary motor cortex; units 2 and 3 were from premotor cortex. At the bottom, the temporal distribution of task events is shown: 100 ms after target onset (TARG, green), end of delay period (GO, blue), and movement onset (MOVE, red).

(C) Scatterplot showing correlation between the hand movement onset on each trial and the learned shift parameter in the shift-only warping model.

(D) Same as (B) but with trials re-ordered by the magnitude of shift warping.

(E) Same as (B) but with the time axis aligned by shift-only warping.

(F) Same as (B) but with the time axis aligned by linear warping.

(G) Same as (B) but with the time axis aligned by piecewise linear warping with one knot.

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80, or 120 cm from the center. We observed similar results on other reach angles (Figure S1), as well as when data was pooled across all reach angles (data not shown). Multunit activity was collected from \( N = 191 \) electrodes across two Utah multielectrode arrays placed in primary motor (M1) and premotor (PMd) cortices (see STAR Methods).

The most dramatic changes in neural firing rates are closely time locked to movement (Kaufman et al., 2016). Thus, it is common to track hand position on a moment-by-moment basis and use these measurements to align spike times to the onset of movement or the peak hand velocity on each trial. We instead examined spike trains aligned to the beginning of the delay period (Figure 4B) and used time warping to infer an alignment without any reference to the animal’s behavior.

As expected, a shift-only warping model closely aligned spike times with the onset of movement. The model’s learned shift parameter on each trial correlated very tightly with movement onset (Figure 4C), achieving a comparable level of performance (\( R^2 = 0.9 \)) to what was recently reported for a complex, nonlinear warping method (Duncker and Sahani, 2018). Furthermore, the shift-only warping model enabled the visualization of movement-related firing rate changes in single-neuron rasters, either by re-sorting the trial order of the raw data (Figure 4D) or by realigning the spike times (Figure 4E).

Thus, learning a single per-trial shift was sufficient to align neural spike times to movement without any reference to hand-tracking data. However, shifting spike times in this manner also destroyed other structure in the data. Namely, a subset of multiunits, mostly in PMd, showed increased firing around \( \sim 100 \) ms into the delay period, i.e., shortly after the reach target was visually presented to the animal (see units 2 and 3 in Figure 4). Due to the variable delay between target onset and movement onset, a shift-only warping model is incapable of simultaneously aligning spikes across these two events.

A linear time warping model more appropriately captures this structure in the data. On each trial, the model utilizes its two free parameters—the slope and intercept of the warping function—to precisely align these two task events (Figure 4F). Thus, these results provide strong evidence, via an unsupervised time warping method, that reliable neural dynamics occur around the time of movement onset and shortly after target onset. Piecewise linear warping functions did not provide substantial benefits over linear warping; however, linear warping provided a reproducible benefit over the shift-only warping model in cross-validated performance (Figure S1).

**Detection of \(~13–40\) Hz Spike-Time Oscillations in Primate Premotor Cortex**

Thus far, we have shown that the temporal alignments learned by simple warping models can closely correlate with behaviors (e.g., movement or sniffing) and sensory cues (e.g., reach target presentation). This agreement demonstrates that time warping models can converge to reasonable and human-interpretable solutions and, conversely, suggests that established alignment practices in these systems are well justified from a statistical perspective. We now show that time warping can also uncover more subtle and unexpected features in spike train data.

LFP signals in primate premotor cortex show oscillations in the beta frequency range (13–40 Hz) during movement preparation, which are correlated with spike timing (Murthy and Fetz, 1992; Sanes and Donoghue, 1993; Reimer and Hatsopoulos, 2010). While recent work has investigated the relationships between LFP and behavior (Khanna and Carmena, 2015, 2017; Chandrasekaran et al., 2019), the impact of beta oscillations on population-level spiking activity is still poorly understood. Recent work used a complex, black box model of neural dynamics to detect oscillatory structure in high-dimensional spike trains (Pandarinath et al., 2018). Here, we show that shift-only or linear time warping models can recover similar oscillations.

We examined premotor cortical data collected from two different monkey subjects (Monkey J and Monkey U) performing point-to-point reaches; one animal performed these reaches with an unrestrained hand, while the other used a manipulandum (see STAR Methods). Since the oscillations are strongest during the pre-movement delay period, we focused on a time window beginning 400 ms prior to and 100 ms after go cue presentation. We found that having a larger number of trials was beneficial, so we pooled trials from all reach angles for this analysis. We analyzed multiunit data for each monkey from \( N = 96 \) electrodes placed in PMd.

No oscillations were visible in spike rasters aligned to go cue (Figure 5A; data from Monkey U). However, re-aligning these spike trains based on a shift-only warping model revealed oscillations at \(~18\) Hz in Monkey U (shown in Figure 5B) and at \(~40\) Hz in Monkey J (Figure S2); these results are within previously reported frequency ranges (Murthy and Fetz, 1992; Sanes and Donoghue, 1993). In Monkey U, these oscillations were more apparent after linear warping (Figure 5C), suggesting that the frequency (in addition to the phase) of the oscillations may be variable. Further, these oscillations were roughly in-phase across multiunits; as a result, averaging spike counts across all multiunits and trials after time warping produced cleaner estimates (Figures 5A–5C, bottom).

To quantify these effects across all multiunits, we compared the PSTHs computed from raw data (blue traces, Figure 5A) to PSTHs computed from data aligned by linear time warping (blue traces, Figure 5C). We used Fourier analysis to estimate the amplitude and phase of the oscillation at 18 Hz and found that time warping increased the strength of the oscillation by 1–2 orders of magnitude in most multiunits (Figure 5D). Furthermore, in the raw PSTHs, the oscillation phases were widely spread across multiunits, consistent with there being no detectable oscillations above background noise (Figure 5E, gray dots); in the aligned PSTHs, the phases were tightly clustered, reflecting that oscillations were detectable and in-phase across nearly all multiunits (Figure 5E, purple dots).

We confirmed that the spike-level oscillations were in-phase with LFP oscillations in Monkey U. To do this, we applied the time warping models fit on spike train data to align bandpass-filtered LFP signals (10–30 Hz). The LFP signal was misaligned across trials in raw data but was accurately aligned by the spike-level time warping models (Figure S3), suggesting that the two signals are coherently time warped (in this case, temporally shifted and/or stretched) on a trial-by-trial basis. On a methodological level, this demonstrates that time warping models...
can generalize and make accurate predictions about other time series (e.g., LFP) with qualitatively distinct statistics from the training data (e.g., spike times).

We wondered whether time warping would fail to recover these oscillations if the movement-related spiking, which occurs at a much higher firing rate than pre-movement activity, was included in the analysis. To examine this, we fit warping models to a larger time window (± 400 ms around go cue presentation). Time warping was still able to extract oscillations under these more challenging circumstances (Figure 5F). Interestingly, a shift-only model was no longer sufficient to capture oscillatory activity, suggesting that the oscillations were not phase locked to movement onset. In contrast, linear or piecewise linear warping functions were able to recover the oscillations (Figure 5F, bottom). Thus, while the shift-only model is simplest to interpret, it may be insufficient to capture certain results. This demonstrates the utility of exploring a range of time warping models with varying levels of complexity on the same dataset.

Detection of ∼6–7 Hz Oscillations in Rat Motor Cortex

We have seen that time warping can reveal interpretable structure, even under well-controlled experimental conditions. Discrete reaching, for example, is arguably the simplest volitional motor behavior that one can study, and yet straightforward behavioral alignments obscure salient spike-time oscillations (Figure 5). To study a more complex behavior, in a different animal model, we analyzed motor cortical activity in rats trained to produce a timed motor sequence (Kawai et al., 2015; Dhawale et al., 2017). Rats were trained to press a lever twice with a target time interval of 780 ms and were rewarded if the sequence was completed within ± 80 ms of this target (Figure 6A). While rats produce stereotyped motor sequences in this setting, the

Figure 5. Time Warping Reveals Spike-Level Oscillations in Primate Premotor Cortex

(A) Multiunit activity aligned to go cue. Four representative multiunits are shown; spike rasters (black) trial-average PSTH (blue) are shown for each multiunit. Go cue onset is shown in red. The average firing rate across all multiunits is shown on the bottom (black trace). Shaded gray region denotes upper and lower quartiles.

(B) Same as (A) but with spike times aligned by shift-only warping model. Each displayed multiunit was held out from the model fit.

(C) Same as (A) but with spike times aligned by linear warping model. Each displayed multiunit was held out from the model fit.

(D) Oscillatory power at 18 Hz in trial-average multiunit activity aligned to go cue (black dots) and after linear warping (purple dots). Each dot represents one of \( N = 96 \) multiunits.

(E) Same as (D) but showing oscillation phase for each multiunit.

(F) Multiunit activity averaged across electrodes and trials in a larger time window (800 ms) around go cue (red arrow). Oscillations are not visible in the raw data (top) or after shift warping (second from top). Oscillations are recovered by either linear (second from bottom) or piecewise linear warping (bottom). Shaded gray region denotes upper and lower quartiles.
duration between lever presses and the timing of intermediate motor actions is variable from trial to trial. We examined a dataset consisting of $N = 30$ neurons and $K = 1,265$ trials (3 recording sessions, collected over a 2-day period). In these sessions, the interval between lever presses ranged from 521 to 976 ms (5th and 95th percentiles) across trials.

This experiment has three obvious alignment procedures: align spike times to the first lever press, align spike times to the second lever press, or linearly stretch and/or compress the spike times to align both lever presses across trials (i.e., human-supervised time warping). Figures 6B–6D show the activity of six example neurons under these alignment strategies. At a high level, these raster plots demonstrate that neurons preferentially respond to different behavioral events within a trial. For example, cell 1 in Figure 6 fires after the second lever press, while cell 6 in Figure 6 fires after the first press. Thus, it is not obvious which alignment is preferable, and indeed different insights may be gained from analyzing each.

Unsupervised time warping revealed structure in the data that is hidden in all three behavioral alignments. A shift-only warping model uncovered strong oscillations in many neurons, as visualized either by re-sorting trials based on the learned shift (Figure 6E, same alignment as Figure 6B), or by using the model to re-align spike times (Figure 6F). These findings are not due to spurious alignments produced by an overfit model; each spike raster in Figure 6F was generated on held-out data as diagrammed in Figure 2K. Linear and piecewise linear time warping recovered qualitatively similar oscillations (data not shown) and were not superior to the shift-only model in terms of cross-validated performance (Figure S4).

These results reveal a partial decoupling of behavioral events (lever presses) with neural firing patterns. After alignment, both the first and second lever presses occur at variable times within each trial (Figure 6F, histograms at bottom), and the learned shift on each trial only loosely correlated with inter-press interval (Figure 6G). Furthermore, while multiple neurons exhibited spike oscillations, these oscillations were out-of-phase across neurons and therefore not visible in single-trial raster plots (Figure S4). Taken together, these features of the data suggest that it would be difficult to discover this oscillatory structure by...
manual alignment, demonstrating the power of unsupervised time warping. While the uncovered oscillations are not phase locked with lever press times, they are nonetheless correlated with certain aspects of the animal’s behavior. In particular, some cells only exhibit the ~6–7 Hz oscillation following the first lever press with remarkable temporal precision (see cells 3 and 6 in Figure 6F). Indeed, multiple cells exhibit non-oscillatory firing prior to the first lever press but rapidly switch to an oscillatory behavior following the lever press (see cell 3 in Figure 6F and Figure S5). Other cells exhibit oscillations prior to the first lever press, but the amplitude and precision of the oscillations appear to improve following the first lever press (see cell 4 in Figure 5F). Still, other cells either do not exhibit oscillations (cell 1 in Figure 6F) or exhibit strong oscillations both prior to and following the first lever press (cell 5 in Figure 6F). Time warping enables us to discover and visualize this full spectrum of functional cell types, which are otherwise difficult to detect and characterize. The presence of oscillations in single neurons can be confirmed by plotting the distribution of inter-spike-intervals (Figure S5); the shift-only model goes beyond this method by demonstrating that a large population of neurons are coherently phase shifted on a trial-by-trial basis and by enabling characterization of the full population dynamics and behavioral events in an aligned temporal space.

We then examined whether these spike-level oscillations were aligned with oscillations in LFP. The average frequency spectrum of the LFP did display a prominent peak at ~6–7 Hz, a very similar frequency range to the spike-level oscillations identified in Figure 6. To characterize the relationship between these two oscillatory signals, we band-pass filtered the LFP between 5 and 9 Hz on each trial and fit a separate shift-only time warping model to the LFP traces. The time warping functions learned on LFP data did not uncover the spike-time oscillations shown in Figure 6, and, likewise, the LFP signals were not well aligned by the time warping functions fit to spike times (Figure S6). Thus, unlike the oscillations identified in primate premotor cortex, the oscillations in rat motor cortex were not aligned with the LFP.

**DISCUSSION**

While the temporal precision of neural coding has been a matter of intense debate, few studies have leveraged statistical alignment methods to investigate this problem. Earlier work incorporated time warping into single neuron encoding and decoding models (Aldworth et al., 2005; Gollisch, 2006; Smith and Paninski, 2013; Lawlor et al., 2018), as well as dimensionality reduction methods (Poole et al., 2017; Duncker and Sahani, 2018). Here, we decoupled time warping from these other modeling objectives to achieve a flexible and simplified framework. We surveyed a broader range of datasets than past work, spanning multiple model organisms, brain areas, and sensory or motor tasks. In all cases, we found that the simplest and most interpretable models—often those with shift-only or linear warping functions—matched the performance of more complex models while uncovering striking and sometimes unanticipated dynamics.

Our results suggest that shift-only and linear warping models can match or even outperform more complex nonlinear warping methods. These simpler models have three attractive properties. First, they manipulate model estimates of single-trial firing rates in a more interpretable manner (see Figure 1), enabling exploratory data analysis and visualization. Second, we developed efficient and scalable optimization methods for this class of models. On a modern laptop, these models can typically be fit to data from 1,000 neurons, 100 time points, and 1,000 trials in 1 min or less. This scalability is of great practical importance given the increasing scale of neural recordings (Stevenson and Kording, 2011) and the growing need for rigorous cross-validation and model comparison methods (Chandrasekaran et al., 2018), which are often computationally intensive, if not prohibitive. Indeed, another contribution of this work is the development of cross-validation procedures to compare the performance of time warping models ranging from simple (shift-only) to more complex (piecewise linear; see Figures 2I–2K). In essence, this enables a data-driven determination of the questions we posed in the introduction of this manuscript: should spike times be reported in absolute clock time, relative time between two behavioral events, or something else? Though the answer to this question will depend on the particulars of the experimental context, time warping may be used to rigorously compare these possibilities on a case-by-case basis.

Time warping also uncovered firing patterns that were not aligned to any stimulus or measured behavior. For example, we observed ~13–40Hz spike time oscillations in primate premotor cortex during movement preparation (see Figure 5), which we then verified were phase aligned with LFP (see Figure S3). Similar non-phase-locked oscillations have been previously observed in functional neuroimaging experiments (Makeig et al., 2002; David et al., 2006). Notably, the time warping models we used did not assume any oscillatory structure in the data and thus provide strong evidence that spike-level oscillations are a salient feature of the dynamics. When an oscillation is uncovered, a linear time warping model considers the activity of the full neural population to estimate the changes in the phase (y intercept) and frequency (slope) of the oscillation on a trial-by-trial basis. This population-level approach can be contrasted with popular frequency-domain statistical measures like coherence, which measures the degree of phase synchronization between two spike trains or between a single spike train and LFP (Fries et al., 1997). These pairwise analysis methods are likely more sensitive to noise in single neurons than the population-level models we investigated here; future work could explore the relative advantages and disadvantages of these approaches and develop new methods that combine desirable elements from both.

Oscillatory spike patterns may not always be synchronized to LFP, as we observed in rat motor cortex (see Figure 6). These ~7 Hz oscillations were sometimes gated by a motor action—specifically, the first lever press—thus suggesting a potential relevance to timed motor production (Figure S5). Another possibility is that orofacial behaviors such as whisking and licking are the primary driver of these oscillations (Hill et al., 2011). Other work has shown that persistent ~7 Hz LFP oscillations may be locked to the respiration cycle (Tort et al., 2018); however, the transient, spike-level oscillations we observed were decoupled from LFP and are thus potentially distinct from this phenomenon.
Regardless of their root cause, this result demonstrates the ability of time warping to extract unexpected features of scientific interest from high-dimensional spike trains. Further work is needed to confirm and fully elucidate the properties of these oscillations.

Time warping is only one form of variability exhibited by single-trial neural dynamics. For simplicity, we examined time warping in the absence of other modeling assumptions, such as trial-to-trial variation in amplitude (Bollimunta et al., 2007; Williams et al., 2018) or condition-specific changes in dynamics (Duncker and Sahani, 2018). Nonetheless, the models described here could be extended in several ways to capture more complex single-trial dynamics. First, we made the restrictive assumption that all neurons share the same time warping function on each trial. Under more complex experimental conditions—e.g., in large-scale recordings of multiple brain regions—it may be fruitful to fit multiple time warping functions associated with different neural sub-populations. Second, we fit only a single response template for each neuron. Relaxing this assumption may be useful in situations where neurons exhibit more than one canonical response pattern. Finally, we assumed that neural data were collected over repeated trials. To accommodate more unstructured experimental designs, future work could incorporate time warping into state space models (Macke et al., 2015) or sequence extraction algorithms (Mackevicius et al., 2019). Despite these exciting prospects for future statistical methodology, our work demonstrates that even a simple time warping framework can provide a rich and practical set of tools for the modern neuroscientist.

While our results already show that averaging over short, stereotyped trials can obscure fine temporal oscillations and firing events, these shortcomings are undoubtedly more severe in behaviors that have longer temporal extents and exhibit more variability. Thus, we expect time warping methods to play an increasingly crucial role in neural data analysis as the field moves to study more complex and unstructured animal behaviors (Krakauer et al., 2017). Furthermore, in complex experimental tasks involving large numbers of conditions and exploratory behaviors, the same motor act or sensory percept may present itself only a small number of times. In this trial-limited regime, precise data alignment may be critical to achieve the necessary statistical power to make scientific claims. We expect simple models, such as linear and piecewise linear warping, to perform best on these emerging datasets due to their interpretability, computational efficiency, and robustness to overfitting.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.neuron.2019.10.020.

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**AUTHOR CONTRIBUTIONS**


**DECLARATION OF INTERESTS**

K.V.S. is a consultant to CTRL-Labs and Neuralink Inc. and on the Scientific Advisory Boards of Cognescent and Heal; these entities in no way influenced or supported this work.

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**REFERENCES**


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KEY RESOURCES TABLE

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LEAD CONTACT AND MATERIALS AVAILABILITY

Further requests for resources should be directed to and will be fulfilled by the Lead Contact, Alex H. Williams (ahwillia@stanford.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mouse Olfactory Task
Experimental subjects were male C57B/6 mice (Jackson Laboratories). All procedures were approved by the Institutional Animal Care and Use Committee of New York University Langone Medical Center.

Primate Motor Task
Experimental subjects were two male rhesus macaque monkeys (Macaca mulatta), denoted monkey J and monkey U. The monkeys were 13 (J) and 7 (U) years old and weighed 16 kg (J) and 13 kg (U) at the time of these experiments. All procedures and experiments were reviewed and approved by the Stanford University Institutional Animal Care and Use Committee.

Rat Motor Task
Experimental subjects were female Long Evans rats, 3-8 months old at the start of the experiment (Charles River). All procedures and experiments were reviewed and approved by the Harvard Institutional Animal Care and Use Committee.

METHOD DETAILS

Mathematical Notation
We follow the same notation introduced in the main text. Matrices are denoted in bold, uppercase fonts, e.g., M, while vectors are denoted in bold, lowercase fonts, e.g., v. Unless otherwise specified, non-boldface letters specify scalar quantities, e.g., S or s. We use $M^T$ and $M^{-1}$ to denote the transpose and inverse of a matrix, respectively.
We consider a dataset consisting of $N$ features over $K$ trials with $T$ timesteps per trial. For simplicity, we refer to $N$ as the number of neurons in the dataset; however, $N$ could also refer to the number of fMRI voxels, multiunits, or regions of interest in imaging data. The full dataset is a third-order tensor (a three-dimensional data array) with dimensions $K \times T \times N$. The $k^{th}$ slice of the data tensor is a $T \times N$ matrix $X_k$, which represents the activity of the neural population on trial $k$. We denote a single element of the tensor as $X_{k,t,n}$, which specifies the activity of neuron $n$ at timebin $t$ on trial $k$.

The time warping model produces an estimate of population activity on each trial. Mirroring standard notation in linear regression, we denote the model estimate on trial $k$ as $\hat{X}_k$ (a $T \times N$ matrix).

**Model Estimate and Template Interpolation Scheme**

The main idea behind time warping is to approximate each trial, $X_k$, as a warped version of a $T \times N$ template, $\bar{X}$, that is shared across all trials. For neuron $n$, at time bin $t$, on trial $k$, the spirit behind the model is:

$$\hat{X}_{k,t,n} = \hat{X}_{\omega_k(t),n}$$  \hspace{1cm} \text{(Equation 3)}

However, this expression is only valid when the warping function, $\omega_k(t)$, produces integer values. To allow the warping functions to produce non-integer values, we adopt a standard linear interpolation scheme. Let $\omega_k(t)$ denote the time warping function for trial $k$. The inputs to the warping function are integer-valued time indices $t$ (clock time), and the outputs are any real number (aligned time). We assume that $\omega_k(t)$ is monotonically increasing and thus invertible. Then, the model estimate for neuron $n$, at time bin $t$, on trial $k$ is given by:

$$\hat{X}_{k,t,n} = (\lfloor \omega_k(t) \rfloor - \omega_k(t)) \hat{X}_{\lfloor \omega_k(t) \rfloor,n} + (\omega_k(t) - \lfloor \omega_k(t) \rfloor) \hat{X}_{\lfloor \omega_k(t) \rfloor + 1,n}$$  \hspace{1cm} \text{(Equation 4)}

where $\lfloor \cdot \rfloor$ represents the “flooring” operation, and $\lceil \cdot \rceil$ represents the “ceiling” operation. Because the model estimate (Equation 4) is a linear combination of $\hat{X}_{\lfloor \omega_k(t) \rfloor,n}$ and $\hat{X}_{\lfloor \omega_k(t) \rfloor + 1,n}$, the warping transformation can be represented as a matrix $W$ with nonzero elements given by:

$$W_{\lfloor \omega_k(t) \rfloor} = \omega_k(t) - \lfloor \omega_k(t) \rfloor$$
$$W_{\lfloor \omega_k(t) \rfloor + 1} = \lceil \omega_k(t) \rceil - \omega_k(t)$$  \hspace{1cm} \text{(Equation 5)}

For each trial, the warping matrix $W_k$ can be uniquely determined from the warping function $\omega_k$. Thus, the model estimate on each trial is given by:

$$\hat{X}_k = W_k \bar{X}$$  \hspace{1cm} \text{(Equation 6)}

**Optimization Strategy**

The model template and warping functions are optimized to minimize an objective function, which we denote as $F(\bar{X}, \omega_1, \omega_2, \ldots, \omega_K)$. We assume that this objective function decomposes across trials as follows:

$$F(\bar{X}, \omega_1, \omega_2, \ldots, \omega_K) = \sum_{k=1}^{K} f_k(\bar{X}, \omega_k) + \rho_1(\bar{X})$$  \hspace{1cm} \text{(Equation 7)}

Here $f_k$ is a function defining the model loss on trial $k$, and $\rho_1$ is a regularization term, penalizing the roughness and size of the template (described in the next section). Our online code package supports least-squares and Poisson loss functions; we adopted the least-squares criterion for the purposes of this paper due to its computational efficiency and its ability to be adapted to non-spike time data (e.g., fMRI or calcium imaging). Under this choice, the per-trial loss function is:

$$f_k(\bar{X}, \omega_k) = \|W_k \bar{X} - X_k\|_F^2 + \rho_2(\omega_k)$$  \hspace{1cm} \text{(Equation 8)}

Here, $\rho_2$ is a regularization term that penalizes the magnitude of warping (described in the next section), and $\| \cdot \|_F^2$ denotes the squared Frobenius norm, which is simply the sum of squared residuals, $\|M\|_F^2 = \sum y_i^2$.

To minimize $F$, we adopt an alternating optimization (block coordinate descent) approach (Wright 2015). First, each warping function is initialized to be the identity, $\omega_k(t) = t$, and the template and warping functions are cyclically updated according to the following sequence of optimization subproblems:

$$\bar{X} \leftarrow \arg \min_{\bar{X}} F(\bar{X}, \omega_1, \ldots, \omega_K)$$
$$\omega_1 \leftarrow \arg \min_{\omega_1} F(\bar{X}, \omega_1, \ldots, \omega_K)$$
$$\vdots$$
$$\omega_K \leftarrow \arg \min_{\omega_K} F(\bar{X}, \omega_1, \ldots, \omega_K)$$  \hspace{1cm} \text{(Equation 9)}

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Here, an underlined variable denotes a dummy variable that is optimized over in each subproblem. This sequence of parameter updates is cyclically repeated until the objective value ceases to improve; by construction, the objective monotonically decreases at each step of the algorithm so convergence is guaranteed under mild assumptions (Wright 2015).

Partitioning the parameter updates in this manner is useful because each subproblem can be solved very efficiently. When the template is considered a fixed variable, the objective function decouples across trials (Equation 8), which simplifies the warping function updates considerably:

$$\omega_k \leftarrow \arg \min_{\omega_k} f_k (\mathbf{X}, \omega_k) \quad \text{(Equation 10)}$$

These parameter updates are entirely independent, with each update only depending on the raw data for trial k, \(\mathbf{X}_k\), and the current warping template \(\mathbf{X}\). Our code package executes them efficiently in parallel across CPU threads. Furthermore, each warping function is controlled by a small number of parameters in our framework—at best a single parameter (shift-only warping) and at worst only a few parameters (piecewise linear warping). Thus, we perform these updates by a brute force random search (see Warping Function Regularization and Update Rule).

The response template is also very simple to update, especially under a least-squares loss criterion. Assume for the moment that the model is not regularized; i.e., \(\rho_1 (\mathbf{X}) = 0\) and \(\rho_2 (\mathbf{W}_k) = 0\). Then, because each \(\mathbf{W}_k\) is held constant, updating the template amounts to a least-squares problem that can be solved in closed form:

$$\begin{align*}
\arg \min_{\mathbf{X}} \sum_{k=1}^{K} \| \mathbf{W}_k \mathbf{X} - \mathbf{X}_k \|^2_F &= \arg \min_{\mathbf{X}} \left\| \left( \sum_{k} \mathbf{W}^T_k \mathbf{W}_k \right) \mathbf{X} - \left( \sum_{k} \mathbf{W}^T_k \mathbf{W}_k \right) \mathbf{X}_k \right\|^2_F \\
&= \left( \sum_{k} \mathbf{W}^T_k \mathbf{W}_k \right)^{-1} \sum_{k} \mathbf{W}^T_k \mathbf{W}_k \mathbf{X}_k
\end{align*} \quad \text{(Equation 11)}$$

Furthermore the matrix \(\sum_{k} \mathbf{W}_k^T \mathbf{W}_k\) is a symmetric, tridiagonal matrix. Intuitively, this tridiagonal structure arises from the constraint that each warping function is monotonically increasing, and the local structure of the linear interpolation scheme: for any warping matrix \(\mathbf{W}\), with an associated warping function \(\omega\), Equation 4 implies that \(\mathbf{W}_i \mathbf{W}_j = 0\) if \(|\omega(t) - i| > 1\) or if \(|\omega(t) - j| > 1\) and thus \(\mathbf{W}^T \mathbf{W}_{ij} = 0\) if \(i - j > 1\).

The tridiagonal structure of \(\sum_{k} \mathbf{W}_k^T \mathbf{W}_k\) enables the template parameters to be updated extraordinarily fast for practical applications. We use a specialized solver for systems of linear equations with banded, symmetric structure (scipy.linalg.solve_banded). This allows the optimization problem in Equation 12 to be solved with \(O(TN)\) operations. If \(\sum_{k} \mathbf{W}_k^T \mathbf{W}_k\) were naively treated as a dense matrix, the solution would instead require \(O(T^3 + T^2 N)\) operations.

**Template Regularization and Update Rule**

We found that introducing regularization (penalties on the magnitude or complexity of model parameters) can improve the interpretability of the model and its ability to predict held out data. First, we found in some datasets that the warping template could exhibit rapid, high-frequency changes in firing rate (see, e.g., the template in Figure 1C, bottom panel, which was fit without regularization). These irregularities likely correspond to the model overfitting to noisy neuronal data, and can be discouraged by penalizing the parameter updates as well. The penalty on second derivative of the response template is:

$$\rho_1 (\mathbf{X}) = \lambda \|D \mathbf{X} \|_F^2 + \gamma \|\mathbf{X}\|_F^2 \quad \text{(Equation 12)}$$

where \(\lambda > 0\) controls the strength of the roughness penalty and \(\gamma > 0\) controls the strength of the Frobenius norm penalty. The matrix \(D\) is a \((T-2) \times T\) matrix that computes second-order finite differences:

$$\begin{pmatrix} 1 & -2 & 1 & 0 & \cdots & 0 \\ 0 & 1 & -2 & 1 & \cdots & \vdots \\ \vdots & & & \ddots & & \vdots \\ 0 & \cdots & 0 & 1 & -2 & 1 \end{pmatrix} \quad \text{(Equation 13)}$$
Incorporating this regularization term into the update of the warping template (Equation (11)), we get:

\[
\arg \min_{X} \left\| \sum_{k} W_{k} W_{k} X - \left( \sum_{k} W_{k} \right) X \right\|_{F}^{2} + \lambda \| DX \|_{F}^{2} + \gamma \| X \|_{F}^{2}
\]

(Equation 14)

\[
= \arg \min_{X} \left\| \left( \sum_{k} W_{k} + \lambda D^{T} D + \gamma I \right) X - \left( \sum_{k} W_{k} \right) X \right\|_{F}^{2}
\]

Which yields the template update rule:

\[
X \leftarrow \left( \sum_{k} W_{k} + \lambda D^{T} D + \gamma I \right) \left( \sum_{k} W_{k} \right) X \]

(Equation 15)

Thus, the solution is the same as before except a term \( \lambda D^{T} D + \gamma I \) is added to the inverted matrix (left-hand side of linear system). These modifications hardly affect the computational complexity of the parameter update since \( \lambda D^{T} D + \gamma I \) is also a symmetric, banded matrix. Furthermore, as long as \( \gamma > 0 \) the overall matrix is positive definite and therefore guaranteed to be invertible.

In practice, we have found that it is simple to hand-tune the regularization strengths for exploratory analysis (though cross-validation procedures, described below, should always be used to monitor for overfitting). We typically set the L2 regularization (\( \gamma \)) to be zero or very small (e.g., \( 1e-4 \)) and do not tune it further. A reasonable value for the roughness penalty scale can be found by visually inspecting the template for various neurons (columns of \( X \)) and increasing \( \lambda \) if these time series appear noisy.

**Warping Function Regularization and Update Rule**

We found that the optimization landscape of linear and piecewise linear warping functions is complex and full of local minima. Thus, gradient-based optimization methods can be ineffective. Thankfully, the warping functions are (a) low-dimensional and (b) entirely decoupled across trials. Thus, when updating the warping functions, we perform a brute force parameter search for each trial in parallel. For shift-only warping models, we perform a dense grid search over the parameter (the magnitude of the shift).

For piecewise linear warping models we perform an annealed random search as follows. Consider a warping function \( \hat{w}_{k}(t) \) for any arbitrary trial. We parameterize the warping function as:

\[
\hat{w}_{k}(t) = 1 + (T - 1) \cdot \left[ \hat{w}_{k} \left( \frac{t - 1}{T - 1} \right) \right] \]

where \( \hat{w}_{k} \) is a piecewise linear function mapping the unit interval \([0, 1]\) to any real number, and \(|z| = \max(\min(z, 1), 0)\) denotes clipping any real number \( z \) to have a value between zero and one.

The piecewise linear function \( \hat{w}_{k}(t) \) intuitively defines the “unclipped” warping function for trial \( k \). We define these functions by a series of \( M \times k \) y coordinates, \( \{ (\alpha_{1}, \beta_{1}), (\alpha_{2}, \beta_{2}), \ldots (\alpha_{M}, \beta_{M}) \} \), where \( 0 = \alpha_{1} < \alpha_{2} < \ldots < \alpha_{M} = 1 \) and \( \beta_{1} \leq \beta_{2} \leq \ldots \leq \beta_{M} \). We refer to these coordinates as the knots of the warping function.

\[
\hat{w}_{k}(t) = \begin{cases} 
\beta_{1} & \text{if } t = \alpha_{1} = 0 \\
\beta_{1} \left( 1 - \frac{t - \alpha_{1}}{\alpha_{2} - \alpha_{1}} \right) + \beta_{2} \left( \frac{t - \alpha_{1}}{\alpha_{2} - \alpha_{1}} \right) & \text{if } \alpha_{1} < t < \alpha_{2} \\
\vdots & \\
\beta_{M-1} \left( 1 - \frac{t - \alpha_{M-1}}{\alpha_{M} - \alpha_{M-1}} \right) + \beta_{M} \left( \frac{t - \alpha_{M-1}}{\alpha_{M} - \alpha_{M-1}} \right) & \text{if } \alpha_{M-1} < t < \alpha_{M} \\
\beta_{M} & \text{if } t = \alpha_{M} = 1
\end{cases}
\]

(Equation 17)

Note that the warping function for each trial has a different set of knots; however, we have suppressed the trial index variable \( k \) on the knot coordinates \( (\alpha_{k} \text{ and } \beta_{k}) \) for notational brevity.

To optimize the warping functions we perform a random search over these coordinates/knots. Let \( \alpha = [\alpha_{1}, \alpha_{2}, \ldots, \alpha_{M}] \) and \( \beta = [\beta_{1}, \beta_{2}, \ldots, \beta_{M}] \) denote the current coordinates for any particular trial. We form a new proposed warping function by:

\[
\alpha' \leftarrow \text{sort}(\alpha + \eta Q) \\
\alpha' \leftarrow \left( \alpha' - \alpha_{i}' \right) / (\alpha_{M}' - \alpha_{i}') \\
\beta' \leftarrow \text{sort}(\beta + \eta Q)
\]

(Equation 18)

where \( Q > 0 \) is a scalar parameter tuning the amount of exploration, and \( \eta \) is a vector of random normal variables with mean zero and unit variance. The procedure “sort(\( \eta \))” re-orders the elements of a vector so that they are in ascending order. If the proposed warping function improves the objective function, we accept the new parameters.
Synthetic data examples

In Figure 1 data from a single neuron was simulated as a difference of two exponential curves. The activity at $T = 100$ equally spaced time points between $[-8, +8]$ was given by:

$$x_k(t) = \begin{cases} 
0 & \text{for } s_k - t < 0 \\
3.3 \cdot \exp((s_k - t)/2) - \exp((s_k - t) + \eta) & \text{for } s_k - t \geq 0
\end{cases}$$

where $s_k$ was a random shift parameter drawn uniformly on the interval $[-5.5, 3.5]$, and $\eta$ was randomly drawn zero-mean Gaussian noise with a standard deviation of 0.15. Unregularized shift-only, linear, and piecewise linear (with 1 knot) models were fit to $K = 100$ simulated trials. DTW-Barycenter Averaging (DBA; Petitjean et al., 2011) was fit to the same data using an open-source package called tsclearn (Tavenard, 2017).

In Figure 2 we simulated random warping functions following the procedure listed in Equation 18, with $Q = .12$. The firing rate template of each neuron was given by a smoothed, sparse sequence of heavy-tailed random variables:

$$x(t) = 0.01 + \text{conv} (b_t \cdot e_t)$$

where $e_t$ were randomly drawn from an exponential distribution (with scale parameter equal to one) and $b_t$ were binary random variables drawn from a Bernoulli distribution (with probability of 0.92 that $b_t = 0$). The conv(·) procedure denotes convolution with a Gaussian smoothing kernel with a standard deviation of 2. Truncated Poisson random variables were then drawn in each timebin; any bins with more than two spikes were truncated to one spike.

Mouse Olfactory Task

We analyzed data from a single recording session collected as part of a previously published study (Wilson et al., 2017). Subjects were implanted with a RIVETS headbar for head-fixation 7 days prior to the experiment (see description in Arneodo et al., 2018). Subjects were water deprived prior to the experiment and were administered water during random odor presentations to acclimate animals to the experimental apparatus.

On the day of experiment, subjects were anesthetized using isoflurane and a ~0.3 mm craniotomy was performed to gain access to the dorsal olfactory bulb. NeuroNexus A2x32 probes were inserted approximately 500 μm into the dorsal bulb to record from the mitral-tufted cell layer. After probe insertion, subjects were allowed to recover from anesthesia for 30 min prior to recording. Electrophysiological and respiration signals were recorded using the HHMI Janelia Whisper recording system at 25000 Hz. Respiration (sniff) was monitored non-invasively using a pressure sensor sampling from the airflow in front of the nose. Action potentials from the recording were identified and classified into units offline using Spyking Circus template-matching software (Yger et al., 2018).

Two odors at 3 concentrations were presented in randomly interleaved trials. Subjects were passively sampling odor during trials. Concentrations covered a range of 2 orders of magnitude of molarity in carrier air. Odorants were diluted in mineral oil, stored in amber volatile organic analysis vials, and delivered via a 8-odor olfactometer. Odorant concentrations were controlled using a combination of gas- and liquid-phase dilution. We restricted our analysis to subsets of trials with odorant concentrations of $10^{-2}$ dilution from standard vapor pressure—the highest concentration analyzed in (Wilson et al., 2017).

Primate Motor Task

Monkeys performed a standard center-out delayed reach task described previously in (Gilja et al., 2012; Ames et al., 2014). Targets were presented at (40, 80, 120) cm and 90 cm from the central starting location for monkeys J and U respectively. For
monkey J, delay periods were evenly distributed between 300 and 700 ms (monkey J) with ~4.5% non-delay trials randomly interleaved. For monkey U, delays were randomly distributed between 350 and 600 ms on 87% trials, between 5 and 350 ms on 10% of trials, with 3% non-delay trials. Monkeys received a liquid reward upon touching and holding the cursor on the target. Movement during the delay period caused a trial failure and provided a brief automated timeout (~1 s). Non-delay trials were not analyzed.

For monkey J, the virtual cursor and targets were presented in a three-dimensional environment (MusculoSkeletal Modeling Software, Medical Device Development Facility, University of Southern California). Hand-position data were measured at 60 Hz with an infrared reflective bead-tracking system (Polaris, Northern Digital). Behavioral control and neural decode were run on separate PCs using the Simulink/xPC platform (Mathworks) with communication latencies of less than 3ms. This system enabled millisecond timing precision for all computations. Visual presentation was provided via two LCD monitors in a Wheatstone stereotax configuration, with refresh rates at 120 Hz, yielding frame updates of 7 ± 4 ms. Two mirrors visually fused the displays into a single three-dimensional percept for the user, as described previously in (Gilja et al., 2012).

For monkey U, the virtual cursor and targets were presented on a standard 2D display. The monkey controlled the position of an onscreen cursor using a haptic manipulandum which applied no additional forces applied to the arm and was only used for positional cursor control. The haptic device was constrained to move within a 2D vertical workspace and cursor position tracks hand position 1:1 without perceptible lag.

Neural recordings were obtained via implanted 96-electrode Utah Microelectrode arrays (Blackrock Microsystems) using standard neurosurgical techniques. Two arrays were implanted in the left hemisphere of Monkey J, one in dorsal premotor cortex (PMd) and one in primary motor cortex (M1). Three arrays were implanted in the left hemisphere of Monkey U, one in PMd, one in medial M1, and one in lateral M1. For both monkeys, implantation location was estimated visually from local anatomical landmarks.

Neural data were band-pass filtered between 250-7500 Hz, and processed to obtain multiunit ‘threshold crossings’ spikes, defined as any time the signal crosses ~3.5 times RMS voltage. We did not perform spike sorting, and instead grouped together the multiunits present on each electrode. As such, we anticipate that these population recordings contain both single and multiunit activity.

For Figure 4 and Figure S2 (Monkey J), each trial was defined as the 1200 ms following the reach target onset. Spike times were binned in 5 ms increments. For Figure 5 (Monkey U) and Figure S3 (Monkey J), we aligned spike times to the go cue instead of target onset. To highlight oscillatory spiking activity, we defined each trial as the period occurring 400 ms prior to go cue and 100 ms after go cue. Spike times were binned in 2.5 ms increments for Monkey J and 5 ms increments for Monkey U; similar results were found for smaller bin sizes, and stronger smoothness regularization. In Figure 5F (Monkey U), we extended each trial duration to ± 400 ms around the go cue, but otherwise kept the same parameters.

Tuning the regularization strength of on template smoothness (λ) and warp magnitude (μ) was important to uncover the oscillations in premotor cortex. We used the cross-validation procedure described above to determine roughly appropriate values for these parameters. To be conservative and to be confident that these results were not due to overfitting, we increased the regularization strength further for the purposes of visualization.

**Rat Motor Task**

We analyzed data that was collected as part of a previously published study (Dhawale et al., 2017), which describes all experimental procedures and data collection protocols in greater detail. Extracellular recordings were obtained from 16 chronically implanted tetrodes in the motor cortex. Signals were amplified and digitized on a customized head-stage, and sampled at 30 kHz. The head stage was attached to a custom-designed tethering system that allowed the animal to move freely within its cage. Before implantation, an automated behavioral training framework (described in Poddar et al., 2013) was used to train the rats on a timed lever-pressing task (described in Kawai et al., 2015) until asymptotic performance was achieved.

The tetrode drive was then surgically implanted and targeted to motor cortex, through a 4-5 mm diameter craniotomy made 2 mm anterior and 3 mm lateral to bregma. The tetrode array was lowered to a target depth of 1.85 mm. At the end of the experiments, the position of the electrodes was verified by standard histological methods—brains were fixed via transcardial perfusion (4% paraformaldehyde in phosphate-buffered saline, Electron Microscopy Sciences) and the location of the electrodes was reconstructed by viewing mounted coronal sections (60 mm).

After 7 days of recovery post-surgery, training on the task resumed in the animal’s home cage. Neural and behavioral data was recorded continuously during this time (12-16 weeks) with only brief interruptions (median time of 0.2 h). Spikes were sorted using Fast Automated Spike Tracker (FAST), a custom algorithm designed for parsing long-term continuous neural recordings (for details, see Dhawale et al., 2017). We examined $K = 1265$ trials, collected over a two day period.

Each trial was defined as the period starting 500 ms prior to the first lever press and 1500 ms after the first lever press. Spike times were binned in 10 ms increments for each unit. Raw spike counts were provided to the time warping algorithm; however, we observed similar results under various normalization schemes, such as soft-normalization (Churchland et al., 2012). All analyses of these data used a shift-only time warping model. The per-trial shift was constrained to be less than 10% of the total trial duration.
QUANTIFICATION AND STATISTICAL ANALYSIS

Cross-validation

As with any statistical method, one must be very careful that time warping does not reveal spurious structure and features of the data. In Figure 1, we saw that even a simple linear warping model can result in noticeable overfitting on a simple synthetic time series. An important technical contribution of our work is a rigorous cross-validation framework for time warping models. This framework, described in detail below, enables us to fine-tune all regularization terms—i.e., the hyperparameters \( \{ \gamma, \lambda, \mu \} \)—across all warping models. That is, we can rigorously compare the performance of shift-only, linear, and piecewise linear time warping models on an even footing, and thus critically examine the degree of nonlinearity in time warping. While cross-validation is a common procedure in statistical modeling and in modern neuroscience, there are subtle pitfalls that must be avoided in unsupervised learning models (Bro et al., 2008; Perry 2009), and in models with temporally correlated noise (Opsomer et al., 2001).

To properly compare the performance of different warping models, it is important to perform nested cross-validation, so that regularization terms are separately tuned for each model. For example, a piecewise linear warping model will often require stronger smoothness and warp regularization terms, compared to a simpler, shift-only warping model. Thus, on each cross-validation run we split the data in three partitions: a training set, a validation set, and a test set. For each model class (shift-only, linear warping, piecewise linear warping, etc.) we fit 100 models with randomized regularization strengths to the training set; we then evaluated all 100 models on the held out validation set; finally, the best-performing model was evaluated on the test set. The test set performance is then compared across model classes. Model performance was measured by computing the coefficient of determination \( (R^2) \), averaged across neurons. Formally, for any partition of neurons \( i \subseteq \{1, 2, \ldots, N\} \) and trials \( k \subseteq \{1, 2, \ldots, K\} \) we define the coefficient of determination as:

\[
R^2 = 1 - \frac{\sum_{n \in i} \sum_{t=1}^{T_{kn}} \sum_{k \in k} \left( X_{ktn} - \tilde{X}_{ktn} \right)^2}{\sum_{n \in i} \sum_{t=1}^{T_{kn}} \sum_{k \in k} \left( X_{ktn} - \bar{X}_n \right)^2}
\]

where \( \bar{X}_n = (1/TK) \sum_{t=1}^{T} \sum_{k=1}^{K} X_{ktn} \) is the average activation for neuron \( n \).

Recall that our dataset consists of \( N \) neurons, \( T \) time bins, and \( K \) trials. The question then arises, should we hold out neurons, time points, or trials during cross-validation? Since the warping functions are assumed to be shared across all neurons, these model parameters can be fit on a subset of neurons (training set), and then evaluated on held out neurons (validation/test sets). However, if we hold out individual neurons entirely, then it is impossible to fit the response template matrix \( X \) for those cells. Conversely, the response template can be fit to a subset of trials (training set) and evaluated on the remaining trials (validation/test sets). However, if we hold out individual trials entirely, then it is impossible to fit the warping functions associated with those held out trials.

To circumvent this problem we adopt a bi-cross-validation hold out pattern (Owen and Perry, 2009). This entails separately and independently partitioning neurons and trials. Thus, we randomly choose training neurons (≈73% of all cells), validation neurons (≈13% of all cells), and testing neurons (the remaining ≈13%). Additionally, we randomly choose training trials, validation trials, and testing trials, according to these same ratios. The model warping functions are fit to all trials, but only on the training neurons; the response template is fit for all neurons, but only on the training trials. When reporting the training performance, we compute the reconstruction loss on the intersection of the validation (or test) neurons and trials. Likewise, when evaluating models on the validation (or test) set, we compute the reconstruction loss on the intersection of the validation (or test) neurons and validation (or test) trials.

Temporal dependencies in model errors can complicate proper cross-validation (Opsomer et al. 2001). To avoid these complications, we leave out entire trials for individual neurons, rather than leaving out a subset of time bins.

Null models and other sanity checks

The cross-validation procedure described above is fully rigorous, but computationally expensive to perform. Even if each optimization run only takes a few seconds to complete, repeating the whole process over \( Q \) randomized folds leads to long run times; for example, ≈26 h for \( M = 5, P = 100, Q = 50 \) and each model taking ≈20 s to optimize. This rather unfavorable scaling underscores why our attention to performance enhancements—e.g. by exploiting banded matrix structure when updating the model template—is critical for practical applications. On the other hand, a full cross-validation run is often unnecessary for exploratory data analysis and visualization. Here we briefly outline two simple procedures for validating time warping visualizations in a more interactive manner. Our online code package also supports both of these options.

First, one can create a very simple null dataset of neural activity that, by construction, contains no warping. By comparing the results of time warping on this null dataset to those achieved on the real data, we gain an informative reference point. For spiking data, we simulate null data by computing the trial-average firing rate of each neuron and then drawing Poisson i.i.d. random spike trains on every trial. That is, on each trial, the spike train for a neuron is drawn from an inhomogeneous Poisson process, with a rate function given by the trial-average firing rate. Similar baselines could be developed for calcium imaging and fMRI studies after specifying an appropriate noise model. We did not apply this method directly in this manuscript, but nonetheless mention it here for completeness.
Second, a key visualization tool enabled by time warping is the alignment of neural activity across trials. This alignment is achieved by applying the inverse warping functions to re-scale the time axis on the raw data; it does not directly rely on the response template, $X$. Thus, one can visualize the aligned activity of an individual neuron in a held out manner—the model is fit to all trials and all other neurons, and the warping functions are applied to the held out cell (see Figure 2K). This can then be repeated for each neuron in the full population. All spike raster plots in the main paper were produced using this procedure.

While these two approaches do not supplant the need for careful cross-validation, they can provide a quick validation for visualizations and presented results.

DATA AND CODE AVAILABILITY

Our code for fitting linear and piecewise linear time warping models is distributed as a GitHub repository (under an MIT license): https://github.com/ahwillia/affinewarp. This repository also contains pre-processed versions of all three experimental datasets detailed in this paper, alongside step-by-step tutorials which reproduce the major results in Figures 2, 3, 5, and 6. Our Python implementation relies on the standard SciPy scientific computing libraries (Jones et al., 2001–; Hunter, 2007). Additionally, we achieved substantial performance enhancements by leveraging numba, a Python library that enables just-in-time (JIT) compilation (Lam et al., 2015).