- 1 A large-scale neural network training framework for generalized estimation of single-trial population dynamics
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22 Abstract

23 Large-scale recordings of neural activity are providing new opportunities to study network-level dynamics. However, the 24 sheer volume of data and its dynamical complexity are critical barriers to uncovering and interpreting these dynamics. 25 Deep learning methods are a promising approach due to their ability to uncover meaningful relationships from large, 26 complex, and noisy datasets. When applied to high-D spiking data from motor cortex (M1) during stereotyped behaviors, 27 they offer improvements in the ability to uncover dynamics and their relation to subjects' behaviors on a millisecond 28 timescale. However, applying such methods to less-structured behaviors, or in brain areas that are not well-modeled by 29 autonomous dynamics, is far more challenging, because deep learning methods often require careful hand-tuning of 30 complex model hyperparameters (HPs). Here we demonstrate AutoLFADS, a large-scale, automated model-tuning 31 framework that can characterize dynamics in diverse brain areas without regard to behavior. AutoLFADS uses distributed 32 computing to train dozens of models simultaneously while using evolutionary algorithms to tune HPs in a completely 33 unsupervised way. This enables accurate inference of dynamics out-of-the-box on a variety of datasets, including data 34 from M1 during stereotyped and free-paced reaching, somatosensory cortex during reaching with perturbations, and 35 frontal cortex during cognitive timing tasks. We present a cloud software package and comprehensive tutorials that 36 enable new users to apply the method without needing dedicated computing resources.

38 Introduction

39 Ongoing advances in neural interfacing technologies are enabling simultaneous monitoring of the activity of large neural 40 populations across a wide array of brain areas and behaviors (1-5). Such technologies may fundamentally change the 41 questions we can address about computations within a neural population, allowing neuroscientists to shift focus from 42 understanding how individual neurons' activity relates to externally-measurable or controllable parameters, toward 43 understanding how neurons within a network coordinate their activity to perform computations underlying those 44 behaviors. A natural method for interpreting these complex, high-dimensional datasets is that of neural population dynamics (6-8). The dynamical systems framework centers on uncovering coordinated patterns of activation across a 45 46 neural population and characterizing how these patterns change over time. Knowledge of these hidden dynamics has 47 provided new insights into how neural populations implement the computations necessary for motor, sensory, and 48 cognitive processes (9-15).

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A focus on population dynamics could also facilitate a shift away from reliance on stereotyped behaviors and trialaveraged neural responses. Standard approaches must typically average activity across trials, sacrificing single trial interpretability for robustness against what is perceived as noise in single trials. However, as articulated by Cunningham and Yu (16): "If the neural activity is not a direct function of externally measurable or controllable variables (for example, if activity is more a reflection of internal processing than stimulus drive or measurable behavior), the time course of neural responses may differ substantially on nominally identical trials." This may be especially true of non-primary cortical
 areas, and cognitively demanding tasks that involve decision-making, allocation of attention, or varying levels of
 motivation.

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59 To move beyond this bottleneck, high-time resolution single-trial analyses are essential. These can be enabled by a 60 combination of neural population recordings and novel analytical tools like those proposed here. Single-trial, population-61 level analyses benefit from two principles of the dynamical systems view: first, that simultaneously recorded neurons are 62 not independent, but rather exhibit coordinated patterns of activation that reflect the state of the overall network rather 63 than individual neurons. Second, the coordinated patterns evolve over time in ways that are largely predictable based 64 on the population's internal dynamics. Thus, while it may be challenging to accurately estimate the network's state based 65 solely on activity observed at a single time point, knowledge of how the state evolves can constrain an estimate at any 66 given time point.

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Several approaches have been developed to infer latent dynamical structure from neural population activity on individual trials, including a growing number that leverage artificial neural networks (17–22). One such method, latent factor analysis via dynamical systems (LFADS) (22,20) achieved precise inference of motor cortical firing rates on single trials of stereotyped behaviors, enabling accurate prediction of subjects' behaviors on a moment-by-moment, millisecond timescale (20). Further, in tasks with unpredictable events, a modified network architecture enabled inference of dynamical perturbations that corresponded to how subjects ultimately responded to the unpredictable events.

75 Though highly effective, artificial neural networks, including LFADS, typically have many thousands of parameters, and 76 potentially dozens of non-trainable hyperparameters (HPs) that need to be tuned to achieve good performance. HPs 77 include architecture parameters like the type, dimensionality, and number of various layers, as well as regularization and 78 optimization parameters. Until recently, the HP optimization problem was typically addressed by an iterative manual 79 process, a random search, or some combination of the two. In the past several years, a host of more advanced 80 approaches promises to eliminate the tedious work and domain knowledge required for manual tuning while performing 81 better and more efficiently than random search (23-25). The form and variety of possible neuroscientific datasets present 82 unique challenges that make HP optimization a particularly impactful problem (26). Thus, bringing efficient HP search 83 algorithms to neuroscience could allow more effective experimentation with models based on artificial neural networks, 84 like LFADS.

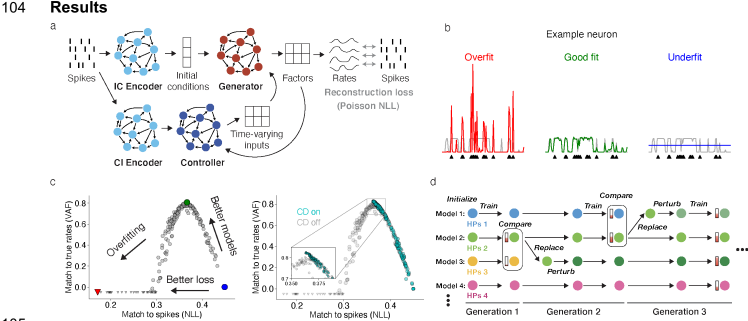
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Here we present AutoLFADS, a framework for large-scale, automated model tuning that enables accurate single-trial inference of neural population dynamics across a range of brain areas and behaviors. We evaluate AutoLFADS using data from three cortical regions: primary motor and dorsal premotor cortex (M1/PMd), somatosensory cortex area 2, and dorsomedial frontal cortex (DMFC). The tasks span a mix of functions where population activity can be well-modeled by autonomous dynamics (e.g., pre-planned reaching movements, estimation of elapsed time) and those for which population activity is responsive to external inputs (e.g., mechanical perturbations, unexpected appearance of reaching targets, variable timing cues).

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94 Using this broad range of datasets, we show that AutoLFADS achieves high-time resolution, single-trial inference of 95 neural population dynamics, surpassing LFADS in all scenarios tested. Remarkably, AutoLFADS does this in a 96 completely unsupervised manner that does not depend on the knowledge of the tasks, subjects' behaviors, or brain 97 areas. In all applications, the method is applied "out of the box" without careful adjustment for each dataset. We believe 98 these capabilities greatly extend the range of neuroscientific applications for which accurate inference of single-trial 99 population dynamics should be achievable, and substantially lower the barrier to entry for applying these methods. 100 Finally, we present a cloud software package and comprehensive tutorials to enable new users without machine learning 101 expertise or dedicated computing resources to apply AutoLFADS successfully.

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106 Fig1 | AutoLFADS combines a novel neural network regularization method with a large-scale framework for automated 107 hyperparameter optimization. (a) Schematic of the LFADS architecture, showing how the generative model infers the firing rates 108 that underlie the observed spikes. (b) Examples of LFADS-inferred rates (colored) and the corresponding synthetic input data (spikes, 109 shown as black triangles) and data-generating distribution (ground truth rates, shown as gray traces) for three fitting modes. (c) Left: 110 performance of 200 LFADS models with random HPs in matching the spikes and the known rates of a synthetic dataset, measured 111 by negative log-likelihood (NLL) and variance accounted for (VAF) respectively. Colored points indicate the models that produced the 112 rates in the previous panel. Right: same as previous, but for models trained with CD. (d) Schematic of the PBT approach to HP 113 optimization. Each colored circle represents an LFADS model with a certain HP configuration and partially filled bars represent model 114 performance. Models are trained for fixed intervals (generations), between which poorly-performing models are replaced by copies of 115 better-performing models with perturbed HPs.

117 LFADS architecture

118 The LFADS architecture (Fig. 1a) has been detailed previously (20,22,26). Briefly, LFADS is based on the idea that the 119 evolution of a neural population's activity in time can be modeled as a non-autonomous dynamical system, i.e., a 120 dynamical system whose state evolution is influenced by both internal dynamics and external inputs. This dynamical 121 system is approximated by a recurrent neural network (RNN) known as the generator. Observed spiking activity from 122 each neuron is assumed to reflect an underlying firing rate that is linked to the state of the generator at each timestep. 123 Separately, to enable modeling of input-driven dynamical systems, time-varying inputs are inferred by a controller RNN, 124 which receives as input an encoding of the spike count data as well as the generator's output at the previous time step. 125 This architecture is a modification of a sequential variational autoencoder (VAE) (22,27,28). When training the model, 126 the objective is to maximize a lower bound on the Poisson likelihood of the observed spiking activity given the inferred 127 rates (see Methods for details).

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129 It is imperative to regularize the model properly in order to extract useful spike rates (Fig. 1b) (26). This can be achieved 130 through HP optimization. The two main classes of LFADS HPs are those that set the network architecture (e.g., number 131 of units in each RNN, dimensionality of initial conditions, inputs, and factors), and those that control regularization and 132 training (e.g., L2 penalties, scaling factors for KL penalties, dropout probability, and learning rate; described in *Methods*). 133 The optimal values of these HPs could depend on various factors such as dataset size, dynamical structure underlying 134 the activity of the brain region being modeled, and the behavioral task.

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A critical challenge for autoencoders is that automatic HP searches face a type of overfitting that is particularly hard to address (26). Given enough capacity, the model can find a trivial solution where it simply passes individual spikes from the input to the output firing rates, akin to an identity transformation of the input without modeling any meaningful structure underlying the data (**Fig. 1b**). Importantly, such pathological overfitting is not detectable by standard validation likelihood, 140 as the failure mode also results in high likelihood and poor modeling of validation data. We performed a 200-model 141 random search over a space of KL, L2, and dropout regularization HPs that was empirically determined to yield both 142 underfitting and overfitting models on a synthetic dataset (see *Methods* for a description of the dataset). Models that 143 appear to have the best likelihoods actually exhibit poor inference of underlying firing rates, indicating a type of 144 pathological overfitting (**Fig. 1c**, left). This phenomenon is also consistently observed on real data throughout this paper: 145 better validation loss did not indicate better performance for any of our decoding or PSTH-based metrics.

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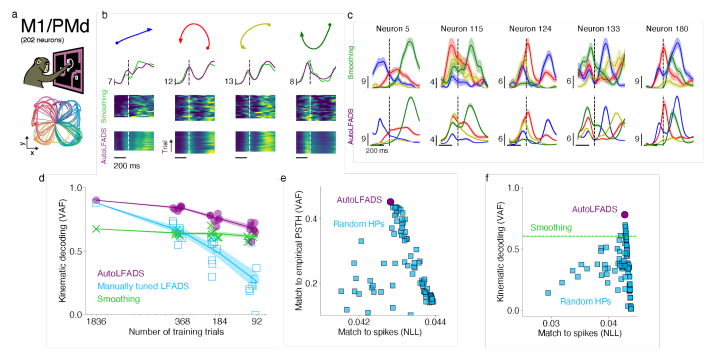
The lack of a reliable validation metric has prevented automated HP searches because it is unclear how one should select between models when underlying firing rates are unavailable or non-existent. To address this issue, we developed a novel regularization technique called coordinated dropout (CD) that forces the network to model only structure that is shared across neurons (26). After applying CD, we repeated the previous test on synthetic data using 200 LFADS models from the same HP search space, and found that they no longer overfit spikes (**Fig. 1c**, right). CD restored the correspondence between model quality assessed from matching spikes (validation likelihood) and matching rates, allowing the former to be used as a surrogate when the latter is not available.

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The premise of this paper is that this reliable validation metric should enable large-scale HP searches and fullyautomated selection of high-performing neuroscientific models despite having no access to ground truth firing rates. To test this, we needed an efficient HP search strategy. We chose a recent method based on parallel search called Population Based Training (PBT; **Fig. 1d**) (25,29). PBT distributes training across dozens of models simultaneously, and uses evolutionary algorithms to tune HPs over many generations. Because PBT distributes model training over many workers, it matches the scalability of parallel search methods such as random or grid search, while achieving higher performance with the same amount of computational resources (25,29).

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163 These two key modifications - a novel regularization strategy (CD) that results in a reliable validation metric, and an 164 efficient approach to HP optimization (PBT) - yield a large-scale, automated framework for model tuning, which we refer 165 to as AutoLFADS. In the following sections, we test the performance of AutoLFADS on previously characterized datasets, 166 as well as novel ones. We start by evaluating AutoLFADS using data from M1/PMd in a structured reaching task to 167 investigate the model's performance on a well-characterized dataset that had been previously used to benchmark the 168 performance of LFADS (20,26). On this data, we demonstrate that proper HP tuning leads to models that consistently 169 outperform LFADS and that this gap grows substantially when data are limited. Next, we move to assessing the ability 170 of AutoLFADS to approximate input-driven dynamics, using data from M1 in a random target task, data from area 2 in a 171 reaching task with mechanical perturbations, and data from DMFC in a cognitive timing task. In each case, by several 172 metrics. AutoLFADS consistently achieves better results than random searches that used three times the computational 173 resources, despite performing model selection in a completely unsupervised fashion. 174



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176 Fig2 | Application of AutoLFADS to data from motor cortex. (a) Schematic of the maze task (top), and representative reach 177 trajectories across 108 total conditions, colored by target location (bottom). (b) Average reach trajectories (top), PSTHs (second row) 178 and single-trial firing rates (bottom) obtained by smoothing (Gaussian kernel, 30 ms s.d.) or AutoLFADS for a single neuron across 4 179 reach conditions. All data is modeled at 2 ms bins. Dashed lines indicate movement onset and vertical scale bars denote rates 180 (spikes/s). (c) PSTHs produced by smoothing spikes (top) or by applying AutoLFADS (bottom), for 5 example neurons. Shaded 181 regions are standard errors. Movement onset and rate scales are denoted as in the previous panel. (d) Performance in decoding 182 reaching kinematics (arm velocities) as a function of training dataset size. Trial counts exclude the 20% of trials for each dataset size 183 that were held-out for model evaluation. We decoded X and Y arm velocities from smoothed spikes, rates inferred by LFADS with 184 manually-tuned hyperparameters (HPs), and rates inferred by AutoLFADS. Accuracy was guantified by VAF. Lines and shading 185 denote mean +/- standard error across 7 models trained on randomly-drawn subsets of the full dataset. (e) Performance in replicating 186 the empirical PSTHs computed on all trials using rates inferred from a 184-trial training set using AutoLFADS and LFADS with random 187 HPs (100 models). (f) Hand velocity decoding performance for firing rates from a 184-trial training set (same models as in (e)).

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189 AutoLFADS outperforms original LFADS when applied on benchmark data from M1/PMd

190 We first evaluated AutoLFADS on data from motor cortex during a highly stereotyped behavior, which was used to 191 assess the original LFADS method (20). We used 202 neurons simultaneously recorded from M1 and PMd during a 192 maze reaching task (see Methods) in which a monkey made a variety of straight and curved reaches after a delay period 193 following target presentation (Fig. 2a; dataset consisted of 2296 individual reach trials spanning 108 reach types). 194 Previous analyses of the delayed reaching paradigm demonstrated that activity during the movement period is well 195 modeled as an autonomous dynamical system (10,20). In this abstract model, the temporal evolution of the neural 196 population's activity is predictable based on the state it reaches during the delay period. Therefore, previous work 197 modeled these data with a simplified LFADS configuration which could only approximate autonomous dynamics (20). 198 However, this simplified model is not applicable more broadly to situations in which both autonomous dynamics and 199 external inputs might be needed to describe neural activity. Therefore, in this paper we do not constrain the network 200 architecture to only model autonomous dynamics for any applications tested, to determine whether AutoLFADS can 201 automatically adjust the degree to which autonomous dynamics and inputs are needed to model the data.

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AutoLFADS operates on unlabeled segments of binned spiking data and infers firing rates for each neuron in an unsupervised manner. Consistent with previous applications of LFADS on this dataset (20,26), the firing rates inferred by AutoLFADS for 2 ms bins exhibited clear and consistent structure on individual trials (**Fig. 2b**, bottom). We also verified that these firing rates captured features of the neural responses revealed by averaging across trials, a common method of de-noising neural activity (**Fig. 2b**, second row, and **Fig. 2c**).

209 A generalizable method should be able to perform well across the broad range of dataset sizes typical of neuroscience 210 experiments. To test this, we compared AutoLFADS and manually-tuned LFADS models that were trained using either 211 the full dataset (2296 trials), or randomly sampled subsets containing 5, 10, and 20% of the trials. We first tested the 212 degree to which the representations produced by the models were informative about observable behavior, which we 213 quantified by decoding the monkey's hand velocity from the inferred rates using optimal linear estimation (Fig 2d). At 214 the largest dataset size, decoding performance for AutoLFADS and manually-tuned LFADS was comparable. This result 215 fits with standard intuition that performance is less sensitive to HPs when sufficient data are available. However, for all 216 three reduced dataset sizes, the AutoLFADS outperformed the manually-tuned model (p<0.05 for all three sizes, paired, 217 one-tailed Student's t-test).

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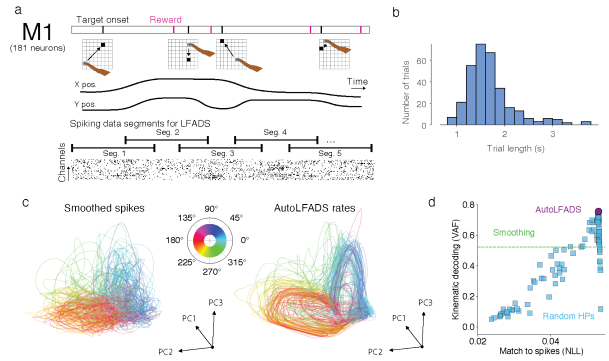
219 While this result is promising, the difference in robustness to dataset size between AutoLFADS and LFADS could have 220 resulted from a particularly poor selection of HPs during manual tuning. To control for this possibility, we chose one of 221 the smaller data subsets (184 trials) and trained 100 additional LFADS models with randomly-selected HPs. We 222 evaluated the models' performance in two ways: how accurately the models replicated the empirical trial-averaged firing 223 rates (PSTHs; Fig. 2e), and how accurately arm velocity could be decoded from inferred rates (Fig. 2f). While the LFADS 224 models achieved a broad range of performance, models with better validation likelihoods did not achieve better inference 225 of firing rates, mirroring our earlier findings with synthetic data (Fig. 1c). Thus it is unclear how one could select amongst 226 the LFADS models with random HPs without some supervised intervention. In contrast, the single AutoLFADS model, 227 chosen in a completely unsupervised fashion, outperformed all LFADS models for both performance metrics.

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Taken together, these results show that even if one performed a random search and then selected a model using a supervised approach (e.g., based on reconstruction of empirical PSTHs or decoding accuracy), its performance would still be substantially lower than that of AutoLFADS. Additionally, this validation - i.e., that the unsupervised approach produces high-performing models - provides evidence that even in cases where such supervision is unavailable (e.g., settings that lack clear task structure or measurement of behavioral variables), AutoLFADS models will still be high performing.

236 AutoLFADS uncovers population dynamics without structured trials

237 To-date, most efforts to tie dynamics to neural computations have used experiments where subjects perform constrained 238 tasks with repeated, highly structured trials. For example, motor cortical dynamics are often framed as a computational 239 engine to link the processes of motor preparation and execution (6-8). To interrogate these dynamics, most studies use 240 a delayed-reaching paradigm that creates explicit pre-movement and movement periods. However, constrained 241 behaviors may have multiple drawbacks in studying dynamics. First, it is unclear whether such artificial paradigms are 242 good proxies for everyday behaviors. Second, highly constrained, repeated behaviors might impose artificial limits on 243 the properties of the uncovered dynamics, such as the measured dimensionality of the neural population activity (30). 244 Even outside of movement neuroscience, the requirement that we conduct many repetitions of constrained tasks 245 significantly hinders our ability to study a rich sample of the dynamics of a given neural population. Accurate inference 246 of neural dynamics without these constraints could facilitate dynamics-based analyses of richer datasets that are more 247 reflective of the brain's natural behavior.



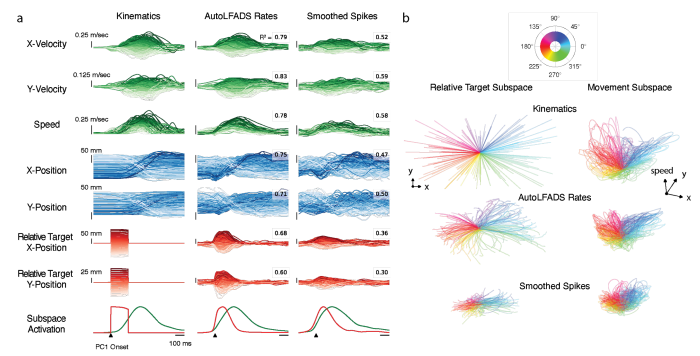
249 Fig3 | Modeling neural activity in M1 without knowledge of trial or task information. (a) Top: Schematic of the random target 250 task, which lacks stereotyped trial structure and delay periods. Bottom: Continuous neural activity (spiking data) recorded during back-251 to-back reaching trials was divided into 600 ms segments with 200 ms of overlap between adjacent segments. After modeling by 252 AutoLFADS, the inferred firing rates from different segments were merged together to create a continuous segment, using a weighted 253 average of data at overlapping timepoints. (b) Distributions of trial lengths (time between onsets of successive targets) for 313 total 254 trials. (c) Subspaces of neural activity extracted using PCA and colored by angle to the target. Left: 3D subspace that captures the 255 most variance in smoothed spiking activity. Center: Subspace that captures the most variance in AutoLFADS rates. (d) Accuracy in 256 decoding hand velocity from firing rates inferred by smoothing, 100 LFADS models with random HPs, and AutoLFADS.

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258 In order to provide access to a much broader range of experimental data, we tested whether AutoLFADS could model 259 data without regard to trial structure. We applied AutoLFADS to neural activity from a monkey performing a continuous, 260 self-paced random target reaching task (Fig. 3a, top) (31), in which each movement started and ended at a random 261 position, and movements were highly variable in duration (Fig. 3b). Analysis of data without consistent temporal structure 262 repeated across trials is challenging, as trial-averaging is not feasible. Even the available single-trial analytical methods 263 have typically relied on strong simplifying assumptions that are not applicable to less-structured tasks. For example, 264 previous efforts to uncover motor cortical dynamics during single reaches have been able to consider only brief data 265 segments that begin with the arm at a consistent starting point, and relied on behavioral events such as target or 266 movement onset to align trials before analysis (17.20.26.32-36). 267

268 Like most machine learning algorithms, AutoLFADS operates on discrete, fixed-length segments of neural data. To 269 create these segments from a task with highly variable timing, we chopped an approximately 9 minute window of 270 continuous neural data into 600 ms segments with 200 ms of overlap (Fig. 3a, bottom) without regard to trial boundaries. 271 After modeling with AutoLFADS, we merged inferred firing rates from individual segments, which yielded inferred rates 272 for the original continuous window. We then analyzed the inferred rates by aligning the data to movement onset for each 273 trial (see Methods). Even though the dataset was modeled without the use of trial information, inferred firing rates during 274 the reconstructed trials exhibited consistent progression in an underlying state space, with clear structure that 275 corresponded with the monkey's reach direction on each trial (Fig. 3c, right). Further, the inferred firing rates were highly 276 informative about moment-by-moment details of the measured reaching movements: AutoLFADS enabled decoding of 277 continuous hand velocities with substantially higher accuracy than did smoothing (R² of 0.76 for AutoLFADS v. 0.52 for 278 smoothing), and it also outperformed all LFADS models with random HPs (Fig. 3d).

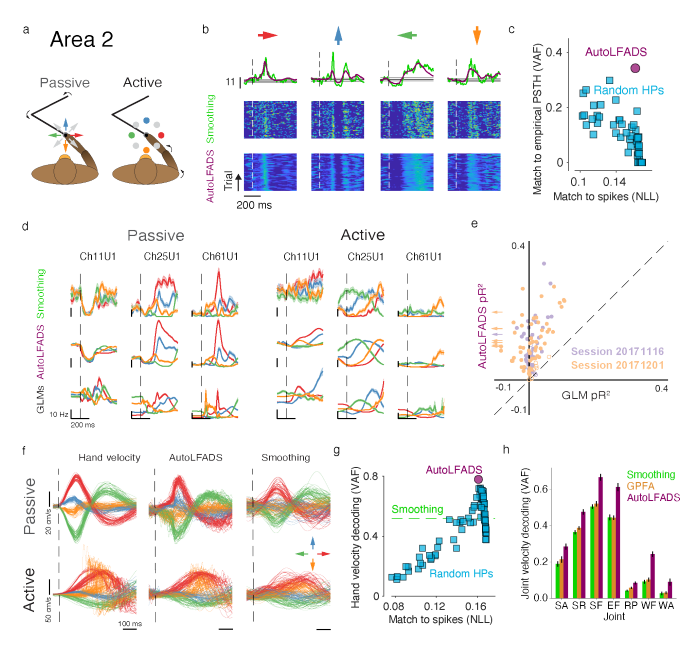


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Fig4 | Inferred firing rates contain neural subspaces that are informative about movement kinematics and reach targets. (a) Kinematic and relative target variables and their corresponding neural representations, uncovered via linear regression. The quality of each projection is quantified by accuracy in decoding kinematic and target variables (R²). Plots are colored by x and y distance to 283 target, except for speed which is colored by peak speed. Bottom row represents the normalized activation of movement (green) and 284 relative target (red) subspaces, illustrating the more transient activation in the target subspace. (b) Movement and relative target 285 subspaces plotted as 3D trajectories and colored by angle to target.

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287 In support of the hypothesis that AutoLFADS is picking up on meaningful dynamics that occurred throughout the session, 288 we found that the firing rates inferred by AutoLFADS were informative of the previously-hypothesized computational role 289 of motor cortical dynamics - i.e., linking the process of movement preparation and execution - despite the model being 290 trained without information about the monkey's behavior (Fig. 4). In particular, firing rates contained subspaces that 291 were highly informative about hand position, hand velocity, and reach target on individual trials (Fig. 4a) and showed 292 clear structure relative to the task (Fig. 4b). To find the subspaces, we used linear regression to project neural activity 293 onto variables related to movement goals (reach target) and movement details (position, velocity and speed). Notably, 294 the subspace reflecting reach target was transiently active around the time of movement execution, consistent with 295 previous studies that have demonstrated the presence of preparatory activity in motor cortex, yet revealed without an 296 explicit preparatory period. It is likely that the rates inferred by AutoLFADS also contain yet undiscovered subspaces 297 and representations that can be explored in this same dataset without experiments explicitly designed to reveal them. 298 Thus, AutoLFADS has the potential to greatly improve the utility and versatility of rich behavioral datasets via a unique 299 unsupervised modeling process.



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301 Fig5 | Application of AutoLFADS to data from somatosensory cortex area 2. (a) Schematic of the center-out, bump task showing 302 passive and active conditions. (b) PSTHs and single-trial firing rates for a single neuron across 4 passive perturbation directions. 303 Smoothing was performed using a Gaussian kernel with 10 ms s.d.. Dashed lines indicate movement onset. (c) Comparison of 304 AutoLFADS vs. random search in matching empirical PSTHs. (d) PSTHs produced by smoothing spikes (top), AutoLFADS (middle), 305 or GLM predictions (bottom) for 3 example neurons. (e) Comparison of spike count predictive performance for AutoLFADS and GLMs. 306 Filled circles correspond to neurons for which AutoLFADS pR² was significantly higher than GLM pR², and open circles correspond 307 to neurons for which there was no significant difference. Arrows (left) indicate neurons for which GLM pR² was outside of the plot 308 bounds. (f) Subspace representations of hand x-velocity during active and passive movements extracted from smoothed spikes and 309 rates inferred by AutoLFADS. (g) Comparison of AutoLFADS vs. random search in decoding hand velocity during active trials. (h) 310 Joint angular velocity decoding performance from firing rates inferred using smoothing, Gaussian process factor analysis (GPFA), 311 and AutoLFADS. Error bars denote standard error of the mean. Joint abbreviations: shoulder adduction (SA), shoulder rotation (SR), 312 shoulder flexion (SF), elbow flexion (EF), wrist radial pronation (RP), wrist flexion (WF), and wrist adduction (WA).

315 AutoLFADS accurately captures single-trial population dynamics in somatosensory cortex

Results from the motor cortical datasets demonstrated that AutoLFADS could produce accurate dynamical models that were robust to training dataset size and generalized well across task conditions, without requiring highly constrained tasks or repeated trials. We next investigated whether AutoLFADS, without manual adjustment, could accurately model dynamics associated with sensory processes. Specifically, we modeled activity in somatosensory area 2 during a reaching task with mechanical perturbation.

Area 2 provides a valuable test case for AutoLFADS generalization. As a sensory area, area 2 receives strong afferent input from cutaneous receptors and muscles and is robustly driven by mechanical perturbations to the arm (37–39). Functionally, area 2 is thought to serve a role in mediating reach-related proprioception (38–41), was recently shown to contain information about whole-arm kinematics (39), and may also receive efferent input from motor areas (38,39,42,43).

328 In the area 2 experiment (Fig. 5a), a monkey used a manipulandum to control a cursor. The task began with a center-329 hold period where the monkey held the cursor in the center of the screen. During half of the center-hold attempts, the 330 manipulandum randomly perturbed the monkey's arm in one of the eight directions, and the monkey had to re-acquire 331 the central target (passive movement trials). Following the center-hold, the monkey moved to acquire one of eight 332 peripheral targets (active movement trials). The single-trial rates inferred by AutoLFADS for passive trials exhibited clear 333 and structured responses to the unpredictable perturbations (Fig. 5b), highlighting the model's ability to approximate 334 input-driven dynamics.

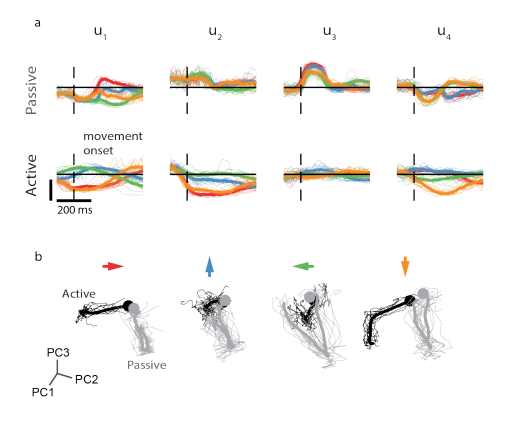
335 336 As for M1/PMd, we verified that the rates inferred by AutoLFADS accurately reproduced empirical PSTHs and were 337 informative of task variables. The inferred rates captured the distinct features of PSTHs during active and passive trials, 338 even though no behavioral or task information was provided to the model (Fig. 5b; top, and Fig. 5c). The rates inferred 339 by AutoLFADS also had a much closer correspondence to the empirical PSTHs during passive trials than LFADS models 340 trained with random HPs (Fig. 5c). However, sensory brain regions like area 2 are typically characterized in terms of 341 how neural activity encodes sensory stimuli (37-39). Thus, we examine whether rates inferred by AutoLFADS explain 342 observed spikes better than a typical area 2 neural encoding model, in which neural activity is fit to some function of the 343 state of the arm. We fit a generalized linear model (GLM) for each neuron over both active and passive movements, 344 where the firing rate was solely a function of the position and velocity of the hand, as well as the contact forces with the 345 manipulandum handle (39) (GLM predictions shown in Fig. 5d). We then compared the ability of the GLM and 346 AutoLFADS to capture each neuron's observed response using pseudo-R² (pR²), a metric similar to R² but adapted for 347 the Poisson statistics of neural firing (44). For the vast majority of neurons across two datasets. AutoLFADS predicted 348 the observed activity significantly better than GLMs (p<0.05 for 110/121 neurons, bootstrap; see *Methods*), and there 349 were no neurons for which the GLM produced better predictions than AutoLFADS (Fig. 5e).

351 We used linear decoding to extract subspaces of neural activity that corresponded to x and y hand velocities for both 352 smoothed spikes and rates inferred by AutoLFADS (Fig. 5f). The AutoLFADS rates contained subspaces that more 353 clearly separated hand velocities for all active conditions and all passive conditions than smoothing, showing that they 354 are better represented in the modeled dynamics of area 2. Further, single-trial hand velocity decoding from rates inferred 355 by AutoLFADS for active trials was substantially more accurate than that of smoothing, and also more accurate than 356 decoding from the output of any random search model (Fig. 5g). On a second dataset that included whole-arm motion 357 tracking, the velocity of all joint angles was decoded from AutoLFADS rates with higher accuracy than from smoothing 358 or GPFA (Fig. 5h, right; p<0.05 for all joints, paired, one-sided Student's t-Test).

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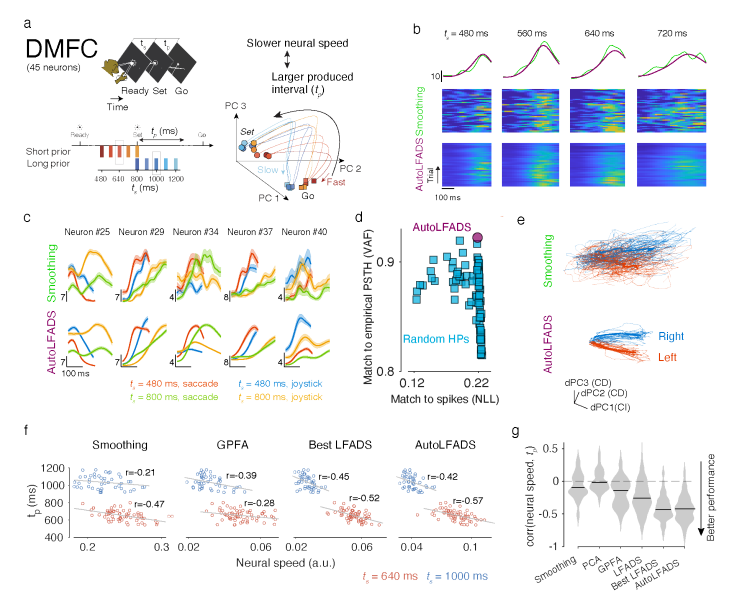


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Fig6 | AutoLFADS-inferred inputs for area 2 neural activity. (a) Time-courses of the four inferred generator input dimensions for passive (top) and active (bottom) conditions. Thick line indicates average input trace for each direction, indicated by color, while thin colored lines show input traces for ten randomly chosen trials. Vertical scale bar is A.U. (b) Projection of four-dimensional inputs, from -100 ms to 200 ms around movement onset, into the top three principal components, with separate plots for each movement direction. Darker lines indicate active trials while lighter lines denote passive trials. Large dots indicate average initial input in PC space. Thick and thin lines follow conventions in (a).

Since area 2 plays a significant role in processing sensory inputs, it stands to reason that the inputs inferred by AutoLFADS are important for successfully modeling the area's activity as a dynamical system. If AutoLFADS is successfully modeling area 2 as an input-driven dynamical system, we should expect the inferred inputs to be consistent across trials with the same behavioral conditions. In these experiments, AutoLFADS models the data as fixed-length segments without regard to trial boundaries, so there is no guarantee of the consistency of the meaning of a given input between different trials of the same condition or even within a single trial.

373 Despite the unsupervised modeling process, AutoLFADS inferred input trajectories that were consistent with the 374 supervised notions of trials, directions, and perturbation types (Fig. 6a). Inputs were continuous over the course of a 375 trial, implying that the model was able to pick up on statistical similarities between adjacent segments. The model also 376 produced similar input patterns within a given condition, showing that it was able to detect the statistical patterns of a 377 given condition from arbitrary segments of time during arbitrary trials. Finally, AutoLFADS produced distinct and logically 378 consistent output patterns for active and passive trials. Inputs for abrupt passive movements generally had a much 379 shorter time course that unfolded post-perturbation, while inputs for active trials began before movement and evolved 380 more slowly. Visualization of these inputs highlights AutoLFADS's ability to infer distinct inputs for distinct subsets of the 381 data (Fig. 6b).



382

383 Fig7 | Application of AutoLFADS to data from dorsomedial frontal cortex (DMFC). (a) Top left: the time interval reproduction 384 task. Bottom left: timing conditions used. Right: schematic illustrating the inverse correlation between neural speed and monkey's 385 produced time (t_p). (b) PSTHs and single-trial firing rates for an example neuron during the Set-Go period of leftward saccade trials 386 across 4 different values of t_s (vertical scale bar: spikes/sec). Smoothing was performed using a Gaussian kernel with 25 ms s.d., (c) 387 PSTHs for 5 example neurons during the Set-Go period of rightward trials for two response modalities and two values of t_s . (d) 388 Performance in replicating the empirical PSTHs. (e) Visualization of low-dimensional trial-averaged and single-trial neural trajectories 389 for the Ready-Set period for left and right joystick trials with t_s of 1000 ms. 30 trials are shown for each condition. dPC: demixed 390 principal component, CI: condition-independent, CD: condition-dependent. (f) Example plots showing correlations between neural 391 speed and behavior (i.e., production time, t_p) for individual trials across two timing intervals (red: 640 ms blue: 1000 ms). Neural speed 392 was obtained based on the firing rates inferred from smoothing, GPFA, the LFADS model with best median speed- t_{ρ} correlation across 393 the 40 different task conditions (Best LFADS), and AutoLFADS. (g) Distributions of correlation coefficients across 40 different task 394 conditions. Horizontal lines denote medians. For LFADS, the distribution includes correlation values for all 96 models with random 395 HPs (40x96 values).

396

398 AutoLFADS accurately captures single-trial dynamics during cognition

399 While activity in M1 and area 2 are largely driven by internal dynamics and inputs, respectively, many brain areas depend 400 critically on the confluence of internal dynamics and inputs. To further test the generality of AutoLFADS to these 401 situations, we applied it to data collected from dorsomedial frontal cortex (DMFC) during a cognitive time estimation task. 402 DMFC comprises the supplementary eye field, dorsal supplementary motor area, and presupplementary motor area. It 403 is often considered an intermediate region in the sensorimotor hierarchy (45), interfacing with both low-level sensory and 404 motor (PMd/M1) areas. DMFC activity is less closely tied to the moment-by-moment details of movements than activity 405 in M1 or area 2 - instead, its activity seems to relate to higher-level aspects of motor control, including motor timing 406 (46,47), planning movement sequences (48), learning sensorimotor associations (49) and context-dependent reward 407 modulation (50). However, population dynamics in DMFC are tied to behavioral correlates such as movement production 408 time (15,47,51). This makes DMFC another excellent test case for unsupervised modeling with AutoLFADS.

409

For this task, the monkey was presented with two visual stimuli ("Ready" and "Set", respectively), separated by sample timing interval t_s . After "Set", the monkey attempted to reproduce the interval by waiting for the same amount of time (t_p) before initiating a movement ("Go") (**Fig. 7a**, left). The movement was either a saccade or joystick manipulation to the left or right depending on the location of a peripheral target. The two response modalities, combined with 10 timing conditions (t_s) and two target locations, led to a total of 40 task conditions.

415

416 Consistent with our observations on M1/PMd and area 2 data, AutoLFADS-inferred rates for this dataset showed 417 consistent, denoised structure at the single-trial level (Fig. 7b, bottom) and recapitulated the features of neural responses 418 uncovered by trial averaging (Fig. 7b, top; Fig. 7c). Quantitative comparison of the PSTHs shows that AutoLFADS-419 inferred rates again achieved a better match to the empirical PSTHs than all of the random search models (Fig. 7d), 420 providing further evidence that AutoLFADS can achieve superior models without expert tuning of regularization HPs or 421 supervised model selection criteria. Additionally, when visualized in a low-dimensional space using demixed principal 422 components analysis (dPCA), the AutoLFADS-inferred firing rates showed much greater consistency across trials of a 423 given condition than firing rates computed by smoothing spikes (Fig. 7e).

424

425 To evaluate the AutoLFADS model beyond its ability to capture trial-averaged responses, we sought to evaluate whether 426 its predicted firing rates were more informative of trial-by-trial timing behaviors than other methods. Previous studies 427 have shown that the monkey's produced time interval (t_{ρ}) is negatively correlated to the speed at which the neural 428 trajectories evolve during the Set-Go period (Fig. 7a, right) (15,51). To evaluate the correspondence between neural 429 activity and behavior, we estimated neural speeds using representations produced by smoothing spikes, GPFA, principal 430 component analysis (PCA), the best random search model ('Best LFADS', see Methods for details), and an AutoLFADS 431 model, and measured the trial-by-trial correlation between the estimated speeds and t_p . Note that selecting the best 432 random search model again required a supervised calculation (t_p correlation) for each model. If a given representation 433 of neural activity is more informative about behavior, we expect a stronger (more negative) correlation between predicted 434 and observed t_{ρ} .

435

436 We show correlation values for individual trials across two different values of t_s (**Fig. 7f**), and summarize across all 40 437 task conditions (**Fig. 7g**). We observed consistent negative correlations between t_p and the estimated neural speed from 438 rates obtained by different methods. Correlations from rates inferred by AutoLFADS were significantly better than all 439 unsupervised approaches (p<0.001, Wilcoxon signed rank test), and comparable with the supervised selection approach 440 ('Best LFADS', p=0.758, Wilcoxon signed rank test), despite using no task information.

441

Taken together, the area 2 and DMFC results demonstrate that the out-of-the-box, automated inference of neural population dynamics provided by AutoLFADS allows modeling of diverse brain areas, with dynamics that span the continuum from autonomous to input-driven. AutoLFADS provides a powerful framework for generalized inference of input-driven dynamics and enables decoding of simultaneously monitored behavioral variables with unprecedented accuracy. Importantly, the unsupervised approach of AutoLFADS avoids the use of any behavioral data and optimizes only for neural modeling. This allows for modeling when behavioral data is not available and also prevents any behavioral biases from being introduced to the firing rates, resulting in better inference of the brain's inherently generalized representations. This is evident in the high performance of AutoLFADS rates in both PSTH reconstruction and variousdecoding tasks.

451

452 Running AutoLFADS in the Cloud

453 A key challenge with emerging, computationally-intensive data analysis methods is that the computational infrastructure 454 and expertise necessary to make effective use of these tools is a significant barrier to widespread adoption (52). For 455 example, many labs do not have the resources necessary to train dozens of models in parallel across many GPUs. To 456 address this hurdle, we provide an open-source implementation of AutoLFADS designed to operate on Google Cloud 457 Platform (GCP). Additionally, we provide a comprehensive tutorial to help novice users get started running AutoLFADS 458 on GCP without expert knowledge of cloud computing or machine learning. The tutorial describes how to set up the 459 framework, prepare input data, set up AutoLFADS runs, and load the final results. Users of AutoLFADS on GCP don't 460 need to worry about the upfront hardware and labor costs associated with maintaining a local computing cluster, yet 461 have access to virtually unlimited computation on demand. This framework allows researchers to spend less time doing 462 non-research tasks like dependency management and hyperparameter optimization, while giving them confidence that 463 their models are performing well, regardless of brain area or task. We include links to the code and tutorial in Code 464 Availabilitv.

465

466 Discussion

467 The original LFADS work (20) provided a method for inferring latent dynamics, denoised firing rates, and external inputs 468 from large populations of neurons, producing representations that were more informative of behavior than previous 469 approaches (33). However, application of LFADS to neural populations with different dynamics, strong external inputs, 470 or unconstrained behavior would have necessitated time-consuming and subjective manual tuning. In the current work, 471 we show that with robust regularization and efficient hyperparameter tuning it is possible to train high-performing LFADS 472 models for neural spiking datasets with arbitrary size, trial structure, and dynamical complexity. We demonstrated several 473 properties of the AutoLFADS training approach which have broad implications. On the maze task, we showed that 474 AutoLFADS models are more robust to dataset size, opening up new lines of inquiry on smaller datasets and reducing 475 the number of trials that must be conducted in future experiments. Using the random target task, we demonstrated how 476 AutoLFADS needs no task information in order to generate rich dynamical models of neural activity. This enables the 477 study of dynamics during richer tasks and reuse of datasets collected for another purpose. With the perturbed reaching 478 task, we demonstrated the first application of dynamical modeling, as opposed to encoder-based modeling, to the highly 479 input-driven somatosensory area 2. Finally, in the timing task, we showed that AutoLFADS found the appropriate balance 480 between inputs and internal dynamics for a cognitive area by modeling DMFC.

481

482 AutoLFADS inherits some of the flaws of the LFADS model. For example, the linear-exponential-Poisson observation 483 model is likely an oversimplification. However, we used this architecture as a starting point to show that a large-scale 484 hyperparameter search is feasible and beneficial. By enabling large-scale searches, we can be reasonably confident 485 that any performance differences achieved by future architecture changes will be due to real differences in modeling 486 capabilities rather than a simple lack of HP optimization.

487

AutoLFADS performed well using a simple binary tournament exploitation and perturbation exploration strategies for PBT (25). Future work might investigate alternate exploitation or exploration strategies, or whether more powerful and efficient PBT variants (53) can increase speed and performance of AutoLFADS while lowering computational cost. A current limitation of AutoLFADS is its inability to explore hyperparameters that modify the underlying model architecture. Thus, another avenue for further work lies in combining AutoLFADS with the recent techniques for automated neural architecture search (54).

494

Though AutoLFADS is much more efficient than previous approaches, it still requires substantial computational resources that may not be available for all potential users. Setting up the requisite software environments can be an additional hurdle. Our GCP implementation allows users to apply AutoLFADS without needing to purchase and maintain a local cluster. We estimate that the compute cost for a typical AutoLFADS run on GCP is between \$5-25, depending on dataset and model sizes. We have created detailed tutorials to guide novice users through the setup, model training, and data retrieval processes, making AutoLFADS accessible to anyone who works with neural spiking data.

501

502

503 Taken together, AutoLFADS provides an accessible and extensible framework for generalized inference of single-trial 504 neural dynamics that has the potential to unify the way we study computation through dynamics across brain areas and 505 tasks.

506

507 Code Availability

508 AutoLFADS for GCP can be downloaded from GitHub at <u>github.com/snel-repo/autolfads</u> and the tutorial is 509 available at <u>snel-repo.github.io/autolfads</u>.

511 Data Availability

512 Data will be made available upon reasonable request from the authors. The random target dataset is publicly available
 513 at http://doi.org/10.5281/zenodo.3854034.

514

510

515 Acknowledgements

516 We thank K. Shenoy, M. Churchland, M. Kaufman, and S. Ryu for sharing the Monkey J Maze dataset. We also thank 517 J. O'Doherty, M. Cardoso, J. Makin, and P. Sabes for making the random target dataset publicly available. This work 518 was supported by the Emory Neuromodulation and Technology Innovation Center (ENTICe), NSF NCS 1835364, 519 DARPA PA-18-02-04-INI-FP-021, NIH Eunice Kennedy Shriver NICHD K12HD073945, the Alfred P. Sloan Foundation, 520 the Burroughs Wellcome Fund, and the Simons Foundation as part of the Simons-Emory International Consortium on 521 Motor Control (CP), NIH NINDS R01 NS053603, R01 NS095251, and NSF NCS 1835345 (LEM), NSF Graduate 522 Research Fellowships DGE-1650044 (ARS) and DGE-1324585 (RHC), the Center for Sensorimotor Neural Engineering 523 and NARSAD Young Investigator grant from the Brain & Behavior Research Foundation (HS), NIH NINDS NS078127, 524 the Sloan Foundation, the Klingenstein Foundation, the Simons Foundation, the McKnight Foundation, the Center for 525 Sensorimotor Neural Engineering, and the McGovern Institute (MJ).

526

527 Author Contributions

	MRK	ARS	RHC	RT	DB	SLN	HS	MJ	LEM	СР
Conceptualization										
Funding acquisition										
Investigation										
Resources										
Software										
Tutorials										
Writing										
Revision										

528

529 **Competing Interests**

530 The authors declare no competing interests.

531

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671 Methods

670

672 LFADS architecture and training

A detailed overview of the LFADS model is given in (20). Briefly: at the input to the model, a pair of bidirectional RNN encoders read over the spike sequence and produce initial conditions for the generator RNN and time-varying inputs for the controller RNN. All RNNs were implemented using gated recurrent unit (GRU) cells. At each time step, the generator state evolves with input from the controller and the controller receives delayed feedback from the generator. The generator states are linearly mapped to factors, which are mapped to the firing rates of the original neurons using a linear mapping followed by an exponential. The optimization objective is to minimize the negative log-likelihood of the data given the inferred firing rates, and includes KL and L2 regularization penalties.

- 681 Identical architecture and training hyperparameter values were used for most runs, with a few deviations. We used a 682 generator dimension of 100, initial condition dimension of 100 (50 for area 2 runs), initial condition encoder dimension 683 of 100, factor dimension of 40, controller and controller input encoder dimension of 80 (64 for DMFC runs), and controller 684 output dimension of 4 (10 for overfitting runs).
- 685

We used the Adam optimizer with an initial learning rate of 0.01 and, for non-AutoLFADS runs, decayed the learning rate by a factor of 0.95 after every 6 consecutive epochs with no improvement to the validation loss. Training was halted for these runs when the learning rate reached 1e-5. The loss was scaled by a factor of 1e4 immediately before optimization for numerical stability. GRU cell hidden states were clipped at 5 and the global gradient norm was clipped at 200 to avoid occasional pathological training.

691

We used a trainable mean initialized to 0 and fixed variance of 0.1 for the Gaussian initial condition prior and set a minimum allowable variance of 1e-4 for the initial condition posterior. The controller output prior was autoregressive with a trainable autocorrelation tau and noise variance, initialized to 10 and 0.1, respectively.

Memory usage for RNNs is highly dependent on the sequence length, so batch size was varied accordingly (100 for maze and random target datasets, 500 for synthetic and area 2 datasets, and 300/400 for the DMFC dataset). KL and L2 regularization penalties were linearly ramped to their full weight during the first 80 epochs for most runs to avoid local minima induced by high initial regularization penalties. Exceptions were the runs on synthetic data, which were ramped over 70 epochs and random searches on area 2 and DMFC datasets, which used step-wise ramping over the first 400 steps.

702

Random searches and AutoLFADS runs used the architecture parameters described above, along with regularization HPs sampled from ranges (or initialized with constant values) given in **Supp. Table 2**. Most runs used a default set of ranges, with a few exceptions outlined in the table. Dropout was sampled from a uniform distribution and KL and L2 weight HPs were sampled from log-uniform distributions.

707

During PBT, weights were used to control maximum and minimum perturbation magnitudes for different HPs (e.g. a weight of 0.3 results in perturbation factors between 0.7 and 1.3). The dropout and CD HPs used a weight of 0.3 and KL and L2 penalty HPs used a weight of 0.8. CD rate, dropout rate, and learning rate were limited to their specified ranges, while the KL and L2 penalties could be perturbed outside of the initial ranges. Each generation of PBT consisted of 50 training epochs. AutoLFADS training was stopped when the best smoothed validation NLL improved by less than 0.05% over the course of four generations.

714

Validation NLL was exponentially smoothed with $\alpha = 0.7$ during training. For non-AutoLFADS runs, the model checkpoint with the lowest smoothed validation NLL was used for inference. For AutoLFADS runs, the checkpoint with the lowest smoothed validation NLL in the last epoch of any generation was used for inference. Firing rates were inferred 50 times for each model using different samples from initial condition and controller output posteriors. These estimates were then averaged, resulting in the final inferred rates for each model.

721 Overfitting on synthetic data

Synthetic data were generated using a 2-input chaotic vanilla RNN ($\gamma = 1.5$) as described in the original LFADS work (20,22). The only modification was that the inputs were white Gaussian noise. In brief, the 50-unit RNN was run for 1 second (100 time steps) starting from 400 different initial conditions to generate ground-truth Poisson rates for each condition. These distributions were sampled 10 times for each condition, resulting in 4000 spiking trials. Of these trials, 80% (3200 trials) were used for LFADS training and the final 20% (800 trials) were used for validation.

- We sampled 200 HP combinations from the distributions specified in **Supp. Table 2** and used them to train LFADS models on the synthetic dataset. We then trained 200 additional models with the same set of HPs using a CD rate of 0.3 (i.e., using 70% of data as input and remaining 30% for likelihood evaluation) (26). The coefficient of determination between inferred and ground truth rates was computed across all samples and neurons on the 800-sample validation set.
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734 M1 maze task

We used the previously-collected maze dataset (55) described in detail in the original LFADS work (20). Briefly, a male macaque monkey performed a two-dimensional center-out reaching task by guiding a cursor to a target without touching any virtual barriers while neural activity was recorded via two 92-electrode arrays implanted into M1 and dorsal PMd. The full dataset consisted of 2,296 trials, 108 reach conditions, and 202 single units.

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The spiking data were binned at 1 ms and smoothed by convolution with a Gaussian kernel (30 ms s.d.). Hand velocities were computed using second order accurate central differences from hand position at 1kHz. An antialiasing filter was applied to hand velocities and all data were then resampled to 2 ms. Trials were created by aligning the data to 250 ms before and 450 ms after movement onset, as calculated in the original paper.

745 Datasets of varying sizes were created for LFADS by randomly selecting trials with 20, 10, and 5% of the original dataset 746 using seven fixed seeds, and then splitting each of these into 80/20 training and validation sets for LFADS (22 total, 747 including the full dataset). As a baseline for each data subset, we trained LFADS models with fixed HPs that had been previously found to result in high-performing models for this dataset, with the exception of controller input encoder and controller dimensionalities (see *LFADS architecture and training* and **Supp. Table 2**). We increased the dimensionality of these components to allow improved generalization to the datasets from more input-driven areas while keeping the architecture consistent across all datasets. We also trained AutoLFADS models (40 workers) on each subset using the search space given in **Supp. Table 2**. Additionally, we ran a random search using 100 HPs sampled from the AutoLFADS search space on one of the 230-trial datasets.

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We used rates from spike smoothing, manually tuned LFADS models, random search LFADS models, and AutoLFADS models to predict x and y hand velocity delayed by 90 ms using ridge regression with a regularization penalty of $\lambda = 1$. Each data subset was further split into 80/20 training and validation sets for decoding. To account for the difficulty of modeling the first few time points of each trial with LFADS, we discarded data from the first 50 ms of each trial and did not use that data for model evaluation. Decoding performance was evaluated by computing the coefficient of determination for predicted and true velocity across all trials for each velocity dimension. The result was then averaged across the two velocity dimensions.

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To evaluate PSTH reconstruction for random search and AutoLFADS models, we first computed the empirical PSTHs by averaging smoothed spikes from the full 2296-trial dataset across all 108 conditions. We then computed model PSTHs by averaging inferred rates across conditions for all trials in the 230-trial subset. We computed the coefficient of determination between model-inferred PSTHs and empirical PSTHs for each neuron across all conditions in the subset. We then averaged the result across all neurons.

769 M1 random target task

The random target dataset consists of neural recordings and hand position data recorded from macaque M1 during a self-paced, sequential reaching task between random elements of a grid (31). For our experiments, we used only the first 30% (approx. 9 minutes) of the dataset recorded from Indy on 04/26/2016.

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774 We started with sorted units obtained from M1 and binned their spike times at 1 ms. To avoid artifacts in which the same 775 spikes appeared on multiple channels, we computed cross-correlations between all pairs of neurons over the first 10 sec 776 and removed individual correlated neurons (n = 34) by highest firing rate until there were no pairs with correlation above 777 0.0625, resulting in 181 uncorrelated neurons. The position data were provided at 250 Hz, so we upsampled these data 778 to 1 kHz using cubic interpolation. We smoothed the spikes by convolving with a Gaussian kernel (50 ms s.d.), applied 779 an antialiasing filter to hand velocities, and downsampled to 2 ms. The continuous neural spiking data were chopped 780 into overlapping segments of length 600 ms, where each segment shared its last 200 ms with the first 200 ms of the 781 next. The resulting 1321 segments were split into 80/20 training and validation sets for LFADS, where the validation 782 segments were chosen in blocks of 3 to minimize the overlap between training and validation subsets.

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The chopped segments were used to train an AutoLFADS model and to run a random search using 100 HPs sampled from the AutoLFADS search space. After modeling, the chopped data were merged using a quadratic weighting of overlapping regions that placed more weight on the rates inferred at the ends of the segments. The merging technique weighted the ends of segments as $w = 1 - x^2$ and the beginnings of segments as 1 - w, with x ranging from 0 to 1 across the overlapping points. After weights were applied, overlapping points were summed, resulting in a continuous ~9-minute stretch of modeled data.

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We computed hand velocity from position using second-order accurate central differences and introduced a 120 ms delay between neural data and kinematics. We used ridge regression ($\lambda = 1e - 5$) to predict hand velocity across the continuous data using smoothed spikes, random search LFADS rates, and AutoLFADS rates. We computed coefficient of determination for each velocity dimension individually and then averaged the two velocity dimensions to compute decoding performance.

To prepare the data for subspace visualization, the continuous activity for each neuron was soft-normalized by subtracting its mean and dividing by its 90th quantile plus an offset of 0.01. Trials were identified in the continuous data as the intervals over which target positions were constant (314 trials). To identify valid trials, we computed the normalized 800 distance from the final position. Trials were removed if the cursor exceeded 5% of this original distance or overshot by 801 5%. Thresholds (n = 100) were also created between 25 and 95% of the distance and trials were removed if they crossed 802 any of those thresholds more than once. We then computed an alignment point at 90% of the distance from the final 803 position for the remaining trials and labeled it as movement onset (227 trials). For each of these trials, data were aligned 804 to 400 ms before and 500 ms after movement onset. The first principal component of AutoLFADS rates during aligned 805 trials was computed and activation during the first 100 ms of each trial was normalized to [0,1]. Trials were rejected if 806 activation peaked after 100 ms or the starting activation was more than 3 standard deviations from the mean. The PC1 807 onset alignment point was calculated as the first time that activity in the first principal component crossed 50% of its 808 maximum in the first 100 ms (192 trials). This alignment point was used for all neural subspace analyses.

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810 Movement-relevant subspaces were extracted by ridge regression from neural activity onto x-velocity, y-velocity, and 811 speed. Similarly, position-relevant subspaces involved regression from neural activity onto x-position and y-position. For 812 movement and position subspaces, neural and behavioral data were aligned to 200 ms before and 1000 ms after PC1 813 onset. Target subspaces were computed by regressing neural activity onto time series that represented relative target 814 positions. As with the movement and position subspaces, the time series spanned 200 ms before to 1000 ms after PC1 815 onset. A boxcar window was used to confine the relative target position information to the time period spanning 0 to 200 816 ms after PC1 onset, and the rest of the window was zero-filled. For kinematic prediction from neural subspaces, we used 817 a delay of 120 ms and 80/20 trial-wise training and validation split. For each behavioral variable and neural data type, a 818 5-fold cross-validated grid search (n = 100) was used on training data to find the best-performing regularization across 819 orders of magnitude between 1e-5 and 1e4.

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Single subspace dimensions were aligned to 200 ms before and 850 ms after PC1 onset for plotting. Subspace activations were calculated by computing the norm of activations across all dimensions of the subspace and then rescaling the min and max activations to 0 and 1, respectively. Multidimensional subspace plots for the movement subspace were aligned to 180 ms before and 620 ms after PC1 onset and for target subspace 180 ms before and 20 ms after.

826 827 Area 2 bump task

The sensory dataset consisted of two recording sessions during which a monkey moved a manipulandum to direct a cursor towards one of eight targets (active trials). During passive trials, the manipulandum induced a mechanical perturbation to the monkey's hand prior to the reach. Activity was recorded via an intracortical electrode array embedded in Brodmann's area 2 of the somatosensory cortex. For the second session, joint angles were calculated from motion tracking data collected throughout the session. The first session was used for PSTH, GLM, subspace, and velocity decoding analyses and the second session was only used for pseudo-R² comparison to GLM and joint angle decoding. More details on the task and dataset are given in the original paper (39).

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836 For both sessions, only sorted units were used. Spikes were binned at 1 ms and neurons that were correlated over the 837 first 1000 sec were removed (n = 2 for each session) as described for the random target task, resulting in 53 and 68 838 neurons in the first and second sessions, respectively. Spikes were then rebinned to 5 ms and the continuous data were 839 chopped into 500 ms segments with 200 ms of overlap. Segments that did not include data from rewarded trials were 840 discarded (kept 9,626 for the first session and 7,038 for the second session). A subset of the segments (30%) were 841 further split into training and validation data (80/20) for LFADS. An AutoLFADS model (32 workers) was trained on each 842 session and a random search (96 models) was performed on the first session. After modeling, LFADS rates were then 843 reassembled into their continuous form, with linear merging of overlapping data points.

844

845 Empirical PSTHs were computed by convolving spikes binned at 1 ms with a half-Gaussian (10 ms s.d.), rebinning to 5 846 ms, and then averaging across all trials within a condition. LFADS PSTHs were computed by similarly averaging LFADS 847 rates. Passive trials were aligned 100 ms before and 500 ms after the time of perturbation, and active trials were aligned 848 to the same window around an acceleration-based movement onset (39). Neurons with firing rates lower than 1 Hz were 849 excluded from the PSTH analysis. To quantitatively evaluate PSTH reconstruction, the coefficient of determination was 850 computed for each neuron and passive condition in the four cardinal directions, and these numbers were averaged for 851 each model. 852

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As a baseline for how well AutoLFADS could reconstruct neural activity, we fit generalized linear models (GLMs) to each individual neuron's firing rate, based on the position and velocity of and forces on the hand (see Chowdhury et al., 2020 for details of the hand kinematic-force GLM). Notably, in addition to fitting GLMs using the concurrent behavioral covariates, we also added 10 bins of behavioral history (50 ms) to the GLM covariates, increasing the number of GLM parameters almost tenfold. Furthermore, because we wanted to find the performance ceiling of a behavioral-encoderbased GLMs to compare with the dynamics-based AutoLFADS, we purposefully did not cross-validate the GLMs. Instead, we simply evaluated GLM fits on data used to train the model.

- 861 To evaluate AutoLFADS and GLMs individually, we used the pseudo-R² (pR²), a goodness-of-fit metric adapted for the 862 Poisson-like statistics of neural activity. Like variance-accounted-for and R², pR² has a maximum value of 1 when a 863 model perfectly predicts the data, and a value of 0 when a model predicts as well as a single parameter mean model. 864 Negative values indicate predictions that are worse than a mean model. For each neuron, we compared the pR² of the 865 AutoLFADS model to that of the GLM (Fig 5e). To determine statistically whether AutoLFADS performed better than 866 GLMs, we used the relative-pR² (rpR²) metric, which compares the two models against each other, rather than to a mean 867 model (see Perich et al., 2018 for full description of pR^2 and rpR^2). In this case, a rpR^2 value above 0 indicated that 868 AutoLFADS outperformed the GLM (indicated by filled circles in Fig 5e). We assessed significance using a bootstrapping 869 procedure, after fitting both AutoLFADS and GLMs on the data. On each bootstrap iteration, we drew a number of trials 870 from the session (with replacement) equal to the total number of trials in the session, evaluating the rpR² on this set of 871 trials as one bootstrap sample. We repeated this procedure 100 times. We defined neurons for which at least 95 of these 872 rpR² samples were greater than 0 as neurons that were predicted better by AutoLFADS than a GLM. Likewise, neurons 873 for which at least 95 of these samples were below 0 would have been defined as neurons predicted better by GLM 874 (though there were no neurons with this result).
- 875

876 For the subspace analysis, spikes were smoothed by convolution with a Gaussian (50 ms s.d.) and then rebinned to 50 877 ms. Neural activity was scaled using the same soft-normalization approach outlined for the random target task subspace 878 analysis. Movement onset was calculated using the acceleration-based movement onset approach for both active and 879 passive trials. For decoder training, trials were aligned to 100 ms before to 600 ms after movement onset. For plotting, 880 trials were aligned to 50 ms before and 600 ms after movement onset. The data for successful reaches in the four 881 cardinal directions was divided into 80/20 trial-wise training and validation partitions. Separate ridge regression models 882 were trained to predict each hand velocity dimension for active and passive trials using neural activity delayed by 50 ms 883 (total 4 decoders). The regularization penalty was determined through a 5-fold cross validated grid search of 25 values 884 from the same range as the random target task subspace decoders. 885

- For hand velocity decoding, spikes during active trials were smoothed by convolution with a half-Gaussian (50 ms s.d.) and neural activity was delayed by 100 ms relative to kinematics. The data were aligned to 200 ms before and 1200 ms after movement onset and trials were split into 80/20 training and validation sets. Simple regression was used to estimate kinematics from neural activity and the coefficient of determination was computed and averaged across x- and y-velocity.
- 891 GPFA was performed on segments from all rewarded trials using a latent dimension of 20 and Gaussian smoothing 892 kernel (30 ms s.d.). Decoding data were extracted by aligning data from active trials to 200 ms before and 500 ms after 893 movement onset. Data were split into 80/20 training and validation sets and neural activity was lagged 100 ms behind 894 kinematics. Ridge regression ($\lambda = 0.001$) was used to decode all joint angle velocities from smoothed spikes (half-895 Gaussian, 50 ms kernel s.d.), rates inferred by GPFA, and rates inferred by AutoLFADS.
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897 DMFC timing task

The cognitive dataset consisted of one session of recordings from the dorsomedial frontal cortex (DMFC) while a monkey performed a time interval reproduction task. The monkey was presented with a "Ready" visual stimulus to indicate the start of the interval and a second "Set" visual stimulus to indicate the end of the sample timing interval, t_s . Following the Set stimulus, the monkey made a response ("Go") so that the production interval (t_p) between Set and Go matches the corresponding t_s . The animal responded with either a saccadic eye movement or a joystick manipulation to the left or right depending on the location of a peripheral target. The two response modalities, combined with 10 timing conditions 904 (t_s) and two target locations, led to a total of 40 task conditions. A more detailed description of the task is available in the 905 original paper (57).

907 To prepare the data for LFADS, the spikes from sorted units were binned at 20 ms. To avoid artifacts from correlated 908 spiking activity, we computed cross-correlations between all pairs of neurons for the duration of the experiment and 909 sequentially removed individual neurons (n = 8) by the number of above-threshold correlations until there were no pairs 910 with correlation above 0.2, resulting in 45 uncorrelated neurons. Data between the "Ready" cue and the trial end was 911 chopped into 2600 ms segments with no overlap. The first chop for each trial was randomly offset by between 0 and 100 912 ms to break any link between trial start times and chop start times. The resulting neural data segments (1659 total) were 913 split into 80/20 training and validation sets for LFADS. An AutoLFADS model (32 workers) and random search (96 914 models) were trained on these segments (see Supp. Table 2).

For all analyses of smoothed spikes, smoothing was performed by convolving with a Gaussian kernel (widths describedbelow) at 1 ms resolution.

Empirical PSTHs were computed by trial-averaging smoothed spikes (25 ms kernel s.d., 20 ms bins) within each of the
40 conditions. LFADS PSTHs were computed by similarly averaging LFADS rates. The coefficient of determination was
computed between inferred and empirical PSTHs across all neurons and time steps during the "Ready-Set" and "SetGo" periods for each condition and then averaged across periods and conditions.

To visualize low-dimensional neural trajectories, demixed principal component analysis (dPCA; Kobak et al., 2016) was performed on smoothed spikes (40 ms kernel s.d., 20 ms bins) and AutoLFADS rates during the "Ready-Set" period. The two conditions used were rightward and leftward hand movements with $t_s = 1000 ms$.

928 Besides LFADS/AutoLFADS, three alternate methods were applied for speed-tp correlation comparisons: spike 929 smoothing, GPFA, and PCA. For spike smoothing, analyses were performed by smoothing with a 40 ms s.d.. For GPFA, 930 a model was trained on the concatenated training and validation sets with a latent dimension of 9. Principal component 931 analysis (PCA) was performed on smoothed spikes (40 ms kernel s.d., 20 ms bins) and 5-7 top PCs that explained more 932 than 75% of data variance across conditions were included in the later analysis.

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934 Neural speed was calculated by computing distances between consecutive time bins in a multidimensional state space 935 and then averaging the distances across the time bins for the production epoch. The number of dimensions used to 936 compute the neural speed was 45, 5-7, 9, and 45 for smoothing, PCA, GPFA and LFADS, respectively. The Pearson's 937 correlation coefficient between neural speed and the produced time interval was computed across trials within each 938 condition.