

Interrogating theoretical models of neural computation with deep learning

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¹ 1 Abstract

² The cornerstone of theoretical neuroscience is the circuit model: a system of equations that captures
³ a hypothesized neural mechanism of scientific importance. At its best, such a model will give rise
⁴ to an experimentally observed phenomenon – whether behavioral or in terms of neural activity –
⁵ and thus can offer insight into neural computation. The operation of these circuits, like all models,
⁶ critically depends on the choices of model parameters. Historically, the gold standard has been
⁷ to analytically derive the relationship between model parameters and computational properties.
⁸ However, this enterprise quickly becomes infeasible as biologically realistic constraints are included
⁹ into the model, often resulting in *ad hoc* approaches to understanding the relationship between
¹⁰ model and computation. We bring recent machine learning techniques – the use of deep generative
¹¹ models for probabilistic inference – to bear on this problem, learning distributions of parameters
¹² that produce the specified properties of computation. Importantly, the techniques we introduce
¹³ offer a logical and unbiased means to understand the implications of model parameter choices
¹⁴ on computational properties of interest. We motivate this methodology with a worked example
¹⁵ analyzing sensitivity in the stomatogastric ganglion. We then use it to generate insights into neuron-
¹⁶ type input-responsivity in primary visual cortex, a new understanding of rapid task switching in
¹⁷ superior colliculus models, and improved attribution of bias in low-rank recurrent neural networks.
¹⁸ More generally, this work moves us away from the tradeoff of biological realism vs analytical
¹⁹ tractability, and towards the use of modern machine learning for sophisticated interrogation of
²⁰ biologically relevant models.

²¹ 2 Introduction

²² The fundamental practice of theoretical neuroscience is to use a mathematical *model* to understand
²³ neural computation, whether that computation enables perception, action, or some intermediate
²⁴ processing [1]. In this field, a neural computation is systematized with a set of equations – the
²⁵ model – and these equations are motivated by biophysics, neurophysiology, and other conceptual
²⁶ considerations. The function of this system is governed by the choice of model parameters, which

27 when configured in some special way, give rise to some measurable signature of a computation. The
28 work of analyzing a model then becomes the inverse problem: given a computation of interest, how
29 can we reason about these special parameter configurations – their likely values, their uniquenesses
30 and degeneracies, their attractor states and phase transitions, and more?

31 Consider the idealized practice: a theorist considers a model carefully and analytically derives how
32 model parameters govern the computation. Seminal examples of this gold standard include our
33 field’s understanding of memory capacity in associative neural networks [2], chaos and autocorrela-
34 tion timescales in random neural networks [3], and the paradoxical effect in excitatory/inhibitory
35 networks [4]. Unfortunately, as circuit models include more biological realism, theory via analytic
36 derivation becomes intractable. This fact creates an unfavorable tradeoff for the theorist. On the
37 one hand, one may tractably analyze systems of equations with unrealistic assumptions (for ex-
38 ample symmetry or gaussianity), producing accurate inferences about parameters of a too-simple
39 model. On the other hand, one may choose a more biologically relevant model at the cost of *ad hoc*
40 approaches to analysis (simply examining simulated activity), producing questionable or partial
41 inferences about parameters of an appropriately complex, scientifically relevant model.

42 Of course, this same tradeoff has been confronted in many scientific fields and engineering problems
43 characterized by the need to do inference in complex models. In response, the machine learning
44 community has made remarkable progress in recent years, via the use of deep neural networks as a
45 powerful inference engine: a flexible function family that can map observed phenomena (in this case
46 the measurable signal of some computation) back to probability distributions quantifying the likely
47 parameter configurations. One celebrated example of this approach from the machine learning
48 community, from which we draw key inspiration for this work, is the variational autoencoder [5, 6],
49 which uses a deep neural network to induce an (approximate) posterior distribution on hidden
50 variables in a latent variable model, given data. Indeed, these tools have been used to great success
51 in neuroscience as well, in particular for interrogating parameters (sometimes treated as hidden
52 states) in models of both cortical population activity [7, 8, 9, 10] and animal behavior [11, 12, 13].
53 These works have used deep neural networks to expand the expressivity and accuracy of statistical
54 models of neural data [14].

55 However, these inference tools have not significantly influenced the study of theoretical neuroscience
56 models for at least three reasons. First, at a practical level, the nonlinearities and dynamics of many
57 theoretical models are such that conventional inference tools (for example mean field variational
58 inference) typically produce a narrow set of insights into these models [15]. Indeed, only in the last

59 few years has the deep learning toolkit expanded to a point of relevance to this class of problem.
60 Second, the object of interest from a theoretical model is not typically data itself, but rather a
61 qualitative phenomenon – inspection of model behavior, or better, a measurable signature of some
62 computation – an *emergent property* of the model. Third, because theoreticians work carefully to
63 construct a model that has biological relevance, such a model as a result often does not fit cleanly
64 into the framing of a statistical model. Technically, because many such models stipulate a noisy
65 system of differential equations that can only be sampled or realized through forward simulation,
66 they lack the explicit likelihood and priors central to the probabilistic modeling toolkit.

67 To address these three challenges, we developed an inference methodology – ‘emergent property
68 inference’ – which learns a distribution over parameter configurations in a theoretical model. Crit-
69 ically, this distribution is such that draws from the distribution (parameter configurations) corre-
70 spond to systems of equations that give rise to a specified emergent property. First, we stipulate a
71 deep neural network that induces a flexible family of probability distributions over model param-
72 eterizations. We will insist on being able to quantify the probability of various model parameter
73 configurations, and thus we choose the deep neural network to be of the (bijective) normalizing flow
74 class [16]. Second, we quantify the notion of emergent properties as a set of moment constraints
75 on datasets generated by the model. Thus an emergent property is not a single data realization,
76 but a phenomenon or a feature of the model, which is the central object of interest to the theorist
77 (unlike say the statistical neuroscientist). The requirement to condition on an emergent property
78 requires the adaptation of deep probabilistic inference methods, and we extend recent tools to do
79 so [17]. Third, because we can not assume the theoretical model has explicit likelihood on data
80 or the emergent property of interest, we use stochastic gradient techniques in the spirit of likeli-
81 hood free variational inference [18]. Taken together, emergent property inference (EPI) provides
82 a methodology for inferring and then reasoning about parameter configurations that give rise to
83 particular emergent phenomena in theoretical models. Emergent property inference is described
84 schematically in Fig 1A.

85 Equipped with this methodology, we investigated four models of historical and current importance
86 in theoretical neuroscience. These models were chosen to demonstrate generality through ranges
87 of biological realism (conductance-based biophysics to recurrent neural networks), neural system
88 function (pattern generation to abstract cognitive function), and network scale (four to infinite
89 neurons). First, to motivate the contribution of emergent property inference, we investigated
90 network syncing in a classic model of the stomatogastric ganglion [19]. Second, we conducted a

91 exploratory analysis of a four neuron-type dynamical model of primary visual cortex resulting in
92 testable predictions of nonlinear population input-responsivity. Third, we demonstrated how the
93 systematic application of EPI to levels of behavioral accuracy can generate experimentally testable
94 hypotheses regarding connectivity in superior colliculus. Fourth, we leveraged the flexibility of EPI
95 to uncover the sources of bias in a low-rank recurrent neural network executing Bayesian inference.
96 The novel scientific insights offered by EPI contextualize and clarify the previous studies exploring
97 these models [19, 20, 21, 22] and more generally offer a quantitative grounding for theoretical
98 models going forward, pointing a way to how rigorous statistical inference can enhance theoretical
99 neuroscience at large.

100 We note that, during our preparation and early presentation of this work [23, 24], another work
101 has arisen with broadly similar goals: bringing statistical inference to mechanistic models of neu-
102 ral circuits [25]. We are excited by this broad problem being recognized by the community, and
103 we emphasize that these works offer complementary neuroscientific contributions and use different
104 technical methodologies. Scientifically, our work has focused primarily on systems-level theoretical
105 models, while their focus is on lower-level cellular models. Secondly, there are several key technical
106 differences in the approaches (see Section A.1.4) perhaps most notably is our focus on the emer-
107 gent property – the measurable signal of the computation in question, vs their focus on observed
108 datasets; both certainly are worthy pursuits. The existence of these complementary methodologies
109 emphasizes the increased importance and timeliness of both works.

110 3 Results

111 3.1 Motivating emergent property inference of theoretical models

112 Consideration of the typical workflow of theoretical modeling clarifies the need for emergent prop-
113 erty inference. First, the theorist designs or chooses an existing model that, it is hypothesized,
114 captures the computation of interest. To ground this process in a well-known example, consider
115 the stomatogastric ganglion (STG) of crustaceans, a small neural circuit which generates multiple
116 rhythmic muscle activation patterns for digestion [?]. A standard model for the STG is shown
117 schematically in Figure 1A, and note that the behavior of this model will be critically dependent
118 on its parameterization – the choices of conductance parameters $z = [g_{el}, g_{synA}]$. Second, the the-
119 orist defines the emergent property, the measurable signal of scientific interest. To continue our
120 running STG example, one such emergent property is the phenomenon of *network syncing* – in

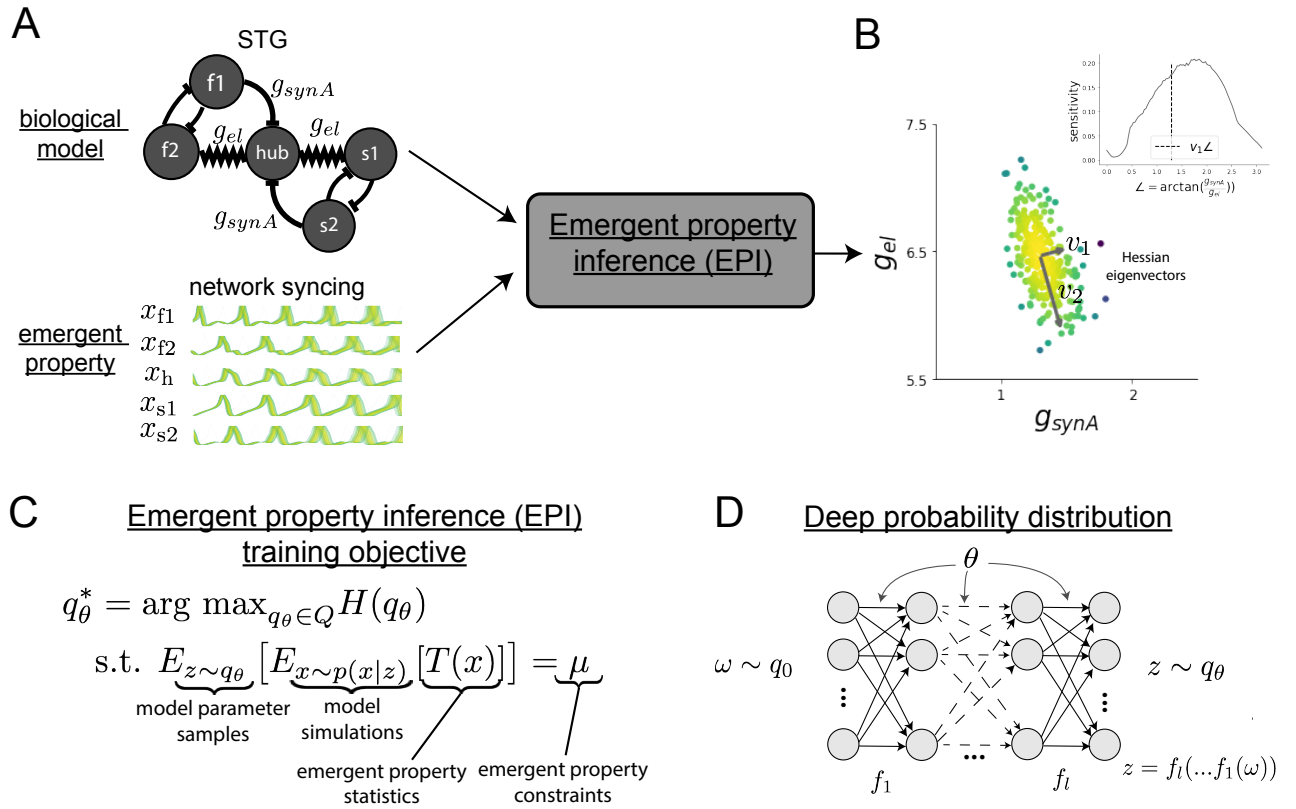


Figure 1: A. For a choice of model (STG) and emergent property (network syncing), emergent property inference (EPI) learns a posterior distribution of the model parameters $z = [g_{el}, g_{synA}]^\top$ conditioned on network syncing. B. An EPI distribution of STG model parameters producing network syncing. (Inset) Sensitivity of the system with respect to network syncing along all dimensions of parameter space away from the mode. C. EPI solves a constrained stochastic optimization, in which emergent property statistics $T(x)$ are fixed in expectation over model simulations $x \sim p(x | z)$ and parameter distribution samples $z \sim q_\theta(z)$ to be a particular value μ . EPI distributions maximize randomness via entropy, although other measures are sensible. D. Degenerate solution networks (DSNs) are deep probability distributions $q_\theta(z)$ of theoretical model parameterizations that produce emergent properties of interest. The stochasticity of a deep probability distribution comes from a simple random variable $\omega \sim q_0$, where q_0 is often chosen to be an isotropic gaussian, and the structure comes from the deterministic transformation made by the deep neural network with optimized parameters θ .

certain parameter regimes, the activity of the hub neuron x_h matches that of the fast (x_{f1}, x_{f2}) and slow (x_{s1}, x_{s2}) populations. This emergent property is shown in Figure 1A. Third, qualitative parameter analysis ensues: since precise mathematical analysis is intractable in this model, a brute force sweep of parameters is done to describe different parameter configurations that lead to the emergent property. In this last step lies the opportunity for modern machine learning. We can precisely quantify the emergent property as a statistical feature of the model, and we can infer a probability distribution over parameter configurations that produce this emergent property.

Before presenting technical details (in the following section), let us understand emergent property inference schematically: the black box in Figure 1A takes, as input, the model and the specified emergent property, and produces as output the parameter distribution shown in Figure 1B. This distribution – represented for clarity as samples from the distribution – is then a scientifically meaningful and mathematically tractable object. It conveys parameter regions critical to the emergent property, directions in parameter space that will be invariant (or not) to that property, and more. In the STG model, this distribution can be specifically queried to determine the prototypical parameter configuration for network syncing (the mode; Figure 1B star), and then how quickly network syncing will decay based on changes away that mode (Figure 1B, inset). Of course, validation of this distribution is critical: we can also plot a metric of network syncing that is independent of this inference process, to show that this EPI distribution indeed captures that structure (Figure 1B, contour lines). Taken together, bringing careful inference to theoretical models offers deeper insight into the behavior of these models, and the opportunity to make rigorous this last analysis step of the practice of theoretical neuroscience.

3.2 A deep generative modeling approach to emergent property inference

Emergent property inference (EPI) systematizes the three-step procedure of the previous section. First, we consider the model as a coupled set of differential (and potentially stochastic) equations. In the running STG example the dynamical state $x = [x_{f1}, x_{f2}, x_{hub}, x_{s1}, x_{s2}]^\top$ is the membrane potential for each neuron, which evolves according to the biophysical conductance-based equation:

$$C_m \frac{\partial x}{\partial t} = -[h_{leak} + h_{Ca} + h_K + h_{hyp} + h_{elec} + h_{syn}] \quad (1)$$

where $C_m=1\text{nF}$, and h_{leak} , h_{Ca} , h_K , h_{hyp} , h_{elec} , h_{syn} are the leak, calcium, potassium, hyperpolarization, electrical, and synaptic currents, all of which have their own complicated dependence on x and θ (see Methods) [THIS NEEDS TO BE CLARIFIED]. Second, we define the emergent

property, which as above is network syncing: the phase locking of the population and its oscillation at an intermediate frequency (Figure 1A bottom) [REPETITIVE]. It is worth noting that theoretical work has characterized how model parameters such as the electrical conductance g_{el} and synaptic conductance g_{synA} govern the production of STG rhythms [19]. In this 5-neuron model, two fast neurons ($f1$ and $f2$) mutually inhibit one another, and oscillate at a faster frequency than the slow neurons ($s1$ and $s2$), which also mutually inhibit each other (Figure 1A). Quantifying this phenomenon is straightforward: we define [XXXXXXXXXX statistics, values, and we insist on these being the measurable signatures of the EP... End up with a constraint function $E(T(X)) = \mu$ XXXXXXXXXXXXXX]. Third, having rationalized the above components, we can introduce deep generative modeling for performing emergent property inference. We seek a distribution over parameter configurations z , and we insist that samples from this distribution produce the emergent property; in other words, they obey the constraints introduced in Equation 1. Thus results the following optimization program:

$$\begin{aligned} & \underset{q_\theta \in \mathcal{Q}}{\operatorname{argmax}} H(q_\theta(z)) \\ & \text{s.t. } E_{z \sim q_\theta} [E_{x \sim p(x|z)} [T(x)]] = \mu \end{aligned} \tag{2}$$

The purpose of each element in this program is detailed in Figure 1D. Stating such a problem is easy enough; finding a tractable and suitably flexible family of probability distributions (\mathcal{Q}) is hard. We use norm flow deep learning [DETAIL HERE], inducing the deep distribution Fig 1E. Finally, we recognize that many distributions will satisfy the emergent property, so we require a normative principle to select amongst them. This principle is captured in Equation [XXX] by the primal objective H . Here we chose Shannon entropy as it has been well used for a variety of things [Elsayed, MEFN, Jaynes, Savin review], but the EPI methods (not the results) offered here are largely unaffected by this choice.

[A few sentences about ... we run optimization... learning.... and then what results is the EPI distribution voila.] Armed with this distribution, we now prove out the value of this technology by investigating a range of models and using EPI to produce novel scientific insights.

3.3 Comprehensive input-responsivity in a nonlinear sensory system

First, dynamical models with two populations (excitatory (E) and inhibitory (I) neurons) of visual processing have been used to reproduce a host of experimentally documented phenomena in primary visual cortex (V1). In particular regimes of excitation and inhibition, these models exhibit the

178 paradoxical effect [4], selective amplification [26], surround suppression [27], and sensory integrative
 179 properties [28]. Since inhibitory neurons mostly fall into one of three classes (parvalbumin (P)-,
 180 somatostatin (S)-, and vasointestinal peptide (V)-expressing neurons) [29, 30], theorists look to
 181 extend these dynamical models to four populations [20] (Fig. 3A).

182 The dynamical state of this model is the firing rate of each neuron type population $x = [x_E \ x_P \ x_S \ x_V]^T$,
 183 which evolves according to rectified, exponentiated dynamics:

$$\tau \frac{dx}{dt} = -x + [Wx + h]_+^n \quad (3)$$

184 with effective connectivity weights W and input h . We set the time constant $\tau = 20ms$ and
 185 dynamics coefficient $n = 2$. We obtained an informative estimate of the effective connectivities
 186 between these neuron types in mice by multiplying their probability of connection by the synaptic
 187 efficacy [?] (see Section A.2.2). Given these fixed parameter choices of W , n , and τ and a baseline
 188 input b , we asked what differential inputs $dh = [dh_E \ dh_P \ dh_S \ dh_V]^T$ cause each neuron type
 189 population to increase its firing rate.

190 Yet, we were at least able to derive the linearized response of the system $\frac{dx_{ss}}{dh}$ at fixed points x_{ss} .
 191 While this linear prediction is accurate for small differential inputs (Fig. 3B, left), it is often mislead-
 192 ing in such nonlinear models as differential input strength increases (Fig. 3B, right). In fact, for a
 193 baseline input of $b = [1 \ 1 \ 1 \ 1]^T$ the linearly predicted response for $dh = [0.5 \ 0.5 \ 0.5 \ 0.5]^T$
 194 was actually in the opposite direction of the true response for the V-population (Fig. 3B, right,
 195 green).

196 To get a more comprehensive picture of the input-responsivity of each neuron type, we used EPI to
 197 learn a distribution of inputs dh that cause the rate of each neuron-type population α to increase by
 198 a value y with some allowed variance. We denote this emergent property of neuron-type responsivity
 199 as $\mathcal{B}(\alpha, y)$. In Fig. 3C, each column visualizes the approximate posterior dh corresponding to a
 200 specific neuron-type increase, while each row corresponds to amounts of increase 0.1 and 0.5. Akin
 201 to the practice of exploratory analysis of neural data sets, in which we search for meaningful
 202 structure, we can do an exploratory analysis of this *model* yielding a meaningful understanding of
 203 its operation.

204 To visualize these four-dimensional distributions, we pairplotted the two-dimensional marginal den-
 205 sities, which yield an insightful picture. As expected, the inferred distributions revealed that each
 206 neuron-type's rate is sensitive to its direct input. The E- and P-population are largely unaffected
 207 by dh_V , and the S-population by dh_P for the rate increases examined. Additionally, we observed

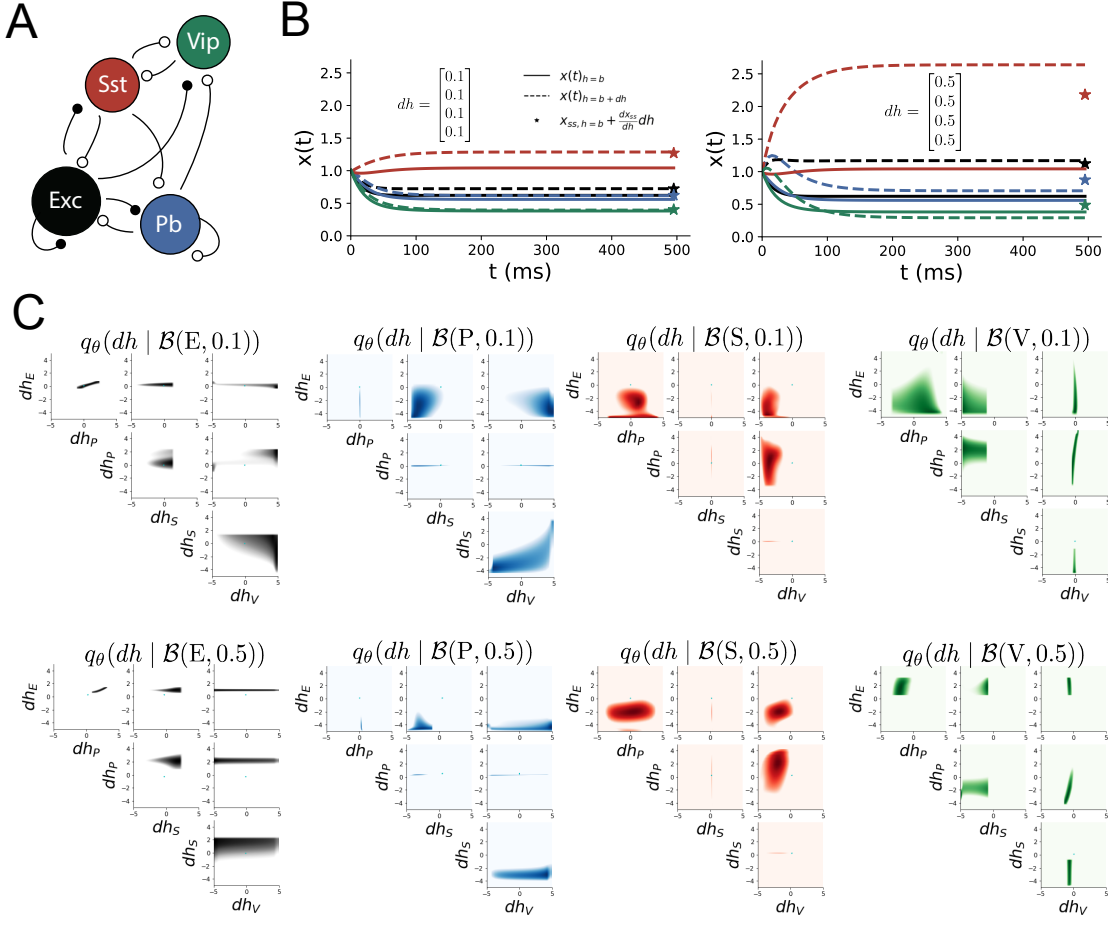


Figure 2: A. Four-population model of primary visual cortex with excitatory (black), parvalbumin (blue), somatostatin (red), and vip (green) neurons. Some neuron types largely do not form synaptic projections to others. Excitatory and inhibitory projections filled and unfilled, respectively. B. Linear response predictions become inaccurate with greater input strength. V1 model simulations for input (solid) $h = b$ and (dashed) $h = b + dh$ with $b = [1, 1, 1, 1]^\top$ and (left) $dh = [0.1, 0.1, 0.1, 0.1]^\top$ (right) $dh = [0.5, 0.5, 0.5, 0.5]^\top$. Stars indicate the linear response prediction. C. EPI distributions on dh conditioned on neuron type α rate increases of y : $\mathcal{B}(\alpha, y)$ (see Section A.2.2).

208 that the P-population likely stabilizes the E- and V-populations (the two it projects to besides
 209 itself). This deduction is supported by the tight positive correlation between dh_P and dh_E for
 210 E-population increases and also between dh_P and dh_V for V-population increases. Finally, EPI
 211 showed that negative dh_E results in small inhibitory firing rate increases. However, for a larger
 212 increase from the V-population, a positive dh_E is required.

213 All of this insight was gained beyond what the analytic linear prediction told us (cyan lines that you
 214 can't see). These analyses can generate experimentally testable hypotheses, that if confirmed, can
 215 be used to build a theory of V1 circuit operation. For example, one could test the P-population's
 216 role in stabilization by optogenetically stimulating the E- and V-populations and measuring the
 217 P-population response. Additionally, one would predict that at some point, the V-population's
 218 response will flip sign as input to the E-population increases.

219 **3.4 Identifying neural mechanisms of behavioral learning.**

220 A key challenge for theorists modeling neural circuits underlying cognitive behavior is the descrip-
 221 tion of sufficient changes to biologically meaningful parameters that result in improved behavoir.
 222 Identifying measurable biological changes that should occur for increased performance is critical
 223 for neuroscience, since they may indicate how the learning brain adapts. We used EPI to learn
 224 connectivities distributions consistent with various levels of rapid task switching accuracy, resulting
 225 in a clear picture of connectivity changes which improve rapid task switching. Furthermore, this
 226 analysis produced experimentally testable predictions regarding effective connectivity throughout
 227 learning of this behavioral paradigm.

228 In a rapid task switching experiment, where rats were to respond right (R) or left (L) to the
 229 side of a light stimulus in the pro (P) task, and oppositely in the anti (A) task predicated by an
 230 auditory cue (Fig. 4A), neural recordings exhibited two population of neurons in each hemisphere
 231 of superior colliculus (SC) that simultaneously represented both task condition and motor response:
 232 the pro/contra and anti/ipsi neurons [21]. Duan et al. proposed a four-population dynamical model
 233 of superior colliculus with a Pro- and Anti-population in each hemisphere, where activities were
 234 bounded from 0-1, and a high output of the Pro population in a given hemisphere corresponds
 235 to the contralateral response. The connectivity matrix is parameterized by the geometry of the
 236 population arrangement (Fig. 4B).

237 We ran EPI to learn appproximate posteriors of SC model weight matrix parameters $z = W$ condi-

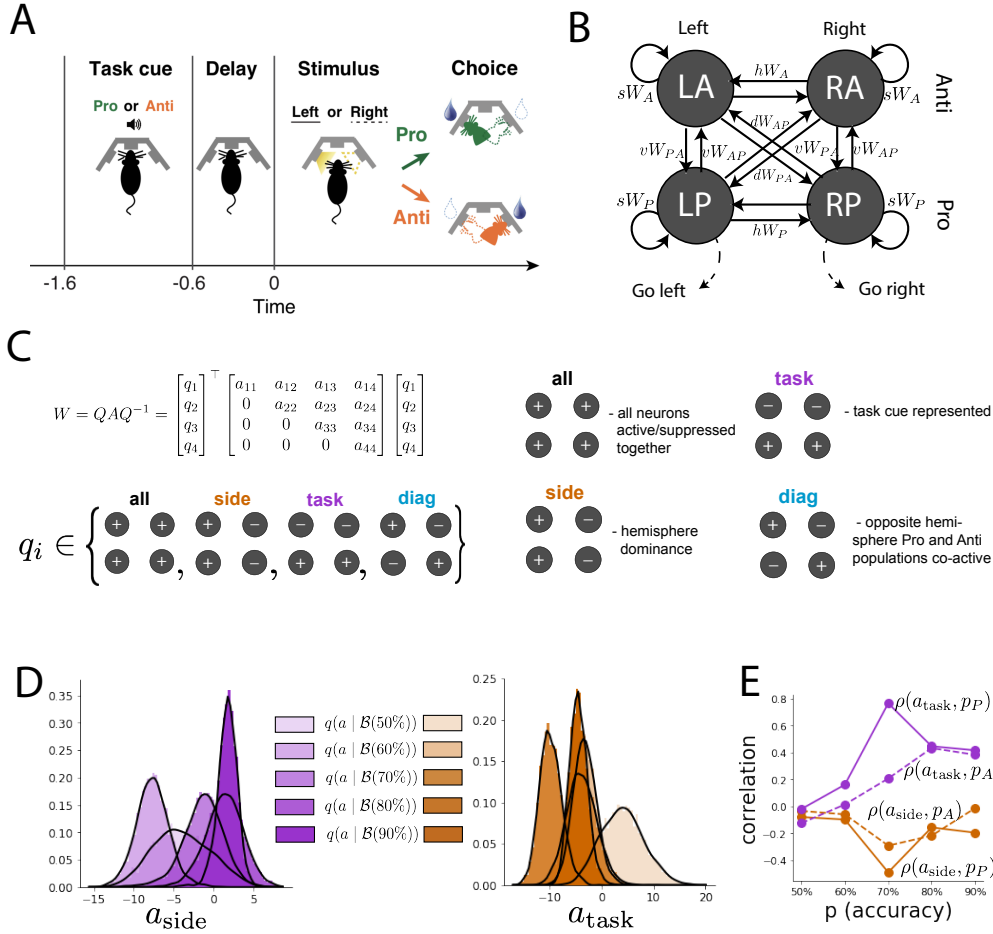


Figure 3: A. Rapid task switching behavioral paradigm. In the Pro (Anti) condition indicated by an auditory cue, the rats are to respond to the same (opposite) side as the light stimulus that is provided after a delay to receive a reward. B. Model of superior colliculus (SC). Neurons: LP - left pro, RP - right pro, LA - left anti, RA - right anti. Parameters: sW - self, hW - horizontal, vW - vertical, dW - diagonal weights. C. The Schur decomposition of the weight matrix. D. The marginal EPI distributions of the Schur eigenvalues at each level of task accuracy. E. The correlation of Schur eigenvalue with task performance.

tioned on of various levels of rapid task switching accuracy $\mathcal{B}(p)$ for $p \in \{50\%, 60\%, 70\%, 80\%, 90\%\}$ (see Section A.2.3). The Schur decomposition of the weight matrix, a unique decomposition revealing the underlying directed structure between modes (the eigenvectors), has the same eigenvectors for all W s under this symmetric parameterization (Fig. 4C). These consistent Schur eigenvectors have intuitive roles in processing for this task, and are accordingly named the *all*, *side*, *task*, and *diag* modes. The corresponding eigenvalues a of each mode (which change according to W indicate the amplification or suppression of activity along that mode).

As learning progresses, the task mode is increasingly amplified, indicating the criticality of a distributed task representation at the time of stimulus presentation, (Fig. 4D, purple). Stepping from task-naive 50% networks to task-performing 60% networks, there is a marked suppression of the side mode (Fig. 4D, orange). Such side mode suppression remains in the regimes of greater accuracy, revealing its importance towards the existence of a distributed task representation. There were no interesting trends with learning in the all or diag mode. We can conclude that side mode suppression allows rapid task switching, and that greater task-mode representation increases accuracy (Fig. 4E). These findings motivate experimental predictions, in which we would expect the effective connectivity between these populations to change throughout learning in a way that increases the task mode and decreases the side mode eigenvalues.

3.5 Characterizing the sources of bias during approximate inference in RNNs

At a more abstract level, recurrent neural networks (RNNs) are high-dimensional models of computation, which have become increasingly popular in systems neuroscience research [32]. Typically, RNNs are trained to do a task from a systems neuroscience experiment, and then the latent factors of the trained RNN are compared to recorded neural activity. Recent theoretical work extends dynamic mean field theory (DMFT) from random [3] to low rank RNNs [22]. This theory establishes a link between interpretable, geometric parameterizations of the RNN connectivity with the emerging dynamics. We used this theory along with EPI to characterize the mechanistic sources of bias during approximate Bayesian inference in rank-1 RNNs.

The connectivity of a rank-1 RNN is the sum of a random component with strength determined by g , and a structured component determined by the outer product of vectors m and n :

$$J = g\chi + \frac{1}{N}mn^\top \quad (4)$$

where $\chi_{ij} \sim \mathcal{N}(0, \frac{1}{N})$ and $m_i \sim \mathcal{N}(M_m, 1)$ and $n_i \sim \mathcal{N}(M_n, 1)$. The rank-1 RNNs were to produce

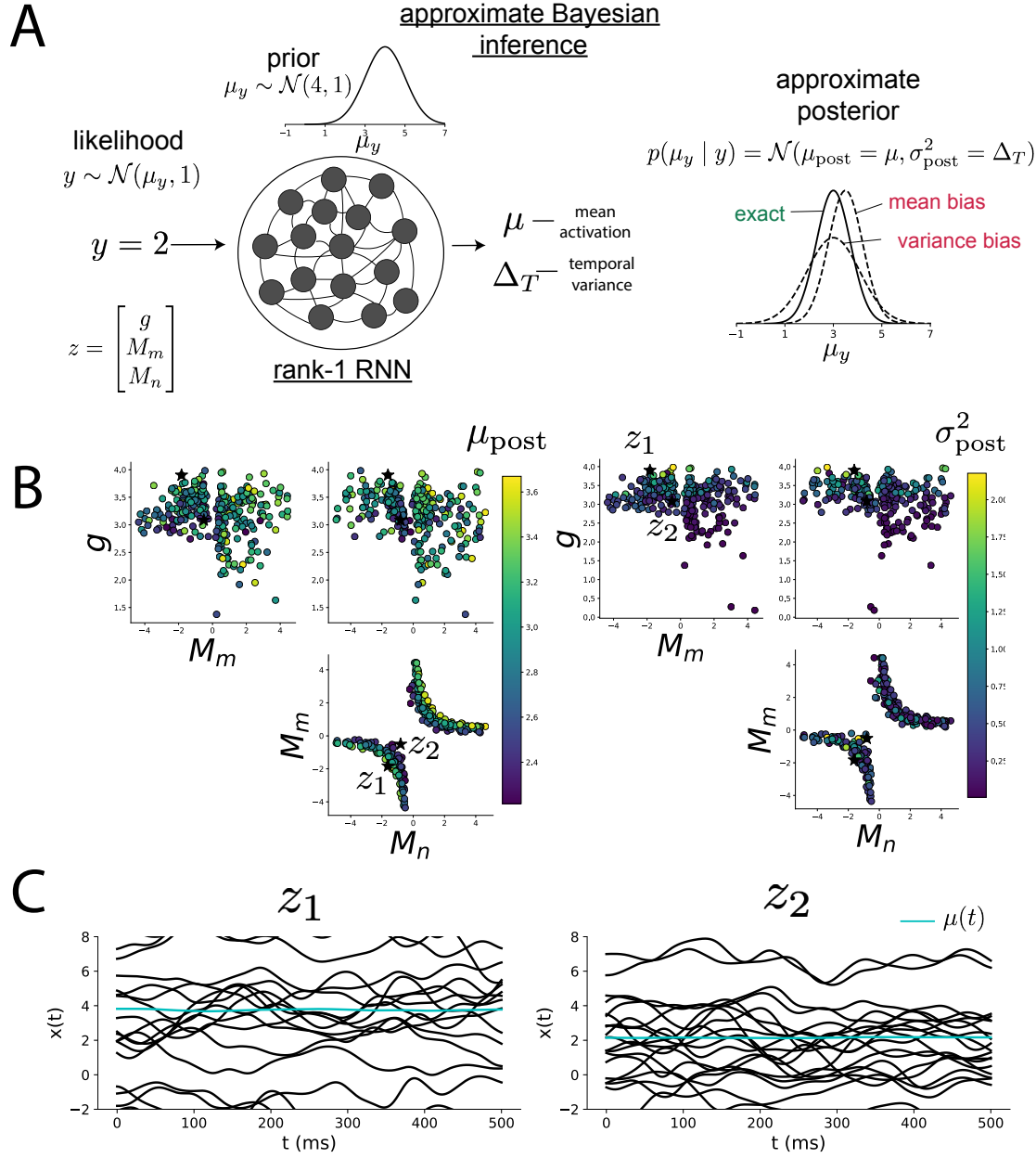


Figure 4: A. We examined a rank-1 RNN running approximate Bayesian inference on μ_y assuming a gaussian likelihood variance of 1 and a prior of $\mathcal{N}(4, 1)$. B. Distribution of rank-1 RNNs executing approximate Bayesian inference. Samples are colored by (left) posterior mean $\mu_{\text{post}} = \mu$ and (right) posterior variance $\sigma_{\text{post}}^2 = \Delta_T$. C. Finite size realizations agree with the DMFT theory.

the posterior mean μ_{post} and variance σ_{post}^2 in their mean activity μ and temporal variance Δ_T (see Section A.2.4). The Bayesian inference problem was to produce the gaussian posterior of the mean μ_y of a gaussian likelihood of observations $y \sim \mathcal{N}(\mu_y, 1)$ given a single observation of $y = 2$ and a prior of $\mu_y \sim \mathcal{N}(4, 1)$ (Fig. 5A). The true posterior to this problem is $\mu_x \sim \mathcal{N}(\mu_{\text{post}} = 3, \sigma_{\text{post}}^2 = 0.5)$, although different parameterizations of the connectivity $z = [g \ M_m \ M_n]^\top$ result in approximate inference procedures of varying biases in μ_{post} and σ_{post}^2 .

Weran EPI on rank-1 RNNs solving this Bayesian inference problem, while allowing a substantial amount of variability in the second moment constraints of the network mean μ and temporal variance Δ_T . This allowed us to study the mechanistic sources of bias in the sampled rank-1 RNNs executing the Bayesian inference computation inexactly. The posterior distribution was roughly symmetric in the M_m - M_n plane which structure suggesting there is a degeneracy in the product of M_m and M_n (Fig. 5B). The product of M_m and M_n almost completely determines the posterior mean (Fig. 5B, left), and the random strength g is the most influential variable on the temporal variance (Fig. 5B, right). Neither of these observations were obvious from the consistency equations afforded by DMFT, the solvers of which we took gradients through to run EPI.

It is good practice to check that finite-size realizations of these infinite-size networks match the predictions from DMFT. A 2,000-neuron network with parameters z_1 produced an overestimate of the posterior mean and variance (mean activity (cyan), temporal variance of traces Fig. 5C, left), while a 2,000-neuron network with parameter z_2 produced underestimates (Fig. 5C, right). This novel procedure of doing inference in interpretable parameterizations of RNNs conditioned on abstract cognitive tasks can be generally executed on other tasks like noisy integration and context-dependent decision making (Fig. S1).

3.6 EPI is a general tool for all theoretical neuroscience

There are many considerations when assessing the feasibility of classical statistical inference in neuroscience such as conjugacy, likelihood tractability, scalability to large data-sets. EPI is no exception, but we emphasize that it is practicable in most settings of theoretical neuroscience. Models of close biological fidelity often have complex nonlinear differential equations, making traditional statistical inference intractable. In contrast, EPI is capable of learning distributions of such model parameters producing low-level signatures of computation as we have shown through our analysis of the STG. This approach is not specific to models of such biological realism, as we have demonstrated its utility studying abstract models like RNNs. We are able to condition both deterministic

298 and stochastic models on all sorts of emergent properties from membrane potential firing to exe-
299 cution of approximate inference. In sum, EPI is feasible when the emergent property statistics are
300 continuously differentiable with respect to the model parameters, which is very often the case. This
301 gradient does not even need to be derived by hand, since we use automatic differentiation tools
302 available in tensorflow[33] and other similar software packages.

303 In this study, we have focused on applying EPI to low dimensional parameter spaces of models
304 with low dimensional dynamical state. These choices were made to present the reader with a
305 series of interpretable conclusions, which is more challenging in high dimensional spaces. In fact,
306 EPI scales well to high dimensional parameter spaces, as the underlying technology has produced
307 state-of-the-art performance on texture generation [17]. However, increasing the dimensionality
308 of the dynamical state of the model makes optimization more expensive, and there is a practical
309 limit there as with any machine learning approach. For systems with high dimensional state, we
310 recommend using theoretical approaches (e.g. [22]) to reason about reduced parameterizations of
311 such high-dimensional (even infinite dimensional) systems.

312 There are additional technical considerations when assessing the suitability of EPI for a particular
313 modeling question. First, one should consider how computationally expensive the gradient of the
314 emergent property statistic is with respect to the model parameters. In the best circumstance, there
315 is a simple, closed form expression (e.g. Section A.1.1) for the emergent property statistic given
316 the model parameters. On the other end of the spectrum, you may require a large number of sim-
317 ulation iterations before a high quality measurement of the emergent property statistic is available
318 (e.g. Section A.2.1). In such cases, optimization will be expensive, and it is worth considering
319 an alternative methodology (see Section A.1.4). Secondly, the defined emergent property should
320 always be appropriately conditioned. Of course, learning can not occur when under- or overcon-
321 strained. When underconstrained, the posterior grows (in entropy) unstably unless mapped to a
322 finite support. If overconstrained, and there is no support producing the emergent property, EPI
323 optimization will never converge.

324 4 Discussion

325 **Draft in progress:**

326 Machine learning has played an effective, multifaceted role in neuroscientific progress. Primarily,
327 has revealed structure in large scale neural data sets [34, 35, 36, 37, 38, 39] (see review, [14]).

328 Secondarily, trained algorithms of varying degrees of biological relevance serve as fully-observable
329 computational systems that are compared to the brain [40, ?]. Theoretical neuroscientists may cur-
330 rently be too focused on this secondary role, and thus missing out on the primary benefit statistical
331 machine learning can offer. Specifically, deep learning for probabilistic inference has matured to a
332 level where theorists can use it to understand their *models* rather than the experimental *data sets*
333 for which it is traditionally purposed.

334 For example, consider the fact that we do not yet understand just a four-dimensional, deterministic
335 model of V1. This should not be surprising, since analytic approaches to studying nonlinear dynam-
336 ical systems complexify greatly when stepping from two-dimensional to three- or four-dimensional
337 systems in the absence of restrictive simplifying assumptions [31]. We are not suggesting to forego
338 the development or application of challenging analytic procedures in theoretical neuroscience. How-
339 ever, we suggest the judicious recognition of arduous mathematical challenges, and alternatively
340 using deep learning through EPI to gain the desired insights. In Section ??, we showed that EPI was
341 far more informative about neuron-type input responsivity than the predictions afforded through
342 analysis. By flexibly conditioning this V1 model on different emergent properties, we performed an
343 exploratory analysis of a *model* rather than a data set, which generated a set of testable predictions.

344 Exploratory analyses in theoretical neuroscience can certainly be less agnostic to the eventual hy-
345 pothesis generated. When interested in the mechanistic changes that occur in the brain throughout
346 learning, one can use EPI to condition on various levels of an emergent property statistic indicative
347 of performance like task accuracy in a behavioral paradigm (see Section ??). This analysis iden-
348 tified experimentally testable predictions of changes in connectivity in SC throughout rapid task
349 switching learning. Precisely, we would predict an initial reduction in side mode eigenvalue, and a
350 steady increase in task mode eigenvalue of such effective connectivity matrices.

351 While experimentally testable predictions are highly valuable, sometimes it is prohibitively chal-
352 lenging to design a biologically realistic model of a neural computation. Thusly, RNNs have become
353 an increasingly popular tool in systems neuroscience research. The scientific philosophy is as fol-
354 lows: optimize an RNN to execute a task from behavioral neuroscience, compare the activity of
355 this optimized system to brain activity from a model organism doing the same task, and lever-
356 age the full observability of the trained RNN to generate hypotheses of the neural mechanisms of
357 computation. While fixed point identification and jacobian measurement yield intuitive, consistent
358 portraits of the implemented computational algorithm [?], there is dizzying degeneracy in the RNN
359 connectivity matrix with respect to these characterizations. Since neural activity generally lies on

360 a low dimensional manifold [?], we may attain an understanding of the neural mechanisms at play
361 in cortical processing by working in a reduced, interpretable parameter setting of these powerfully
362 general models [42].

363 In our final analysis, we present a novel procedure for doing statistical inference on interpretable
364 parameterizations of RNNs executing tasks from behavioral neuroscience. This methodology relies
365 on recently extended theory of responses in random neural networks with minimal structure [22].
366 This theory makes a direct link between a geometric description of the connectivity and the emerging
367 dynamics. These emerging dynamics in response to various inputs can be cast to perform noisy
368 detection, context-dependent evidence integration and more. With this methodology, we can finally
369 open the probabilistic model selection toolkit reasoning about the connectivity of RNNs solving
370 tasks.

371 Some statements to inject somewhere

- 372 • If theory/practice of deep learning improves to a point where we converge to global optima
373 more regularly, we could do cool reasoning about models (the hypothetico-deductive stuff
374 from Gelman/Shalizi) using these max-ent distributions.
- 375 • We can think of the probability in these models as deviation from the mean constraint in the
376 sufficient statistics. See this proved out with the 2D LDS example.

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501 **A Methods**

502 **A.1 Emergent property inference (EPI)**

503 **Draft in progress:**

504 Emergent property inference (EPI) learn distributions of theoretical model parameters that produce
 505 emergent properties of interest. They combine ideas from likelihood-free variational inference [18]
 506 and maximum entropy flow networks [17]. A maximum entropy flow network is used as a deep
 507 probability distribution for the parameters, while these samples are passed through a differentiable
 508 model simulator, which may lack a tractable likelihood function.

509 Consider model parameterization z and data x generated from some theoretical model simulator
 510 represented as $p(x | z)$, which may be deterministic or stochastic. Theoretical models usually have
 511 known sampling procedures for simulating activity given a circuit parameterization, yet often lack
 512 an explicit likelihood function due to the nonlinearities and dynamics. With EPI, a distribution
 513 on parameters z is learned, that yields a behavior of interest \mathcal{B} ,

$$\mathcal{B} : E_{z \sim q_\theta} [E_{x \sim p(x|z)} [T(x)]] = \mu \quad (5)$$

514 by making an approximation $q_\theta(z)$ to $p(z | \mathcal{B})$ (see Section A.1.5). So, over the DSN distribution
 515 $q_\theta(z)$ of model $p(x | z)$ for behavior \mathcal{B} , the emergent properties $T(x)$ are constrained in expectation
 516 to μ .

517 In deep probability distributions, a simple random variable $w \sim p_0$ is mapped deterministically
 518 via a function f_θ parameterized by a neural network to the support of the distribution of interest
 519 where $z = f_\theta(w) = f_l(\dots f_1(w))$. Given a theoretical model $p(x | z)$ and some behavior of interest
 520 \mathcal{B} , the deep probability distributions are trained by optimizing the neural network parameters θ to
 521 find a good approximation q_θ^* within the deep variational family Q to $p(z | \mathcal{B})$.

522 In most settings (especially those relevant to theoretical neuroscience) the likelihood of the behavior
 523 with respect to the model parameters $p(T(x) | z)$ is unknown or intractable, requiring an alternative
 524 to stochastic gradient variational bayes [5] or black box variational inference[43]. These types of
 525 methods called likelihood-free variational inference (LFVI, cite Tran) skate around the intractable
 526 likelihood function in situations where there is a differentiable simulator. Akin to LFVI, DSNs are
 527 optimized with the following objective for a given generative model and statistical constraints on
 528 its produced activity:

$$\begin{aligned} q_\theta^*(z) &= \underset{q_\theta \in Q}{\operatorname{argmax}} H(q_\theta(z)) \\ \text{s.t. } E_{z \sim q_\theta} [E_{x \sim p(x|z)} [T(x)]] &= \mu \end{aligned} \tag{6}$$

529 TODO expand on this in terms of Fig. 2, in a way that complements 3.1.

530 **A.1.1 Example: 2D LDS**

531 **Draft in progress:**

532 To gain intuition for EPI, consider two-dimensional linear dynamical systems, $\tau \dot{x} = Ax$ with

$$A = \begin{bmatrix} a_1 & a_2 \\ a_3 & a_4 \end{bmatrix}$$

533 that produce a band of oscillations. To do EPI with the dynamics matrix elements as the free
 534 parameters $z = [a_1, a_2, a_3, a_4]$, and fixing $\tau = 1$, such that the posterior yields a band of oscillations,
 535 the emergent property statistics $T(x)$ are chosen to contain the first- and second-moments of the
 536 oscillatory frequency Ω and the growth/decay factor d of the oscillating system (the real part of the
 537 complex conjugate pairs of eigenvalues). To learn the distribution of real entries of A that yield a
 538 distribution of d with mean zero with variance 1, and oscillation frequency Ω with mean 1 Hz with
 539 variance 1, the emergent property values would be set to:

$$\mu = E \begin{bmatrix} d \\ \Omega \\ (d - 0)^2 \\ (\Omega - 1)^2 \end{bmatrix} = \begin{bmatrix} 0.0 \\ 1.0 \\ 1.0 \\ 1.025 \end{bmatrix} \tag{7}$$

540 To obtain a differentiable estimate of the oscillation frequency with respect to the dynamics matrices,
 541 we could simulate system activity x from $z = A$ for some finite number of time steps, and estimate
 542 Ω by e.g. taking the peak of the discrete Fourier transform. Instead, the emergent property
 543 statistics for this oscillating behavior are computed through a closed form function $g(z)$ by taking
 544 the eigendecomposition of the dynamics matrix

$$g(z) = E_{x \sim p(x|z)} [T(x)] = \begin{bmatrix} \operatorname{real}(\lambda_1) \\ \frac{\operatorname{imag}(\lambda_1)}{2\pi} \\ \operatorname{real}(\lambda_1)^2 \\ \left(\frac{\operatorname{imag}(\lambda_1)}{2\pi}\right)^2 \end{bmatrix} \tag{8}$$

545

$$\lambda = \frac{\left(\frac{a_1+a_4}{\tau}\right) \pm \sqrt{\left(\frac{a_1+a_4}{\tau}\right)^2 + 4\left(\frac{a_2a_3-a_1a_4}{\tau}\right)}}{2} \quad (9)$$

546 where λ_1 is the eigenvalue of $\frac{1}{\tau}A$ with greatest real part. Even though $E_{x \sim p(x|z)}[T(x)]$ is calculable
 547 directly via $g(z)$, we cannot derive the distribution q_θ^* , since the backward mapping from the mean
 548 parameters μ to the natural parameters η of his exponential family is unknown [44]. Instead, we
 549 can use EPI to learn the linear system parameters producing such a band of oscillations (Fig. S2B).

550 Even this relatively simple system has nontrivial (though intuitively sensible) structure in the
 551 parameter distribution. The contours of the probability density can be derived from the emergent
 552 property statistics and values (Fig. S3). In the $a_1 - a_4$ plane, is a black line at $\text{real}(\lambda_1) = \frac{a_1+a_4}{2} = 0$,
 553 a dotted black line at the standard deviation $\text{real}(\lambda_1) = \frac{a_1+a_4}{2} \pm 1$, and a grey line at twice the
 554 standard deviation $\text{real}(\lambda_1) = \frac{a_1+a_4}{2} \pm 2$ (Fig. S3A). Here the lines denote the set of solutions at
 555 fixed behaviors, which overlay the posterior obtained through EPI. The learned DSN distribution
 556 precisely reflects the desired statistical constraints and model degeneracy in the sum of a_1 and
 557 a_4 . Intuitively, the parameters equivalent with respect to emergent property statistic $\text{real}(\lambda_1)$ have
 558 similar log densities.

559 To explain the structure in the bimodality of the DSN posterior, we can look at the imaginary
 560 component of λ_1 . When $\text{real}(\lambda_1) = \frac{a_1+a_4}{2} = 0$, we have

$$\text{imag}(\lambda_1) = \begin{cases} \sqrt{\frac{a_1a_4-a_2a_3}{\tau}}, & \text{if } a_1a_4 < a_2a_3 \\ 0 & \text{otherwise} \end{cases} \quad (10)$$

561 When $\tau = 1$ and $a_1a_4 > a_2a_3$ (center of distribution above), we have the following equation for the
 562 the other two dimensions:

$$\text{imag}(\lambda_1)^2 = a_1a_4 - a_2a_3 \quad (11)$$

563 Since we constrained $E_{q_\theta}[\text{imag}(\lambda)] = 2\pi$ (with $\omega = 1$), we can plot contours of the equation
 564 $\text{imag}(\lambda_1)^2 = a_1a_4 - a_2a_3 = (2\pi)^2$ for various a_1a_4 (Fig. S3A). If $\sigma_{1,4} = E_{q_\theta}(|a_1a_4 - E_{q_\theta}[a_1a_4]|)$,
 565 then we plot the contours as $a_1a_4 = 0$ (black), $a_1a_4 = -\sigma_{1,4}$ (black dotted), and $a_1a_4 = -2\sigma_{1,4}$ (grey
 566 dotted) (Fig. S3B). We take steps in negative standard deviation of a_1a_4 (dotted and gray lines),
 567 since there are few positive values a_1a_4 in the posterior. More subtle model-behavior combinations
 568 will have even more complexity, further motivating the use of EPI for understanding these systems.

569 For futher validation of the underlying technology, see recovery of ground truth distributions with
 570 maximum entropy flow networks [17].

571 **A.1.2 Augmented Lagrangian optimization**

572 **Draft in progress:**

573 To optimize $q_\theta(z)$ in equation 1, the constrained optimization is performed using the augmented
 574 Lagrangian method. The following objective is minimized:

$$L(\theta; \alpha, c) = -H(q_\theta) + \alpha^\top \delta(\theta) + \frac{c}{2} \|\delta(\theta)\|^2 \quad (12)$$

575 where $\delta(\theta) = E_{z \sim q_\theta} [E_{x \sim p(x|z)} [T(x) - \mu]]$, $\alpha \in \mathcal{R}^m$ are the Lagrange multipliers and c is the
 576 penalty coefficient. For a fixed (α, c) , θ is optimized with stochastic gradient descent. A low
 577 value of c is used initially, and increased during each augmented Lagrangian epoch – a period
 578 of optimization with fixed *alpha* and c for a given number of stochastic optimziation iterations.
 579 Similarly, α is tuned each epoch based on the constraint violations. For the linear 2-dimensional
 580 system (Fig. S4C) optimization hyperparameters are initialized to $c_1 = 10^{-4}$ and $\alpha_1 = 0$. The
 581 penalty coefficient is updated based on a hypothesis test regarding the reduction in constraint
 582 violation. The p-value of $E[\|\delta(\theta_{k+1})\|] > \gamma E[\|\delta(\theta_k)\|]$ is computed, and c_{k+1} is updated to βc_k
 583 with probability $1 - p$. Throughout the project, $\beta = 4.0$ and $\gamma = 0.25$ is used. The other update
 584 rule is $\alpha_{k+1} = \alpha_k + c_k \frac{1}{n} \sum_{i=1}^n (T(x^{(i)}) - \mu)$. In this example, each augmented Lagrangian epoch ran
 585 for 5,000 iterations. We consider the optimization to have converged when a null hypothesis test
 586 of constraint violations being zero is accepted for all constraints at a significance threshold 0.05.
 587 This is the dotted line on the plots below depicting the optimization cutoff of EPI optimization for
 588 the 2-dimensional linear system. If the optimization is left to continue running, entropy usually
 589 decreases, and structural pathologies in the distribution may be introduced.

590 The intention is that c and λ start at values encouraging entropic growth early in optimization.
 591 Then, as they increase in magnitude with each training epoch, the constraint satisfaction terms are
 592 increasingly weighted, resulting in a decrease in entropy. Rather than using a naive initialization,
 593 before EPI, we optimize the deep probability distribution parameters to generate samples of an
 594 isotropic gaussian of a selected variance, such as 1.0 for the 2D LDS example. This provides a
 595 convenient starting point, whose level of entropy is controlled by the user.

596 **A.1.3 Normalizing flows**597 **Draft in progress:**

598 Since we are optimizing parameters θ of our deep probability distribution with respect to the
 599 entropy, we will need to take gradients with respect to the log-density of samples from the deep
 600 probability distribution.

$$H(q_\theta(z)) = \int -q_\theta(z) \log(q_\theta(z)) dz = E_{z \sim q_\theta} [-\log(q_\theta(z))] = E_{\omega \sim q_0} [-\log(q_\theta(f_\theta(\omega)))] \quad (13)$$

$$\nabla_\theta H(q_\theta(z)) = E_{\omega \sim q_0} [-\nabla_\theta \log(q_\theta(f_\theta(\omega)))] \quad (14)$$

601 Deep probability models typically consist of several layers of fully connected neural networks.
 602 When each neural network layer is restricted to be a bijective function, the sample density can be
 603 calculated using the change of variables formula at each layer of the network. For $z' = f(z)$,

$$q(z') = q(f^{-1}(z')) \left| \det \frac{\partial f^{-1}(z')}{\partial z'} \right| = q(z) \left| \det \frac{\partial f(z)}{\partial z} \right|^{-1} \quad (15)$$

604 However, this computation has cubic complexity in dimensionality for fully connected layers. By
 605 restricting our layers to normalizing flows [16] – bijective functions with fast log determinant
 606 jacobian computations, we can tractably optimize deep generative models with objectives that are
 607 a function of sample density, like entropy. Most of our analyses use real NVP [45], which have
 608 proven effective in our architecture searches, and have the advantageous features of fast sampling
 609 and fast density evaluation.

611 **A.1.4 Related work**612 **Draft in progress:**

613

614 **A.1.5 Emergent property inference as variational inference in an exponential family**615 **Draft in progress:**

616 Consider the goal of doing variational inference (VI) in with an exponential family posterior dis-
 617 tribution $p(z | x)$. We'll use the following abbreviated notation to collect the base measure and

618 sufficient statistics into $\tilde{T}(z)$ and likewise concatenate a 1 onto the end of the natural parameter
 619 $\tilde{\eta}(x)$. The log normalizing constant $A(\eta(x))$ will remain unchanged.

$$\begin{aligned} p(z | x) &= b(z) \exp \left(\eta(x)^\top T(z) - A(\eta(x)) \right) = \exp \left(\begin{bmatrix} \eta(x) \\ 1 \end{bmatrix}^\top \begin{bmatrix} T(z) \\ b(z) \end{bmatrix} - A(\eta(x)) \right) \\ &= \exp \left(\tilde{\eta}(x)^\top \tilde{T}(z) - A(\eta(x)) \right) \end{aligned} \quad (16)$$

620 VI looks with an exponential family posterior distribution uses optimization to minimize the fol-
 621 lowing divergence [15]:

$$q_\theta^* = \underset{q_\theta \in Q}{\operatorname{argmin}} KL(q_\theta || p(z | x)) \quad (17)$$

622 $q_\theta(z)$ is the variational approximation to the posterior with variational parameters θ . We can write
 623 this KL divergence in terms of entropy of the variational approximation.

$$KL(q_\theta || p(z | x)) = E_{z \sim q_\theta} [\log(q_\theta(z))] - E_{z \sim q_\theta} [\log(p(z | x))] \quad (18)$$

624

$$= -H(q_\theta) - E_{z \sim q_\theta} [\tilde{\eta}(x)^\top \tilde{T}(z) - A(\eta(x))] \quad (19)$$

625 As far as the variational optimization is concerned, the log normalizing constant is independent of
 626 $q_\theta(z)$, so it can be dropped.

$$\underset{q_\theta \in Q}{\operatorname{argmin}} KL(q_\theta || p(z | x)) = \underset{q_\theta \in Q}{\operatorname{argmin}} -H(q_\theta) - E_{z \sim q_\theta} [\tilde{\eta}(x)^\top \tilde{T}(z)] \quad (20)$$

627 Further, we can write the objective in terms of the first moment of the sufficient statistics $\mu =$
 628 $E_{z \sim p(z|x)} [T(z)]$.

$$= \underset{q_\theta \in Q}{\operatorname{argmin}} -H(q_\theta) - E_{z \sim q_\theta} [\tilde{\eta}(x)^\top (\tilde{T}(z) - \mu)] + \tilde{\eta}(x)^\top \mu \quad (21)$$

629

$$= \underset{q_\theta \in Q}{\operatorname{argmin}} -H(q_\theta) - E_{z \sim q_\theta} [\tilde{\eta}(x)^\top (\tilde{T}(z) - \mu)] \quad (22)$$

630 In emergent property inference (EPI), we're solving the following problem.

$$q_\theta^*(z)y = \underset{q_\theta \in Q}{\operatorname{argmax}} H(q_\theta(z)), \text{ s.t. } E_{z \sim q_\theta} [E_{x \sim p(x|z)} [T(x)]] = \mu \quad (23)$$

631 The lagrangian objective is

$$q_\theta^* = \underset{q_\theta \in Q}{\operatorname{argmin}} -H(q_\theta) + \alpha^\top (E_{z \sim q_\theta} [\tilde{T}(z)] - \mu) \quad (24)$$

632 As the lagrangian optimization proceeds, α should converge to $\tilde{\eta}(x)$ through its adaptations in
 633 each epoch. More formally, $\tilde{\eta}(x) \leftrightarrow \tilde{\eta}(\mu)$ is referred to as the backward mapping, and is formally
 634 hard to identify [44]. Since this backward mapping is deterministic, conceptually, we can replace
 635 $p(z | x)$ with $p(z | \mu)$. More commonly, we write $p(z | \mathcal{B})$ for clarity where \mathcal{B} more explicitly
 636 captures the moment constraints of the sufficient statistics.

637 **A.2 Theoretical models**

638 **Draft in progress:**

639 In this study, we used emergent property inference to examine several models relevant to theoretical
640 neuroscience. Here, we provide the details of each model and the related analyses.

641 **A.2.1 Stomatogastric ganglion**

642 **Draft in progress:**

643 Each neuron's membrane potential is the solution of the following differential equation.

$$C_m \frac{\partial x_m}{\partial t} = -[h_{leak} + h_{Ca} + h_K + h_{hyp} + h_{elec} + h_{syn}] \quad (25)$$

644 The membrane potential of each neuron is affected by the leak, Calcium, Potassium, hyperpo-
645 larization, electrical and synaptic currents, respectively. The capacitance of the circuit is set to
646 $C_m = 1nF$. Each current has an associated reversal potential: $V_{leak} = -40mV$, $V_{Ca} = 100mV$,
647 $V_K = -80mV$, $V_{hyp} = -20mV$, and $V_{syn} = -75mV$. Each current is a function of the difference
648 in membrane and reversal potential multiplied by a conductance:

$$h_{leak} = g_{leak}(x_m - V_{leak}) \quad (26)$$

$$h_{elec} = g_{el}(x_m^{post} - x_m^{pre}) \quad (27)$$

$$h_{syn} = g_{syn}S_\infty^{pre}(x_m^{post} - V_{syn}) \quad (28)$$

$$h_{Ca} = g_{Ca}M_\infty(x_m - V_{Ca}) \quad (29)$$

$$h_K = g_KN(x_m - V_K) \quad (30)$$

$$h_{hyp} = g_hH(x_m - V_{hyp}) \quad (31)$$

654 where g_{el} and g_{syn} are DSN-focused parameters, $g_{leak} = 1 \times 10^{-4}\mu S$, and g_{Ca} , g_K , and g_{hyp}
655 have different values based on fast, intermediate (hub) or slow neuron. Fast: $g_{Ca} = 1.9 \times 10^{-2}$,
656 $g_K = 3.9 \times 10^{-2}$, and $g_{hyp} = 2.5 \times 10^{-2}$. Intermediate: $g_{Ca} = 1.7 \times 10^{-2}$, $g_K = 1.9 \times 10^{-2}$, and
657 $g_{hyp} = 8.0 \times 10^{-3}$. Intermediate: $g_{Ca} = 8.5 \times 10^{-3}$, $g_K = 1.5 \times 10^{-2}$, and $g_{hyp} = 1.0 \times 10^{-2}$.

658 The Calcium, Potassium, and hyperpolarization channels have time-dependent gating dynamics
659 dependent on steady-state gating variables M_∞ , N_∞ and H_∞ , respectively.

$$M_\infty = 0.5 \left(1 + \tanh \left(\frac{x_m - v_1}{v_2} \right) \right) \quad (32)$$

$$\frac{\partial N}{\partial t} = \lambda_N(N_\infty - N) \quad (33)$$

$$N_\infty = 0.5 \left(1 + \tanh \left(\frac{x_m - v_3}{v_4} \right) \right) \quad (34)$$

$$\lambda_N = \phi_N \cosh \left(\frac{x_m - v_3}{2v_4} \right) \quad (35)$$

$$\frac{\partial H}{\partial t} = \frac{(H_\infty - H)}{\tau_h} \quad (36)$$

$$H_\infty = \frac{1}{1 + \exp\left(\frac{x_m + v_5}{v_6}\right)} \quad (37)$$

$$\tau_h = 272 - \left(\frac{-1499}{1 + \exp\left(\frac{-x_m + v_7}{v_8}\right)} \right) \quad (38)$$

666 where $v_1 = 0mV$, $v_2 = 20mV$, $v_3 = 0mV$, $v_4 = 15mV$, $v_5 = 78.3mV$, $v_6 = 10.5mV$, $v_7 = -42.2mV$,
 667 $v_8 = 87.3mV$, $v_9 = 5mV$, and $v_{th} = -25mV$.

668 Finally, there is a synaptic gating variable as well:

$$S_\infty = \frac{1}{1 + \exp\left(\frac{v_{th} - x_m}{v_9}\right)} \quad (39)$$

When the dynamic gating variables are considered, this is actually a 15-dimensional nonlinear dynamical system.

In order to measure the frequency of the hub neuron during EPI, the STG model was simulated for $T = 500$ time steps of $dt = 25ms$. In EPI, since gradients are taken through the simulation process, the number of time steps are kept modest if possible. The chosen dt and T were the most computationally convenient choices yielding accurate frequency measurement.

Our original approach to measuring frequency was to take the max of the fast Fourier transform (FFT) of the simulated time series. There are a few key considerations here. One is resolution in frequency space. Each FFT entry will correspond to a signal frequency of $\frac{F_s k}{N}$, where N is the number of samples used for the FFT, $F_s = \frac{1}{dt}$, and $k \in [0, 1, \dots, N - 1]$. Our resolution is improved by increasing N and decreasing dt . Increasing $N = T - b$, where b is some fixed number of buffer burn-in initialization samples, necessitates an increase in simulation time steps T , which directly increases computational cost. Increasing F_s (decreasing dt) increases system approximation accuracy, but requires more time steps before a full cycle is observed. At the level of $dt = 0.025$, thousands of temporal samples were required for resolution of .01Hz. These challenges in frequency resolution with the discrete Fourier transform motivated the use of an alternative basis of complex

exponentials. Instead, we used a basis of complex exponentials with frequencies from 0.0-1.0 Hz at 0.01Hz resolution, $\Phi = [0.0, 0.01, \dots, 1.0]^\top$

Another consideration is that the frequency spectra of the hub neuron has several peaks. This is due to high-frequency sub-threshold activity. The maximum frequency was often not the firing frequency. Accordingly, subthreshold activity was set to zero, and the whole signal was low-pass filtered with a moving average window of length 20. The signal is subsequently mean centered. After this pre-processing, the maximum frequency in filter bank accurately reflected the firing frequency.

Finally, to differentiate through the maximum frequency identification step, we used a sum-of-powers normalization strategy: Let $\mathcal{X}_i \in \mathcal{C}^{|\Phi|}$ be the complex exponential filter bank dot products with the signal $x_i \in \mathcal{R}^N$, where $i \in \{\text{f1}, \text{f2}, \text{hub}, \text{s1}, \text{s2}\}$. The “frequency identification” vector is

$$u_i = \frac{|\mathcal{X}_i|^\alpha}{\sum_{k=1}^N |\mathcal{X}_i(k)|^\alpha}$$

. The frequency is then calculated as $\Omega_i = u_i^\top \Phi$ with $\alpha = 100$.

Network syncing, like all other emergent properties in this work, are defined by the emergent property statistics and values. The emergent property statistics are the first- and second-moments of the firing frequencies. The first moments are set to 0.55Hz, while the second moments are set to 0.025Hz².

$$E \begin{bmatrix} \Omega_{\text{f1}} \\ \Omega_{\text{f2}} \\ \Omega_{\text{hub}} \\ \Omega_{\text{s1}} \\ \Omega_{\text{s2}} \\ (\Omega_{\text{f1}} - 0.55)^2 \\ (\Omega_{\text{f2}} - 0.55)^2 \\ (\Omega_{\text{hub}} - 0.55)^2 \\ (\Omega_{\text{s1}} - 0.55)^2 \\ (\Omega_{\text{s2}} - 0.55)^2 \end{bmatrix} = \begin{bmatrix} 0.55 \\ 0.55 \\ 0.55 \\ 0.55 \\ 0.55 \\ 0.025^2 \\ 0.025^2 \\ 0.025^2 \\ 0.025^2 \\ 0.025^2 \end{bmatrix} \quad (40)$$

For EPI in Fig 2C, we used a real NVP architecture with two coupling layers. Each coupling layer had two hidden layers of 10 units each.

703 **A.2.2 Primary visual cortex**704 **Draft in progress:**

705 The dynamics of each neural populations average rate $x = \begin{bmatrix} x_E \\ x_P \\ x_S \\ x_V \end{bmatrix}$ are given by:

$$\tau \frac{dx}{dt} = -x + [Wx + h]_+^n \quad (41)$$

706 Some neuron types largely lack synaptic projections to other neuron types [46], and it is popular
707 to only consider a subset of the effective connectivities [20].

$$W = \begin{bmatrix} W_{EE} & W_{EP} & W_{ES} & 0 \\ W_{PE} & W_{PP} & W_{PS} & 0 \\ W_{SE} & 0 & 0 & W_{SV} \\ W_{VE} & W_{VP} & W_{VS} & 0 \end{bmatrix} \quad (42)$$

708 (TODO: ask Ken about how to introduce these values and what to ref). Estimates of the of the
709 probability of connection and strength of connection from the Allen institute result in an estimate
710 of the effective connectivity:

$$W = \begin{bmatrix} 0.0576 & 0.19728 & 0.13144 & 0 \\ 0.58855 & 0.30668 & 0.4285 & 0 \\ 0.15652 & 0 & 0 & 0.2 \\ 0.13755 & 0.0902 & 0.4004 & 0 \end{bmatrix} \quad (43)$$

711 We look at how this four-dimensional nonlinear dynamical model of V1 responds to different inputs,
712 and compare the predictions of the linear response to the approximate posteriors obtained through
713 EPI. The input to the system is the sum of a baseline input $b = [1 \ 1 \ 1 \ 1]^\top$ and a differential
714 input dh :

$$h = b + dh \quad (44)$$

715 All simulations of this system had $T = 100$ time points, a time step $dt = 5\text{ms}$, and time constant
716 $\tau = 20\text{ms}$. And the system was initialized to a random draw $x(0)_i \sim \mathcal{N}(1, 0.01)$.

717 We can describe the dynamics of this system more generally by

$$\dot{x}_i = -x_i + f(u_i) \quad (45)$$

718 where the input to each neuron is

$$u_i = \sum_j W_{ij}x_j + h_i \quad (46)$$

719 Let $F_{ij} = \gamma_i \delta(i, j)$, where $\gamma_i = f'(u_i)$. Then, the linear response is

$$\frac{\partial x_{ss}}{\partial h} = F(W \frac{\partial x_{ss}}{\partial h} + I) \quad (47)$$

720 which is calculable by

$$\frac{\partial x_{ss}}{\partial h} = (F^{-1} - W)^{-1} \quad (48)$$

721 The emergent property we considered was the first and second moments of the change in rate dr
 722 between the baseline input $h = b$ and $h = b + dh$. We use the following notation to indicate that
 723 the emergent property statistics were set to the following values:

$$\mathcal{B}(\alpha, y) \leftrightarrow E \begin{bmatrix} dr_\alpha \\ (dr_\alpha - y)^2 \end{bmatrix} = \begin{bmatrix} y \\ 0.01^2 \end{bmatrix} \quad (49)$$

724 For each $\mathcal{B}(\alpha, y)$ with $\alpha \in \{E, P, S, V\}$ and $y \in \{0.1, 0.5\}$, we ran EPI with five different random
 725 initial seeds using an architecture of four coupling layers, each with two hidden layers of 10 units.

726 We set $c_0 = 10^5$.

727 A.2.3 Superior colliculus

728 **Draft in progress:**

729 There are four total units: two in each hemisphere corresponding to the PRO/CONTRA and
 730 ANTI/IPSI populations. Each unit has an activity (x_i) and internal variable (u_i) related by

$$x_i(t) = \left(\frac{1}{2} \tanh \left(\frac{u_i(t) - \epsilon}{\zeta} \right) + \frac{1}{2} \right) \quad (50)$$

731 $\epsilon = 0.05$ and $\zeta = 0.5$ control the position and shape of the nonlinearity, respectively.

732 We can order the elements of x_i and u_i into vectors x and u with elements

$$x = \begin{bmatrix} x_{LP} \\ x_{LA} \\ x_{RP} \\ x_{RA} \end{bmatrix} \quad u = \begin{bmatrix} u_{LP} \\ u_{LA} \\ u_{RP} \\ u_{RA} \end{bmatrix} \quad (51)$$

733 The internal variables follow dynamics:

$$\tau \frac{\partial u}{\partial t} = -u + Wx + h + \sigma \partial B \quad (52)$$

734 with time constant $\tau = 0.09s$ and gaussian noise $\sigma \partial B$ controlled by the magnitude of $\sigma = 1.0$. The
 735 weight matrix has 8 parameters sW_P , sW_A , vW_{PA} , vW_{AP} , hW_P , hW_A , dW_{PA} , and dW_{AP} (Fig.
 736 4B).

$$W = \begin{bmatrix} sW_P & vW_{PA} & hW_P & dW_{PA} \\ vW_{AP} & sW_A & dW_{AP} & hW_A \\ hW_P & dW_{PA} & sW_P & vW_{PA} \\ dW_{AP} & hW_A & vW_{AP} & sW_A \end{bmatrix} \quad (53)$$

737 The system receives five inputs throughout each trial, which has a total length of 1.8s.

$$h = h_{\text{rule}} + h_{\text{choice-period}} + h_{\text{light}} \quad (54)$$

738 There are rule-based inputs depending on the condition,

$$h_{P,\text{rule}}(t) = \begin{cases} I_{P,\text{rule}} \begin{bmatrix} 1 & 0 & 0 & 1 \end{bmatrix}^\top, & \text{if } t \leq 1.2s \\ 0, & \text{otherwise} \end{cases} \quad (55)$$

$$h_{A,\text{rule}}(t) = \begin{cases} I_{A,\text{rule}} \begin{bmatrix} 0 & 1 & 1 & 0 \end{bmatrix}^\top, & \text{if } t \leq 1.2s \\ 0, & \text{otherwise} \end{cases} \quad (56)$$

740 a choice-period input,

$$h_{\text{choice}}(t) = \begin{cases} I_{\text{choice}} \begin{bmatrix} 1 & 1 & 1 & 1 \end{bmatrix}^\top, & \text{if } t > 1.2s \\ 0, & \text{otherwise} \end{cases} \quad (57)$$

741 and an input to the right or left-side depending on where the light stimulus is delivered.

$$h_{\text{light}}(t) = \begin{cases} I_{\text{light}} \begin{bmatrix} 1 & 1 & 0 & 0 \end{bmatrix}^\top, & \text{if } t > 1.2s \text{ and Left} \\ I_{\text{light}} \begin{bmatrix} 0 & 0 & 1 & 1 \end{bmatrix}^\top, & \text{if } t > 1.2s \text{ and Right} \\ 0, & t \leq 1.2s \end{cases} \quad (58)$$

742 The input parameterization was fixed to $I_{P,\text{rule}} = 10$, $I_{A,\text{rule}} = 10$, $I_{\text{choice}} = 2$, and $I_{\text{light}} = 1$

743 TODO: this is probably a good place to explain the intuition behind the naming of the Schur
 744 eigenmodes.

745 To produce a Bernoulli rate of p_{LP} in the Left, Pro condition (we can generalize this to either cue,
 746 or stimulus condition), let \hat{p}_i be the empirical average steady state (ss) response (final V_{LP} at end
 747 of task) over M=500 gaussian noise draws for a given SC model parameterization z_i :

$$\hat{p}_i = E_{\sigma \partial B} [x_{LP,ss} | s = L, c = P, z_i] = \frac{1}{M} \sum_{j=1}^M x_{LP,ss}(s = L, c = P, z_i, \sigma \partial B_j) \quad (59)$$

748 For the first constraint, the average over posterior samples (from $q_\theta(z)$) to be p_{LP} :

$$E_{z_i \sim q_\phi} [E_{\sigma \partial B} [x_{LP,ss} | s = L, c = P, z_i]] = E_{z_i \sim q_\phi} [\hat{p}_i] = p_{LP} \quad (60)$$

749 We can then ask that the variance of the steady state responses across gaussian draws, is the
750 Bernoulli variance for the empirical rate \hat{p}_i .

$$E_{z \sim q_\phi} [\sigma_{err}^2] = 0 \quad (61)$$

751

$$\sigma_{err}^2 = Var_{\sigma \partial B} [x_{LP,ss} | s = L, c = P, z_i] - \hat{p}_i(1 - \hat{p}_i) \quad (62)$$

752 We have an additional constraint that the Pro neuron on the opposite hemisphere should have the
753 opposite value. We can enforce this with a final constraint:

$$E_{z \sim q_\phi} [d_P] = 1 \quad (63)$$

754

$$E_{\sigma \partial W} [(x_{LP,ss} - x_{RP,ss})^2 | s = L, c = P, z_i] \quad (64)$$

755 We refer to networks obeying these constraints as Bernoulli, winner-take-all networks. Since the
756 maximum variance of a random variable bounded from 0 to 1 is the Bernoulli variance ($\hat{p}(1 - \hat{p})$),
757 and the maximum squared difference between two variables bounded from 0 to 1 is 1, we do not
758 need to control the second moment of these test statistics. In reality, these variables are dynamical
759 system states and can only exponentially decay (or saturate) to 0 (or 1), so the Bernoulli variance
760 error and squared difference constraints can only be undershot. This is important to be mindful
761 of when evaluating the convergence criteria. Instead of using our usual hypothesis testing criteria
762 for convergence to the emergent property, we set a slack variable threshold for these technically
763 infeasible constraints to 0.05.

764 Training DSNs to learn distributions of dynamical system parameterizations that produce Bernoulli
765 responses at a given rate (with small variance around that rate) was harder to do than expected.
766 There is a pathology in this optimization setup, where the learned distribution of weights is bimodal
767 attributing a fraction p of the samples to an expansive mode (which always sends x_{LP} to 1), and a
768 fraction $1 - p$ to a decaying mode (which always sends x_{LP} to 0). This pathology was avoided using
769 an inequality constraint prohibiting parameter samples that resulted in low variance of responses
770 across noise.

771 In total, the emergent property of rapid task switching accuracy at level p was defined as

$$\mathcal{B}(p) \leftrightarrow \begin{bmatrix} \hat{p}_P \\ \hat{p}_A \\ (\hat{p}_P - p)^2 \\ (\hat{p}_A - p)^2 \\ \sigma_{P,err}^2 \\ \sigma_{A,err}^2 \\ d_P \\ d_A \end{bmatrix} = \begin{bmatrix} p \\ p \\ 0.15^2 \\ 0.15^2 \\ 0 \\ 0 \\ 1 \\ 1 \end{bmatrix} \quad (65)$$

772 For each accuracy level p , we ran EPI for 10 different random seeds and selected the maximum
 773 entropy solution using an architecture of 10 planar flows with $c_0 = 2$.

774 **A.2.4 Rank-1 RNN**

775 **Draft in progress:**

776 The network dynamics of neuron i 's rate x evolve according to:

$$\dot{x}_i(t) = -x_i(t) + \sum_{j=1}^N J_{ij} \phi(x_j(t)) + I_i \quad (66)$$

777 where the connectivity is comprised of a random and structured component:

$$J_{ij} = g\chi_{ij} + P_{ij} \quad (67)$$

778 The random all-to-all component has elements drawn from $\chi_{ij} \sim \mathcal{N}(0, \frac{1}{N})$, and the structured
 779 component is a sum of r unit rank terms:

$$P_{ij} = \sum_{k=1}^r \frac{m_i^{(k)} n_j^{(k)}}{N} \quad (68)$$

780 We use this theory to compute $T(x)$ when running EPI.

781 Rank-1 vectors m and n have elements drawn

$$m_i \sim \mathcal{N}(M_m, \Sigma_m)$$

782

$$n_i \sim \mathcal{N}(M_n, \Sigma_n)$$

783 The current has the following statistics:

$$I = M_I + \frac{\Sigma_{mI}}{\Sigma_m} x_1 + \frac{\Sigma_{nI}}{\Sigma_n} x_2 + \Sigma_{\perp} h$$

784 where x_1 , x_2 , and h are standard normal random variables.

785 The $\ddot{\Delta}$ equation is broken into the equation for Δ_0 and Δ_∞ by the autocorrelation dynamics
786 assertions.

$$\ddot{\Delta}(\tau) = -\frac{\partial V}{\partial \Delta}$$

$$787 \quad \ddot{\Delta} = \Delta - \{g^2 \langle [\phi_i(t)\phi_i(t+\tau)] \rangle + \Sigma_m^2 \kappa^2 + 2\Sigma_{mI}\kappa + \Sigma_I^2\}$$

788 We can write out the potential function by integrating the negated RHS.

$$V(\Delta, \Delta_0) = \int \mathcal{D}\Delta \frac{\partial V(\Delta, \Delta_0)}{\partial \Delta}$$

$$789 \quad V(\Delta, \Delta_0) = -\frac{\Delta^2}{2} + g^2 \langle [\Phi_i(t)\Phi_i(t+\tau)] \rangle + (\Sigma_m^2 \kappa^2 + 2\Sigma_{mI}\kappa + \Sigma_I^2)\Delta + C$$

790 We assume that as time goes to infinity, the potential relaxes to a steady state.

$$\frac{\partial V(\Delta_\infty, \Delta_0)}{\partial \Delta} = 0$$

$$791 \quad \frac{\partial V(\Delta_\infty, \Delta_0)}{\partial \Delta} = -\Delta + \{g^2 \langle [\phi_i(t)\phi_i(t+\infty)] \rangle + \Sigma_m^2 \kappa^2 + 2\Sigma_{mI}\kappa + \Sigma_I^2\} = 0$$

$$792 \quad \Delta_\infty = g^2 \langle [\phi_i(t)\phi_i(t+\infty)] \rangle + \Sigma_m^2 \kappa^2 + 2\Sigma_{mI}\kappa + \Sigma_I^2$$

$$793 \quad \Delta_\infty = g^2 \int \mathcal{D}z \left[\int \mathcal{D}x \phi(\mu + \sqrt{\Delta_0 - \Delta_\infty}x + \sqrt{\Delta_\infty}z) \right]^2 + \Sigma_m^2 \kappa^2 + 2\Sigma_{mI}\kappa + \Sigma_I^2$$

794 Also, we assume that the energy of the system is perserved throughout the entirety of its evolution.

$$V(\Delta_0, \Delta_0) = V(\Delta_\infty, \Delta_0)$$

$$795 \quad -\frac{\Delta_0^2}{2} + g^2 \langle [\Phi_i(t)\Phi_i(t)] \rangle + (\Sigma_m^2 \kappa^2 + 2\Sigma_{mI}\kappa + \Sigma_I^2)\Delta_0 + C = -\frac{\Delta_\infty^2}{2} + g^2 \langle [\Phi_i(t)\Phi_i(t)] \rangle + (\Sigma_m^2 \kappa^2 + 2\Sigma_{mI}\kappa + \Sigma_I^2)\Delta_\infty + C$$

$$796 \quad \frac{\Delta_0^2 - \Delta_\infty^2}{2} = g^2 (\langle [\Phi_i(t)\Phi_i(t)] \rangle - \langle [\Phi_i(t)\Phi_i(t)] \rangle) + (\Sigma_m^2 \kappa^2 + 2\Sigma_{mI}\kappa + \Sigma_I^2)(\Delta_0 - \Delta_\infty)$$

$$797 \quad \frac{\Delta_0^2 - \Delta_\infty^2}{2} = g^2 \left(\int \mathcal{D}z \Phi^2(\mu + \sqrt{\Delta_0}z) - \int \mathcal{D}z \int \mathcal{D}x \Phi(\mu + \sqrt{\Delta_0 - \Delta_\infty}x + \sqrt{\Delta_\infty}z) \right) \\ + (\Sigma_m^2 \kappa^2 + 2\Sigma_{mI}\kappa + \Sigma_I^2)(\Delta_0 - \Delta_\infty)$$

798 **Consistency equations:**

799

$$\begin{aligned}
 \mu &= F(\mu, \kappa, \Delta_0, \Delta_\infty) = M_m \kappa + M_I \\
 \kappa &= G(\mu, \kappa, \Delta_0, \Delta_\infty) = M_n \langle [\phi_i] \rangle + \Sigma_{nI} \langle [\phi'_i] \rangle \\
 \frac{\Delta_0^2 - \Delta_\infty^2}{2} &= H(\mu, \kappa, \Delta_0, \Delta_\infty) = g^2 \left(\int \mathcal{D}z \Phi^2(\mu + \sqrt{\Delta_0} z) - \int \mathcal{D}z \int \mathcal{D}x \Phi(\mu + \sqrt{\Delta_0 - \Delta_\infty} x + \sqrt{\Delta_\infty} z) \right) \\
 &\quad + (\Sigma_m^2 \kappa^2 + 2\Sigma_{mI} \kappa + \Sigma_I^2)(\Delta_0 - \Delta_\infty) \\
 \Delta_\infty &= L(\mu, \kappa, \Delta_0, \Delta_\infty) = g^2 \int \mathcal{D}z \left[\int \mathcal{D}x \phi(\mu + \sqrt{\Delta_0 - \Delta_\infty} x + \sqrt{\Delta_\infty} z) \right]^2 + \Sigma_m^2 \kappa^2 + 2\Sigma_{mI} \kappa + \Sigma_I^2
 \end{aligned} \tag{69}$$

800 We can solve these equations by simulating the following Langevin dynamical system.

$$\begin{aligned}
 x(t) &= \frac{\Delta_0(t)^2 - \Delta_\infty(t)^2}{2} \\
 \Delta_0(t) &= \sqrt{2x(t) + \Delta_\infty(t)^2} \\
 \dot{\mu}(t) &= -\mu(t) + F(\mu(t), \kappa(t), \Delta_0(t), \Delta_\infty(t)) \\
 \dot{\kappa}(t) &= -\kappa + G(\mu(t), \kappa(t), \Delta_0(t), \Delta_\infty(t)) \\
 \dot{x}(t) &= -x(t) + H(\mu(t), \kappa(t), \Delta_0(t), \Delta_\infty(t)) \\
 \dot{\Delta}_\infty(t) &= -\Delta_\infty(t) + L(\mu(t), \kappa(t), \Delta_0(t), \Delta_\infty(t))
 \end{aligned} \tag{70}$$

801 Then, the temporal variance is simply calculated via

$$\Delta_T = \Delta_0 - \Delta_\infty \tag{71}$$

802 TODO Need to explain the warm starting for the aficionados.

803 TODO explain the density network architectures used.

804 **A.3 Supplementary Figures**

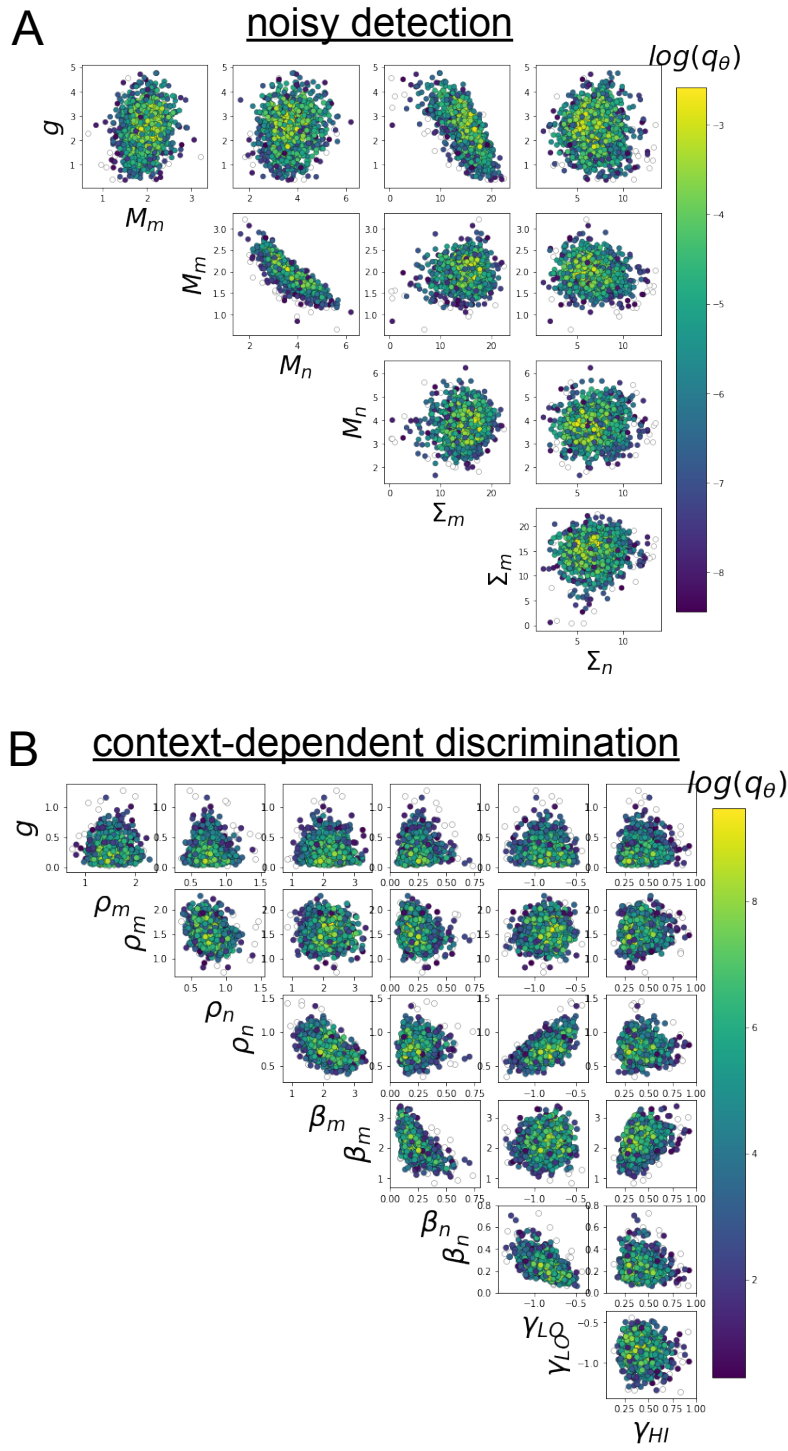


Fig. S1: A. EPI for rank-1 networks doing discrimination. B. EPI for rank-2 networks doing context-dependent discrimination.

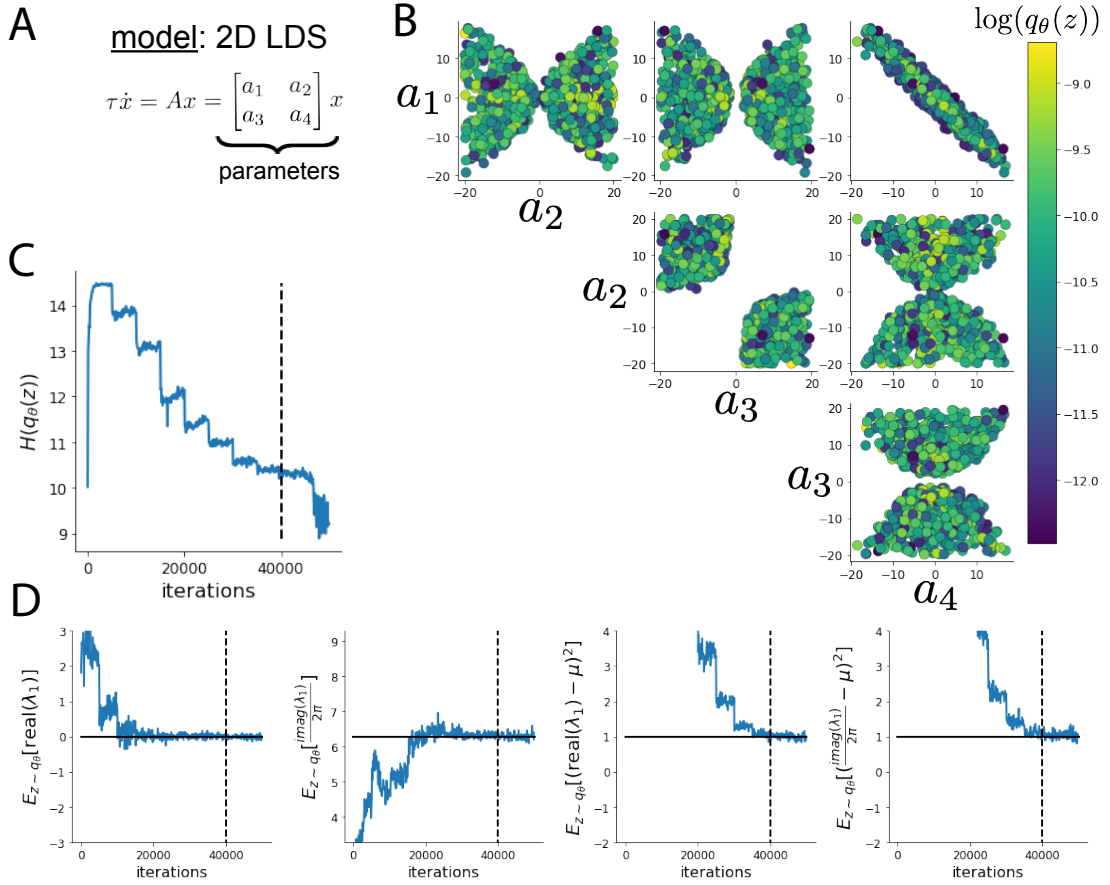


Fig. S2: A. Two-dimensional linear dynamical system model, where real entries of the dynamics matrix A are the parameters. B. The DSN distribution for a 2D LDS with $\tau = 1$ that produces an average of 1Hz oscillations with some small amount of variance. C. Entropy throughout the optimization. At the beginning of each augmented lagrangian epoch (5,000 iterations), the entropy dips due to the shifted optimization manifold where emergent property constraint satisfaction is increasingly weighted. D. Emergent property moments throughout optimization. At the beginning of each augmented lagrangian epoch, the emergent property moments move closer to their constraints.

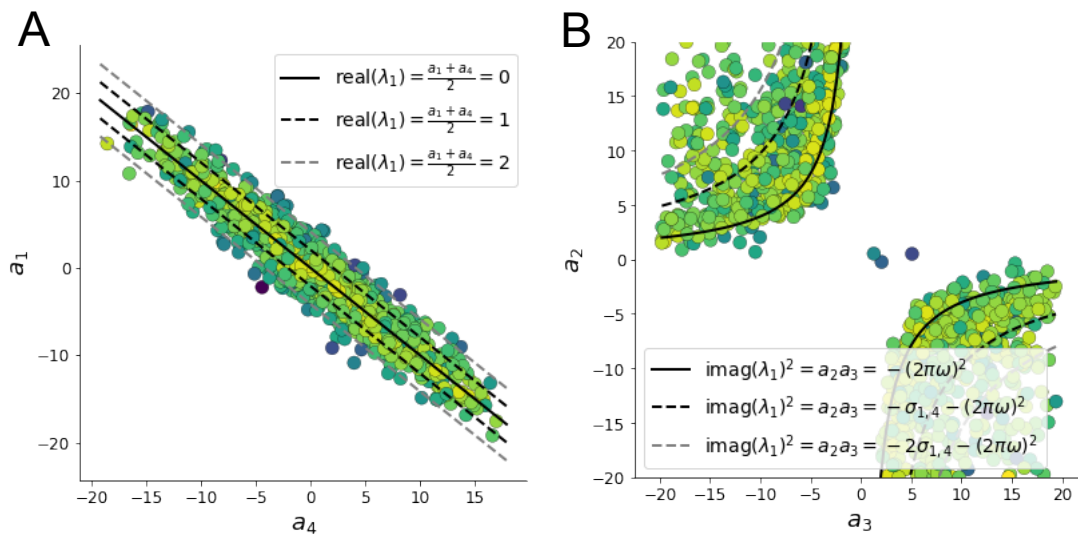


Fig. S3: A. Probability contours in the $a_1 - a_4$ plane can be derived from the relationship to emergent property statistic of growth/decay factor. B. Probability contours in the $a_2 - a_3$ plane can be derived from relationship to the emergent property statistic of oscillation frequency.