# Identifying neurons in *C. elegans* with continuous relaxations for Bayesian permutation inference

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#### **Abstract**

The nematode C. elegans is a unique model organism for neuroscientists as its connectome, or neural wiring diagram, has been known for at least three decades. Despite this knowledge, an understanding of the functional significance of these synaptic connections has remained elusive. Now several groups can routinely image the activity of a large fraction of neurons in the head of the worm, providing a unique opportunity to probe this organism. We propose a hierarchical Bayesian framework that combines strong prior information with data from many experiments to estimate posteriors over the functional connectivity weights. However, these attempts are stifled by a major obstacle: in many cases it is not clear exactly which neurons are being imaged, so to combine information across experiments one must solve a matching, or permutation inference, problem.

In this work we introduce new variational methods that enable the joint inference of connectivity weights and neural identity. Working with actual permutations would involve evaluating and differentiating an intractable partition function. As an alternative, we build upon recent continuous relaxation techniques [Jang et al., 2016, Maddison et al., 2016], extending them from the original case of the probability simplex, to the Birkhoff polytope, the convex hull of permutation matrices. We test our method with simulated data from the true connectome and known covariates (neural position) and show that our approach outperforms many alternatives in identifying neurons.

## 1 Introduction

- 22 The nematode C. elegans plays a special role as a model organism in neuroscience since its neural
- 23 network is stereotyped from animal to animal and its complete neural wiring diagram is known [Varsh-
- 24 ney et al., 2011]. Modern calcium imaging technology enables measurements of hundreds of these
- 25 neurons simultaneously [Kato et al., 2015, Nguyen et al., 2016]. The time is right to employ mod-
- 26 ern statistical methods to learn about the functional connectome in this system and suggest new
- 27 experiments.

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- 28 Ultimately, we are interested the dynamical system that governs how neural activity evolves given its
- 29 history and sensory inputs. Bayesian methods are ideally suited to this goal, allowing us to represent
- 30 hierarchical probabilistic structures and integrate our prior knowledge about the connectome, the
- 31 locations of neurons, etc. Bayesian learning and inference in dynamical systems with MCMC
- methods is well-studied, even for complicated models [De Freitas et al., 2001, Paninski et al., 2010].
- 33 Furthermore, hierarchical models to incorporate information from many worms are easily constructed
- in a Bayesian framework [Gelman et al., 2014].
- 35 However, our efforts to integrate information across worms are complicated by a major hurdle: in
- 36 practice, associating recorded traces to neuron names is a painstaking, manual process. Experimenters

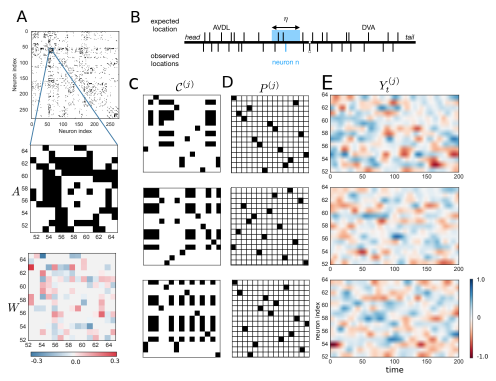


Figure 1: Hierarchical Bayesian framework. A We are given the actual adjaency matrix A from [Varshney et al., 2011]. The full matrix is shown (top) along with a zoom-in to 14 neurons (center). We wish to infer the corresponding weight matrix W, an example of which is shown below. B We also know the typial locations of the neurons [White et al., 1986, Lints et al., 2005]. Given observed locations, we constrain possible assignments to neuron identities within  $\eta$  of the observed location. C These constraints are represented as a matrix  $\mathcal{C}^{(j)}$  for worm j which specifies possible assignments of observed neurons to known identities. This illustration shows three worms. D To infer the weights, we must first infer the permutation  $P^{(j)}$  that matches the observed neurons in worm j to the set of known identities. E The observed data is a matrix  $Y^{(j)}$  whose rows are ordered according to the order in which neurons were observed in that worm. The permutation matrix maps this to the canonical ordering of the adjacency and weight matrices. Given  $\{Y^{(j)}\}_{j=1}^J$  and A, we infer  $\{P^{(j)}\}_{j=1}^J$  and W.

consider the location of the neuron along with its pattern of activity to perform this matching, but the process is laborious and the results are prone to error. Without neuron names, we cannot represent recordings canonically or learn about how one neuron influences another. This technical problem prevents the automatic use of hierarchical methods.

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We present a method for overcoming this hurdle by incorporating inference over permutations that match observed neurons (*neuron 1*, *neuron 2*, ..., *neuron N*) to known names (*AVAL*, *AVAR*, ..., *SMDR*). Once the observed neurons have been mapped to canonical names, we can learn about the shared dynamical system. To start, we focus on a simple linear autoregressive model for neural dynamics,

$$\widetilde{Y}_t^{(j)} = (W \odot A)\widetilde{Y}_{t-1}^{(j)} + \epsilon_t^{(j)}, \tag{1}$$

where  $W \in \mathbb{R}^{N \times N}$  is the weight matrix we wish to infer;  $\odot$  denotes elementwise multiplication;  $A \in \{0,1\}^{N \times N}$  is the known adjacency matrix or connectome; and  $\widetilde{Y}_t^{(j)} \in \mathbb{R}^N$  is the measured neural activity at time t in worm j. The catch is that  $\widetilde{Y}_t^{(j)}$  is assumed to be in canonical order; i.e. in the same order as the rows and columns of W and A. What we actually observe is,

$$Y_t^{(j)} = P^{(j)}\widetilde{Y}_t^{(j)},\tag{2}$$

vectors that are permuted by matrix  $P^{(j)}$ . In order to learn about W, we must also infer the permutation matrices. We assume  $\epsilon_t^{(j)} \sim \mathcal{N}(0, \sigma^2 I)$  with known variance, and we place a Gaussian prior on W.

- The permutation matrices are constrained by side information. Specifically, we use neural position
- along the worm's body to constrain the possible neural identities for a given recorded neuron. We only
- 55 allow an observed neuron to be mapped to a known identity if the observed location is within  $\eta$  of the
- expected location. This is illustrated in Fig. 1B. We represent these constraints with the matrix  $\mathcal{C}^{(j)}$ 56
- so that  $C_{mn}^{(j)}=1$  if and only if observed neuron m is within  $\eta$  of canonical neuron n's expected location. An example is shown in Figure 1C. We let  $P^{(j)}$  have a uniform prior over the set of matrices
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- allowable under the given constraints.
- We need to perform posterior inference of  $p(\{W, P^{(j)}\} | A, \{Y^{(j)}\})$ . MCMC with simple Metropolis-
- Hastings proposals is straightforward, but we found this mixed poorly in practice. Motivated by recent 61
- advances in automatic variational inference [Blei et al., 2017], we considered ways of extending this 62
- technique to permutation inference. In Section 2 we detail our VI formulation and summarize the
- methods we developed. Then in Section 3 we show that these methods outperform alternatives.

#### New methods for variational inference of latent permutations 65

- Consider a latent variable model determined by a prior over the latent  $z \sim p(z)$  and a likelihood
- 67 p(y|z) for the observed data y. In the VI framework, we approximate the intractable posterior p(z|y)
- with the distribution  $q \in \mathcal{Q}$  that best approximates the posterior. For tractability, we assume  $\mathcal{Q}$  is 68
- indexed by a parameter  $\nu$ ; i.e.  $\mathcal{Q} = \{q(z; \nu) : \nu \in \mathcal{V}\}$ . The approximation is typically assessed by
- the Kullback-Leibler divergence between the true posterior and variational approximation,

$$\nu^* = \operatorname*{arg\,min}_{\nu \in \mathcal{V}} \operatorname{KL} \left( q(z; \nu) \parallel p(z \mid y) \right). \tag{3}$$

The above problem is equivalent to maximizing the evidence lower bound (ELBO)

$$\mathcal{L}(\nu) \triangleq \mathbb{E}_{q(z;\nu)}[\log p(y|z)] - \text{KL}(q(z;\nu) \parallel p(z)), \tag{4}$$

- with respect to  $\nu$ . We typically maximize equation (4) with stochastic optimization methods [Kushner
- 73 and Yin, 1987]: specifically, we approximate the expectations in (4) with Monte Carlo estimates
- 74 and optimize the ELBO with stochastic gradient ascent. One critical component is the choice of the
- Monte Carlo approximation. Perhaps the most common choice is through the so called *score function*
- estimator. Unfortunately, this estimator, also referred to as REINFORCE [Williams, 1992], cannot
- 77 be applied to permutations, since it involves the evaluation and differentiation of a likelihood which
- 78 is intractable for any non-trivial distribution over permutations (computing the partition function
- 79 involves a summation over N! terms).
- 80 The re-parameterization trick Kingma and Welling [2013] offers an appealing alternatives. If z can
- 81 be written as a differentiable function of a noise distribution and the parameters—i.e. if for certain
- f and  $\xi \sim p(\xi)$  one has  $z = f(\xi, \nu)$ —then we can write the expectation with respect to q(z) as an expectation with respect to  $p(\xi)$  and bring the gradient inside the expectation. In the case of discrete
- random variables a re-parameterization always exists and it is given by the Gumbel trick [Papandreou
- 85 and Yuille, 2011, Balog et al., 2017], which states that one can sample from any discrete distribution 86
- by perturbing each potential with Gumbel i.i.d noise, and then finding the configuration with the 87 maximum value. Unfortunately, the underlying f to this re-parameterization is the non-differentiable
- arg max operator, precluding the use of gradient descent methods.
- Jang et al. [2016] and Maddison et al. [2016] proposed a solution to this problem, replacing the
- arg max by a temperature-dependent softmax approximation, which in the limit converges to the
- 91 original arg max. By combining the Gumbel trick with the softmax approximation, they conceived
- the Concrete or Gumbel-Softmax distribution, and obtained explicit distribution formulae. Then, 92
- they showed how to perform variational inference in discrete latent variable models using the re-
- 94 parameterization trick and gradient descent. They replaced the original ELBO with a surrogate
- appropriate for their continuous relaxation. As long as the temperature is chosen in a reasonable
- 96 range—not too high as it would lead to a degenerate distribution in the simplex; but also not too low,
- to avoid too high variances of the gradients—the method works well.
- We developed three methods for extending the Gubmel-softmax method to permutations. We name
- then stick-breaking, rounding and Gumbel-Sinkhorn methods. We refer the reader to sections 3.1 99
- 100 and 3.2 of Linderman et al. [2017] and section 4 of Anonymous [2018] for details, respectively.
- Here we briefly summarize them: the primary geometric object is the Birkhoff polytope, the convex

*Table 1:* Accuracy in the C.elegans neural identification problem, for varying mean number of candidate neurons (10, 30, 45, 60) and number of worms.

( 1,11, 1,11,	10		30		45		60	
	1 worm	4 worms	1 Worm	4 worms	1 worm	4 worms	1 worms	4 worms
NAIVE VI	.34	.32	.16	.16	.13	.12	.11	.12
MAP	.34	.32	.17	.17	.14	.13	.13	.12
MCMC	.34	.65	.18	.28	.14	.17	.13	.15
VI	.79	.94	.4	.69	.25	.51	.21	.44

Table 2: Accuracy in inferring true neural identity for different of proportion of known neurons, and two values of n

01 //.	40.%		30.%		20.%		10.%	
	$ \overline{\eta = 0.1} $	$\eta = 0.2$	$ \overline{\eta = 0.1} $	$\eta = 0.2$	$ \overline{\eta = 0.1} $	$\eta = 0.2$	$\overline{\eta = 0.1}$	$\eta = 0.2$
Naive VI	.43	.41	.33	.31	.23	.22	.12	.1
MAP	.42	.41	.33	.32	.23	.22	.12	.11
MCMC	.85	.80	.52	.46	.3	.26	.15	.12
VI	.97	.96	.92	.84	.74	.58	.44	.23

hull of permutation matrices, and the analog of the probability simplex in the discrete case. We 102 generalize the standard stick-breaking construction on the simplex to stick-breaking of the Birkhoff 103 polytope. We show how to consistently "break the stick" while satisfying both the row and column 104 constrains that characterize a doubly stochastic matrix. For the rounding construction, we start 105 106 with a noise distribution and force it to be close to permutation matrices by rounding them towards 107 the extreme-points of the Birkhoff polytope (i.e. permutation matrices). Finally, for the Gumbel-108 Sinkhorn method we notice that the so-called Sinkhorn operator, or infinite and successive row 109 and column normalization of a matrix, is a a natural extension of the softmax operator. With this, 110 we are able to conceive the Gumbel-Sinkhorn distribution, which approximates the sampling of a relevant discrete distribution. Importantly, while stick-breaking and rounding yield explicit densities, 111 Gumbel-Sinkhorn does not. However, there are ways to circumvent this difficulty, and overall we 112 113 observe the latter performs the best.

# 3 Results

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We compared against three methods: (i) naïve variational inference, where we do not enforce the 115 constraint that  $P^{(j)}$  be a permutation and instead treat each row of  $P^{(j)}$  as a Dirichlet distributed 116 vector; (ii) MCMC, where we alternate between sampling from the conditionals of W (Gaussian) 117 and  $P^{(j)}$ , from which one can sample by proposing local swaps, as described in Diaconis [2009], 118 and (iii) maximum a posteriori estimation (MAP). Our MAP algorithm alternates between the 119 optimizing estimate of W given  $\{P^{(j)}, Y^{(j)}\}\$  using linear regression and finding the optimal  $P^{(j)}$ . The second step requires solving a quadratic assignment problem (QAP) in  $P^{(j)}$ ; that is, it can be 121 expressed as  $Tr(APBP^T)$  for matrices A, B. We used the QAP solver proposed by Vogelstein et al. 122 123 [2015].

We found that our method outperforms these alternative approaches. When there are many possible candidates (Table 1) and when only a small proportion of neurons are known with certitude (Table 2), variational inference via continuous relaxation with the Gumbel-Sinkhorn method performs best. Altogether, these results indicate our method enables a more efficient use of information than its alternatives. This is consistent with other results showing faster convergence of variational inference over MCMC [Blei et al., 2017], especially with simple Metropolis-Hastings proposals. We conjecture that MCMC could eventually obtain similar if not better results, if current local proposals—swapping pairs of labels— were replaced by more sophisticated maneuvers, but fundamentally, it seems the hard assignments in the MCMC algorithm lead to poor mixing. We expect that the benefits of VI stem from the continuous relaxation, which enables soft assignments of neurons to identities.

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