Clustering Neural Populations by Poisson Dynamic Factor Model

# Abstract

# Introduction

# Method

## Poisson Dynamic Factor Model

Denote the observed spike-count of neuron at time bin as , and let . Further, let denote the cluster indicator of neuron . To facilitate clustering, we reparametrize the regular Poisson linear dynamical system (PLDS) model to separate the mean log-firing-rate out. Assume neurons are independently Poisson distributed, conditional on the low-dimensional latent state as follows:

, with . We further assume the intercept and the latent state progress linearly with Gaussian noise as

If we denote, and , the model can be equivalently written as regular Poisson factor model as

However, since the condition doesn’t hold, the latent state cannot be consistently estimated, and assuming linear dynamics of resolves the problem. Note that when , the model is not unique, since also satisfies the equation for any orthogonal matrix of order . To ensure the model indefinability, we simply assume and are both diagonal for convenience. Some more careful constraints, such as let be diagonal, with constraints can be implemented, but the results are similar. See more detailed discussions of the constraints in discussion. Overall, denote the cluster-related parameters of cluster as and the spike counts of neuron is generated by Poisson dynamic factor model (PDFM) as , with prior of as . The priors for are regular non-informative normal and inverse-gamma distribution.

The marginal likelihood of neuron is

The marginal likelihood has no closed form and will be used for clustering. To help with fast clustering, instead of doing the Laplace approximation, we choose to make use of the Poisson-Gamma conjugacy [reference]. This leads to the closed form approximation. By the conditional independency assumption, . Since , . Approximate the lognormal distribution by Gamma distribution:

, where and . Then, by Poisson-Gamma conjugacy,

, with and .

Another more general idea is to approximate the log-likelihood by second-order polynomials, with coefficients determined by Chebyshev polynomial approximation [Refer to PAL]. However, the approximation doesn’t work well in practice, especially when the neural spike counts have a wide range. When doing the integration, we need to exponentiate the log-likelihood and this will exaggerate the approximation error.

## Cluster by Mixture of Finite mixture model

When the label is unknown, we cluster the neurons by mixture models. In practice, it’s usually impossible to know the number of neural population. One potential method is to do clustering by Dirichlet process mixtures (DPM) model. However, this is conceptually incorrect, since the number of neural populations is finite but unknown. Besides the conceptual incorrectness, using DPM is not easy to integrate the field knowledge about the number of neural populations. Here, we choose to use the mixture of finite mixtures (MFM) model as follows

whereis a p.m.f. on

given

given

given

independently for , given and

Besides the conceptual correctness, using MFM model allows us integrate the prior knowledge easily. Moreover, compared to DPM, MFM has some better properties for clustering, for example, MFM posterior on number of cluster is more concentrated and consistent, and MFM tend to give clusters size at the same order of magnitude while DPM may lead to a few large clusters and many small clusters. See [reference] for detailed discussion.

# Inference

In this paper, we choose to do inference by MCMC. Because of the Poisson likelihood, the latent has no closed full conditional distribution. We can sample the posterior by particle MCMC directly, but this can be slow. However, due to the Markovian structure of the model, the conditional log-posterior is concave and its Hessian is block-tridiagonal. Thus, we can do the global Laplace approximation efficiently in . The cluster index and number of cluster are sampled by the analog of partition-based algorithm in DPM. See details of the MCMC in appendix.

In practice, using variational Bayes (VB) instead of MCMC may be more favorable. The PLDS can be updated by variational EM. Using the stick-breaking representation of MFM model, we can do VB easily similar to [reference David Blei & Michael I Jordan]. However, checking by the “gold standard” MCMC before doing VB is always a good choice.

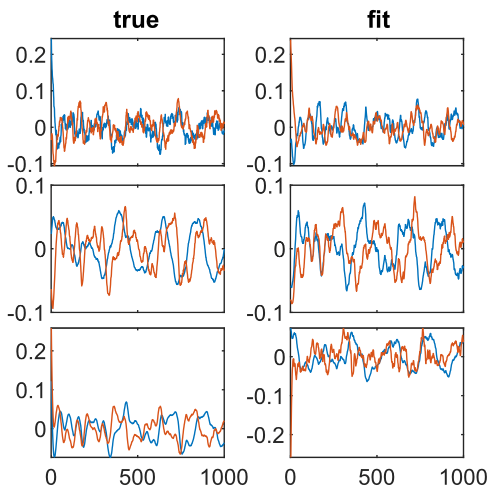
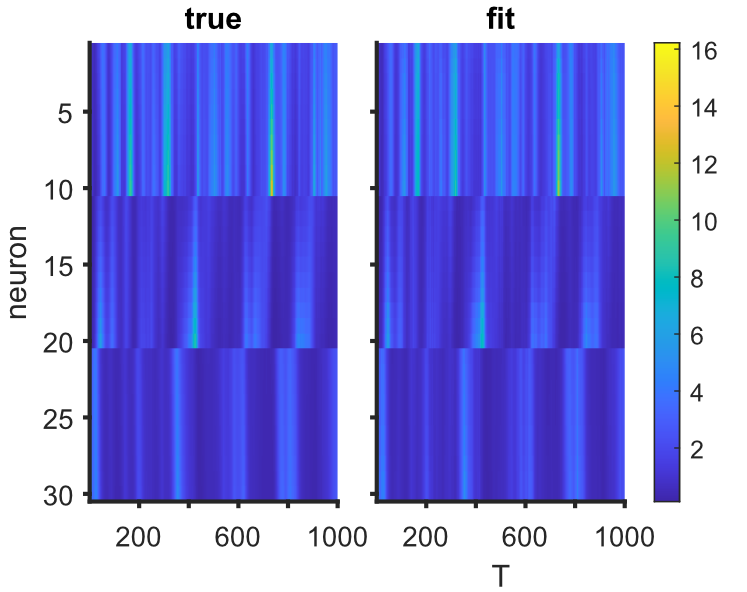
# Simulation

## Model global non-linearity by clustering

There were a rich research results for single PLDS model, but it provides only a global linear model to represent the data in a lower dimensional subspace, which makes the application scope limited. When the input space is nonhomogeneous, a large dimension of latent state is needed and this may lead to the overfitting and poor performance. Mixture of PLDS/ PDFM models allow us to partition the input space into clusters and can therefore capture global nonlinearity by combining local linear models.

We first simulated three neural populations, with 10 neurons in each. We set for each cluster and the recording length is . [Since this model has been studied thoroughly in neuroscience, I didn’t show the trace plots for all parameters. Maybe I just show the trace plots for log-likelihood? ] After checking the trace plots up to 10,000 iterations, the convergence achieved after several steps.

The panel A and B in figure 1 show the posterior mean firing rate and fitted latent state, averaging from iteration 500 to 1000. The latent state is transformed into the commonly used PLDS model: , where which has one more dimension than the parametrization in this paper, i.e. . In the figure 1, has mean zero columns and is orthogonal.

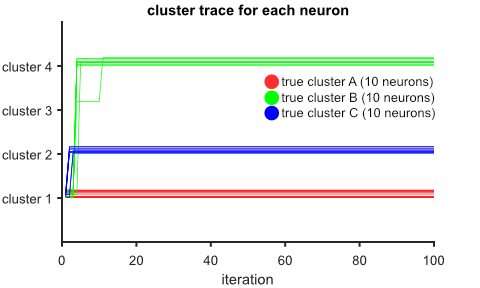


Chart, box and whisker chart

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Then we held out 1/4 and 1/2 data as test set in a “speckled” pattern, i.e. randomly select subset of data for each neuron as held-out dataset. Then we fit the model with and without clusters, keeping the same latent dimension, i.e. (1) 3 clusters with for each and (2) 1 cluster with . The procedure is replicated for 100 times for two proportions, and the difference of held-out likelihood per spike between 2 fittings are shown in panel C in figure 1. The difference between (1) and (2) are always positive, and this shows doing the mixture of PDFM performs better than single PDFM. Moreover, as the proportion of training decrease (less data), the benefit of clustering becomes more significant. This suggests that doing clustering is necessary.

## Clustering

Then use the same setting, we remove the label and use the mixture model to do the clustering by MFM. The prior of the cluster number is . The panel A of figure 2 shows the trace of label for each neuron for the first 100 iterations. Chart

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In this toy simulation example, the signal is strong and the spiking amplitudes of neurons in each population are similar. This might be too simplified for real situation.

In practice, the neural activity is usually sparse and can be recorded in high resolution. Moreover, the spiking amplitude for neuron in one population vary a lot. Therefore, we simulate another more realistic example. In this simulation, there are 3 clusters and the dimension of latent state is for each. However, there are 20 neurons with wider range but smaller value of loading . The simulated firing rate is shown in panel B in figure 2. The trace plot of number of cluster (panel C in figure 2) shows that the model tend to further split some clusters into sub-population, based on the spiking amplitude. The similarity matrix (panel D in figure 2) of posterior, averaging from iteration 500 to 1000, shows that the model tend to split cluster 1 and 3 into two sub-clusters.

# Application

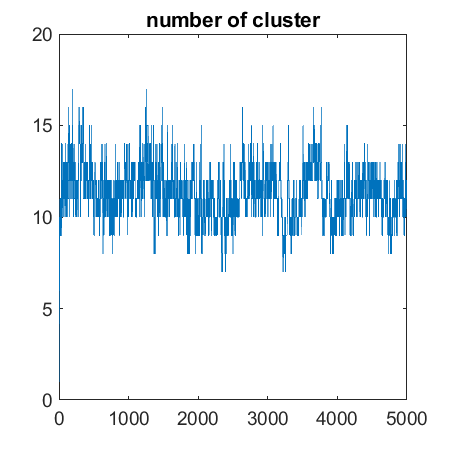
## Neuropixels data

TODO: brief introduction of Neuropixels dataset

The Neuropixels dataset contains recording of neural activities in different brain regions, when doing the drift-grating experiment. Here, we only consider neurons with SNR > 3 and the spiking counts > 1000. Then, we use the recording activity from Lateral posterior nucleus of the thalamus (LP, 20 neurons), anteromedial visual area (VISam, 12 neurons) and ventral posteromedial nucleus of the thalamus (VPM, 14 neurons) during the spontaneous period. The bin size is 0.1s and truncate the data up to 1000 steps (100s recording) for convenience.

Panel A in figure 3 shows the spiking counts of these 46 neurons. Here we first fit model using all data, with and . For the number of clusters, panel B in figure 2 shows the trace plot for the first 5000 iterations and panel C shows the histogram of iteration 2500 to 5000. These plots show that these neurons are quite non-homogenous and they tend to form many sub-populations. Panel D (E sorted) gives the similarity matrix from posterior (iteration 2500 to 5000).

Chart

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Chart, histogram

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Diagram

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I tried to held-out half of the data in a speckled pattern and fit the model with clustering on and off (single population). The dimension is selected by held-out likelihood, which is for clustering on and for single population analysis. The trace plots of the held-out log-likelihood per spike (starting from iteration 2) shows that doing clustering doesn’t improve things…

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# Discussion

# Reference

# Appendix