

Sede Amministrativa: Università degli Studi di Padova

#### Dipartimento di Scienze Statistiche Corso di Dottorato di Ricerca in Scienze Statistiche Ciclo XXXIV

### **Thesis Title**

Coordinatore del Corso: Prof. Nicola Sartori

Supervisore: Prof. Antonio Canale

Co-supervisore: Prof. Michele Guindani

Dottoranda: Laura D'Angelo

Day Month Year of submission

## **Abstract**

Abstract content.

## Sommario

Contenuto del sommario.

# **Contents**

Li	st of	Figures	<b>;</b>	vi
Li	st of	Tables		vii
In	trodu	ıction		1
	Ove	rview		1
	Mai	n contr	ibutions of the thesis	2
1	Bac	kgroun	d: statistical modeling of calcium imaging data	3
	1.1	Overv	riew of calcium imaging data	
		1.1.1	Deconvolution methods	4
		1.1.2	Spike train data analysis	5
	1.2	Data s	sets	5
		1.2.1	Allen Brain Observatory data	
		1.2.2	Altri dati?	7
	1.3	A brie	ef review of some Bayesian nonparametric models	7
		1.3.1	Finite mixture models	8
		1.3.2	Dirichlet process mixture models	10
		1.3.3	Mixtures of finite mixtures	14
		1.3.4	Bayesian nonparametric models for nested data	17
2	Effic	cient po	osterior sampling for Bayesian log-linear models	21
	2.1	Algor	ithm	21
3	Bay	esian n	onparametric analysis for partially exchangeable single-neuron data	23
	3.1	Mode	l specification	23
	3.2		rior computation	
Bi	hling	raphy		25

# **List of Figures**

1.1	Allen Brain Observatory data: calcium traces of two neurons located in the	
	primary visual area during Session A and Session B of the experiment	7

# **List of Tables**

## Introduction

#### **Overview**

A fundamental but unsolved problem in neuroscience is understanding the functioning of neurons and neuronal networks in processing sensory information, generating locomotion, and mediating learning and memory. The investigation of the structure and function of the nervous system can be dated back to the nineteenth century with the invention of the technique of silver impregnation by Camillo Golgi in 1873, which allowed the visualization of individual neurons (Drouin et al., 2015). The technique initiated the study of the microscopic anatomy of the nervous system, and the investigation of how neurons organize to form the brain. Ever since there has been a significant research effort both to discover the cellular properties of the nervous system, and to characterize behaviors and correlate them with activity imaged in different regions of the brain. However, many scientists recognize that despite the innovative techniques developed to observe and analyze neurons, we are still facing an "explanatory gap" between the understanding of elemental components and the outputs that they produce (Parker, 2006, 2010; Dudai, 2004; Paninski and Cunningham, 2018). That is, we know a lot about the components of the nervous system, but still we have little insight into how these components work together to enable us to think, remember, or behave. One of the reasons of this gap is the availability of a huge quantity of data, but a lack of tools to integrate these data in order to obtain a coherent picture of the brain functioning (Parker, 2010).

The technological developments of the last few decades have opened fundamentally new opportunities to investigate the nervous system. Large neuronal networks can now be visualized using *in vivo* high-resolution imaging techniques, which permit to record the neuronal activity in freely moving animals over long periods of time. In this thesis, we focus on data resulting from the application of the two-photon calcium imaging technique. Calcium ions generate intracellular signals that determine a large variety of functions in all neurons: when a neuron fires, calcium floods the cell and produces a transient spike in its concentration (Grienberger and Konnerth, 2012). By using genetically encoded calcium indicators, which are fluorescent molecules that react when binding to the calcium ions, it is possible to optically measure the level of calcium by analyzing the observed fluorescence trace. However,

extracting these fluorescent calcium traces is just the first step towards the understanding of brain circuits: how to relate the observed pattern of neuronal activity with its output remains an open problem of research.

#### Main contributions of the thesis

## Chapter 1

# Background: statistical modeling of calcium imaging data

#### 1.1 Overview of calcium imaging data

Calcium ions generate intracellular signals that control key functions in all types of neurons. At rest, most neurons have an intracellular calcium concentration of about 100 nm; however, during electrical activity, the concentration can rise transiently up to levels around 1000 nm (Berridge *et al.*, 2000). The development of techniques that enable the visualization and quantitative estimation of the intracellular calcium signals have thus greatly enhanced the investigation of neuronal functioning. The development of calcium imaging techniques involved two parallel processes: the development of calcium indicators, which are fluorescent molecules that react when binding to the calcium ions, and the implementation of the appropriate imaging instrumentation, in particular, the introduction of two-photon microscopy (Denk *et al.*, 1990). In recent years, the innovation achieved in these two fields has allowed for real-time observation of biological processes at the single-cell level simultaneously for large groups of neurons (Grienberger and Konnerth, 2012).

The output two-photon calcium imaging is a movie of time-varying fluorescence intensities, and a first complex pre-processing phase deals with the identification of the spatial location of each neuron in the optical field and source extraction (Mukamel *et al.*, 2009; Dombeck *et al.*, 2010). The resulting processed data consist of a fluorescent calcium trace for each observable neuron in the targeted area which, however, is only a proxy of the underlying neuronal activity. Hence further analyses are needed to deconvolve the fluorescence trace to extract the spike train (i.e. the series of recorded firing times), and to try to explain how these firing events are linked with the experiment that generated that particular pattern of activity.

#### 1.1.1 Deconvolution methods

There is currently a rich literature of methods addressing the issue of deconvolving the raw fluorescent trace to extract the spike train. A successful approach is to assume a biophysical model to relate the spiking activity to the calcium dynamics, and to the observed fluorescence. Vogelstein  $et\ al.\ (2010)$  proposed a simple but effective model that has later been adopted by several authors (Pnevmatikakis  $et\ al.\ (2016)$ ; Friedrich and Paninski, 2016; Friedrich  $et\ al.\ (2017)$ ; Jewell and Witten, 2018; Jewell  $et\ al.\ (2019)$ . The model considers the observed fluorescence as a linear (and noisy) function of the intracellular calcium concentration; the calcium dynamics is then modeled using an autoregressive process with jumps in correspondence of the neuron's firing events. Denoting with  $y_t$  the observed fluorescence trace of a neuron and with  $c_t$  the underlying calcium concentration, for time  $t=1,\ldots,T$ , the model can be written as

$$y_t = b + c_t + \epsilon_t, \quad \epsilon_t \sim \mathcal{N}(0, \sigma^2),$$

$$c_t = \gamma c_{t-1} + A_t + w_t, \quad w_t \sim \mathcal{N}(0, \tau^2),$$

$$(1.1)$$

where b models the baseline level of the observed trace and  $\epsilon_t$  is a Gaussian measurement error. In the absence of neuronal activity, the true calcium concentration  $c_t$  is considered to be centered around zero. The parameter  $A_t$  captures the neuronal activity: in the absence of a spike  $(A_t = 0)$ , the calcium level follows a AR(1) process controlled by the parameter  $\gamma$ ; when a spike occurs, the concentration increases instantaneously of a value  $A_t > 0$ . A challenge remains estimating the neuronal activity  $A_t$  in a precise and computationally efficient way.

Vogelstein  $et\ al.\ (2010)$  assume that all spikes have a fixed amplitude, and interpret the parameter  $A_t$  as the number of spikes at time t. Following this definition, they place a Poisson prior distribution on  $A_t$ ; however, the maximum a posteriori estimation of the spike train using a Poisson distribution is computationally intractable. Hence they search an approximate solution by replacing the Poisson distribution with an exponential distribution of the same mean. This leads to some loss of interpretation of the parameters  $A_t$ , as now they are no longer integer values but rather non-negative real numbers, but turns the problem into a convex optimization, which can be solved efficiently. Adopting this approach leads to solving a non-negative lasso problem for estimating the calcium concentration, where the  $L_1$  penalty enforces sparsity of the neural activity. Efficient algorithms to obtain a solution of this problem were also proposed by Pnevmatikakis  $et\ al.\ (2016)$ , Friedrich and Paninski (2016), and Friedrich  $et\ al.\ (2017)$ .

A different perspective is instead proposed by Jewell and Witten (2018) and Jewell *et al.* (2019): rather than interpreting  $A_t$  in model (1.1) as the number of spikes at the t-th timestep, they interpret its sign as an indicator for whether or not *at least one* spike occurred, that is,  $A_t = 0$  indicates no spikes at time t, and  $A_t > 0$  indicates the occurrence of at least one spike. The model so formulated includes an indicator variable, which corresponds to using an  $L_0$  penalization and which makes the optimization problem highly non-convex. In their work,

Jewell and Witten (2018) and Jewell *et al.* (2019) develop fast algorithms to compute the spike trains under these assumptions. Jewell and Witten (2018) assert that the solutions discussed by Vogelstein *et al.* (2010), Friedrich and Paninski (2016), and Friedrich *et al.* (2017) can actually be seen as convex relaxations of this optimization problem, to overcome the computational intractability of the  $L_0$  penalization.

Finally, Pnevmatikakis  $et\ al.\ (2013)$  propose a fully Bayesian approach. Although less computationally efficient than optimization methods, it allows to obtain a posterior distribution of all model parameters instead of just a point estimate, hence improving uncertainty quantification. Differently from previous models, they define the parameter  $A_t$  as the *amplitude* of a spike at time t, taking values in the non-negative real numbers. They formulate the presence/absence of a spike and its amplitude by using the product of a Bernoulli random variable (taking value 0 if there is no spike at time t, and 1 otherwise) with a half-Gaussian random variable (modeling the positive amplitudes). However, they do not explicitly assume sparsity of the spikes.

#### 1.1.2 Spike train data analysis

Standard methods to analyze calcium imaging data rely on a two-step approach: in a first phase, some deconvolution method such as those just described is applied to identify the spikes, then, a different method is used on the deconvolved output to analyze it and, possibly, to relate it with some covariates. However, while there is a rich literature on deconvolution methods, there is still little research on methods that try to derive inferential results from their output.

Concerning encoding models, i.e. models that try to understand how the brain encodes external variables into spike train, Paninski *et al.* (2007) focus on the use of generalized linear models (GLMs). GLMs allow spike trains to be regressed against a potentially large number of covariates such as behavioral parameters, experimental conditions and other relevant factors. Moreover, they provide simple and interpretable results of the effect of each covariate on the neuronal response.

A different approach is proposed by Wei et al. (2019)

#### 1.2 Data sets

Una frase introduttiva? Parlo dei dati dell'Allen Brain Observatory e poi ci sarebbe da mettere i nuovi dati se riesco a fare qualcosa del progetto 3...

#### 1.2.1 Allen Brain Observatory data

The Allen Brain Observatory (Allen Institute for Brain Science, 2016) is a large public data repository for investigating how sensory stimulation is represented by neural activity in the

6 Section 1.2 - Data sets

mouse visual cortex in both single cells and populations. The project aims to provide a standardized and systematic survey to measure and analyze visual responses from neurons across cortical areas and layers, utilizing transgenic Cre lines to drive expression of genetically encoded fluorescent calcium indicators, and measured by *in vivo* two-photon calcium imaging.

The study is an extended survey of physiological activity in the mouse visual cortex in response to a range of visual stimuli (Allen Brain Observatory, 2017). Each mouse is placed in front of a screen where different types of visual stimuli are shown, while the mouse's neuronal activity is recorded. The stimuli vary from simple synthetic images such as locally sparse noise or static gratings, to complex natural scenes and movies. The goal of the study is to investigate how neurons at different depths and in different areas of the visual cortex respond to stimuli of varying complexity, to understand their functional properties. Specifically, each neuron in the visual cortex can be characterized by their receptive field, i.e. the features of the visual stimulus that trigger the signaling of that neuron. An important finding from mammalian is that higher visual areas tend to respond to more complex stimuli relative to lower areas. These differences indicate that the different neurons and visual areas have distinct functional properties. Hence, it is of critical interest to devise methods that allow inferring how the neuronal response varies under the different types of visual stimuli. As an example, Figure 1.1 shows the calcium traces of two neurons recorded during two different experiment sessions from the Allen Brain Observatory study. Each experiment comprises three types of visual stimuli, and has a duration of approximately one hour. These plots highlight that the neuronal response is highly variable, both across experimental conditions and between neurons.

The Allen Brain Observatory study comprises records of neuronal activity from over 60000 cells from six visual areas (VISp, VISl, VISal, VISrl, VISam, and VISpm) and different imaging depths (ranging from 175 to 625 microns). The data were collected analyzing the brain activity of several genetically engineered mice, using different transgenic Cre lines. The neuronal response of each mouse was recorded during three experimental sessions: specifically, each session was made up of different types of visual stimuli displayed sequentially. Session "A" comprises two natural movies and a drifting gratings stimulus; session "B" comprises both natural movies and natural scenes, and a static gratings stimulus; finally, session "C" again includes two natural movies, and a locally sparse noise. Moreover, in all sessions a period of absence of stimuli was used to evaluate the baseline response during spontaneous activity. The synthetic stimuli are used to investigate the parameters that trigger the neuronal response: the locally sparse noise displays white and black spots in different parts of the visual space, and allows to map the spatial size and shape of the receptive field. The grating stimulus is a simple pattern whose intensity varies periodically along one dimension, and is constant in the other dimension. Different spatial frequencies, temporal frequencies and orientations are considered in order to further characterize the receptive fields. A detailed description of the visual stimuli can be found in a technical report (Allen Brain Observatory, 2017).

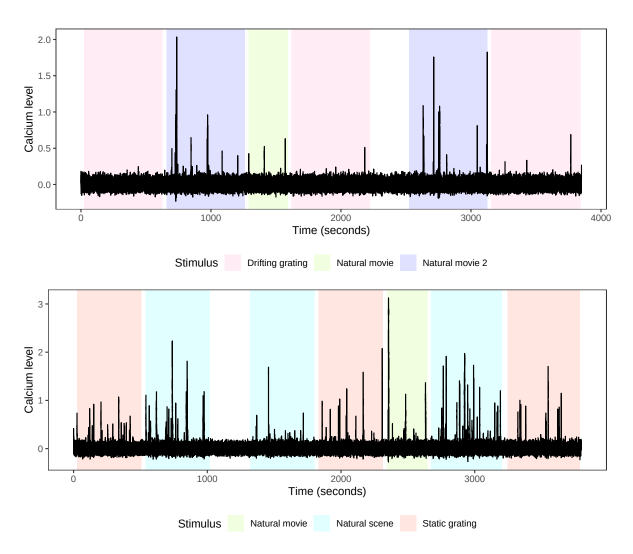


Figure 1.1: Allen Brain Observatory data: calcium traces (black line) of two neurons located in the primary visual area during Session A (upper plot) and Session B (bottom plot) of the experiment. The background colors denote the visual stimulus displayed at each time.

#### 1.2.2 Altri dati?

Paragrafo qui.

### 1.3 A brief review of some Bayesian nonparametric models

In this section we review some statistical tools that will be employed in this thesis in the analysis of calcium imaging data. The purpose of this section is not to provide a comprehensive review, but rather to outline the theoretical framework we adopted and fix some notation. The core topic will be the Bayesian methodology, with a focus on Bayesian nonparametric models.

#### 1.3.1 Finite mixture models

We start our discussion by reviewing finite mixtures. Although they are not part of the Bayesian nonparametric methodology, they provide the starting point for many models that we will review in the following. The content of this brief overview on finite mixtures is largely based on the dedicated chapter in Gelman *et al.* (2013).

#### Definition and hierarchical representations

Mixtures are a popular tool to model heterogeneous data, characterized by the presence of subpopulations within the overall population. In many practical problems the data are collected under different conditions – unfortunately, it is not always possible to have information on the subpopulation to which each individual observation belongs. Mixture models can be used in problems of this type, where the population consists of a number of latent subpopulations, and within each of them a relatively simple model can be applied.

Denote the observed data as a vector of n units  $\mathbf{y}=(y_1,\ldots,y_n)$ ; also, assume that the n observations are exchangeable, meaning that the joint probability density  $p(y_1,\ldots,y_n)$  is invariant to permutations of the indices. In the framework of finite mixtures, we assume that the population is made of  $K \leq n$  subpopulations, with K known and fixed. We assume that within each of these groups, the distribution of  $y_i$ ,  $i=1,\ldots,n$ , can be modeled as  $f(y_i\mid\theta_k^*)$ , for  $k=1,\ldots,K$ . Usually a common parametric family is assumed for all these component distributions, which however depend on specific parameter vectors  $\theta_k^*$ . The last missing piece to construct a mixture model is the parameter describing the proportion of population from each component k: we denote this parameter with  $\pi_k$ , satisfying  $\sum_{k=1}^K \pi_k = 1$ . Denoting the full vectors of parameters as  $\mathbf{\theta}^* = (\theta_1^*,\ldots,\theta_K^*)$  and  $\mathbf{\pi} = (\pi_1,\ldots,\pi_K)$ , the data distribution for observation i can be formulated as

$$p(y_i \mid \boldsymbol{\theta}^*, \boldsymbol{\pi}) = \pi_1 f(y_i \mid \theta_1^*) + \dots + \pi_K f(y_i \mid \theta_K^*).$$

In mixture models it is convenient to think of the component indicators as missing data, and to impute them to obtain a much simpler form of the data distribution. Hence we introduce the indicator  $S_{ik}$  of component k for observation i, with

$$S_{ik} = \begin{cases} 1 & \text{if } y_i \text{ is drawn from component } k \\ 0 & \text{otherwise.} \end{cases}$$

Given  $\pi$ , the distribution of  $S_i = (S_{i1}, \dots, S_{iK})$  is Multinomial $(1; \pi_1, \dots, \pi_K)$ . Conditionally on  $S_i$ , the data distribution of  $y_i$  is simply  $p(y_i \mid S_i, \theta^*) = \prod_{k=1}^K f(y_i \mid \theta_k^*)^{S_{ik}}$ ; moreover, given  $S = (S_1, \dots, S_n)$ , the  $y_i$  are assumed to be independent. The joint density of the observed data

and the unobserved indicators, conditionally on the model parameters, can now be written as

$$p(\boldsymbol{y}, \boldsymbol{S} \mid \boldsymbol{\theta}^*, \boldsymbol{\pi}) = p(\boldsymbol{y} \mid \boldsymbol{S}, \boldsymbol{\theta}^*) p(\boldsymbol{S} \mid \boldsymbol{\pi}) = \prod_{i=1}^n \prod_{k=1}^K \left\{ \pi_k f(y_i \mid \boldsymbol{\theta}_k^*) \right\}^{S_{ik}}.$$

Having defined the data distribution, we need to specify adequate prior distributions on the model parameters  $\pi$  and  $\theta^*$ . The prior  $G_0$  on  $\theta^*$  is usually chosen depending on the specific application and on the basis of the component distribution f. For the mixture proportions  $\pi_k$ , the conjugate and most natural prior distribution is the Dirichlet distribution,  $\pi \sim \text{Dir}_K(\alpha_1, \dots, \alpha_K)$ .

This model also admits a useful hierarchical representation. Rewriting the latent allocation variables using the cluster indicators  $c_i \in \{1, ..., K\}$ , with  $c_i = k$  if  $y_i$  belongs to the k-th mixture component (i.e.  $S_{ik} = 1$ ), the model is, for i = 1, ..., n

$$y_{i} \mid c_{i} = k, \theta_{k}^{*} \sim f(y_{i} \mid \theta_{k}^{*})$$

$$\Pr(c_{i} = k \mid \pi_{1}, \dots, \pi_{K}) = \pi_{k} \quad \text{for } k = 1, \dots, K$$

$$\theta_{1}^{*}, \dots, \theta_{K}^{*} \sim G_{0}$$

$$\pi_{1}, \dots, \pi_{K} \sim \operatorname{Dir}_{K}(\alpha_{1}, \dots, \alpha_{K}).$$

$$(1.2)$$

It is possible to rewrite Equation (1.2) in a slightly different way by thinking that each observation  $y_i$  is associated with a parameter  $\theta_i$ , where these parameters are drawn from a discrete distribution G with support on the K locations  $\{\theta_1^*, \dots, \theta_K^*\}$ . The model then becomes, for  $i = 1, \dots, n$ 

$$y_{i} \mid \theta_{i} \sim f(y_{i} \mid \theta_{i})$$

$$\theta_{i} \mid \boldsymbol{\theta}^{*}, \boldsymbol{\pi} \sim G = \sum_{k=1}^{K} \pi_{k} \delta_{\theta_{k}^{*}}$$

$$\theta_{1}^{*}, \dots, \theta_{K}^{*} \sim G_{0}$$

$$\pi_{1}, \dots, \pi_{K} \sim \operatorname{Dir}_{K}(\alpha_{1}, \dots, \alpha_{K}).$$

$$(1.3)$$

#### Posterior inference for finite mixture models

Posterior inference for mixture models is usually performed through Markov Chain Monte Carlo (MCMC) methods and, in particular, the Gibbs sampler, as the full conditionals after imputing the cluster indicators  $\mathcal{C} = \{c_1, \ldots, c_n\}$  are greatly simplified. Moreover, for the distribution of the mixture weights it is possible to exploit the conjugacy of the Dirichlet distribution with the multinomial model. A Gibbs sampler then simply iterates these three steps:

1. Update the cluster-specific parameters  $\theta_k^*$ , for k = 1, ..., K, from

$$p(\theta_k^* \mid \mathcal{C}, \boldsymbol{y}) \propto G_0(\theta_k^*) \prod_{i:c_i=k} f(y_i \mid \theta_k^*).$$

2. Update the weights  $\pi_1, \dots, \pi_K$  by sampling from a Dirichlet distribution with updated parameters

$$\pi_1, \ldots, \pi_K \mid \mathcal{C} \sim \operatorname{Dir}_K(\alpha_1 + n_1, \ldots, \alpha_K + n_K)$$

where  $n_k$  is the number of observations allocated to cluster k, for k = 1, ..., K.

3. Update the cluster indicators: for i = 1, ..., n and k = 1, ..., K,

$$\Pr(c_i = k \mid \boldsymbol{\pi}, \boldsymbol{\theta}^*, y_i) \propto \pi_k f(y_i \mid \theta_k^*).$$

#### 1.3.2 Dirichlet process mixture models

Nonparametric mixtures extend model (1.3) by placing a nonparametric prior on G. The most common prior on random probability measures is the Dirichlet process (DP), introduced by Ferguson (1973, 1974). Draws from a DP are discrete distributions with probability one, hence they turned out useful as flexible mixing measures in discrete mixtures.

#### The Dirichlet process

The Dirichlet process is a stochastic process whose realizations are probability distributions with probability one. Stochastic processes are distributions over function spaces, with their realizations being random functions. In the case of the DP, it is a distribution over the space of probability measures, which are real-valued functions with particular properties, which can be interpreted as distributions over some probability space. For this short review of some of the main properties of the DP we followed the unpublished report by Yee Whye Teh, which illustrates in a clear and simple way the fundamental properties of this process.

Formally, a random distribution G on some probability space  $\Theta$  is said to follow a DP prior with base measure  $G_0$  and concentration parameter  $\alpha$ , denoted  $G \sim \mathrm{DP}(\alpha, G_0)$ , if for any partition  $\{B_1, \ldots, B_H\}$  of  $\Theta$ 

$$(G(B_1),\ldots,G(B_H)) \sim \operatorname{Dir}_H(\alpha G_0(B_1),\ldots,\alpha G_0(B_H)).$$

That is, the finite-dimensional marginal distributions of a DP are Dirichlet distributions.

The success of the DP mainly arises from two appealing characteristics: its large support, with respect to the space of probability distributions, and tractability of the posterior distribution. Closely related to this last aspect is the conjugacy property of the DP: as the finite dimensional Dirichlet distribution is conjugate to the multinomial likelihood, the DP is conjugate with respect to i.i.d. sampling, that is, with respect to a completely unknown distribution from i.i.d. data. More precisely, if we take  $\{\theta_1,\ldots,\theta_n\}$  a sequence of independent draws from  $G \sim \mathrm{DP}(\alpha,G_0)$ , then the posterior distribution of G given these observed values is still a DP. Letting again  $\{B_1,\ldots,B_H\}$  be a finite measurable partition of  $\Theta$ , and letting  $n_h$  be the number

of observed values in  $B_h$ , for h = 1, ..., H, the posterior distribution is given by

$$(G(B_1),\ldots,G(B_H)) \mid \theta_1,\ldots,\theta_n \sim \operatorname{Dir}_H(\alpha G_0(B_1) + n_1,\ldots,\alpha G_0(B_H) + n_H).$$

In other terms, the posterior distribution is still DP with updated parameters:

$$G \mid \theta_1, \dots, \theta_n \sim \mathrm{DP}\left(\alpha + n, \frac{\alpha G_0 + \sum_{i=1}^n \delta_{\theta_i}}{\alpha + n}\right)$$

where the posterior base measure is a weighted average between the prior base measure  $G_0$  and the empirical distribution  $\sum_{i=1}^{n} \delta_{\theta_i}/n$ . The weight associated with the prior base distribution is proportional to  $\alpha$ , while the empirical distribution has weight proportional to the number of observations.

Another useful result, which allows to get a better understanding of the effect of using a DP as mixing measure, is the represented by Blackwell-MacQueen urn scheme (Blackwell and MacQueen, 1973), which describes the predictive distribution of draws from a DP. Consider again a sequence  $\{\theta_1,\ldots,\theta_n\}$  of independent draws from  $G \sim \mathrm{DP}(\alpha,G_0)$ . The predictive distribution of  $\theta_{n+1}$  conditioned on these values, and with G marginalized out is given by

$$\theta_{n+1} \mid \theta_1, \dots, \theta_n \sim \frac{1}{\alpha + n} \left( \alpha G_0 + \sum_{i=1}^n \delta_{\theta_i} \right).$$

Therefore the posterior base measure given  $\{\theta_1,\ldots,\theta_n\}$  is also the predictive distribution of  $\theta_{n+1}$ . This distribution highlights the discreteness of draws from a DP, and allows to investigate the clustering structure induced by the DP when is used as mixing measure in mixture models. Since the distribution is discrete, it is possible that some of the values  $\{\theta_1,\ldots,\theta_n\}$  will be repeated. In particular, the unique values of  $\{\theta_1,\ldots,\theta_n\}$  induce a partition of the set  $\{1,\ldots,n\}$  into clusters defined by observations with the same value. Denoting with  $\{\theta_1^*,\ldots,\theta_K^*\}$  the unique values among the  $\theta_i$ , and letting  $n_k$  be the number of  $\theta_i$  equal to  $\theta_k^*$ , for  $i=1,\ldots,n$  and  $k=1,\ldots,K$ , the predictive distribution can be written as

$$\theta_{n+1} \mid \theta_1, \dots, \theta_n \sim \frac{1}{\alpha + n} \left( \alpha G_0 + \sum_{k=1}^K n_k \delta_{\theta_k^*} \right).$$

From this equation, it is possible to notice that  $\theta_{n+1}$  will take a value  $\theta_k^*$  with a probability proportional to  $n_k$ , the number of times it has already been observed. Hence, the larger  $n_k$  is, the higher the probability that it will grow. This is a rich-gets-richer phenomenon, where large clusters grow larger faster.

The DP admits several nice representations. An intuitive constructive definition of a DP random probability measure is given by Sethuraman (1994) and is based on the discrete nature of the process, which can be represented as a weighted sum of point masses. This

definition states that if  $G \sim DP(\alpha, G_0)$ , then it can be expressed as follows:

$$v_k \sim \text{Beta}(1, \alpha), \quad \theta_k^* \sim G_0 \quad \text{for } k \ge 1$$

$$\pi_1 = v_1, \quad \pi_k = v_k \prod_{h=1}^{k-1} (1 - v_h) \quad \text{for } k \ge 2$$

$$G = \sum_{k=1}^{\infty} \pi_k \delta_{\theta_k^*}.$$

$$(1.4)$$

The construction of the weights  $\{\pi_k\}_{k=1}^{\infty}$  by means of beta random variable is usually called stick-breaking process, also denoted as  $\pi \sim \operatorname{GEM}(\alpha)$  after the names of Griffiths, Engen and McCloskey. The name arises from a metaphor for this construction, where a unit stick is broken in infinitely many parts, and each piece is used to define a weight. Because of its simplicity, this representation has motivated extensions of the process as well as new inference procedures, as for example the algorithms described in the following to sample from the posterior distribution of Dirichlet process mixtures.

#### Dirichlet process mixtures

Getting back to the framework of mixture models, from the representation of the process as discrete measure it is straightforward to extend the finite mixtures described in the previous section to nonparametric mixtures: following the structure of Equation (1.3), DP mixtures (DPMs) can be written as

$$y_i \mid \theta_i \sim f(y_i \mid \theta_i)$$
  
 $\theta_i \mid G \sim G$   
 $G \sim DP(\alpha, G_0).$ 

Alternatively, making use of the set of unique values  $\{\theta_k^*\}_{k=1}^{\infty}$  and of Sethuraman's representation, the model can be expressed as

$$p(y_i \mid G) = \sum_{k=1}^{\infty} \pi_k f(y_i \mid \theta_k^*)$$
(1.5)

where the weights  $\{\pi_k\}_{k=1}^{\infty}$  follow a stick-breaking construction and the  $\{\theta_k^*\}_{k=1}^{\infty}$  are i.i.d. samples from the base measure  $G_0$ .

Differently from finite mixtures, DPMs are infinite mixture models, as they assume a countably infinite number of components. However, because the  $\pi_k$ 's decrease exponentially quickly, only a small number of components will be used to model the data a priori: in the following, we will define *clusters* these non-empty components. In general, the prior expected number of clusters in a sample of size n is approximately equal to  $\alpha \log(1 + n/\alpha)$ . This means that while in finite mixture models the number of clusters has to be fixed in advance, in DPMs the number of clusters is determined by the data and can be inferred.

#### Posterior inference for DPMs

Applying MCMC techniques to DPMs directly is not feasible, as it would require imputing the infinite-dimensional distribution G. Instead, successful algorithms to perform posterior inference on DP mixtures have been developed using the particular representations that this process admits. It is common to divide these methods into two classes depending on the strategy they adopt to deal with the infinite-dimensional component of the model: marginal algorithms are based on model representations with the DP is integrated out, while conditional algorithms explicitly represent the measure generated by the process using its stick-breaking construction. Here we will focus on conditional methods. Avoiding marginalization of the random measure allows to perform inference on it directly, moreover, these methods are often computationally more efficient than marginal ones; however, they require to devise good strategies to deal with the infinite dimension of the process.

In the blocked Gibbs sampler of Ishwaran and James (2001) the infinite random measure is truncated to an upper bound K for the number of clusters. The motivation that justifies this approach is that the weights  $\{\pi_k\}_{k=1}^{\infty}$  determined by the stick-breaking construction are stochastically decreasing in k. By choosing a sufficiently large value K one can assume that  $\sum_{k=K+1}^{\infty} \pi_k$  has a distribution concentrated near zero. Adopting such truncation leads to a representation of the model as finite mixture, so the sampler described in Section 1.3.1 can be applied with just few changes on the weights distribution. In particular, the weights are now sampled from a stick-breaking process with  $v_k \sim \text{Beta}(1+n_k,\alpha+\sum_{h=k+1}^K n_h)$  for  $k=1,\ldots,K-1$ . Finally, letting  $v_K=1$  ensures that the first K weights sum to one. Adopting this approach leads to a straightforward MCMC algorithm for posterior inference, however, in some applications K needs to be set to very large values in order to obtain an adequate approximation.

The slice sampler (Walker, 2007; Kalli *et al.*, 2011) has been adopted as an alternative "dynamic" truncation method to automatically select the active components of the mixture. The slice sampler relies on the introduction of latent uniform random variables  $u_i$ ,  $i=1,\ldots,n$ , such that marginalizing the joint density of  $(y_i,u_i)$  still returns the desired density in Eq. (1.5), and such that the conditional density of  $y_i \mid u_i$  only involves a finite number of mixture components. Specifically, model (1.5) can be obtained as marginal density with respect to  $u_i$  of

$$p(y_i, u_i \mid G) = \sum_{k=1}^{\infty} \mathbb{I}(u_i < \pi_k) f(y_i \mid \theta_k^*),$$

where  $\mathbb{I}(A)$  is the indicator variable, taking value 1 if event A occurs and 0 otherwise. The key of introducing this variable is that, conditionally on  $u_i$ , the density can be written as

$$p(y_i \mid u_i, G) = N_u^{-1} \sum_{k \in A_u} f(y_i \mid \theta_k^*)$$

where  $A_u$  is the set of indices of active components  $A_u = \{k : \pi_k > u\}$  and  $N_u = \sum_{k \in A_u} \pi_k$ . This model defines a finite mixture with equal weights  $N_u^{-1}$ . Introducing the cluster indicators  $\mathcal{C} = \{c_1, \ldots, c_n\}$  further simplifies the density, similarly to the case of finite mixtures. Finally, Kalli *et al.* (2011) also introduce a non-stochastic positive sequence  $\{\xi_1, \xi_2, \ldots\}$  in order to improve mixing (we refer to the original paper for a discussion on the choice of the sequence). With all these elements, the joint density of  $(y, u, \mathcal{C})$  conditioned on  $\pi$  and  $\theta^*$  becomes

$$p(\boldsymbol{y}, \boldsymbol{u}, \mathcal{C} \mid \boldsymbol{\pi}, \boldsymbol{\theta}^*) = \prod_{i=1}^n \mathbb{I}(u_i < \xi_{c_i}) \frac{\pi_{c_i}}{\xi_{c_i}} f(y_i \mid \theta_{c_i}^*).$$
(1.6)

The slice sampler algorithm iterates the following steps:

1. Update the cluster-specific parameters  $\theta_k^*$ , for k = 1, ..., K, where K is defined as the maximum index h such that  $\xi_h > u_i$  for all i = 1, ..., n, from

$$p(\theta^* \mid \mathcal{C}) \propto G_0(\theta_k^*) \prod_{i:c_i=k} f(y_i \mid \theta_k^*).$$

- 2. Update the weights  $\pi_k$  for k = 1, ..., K using the stick-breaking construction (1.4) with beta random variables  $v_k \sim \text{Beta}(1 + n_k, \alpha + \sum_{h=k+1}^K n_h)$ .
- 3. Sample the latent variables  $u_i$  from a uniform distribution  $u_i \mid c_i \sim \text{Unif}(0, \xi_{c_i})$
- 4. Update the cluster allocations  $c_i$  for i = 1, ..., n from

$$\Pr(c_i = k \mid u_i, y_i, \theta_k^*) \propto \mathbb{I}(k : \xi_k > u_i) \frac{\pi_k}{\xi_k} f(y_i \mid \theta_k^*).$$

#### 1.3.3 Mixtures of finite mixtures

To avoid fixing the number of clusters, a different approach to DP mixtures is to consider finite mixtures with a prior on the number of components. Although this may seem as the most natural way to infer the number of groups, application of mixtures of finite mixtures (MFMs) has long been hindered by the difficulty of performing posterior inference. Several inference methods have been proposed for this type of models (McCullagh and Yang, 2008; Nobile, 2004; Nobile and Fearnside, 2007; Richardson and Green, 1997) often exploiting the reversible jump Markov chain Monte Carlo techniques. However, applying the reversible jump in new situations can be hard, as it requires designing new reversible jump moves.

More recently, different models have been proposed inspired by nonparametric mixtures. Miller and Harrison (2018) explicitly derived MFMs counterparts for several properties exhibited by DPMs. They considered a finite mixture model similar to the one defined in section 1.3.1, where however the number of clusters is random and is assigned a prior distribution. A main limitation of their approach is that it assumes a fixed parameter  $\alpha$  for the Dirichlet distribution, regardless of the dimension K, which can lead to undesired

Another approach is discussed by Malsiner-Walli *et al.* (2016) and Frühwirth-Schnatter and Malsiner-Walli (2019), based on the use of sparse mixtures. Similarly to DPMs, in this formulation they distinguish between the mixture components and the clusters, which are defined as the components actually used by the data. In their approach, the number of components K is fixed to a large and clearly overfitting value, and a Dirichlet prior with small parameter on the mixture weights ensures that the (random) number of clusters  $K_+$  will take values smaller than K with high probability a priori and also a posteriori.

#### Generalized mixtures of finite mixtures

A general formulation that includes all three models described above as special cases (finite mixtures with a fixed number of clusters, MFMs and overfitted mixtures) has recently been described by Frühwirth-Schnatter *et al.* (2020): they call this specification generalized mixture of finite mixtures (gMFM). Similarly to overfitted mixtures, a key aspect of this approach is the distinction between the number of components K and the number of clusters  $K_+$ ; however, here, the number of components is also random. In the following we will review some aspects of these models that will be used later in this thesis. The gMFM model can be defined in a hierarchical form analogous to Equation (1.2) as

$$y_i \mid K, c_i = k, \theta_k^* \sim f(y_i \mid \theta_k^*)$$

$$\Pr(c_i = k \mid K, \pi_1, \dots, \pi_K) = \pi_k \quad \text{for } k = 1, \dots, K$$

$$\theta_1^*, \dots, \theta_K^* \sim G_0$$

$$\pi_1, \dots, \pi_K \mid K, \alpha \sim \text{Dir}_K(\alpha/K, \dots, \alpha/K)$$

$$K \sim p(K)$$

Including a prior on K also induces a prior on the number of clusters  $K_+$ : here a crucial role is played by the sequence of concentration parameters of the Dirichlet distribution. Considering fixed parameters as in Miller and Harrison (2018), where a  $\operatorname{Dir}_K(1,\ldots,1)$  is used regardless of the value of K, leads to a prior expected number of clusters  $E(K_+)$  close to E(K) for many priors p(K). Having instead concentration parameters that decrease with increasing K induces randomness in the prior distribution of  $K_+ \mid K$ , allowing for a gap between  $K_+$  and K. In this formulation the parameters decrease linearly with K, and a K prior distribution is used for the hyperparameter K. The specification of a gMFM is completed with a suitable prior K0 on the number of components. In their work, Frühwirth-Schnatter K1 desirable prior on K2 should be weakly informative on the number of clusters. A desirable prior on K3 should be weakly informative on the number of clusters, and should lead to a prior on K4 which is concentrated on moderate number of clusters, with fat tails to ensure that also a high number of clusters may be estimated. They suggest to use a translated prior for K5, where

K-1 follows a beta-negative-binomial distribution, which is a hierarchical generalization of the Poisson, the geometric and the negative-binomial distribution.

#### Posterior inference for gMFMs

An important contribution of the work of Frühwirth-Schnatter et~al.~(2020) is the introduction of a new inference algorithm "telescoping sampler" to obtain the posterior distribution of all model parameters without resorting to reversible jump MCMC techniques. Their algorithm is a trans-dimensional Gibbs sampler which at each iteration alternates two key steps: first, it updates the partition of the observations  $\mathcal{C} = \{c_1, \ldots, c_n\}$  conditionally on the number of components, then, it samples a new value for K on the basis of this partition. Explicitly sampling the number of mixture components greatly simplifies inference as, conditionally on K, the updates of the partition and of the model parameters are brought back to standard steps. Hence the crucial aspect is sampling from the full conditional of the number of components. This is achieved by combining the conditional exchangeable partition probability function  $p(\mathcal{C} \mid n, K, \alpha)$ , derived in Section 2.2 of Frühwirth-Schnatter et~al.~(2020), with the prior p(K). The algorithm performs the following steps:

- 1. Update the partition C:
  - (a) Sample  $c_i$ , for i = 1, ..., n from  $\Pr(c_i = k \mid \boldsymbol{\pi}, \boldsymbol{\theta}^*, y) \propto \pi_k f(y_i \mid \theta_k^*)$ ;
  - (b) Determine the number  $K_+$  of non-empty clusters and relabel the components such that the first  $K_+$  clusters are non-empty.
- 2. Conditional on C, update the parameters of the non-empty components (and eventual hyperparameters):

$$p(\theta_k^* \mid \mathcal{C}, \boldsymbol{y}) \propto G_0(\boldsymbol{\theta}^*) \prod_{i:c_i=k} f(y_i \mid \theta_k^*) \text{ for } k = 1, \dots, K.$$

3. Conditional on C, draw a new value for K from

$$p(K \mid \mathcal{C}, \alpha) \propto p(\mathcal{C} \mid n, K, \alpha) p(K) = p(K) \frac{K! \, \alpha^{K_+}}{(K - K_+)! \, K^{K_+}} \prod_{k=1}^{K_+} \frac{\Gamma(n_k + \alpha/K)}{\Gamma(1 + \alpha/K)}$$

for  $K = K_+, K_+ + 1, ...$ ; where  $n_k$  is the number of observations in cluster k.

4. Update  $\alpha$  by performing a Metropolis-Hastings step to sample  $\alpha$  from its full conditional

$$p(\alpha \mid \mathcal{C}, K) \propto p(\alpha) \frac{\alpha^{K_+} \Gamma(\alpha)}{\Gamma(n+\alpha)} \prod_{k=1}^{K_+} \frac{\Gamma(n_k + \alpha/K)}{\Gamma(1+\alpha/K)}$$

5. Add  $K - K_+$  empty components,

- (a) if  $K > K_+$ , sample  $K K_+$  new values  $\theta_k^*$  from the prior,  $k = K_+ + 1, \dots, K$ ;
- (b) update the weight vector as

$$\pi_1, \ldots, \pi_K \mid K, \alpha, \mathcal{C} \sim \text{Dir}_K(\alpha/K + n_1, \ldots, \alpha/K + n_K).$$

#### 1.3.4 Bayesian nonparametric models for nested data

All models described so far assumed that the data were exchangeable, that is, they arise from one unknown distribution. However, there is growing interest in modeling scenarios where the data come from different but related groups, as for example different populations or experiments. In these cases it is desirable to individually model the distribution of each group, while also borrowing information between them. When data are collected from individuals in multiple groups, the exchangeability assumption is no longer valid: in these cases, observations are said to be partially exchangeable, meaning that they are exchangeable *within* groups.

Suppose that in addition to each  $y_i$ ,  $i=1,\ldots,n$ , we also observe a categorical variable  $g_i$  with values in  $\{1,\ldots,J\}$  indicating the population to which  $y_i$  belongs, so that when  $g_i=j$  means that  $y_i$  comes from the j-th group. In the Bayesian nonparametric framework, several approaches have been proposed to deal with these nested data sets.

The hierarchical Dirichlet process (Teh  $et\,al.$ , 2006) arises from the desire to flexibly model the distributions of the observations of each group, while also performing clustering to capture latent structures among all individuals. Each group-specific distribution is modeled with a mixture model which uses a random probability measure  $G_j$  as mixing measure, where the  $G_j$ 's are distributed according to a DP: this allows to obtain a partition of individuals within each group. In order to borrow information across groups, they propose a hierarchical formulation where one draw from a Dirichlet process is used as the base measure  $G_0$  of the Dirichlet process generating the individual  $G_j$ 's. This construction implies that the distributions  $\{G_1,\ldots,G_J\}$  share the same atoms (the atoms of  $G_0$ ), thus the model yields a clustering of the individuals also across groups. However, as these  $G_j$ 's are independent draws, in general, they will have different weights: as a result, there is no clustering of these group-specific distributions. The hierarchical DP hence does not allow to investigate similarities between the distributions, but only to cluster individuals.

To overcome this limitation, Rodríguez *et al.* (2008) introduced the nested Dirichlet process, which allows to obtain a clustering both of the observations within each group, and of the groups themselves. Consider again a collection of distributions  $\{G_1, \ldots, G_J\}$  serving as group-specific mixing measures of a nonparametric mixture. The nested DP assumes that

 $G_j \mid Q \sim Q$  for j = 1, ..., J and  $Q \sim DP(\alpha, DP(\beta, G_0))$ . Using the stick-breaking representation of the DP, this model can be expressed as

$$G_1, \dots, G_J \mid Q \sim Q, \qquad Q = \sum_{k=1}^{\infty} \pi_k \delta_{G_k^*}$$

$$G_k^* = \sum_{l=1}^{\infty} \omega_{l,k} \delta_{\theta_{l,k}^*}$$

$$(1.7)$$

where the sequences of weights  $\{\pi_k\}_{k=1}^{\infty}$  and  $\{\omega_{l,k}\}_{l=1}^{\infty}$  for  $k \geq 1$  follow a stick breaking construction,  $\pi \sim \operatorname{GEM}(\alpha)$  and  $\omega_k \sim \operatorname{GEM}(\beta)$ , and the parameters  $\theta_{l,k}^*$  are i.i.d. samples from  $G_0$ . A main difference from the hierarchical DP is that this formulation allows to cluster the group-specific distributions: indeed, if two  $G_j$ 's are assigned to the same  $G_k^*$ , then the observations in the two groups have exactly the same generating distribution. When two groups share the same distribution, they are said to belong to the same distributional cluster.

In a recent paper by Camerlenghi *et al.* (2019), however, they noted a degeneracy property that occurs in the nested DP in case of ties across samples at the observed or latent level. In particular, if two distributions  $G_1$  and  $G_2$  share at least one atom, then their posterior distribution degenerates on  $\{G_1 = G_2\}$ , forcing homogeneity across the two samples. To overcome this drawback, they introduce a novel class of latent nested processes. These processes are based on a mixture of two random distributions: an idiosyncratic component and a shared component. The shared random distribution explicitly accounts for the possibility of observing some common atoms, while the idiosyncratic one allows some atoms to be distinct and specific to each subpopulation. However, implementation of this model becomes challenging and computationally burdensome when the number of groups increases.

#### The common atoms model

Another nested nonparametric prior, more suited to practical applications, is proposed by Denti  $et\ al.\ (2021)$ . They formulate a constrained modification of the nested DP which, however, does not suffer from the degeneracy issue. Moreover, compared to the models introduced by Camerlenghi  $et\ al.\ (2019)$ , it is computationally more efficient and allows to perform posterior inference in a quite straightforward way even when the number of groups is moderate. The first level of their common atoms model (CAM) is analogous to the nested DP (Eq. 1.7), however, the specification of the distributional atoms  $G_k^*$  makes use of a common set of atoms for all k. Hence the distributions  $G_k^*$  can be seen as realizations of a single-atom dependent DP,

$$G_k^* = \sum_{l=1}^{\infty} \omega_{l,k} \delta_{\theta_l^*}$$

where the sequences of weights  $\{\omega_{l,k}\}_{l=1}^{\infty}$  for  $k \geq 1$  again follow a stick breaking construction, and the common atoms  $\{\theta_l^*\}_{l=1}^{\infty}$  are independent draws from a base measure  $G_0$ . Similarly

to the nested DP, the first mixture level of the CAM allows to perform a clustering of the group-specific distributions  $G_j$ ; however, as the set of atoms defining the  $G_k^*$  is common across distributions, the CAM also allows to obtain a clustering of individuals both within each group and across them, in a similar fashion to the hierarchical DP.

Convoluting this nested prior with a kernel we obtain a nested infinite mixture model: the density for observations  $y_j = \{y_i : g_i = j; i = 1, ..., n\}$  belonging to group j can be written as

$$p(\mathbf{y}_j \mid \boldsymbol{\theta}^*, \boldsymbol{\pi}, \boldsymbol{\omega}) = \sum_{k=1}^{\infty} \pi_k \prod_{i:g_i=j} \sum_{l=1}^{\infty} \omega_{l,k} f(y_i \mid \boldsymbol{\theta}_l^*).$$

Introducing two sequences of latent cluster allocations  $C^D = \{c_{g_i}^D\}$  and  $C = \{c_i\}$  for i = 1, ..., n, corresponding respectively to the distributional cluster allocation of the J groups and to the observational cluster allocation of individuals, the model admits a hierarchical representation as

$$y_i \mid c_i, \boldsymbol{\theta}^* \sim f(y_i \mid \boldsymbol{\theta}_{c_i}^*)$$

$$c_i \mid c_{g_i}^D = k, \boldsymbol{\omega}_k \sim \sum_{l=1}^{\infty} \omega_{l,k} \delta_l \qquad \boldsymbol{\omega}_k \sim \text{GEM}(\beta) \quad \text{for } k \ge 1$$

$$c_{g_i}^D \mid \boldsymbol{\pi} \sim \sum_{k=1}^{\infty} \pi_k \delta_k \qquad \boldsymbol{\pi} \sim \text{GEM}(\alpha)$$

$$\boldsymbol{\theta}_l^* \sim G_0 \quad \text{for } l \ge 1.$$

#### Posterior inference for the CAM

To infer on the posterior distribution of the CAM model, Denti et~al.~(2021) develop a nested version of the slice sampler of Kalli et~al.~(2011) (described in the last part of Section 1.3.2). This sampler relies on two sequences of latent uniform variables  $\boldsymbol{u}^D = \{u_{g_i}\}_{g_i=1}^J$  (on the distributional level) and  $\boldsymbol{u}^O = \{u_i^O\}_{i=1}^n$  (on the observational level) pointing to the active mixture components. Moreover, they also introduce two deterministic sequences  $\boldsymbol{\xi}^D = \{\xi_k\}_{k=1}^\infty$  and  $\boldsymbol{\xi}^O = \{\xi_{l,k}^O\}_{l=1}^\infty$  for  $k \geq 1$ . Similarly to Equation 1.6, the model can be written as

$$p(\boldsymbol{y}, \boldsymbol{u}^{D}, \boldsymbol{u}^{O}, \mathcal{C}^{D}, \mathcal{C} \mid \boldsymbol{\theta}^{*}, \boldsymbol{\pi}, \boldsymbol{\omega}) = \prod_{j=1}^{J} \mathbb{I}(u_{j}^{D} < \xi_{c_{j}^{D}}^{D}) \frac{\pi_{c_{j}^{D}}}{\xi_{c_{j}^{D}}^{D}} \prod_{i:g_{i}=j} \mathbb{I}(u_{i}^{O} < \xi_{c_{i}, c_{j}^{D}}^{O}) \frac{\omega_{c_{i}, c_{j}^{D}}}{\xi_{c_{i}, c_{j}^{D}}^{O}} f(y_{i} \mid \boldsymbol{\theta}_{c_{i}}^{*}).$$

Then, the algorithm iterates the following steps

- 1. Sample the latent uniform random variables
  - (a) At the distributional level, for  $j=1,\ldots,J$ , sample  $u^D_j \sim \mathrm{Unif}(0,\xi^D_{c^D_j})$
  - (b) At the observational level, for  $i=1,\ldots,n$ , sample  $u_i^O\sim \mathrm{Unif}(0,\xi^O_{c_i,c^D_{q_i}})$
- 2. Update the weight vectors  $\boldsymbol{\pi}$  and  $\boldsymbol{\omega}_k$ , for  $k \geq 1$

- (a) At the distributional level, sample the distributional stick-breaking proportions  $v_k \sim \text{Beta}(1+m_k,\alpha+\sum_{h=k+1}^{K^*}m_h)$ , where  $m_k$  is the number of groups in distributional cluster k.
- (b) At the observational level, for each  $k=1,\ldots,K^*$ , sample the stick-breaking proportions  $u_{l,k} \sim \text{Beta}(1+n_l^k,\beta+\sum_{h=l+1}^{L^*}n_h^k)$ , where  $n_l^k$  is the number of individuals assigned to observational cluster l and distributional cluster k.
- 3. Update the cluster indicators
  - (a) For the distributional clusters, sample the variables  $c_i^D$  from

$$\Pr(c_j^D = k \mid \boldsymbol{u}^D, \boldsymbol{\pi}, \boldsymbol{\omega_k}, \mathcal{C}) \propto \mathbb{I}(u_j^D < \xi_{c_j^D}^D) \frac{\pi_{c_j^D}}{\xi_{c_j^D}^D} \prod_{i:g_i = j} \omega_{c_i, k}$$

(b) For the observational clusters, sample the variables  $c_i$  from

$$\Pr(c_i = l \mid y_i, g_i = j, c_j^D, \mathbf{u}^O, \omega_{l, c_j^D}) \propto \mathbb{I}(u_i^O < \xi_{l, c_j^D}^O) \frac{\omega_{l, c_j^D}}{\xi_{l, c_j^D}^O} f(y_i \mid \theta_l^*).$$

4. Conditional on the observational clusters, sample the cluster-specific parameters  $\theta_l^*$  from

$$p(\theta^* \mid \mathcal{C}) \propto G_0(\theta_l^*) \prod_{i:c_i=l} f(y_i \mid \theta_l^*).$$

We refer to the Supplementary Material of the original paper for details about the specific sequences  $\xi^{O}$  and  $\xi^{D}$  and computation of the upper bounds  $K^*$  and  $L^*$  involved at step 2.

# Chapter 2

# Efficient posterior sampling for Bayesian log-linear models

2.1 Algorithm

## **Chapter 3**

Bayesian nonparametric analysis for partially exchangeable single-neuron data

- 3.1 Model specification
- 3.2 Posterior computation

- Allen Brain Observatory (2017) Technical whitepaper: stimulus set and response analyses. Available at: http://help.brain-map.org/display/observatory/Documentation.
- Allen Institute for Brain Science (2016) Allen brain observatory. Available at: http://observatory.brain-map.org/visualcoding.
- Berridge, M. J., Lipp, P. and Bootman, M. D. (2000) The versatility and universality of calcium signalling. *Nature Reviews Molecular Cell Biology* **1**, 11 21.
- Blackwell, D. and MacQueen, J. B. (1973) Ferguson distributions via Polya urn schemes. *The Annals of Statistics* **1**(2), 353 355.
- Camerlenghi, F., Dunson, D. B., Lijoi, A., Prünster, I. and Rodríguez, A. (2019) Latent Nested Nonparametric Priors (with Discussion). *Bayesian Analysis* **14**(4), 1303 1356.
- Denk, W., Strickler, J. H. and Webb, W. W. (1990) Two-photon laser scanning fluorescence microscopy. *Science* **248**, 73–76.
- Denti, F., Camerlenghi, F., Guindani, M. and Mira, A. (2021) A common atoms model for the bayesian nonparametric analysis of nested data. *Journal of the American Statistical Association* pp. 1–12.
- Dombeck, D. A., Harvey, C. D., Tian, L., Looger, L. L. and Tank, D. W. (2010) Functional imaging of hippocampal place cells at cellular resolution during virtual navigation. *Nature Neuroscience* **13**, 1433 1440.
- Drouin, E., Piloquet, P. and Péréon, Y. (2015) The first illustrations of neurons by Camillo Golgi. *The Lancet Neurology* **14**(6), 567.
- Dudai, Y. (2004) The neurosciences: the danger that we will think that we have understood it all. In *The new brain sciences: perils and prospects*, eds S. Rose and D. Rees, pp. 167 180. Cambridge University Press.
- Ferguson, T. S. (1973) A Bayesian analysis of some nonparametric problems. *The Annals of Statistics*  $\mathbf{1}(2)$ , 209 230.

Ferguson, T. S. (1974) Prior distributions on spaces of probability measures. *The Annals of Statistics* **2**(4), 615 – 629.

- Friedrich, J. and Paninski, L. (2016) Fast active set methods for online spike inference from calcium imaging. In *Advances In Neural Information Processing Systems*, eds D. Lee, M. Sugiyama, U. Luxburg, I. Guyon and R. Garnett, pp. 1984 1992.
- Friedrich, J., Zhou, P. and Paninski, L. (2017) Fast online deconvolution of calcium imaging data. *PLOS Computational Biology* **13**(3), 1 26.
- Frühwirth-Schnatter, S. and Malsiner-Walli, G. (2019) From here to infinity: Sparse finite versus Dirichlet process mixtures in model-based clustering. *Advances in Data Analysis and Classification* **13**, 33 64.
- Frühwirth-Schnatter, S., Malsiner-Walli, G. and Grün, B. (2020) Generalized mixtures of finite mixtures and telescoping sampling.
- Gelman, A., Carlin, J., Stern, H., Dunson, D., Vehtari, A. and Rubin, D. (2013) *Bayesian Data Analysis, Third Edition*. Chapman & Hall/CRC Texts in Statistical Science. Taylor & Francis.
- Grienberger, C. and Konnerth, A. (2012) Imaging calcium in neurons. *Neuron* 73(5), 862 885.
- Ishwaran, H. and James, L. F. (2001) Gibbs sampling methods for stick-breaking priors. *Journal of the American Statistical Association* **96**(453), 161–173.
- Jewell, S. and Witten, D. (2018) Exact spike train inference via L0 optimization. *The Annals of Applied Statistics* **12**(4), 2457 2482.
- Jewell, S. W., Hocking, T. D., Fearnhead, P. and Witten, D. M. (2019) Fast nonconvex deconvolution of calcium imaging data. *Biostatistics* **21**(4), 709–726.
- Kalli, M., Griffin, J. and Walker, S. (2011) Slice sampling mixture models. *Statistics and Computing* **21**, 93 105.
- Malsiner-Walli, G., Frühwirth-Schnatter, S. and Grün, B. (2016) Model-based clustering based on sparse finite Gaussian mixtures. *Statistics and Computing* **26**, 303 324.
- McCullagh, P. and Yang, J. (2008) How many clusters? *Bayesian Analysis* 3(1), 101 120.
- Miller, J. W. and Harrison, M. T. (2018) Mixture models with a prior on the number of components. *Journal of the American Statistical Association* **113**(521), 340 356.
- Mukamel, E. A., Nimmerjahn, A. and Schnitzer, M. J. (2009) Automated analysis of cellular signals from large-scale calcium imaging data. *Neuron* **63**(6), 747 760.
- Nobile, A. (2004) On the posterior distribution of the number of components in a finite mixture. *The Annals of Statistics* **32**(5), 2044 2073.

Nobile, A. and Fearnside, A. (2007) Bayesian finite mixtures with an unknown number of components: the allocation sampler. *Statistics and Computing* **17**, 147 – 162.

- Paninski, L. and Cunningham, J. P. (2018) Neural data science: accelerating the experiment-analysis-theory cycle in large-scale neuroscience. *Current opinion in neurobiology* **50**, 232 241.
- Paninski, L., Pillow, J. and Lewi, J. (2007) Statistical models for neural encoding, decoding, and optimal stimulus design. In *Computational Neuroscience: Theoretical Insights into Brain Function*, eds P. Cisek, T. Drew and J. F. Kalaska, volume 165 of *Progress in Brain Research*, pp. 493–507. Elsevier.
- Parker, D. (2006) Complexities and uncertainties of neuronal network function. *Philosophical transactions of the Royal Society of London. Series B, Biological sciences* **361**(1465), 81 99.
- Parker, D. (2010) Neuronal network analyses: premises, promises and uncertainties. *Philosophical transactions of the Royal Society of London. Series B, Biological sciences* **365**(1551), 2315 2328.
- Pnevmatikakis, E., Merel, J., Pakman, A. and Paninski, L. (2013) Bayesian spike inference from calcium imaging data. In *In Signals, Systems and Computers*, pp. 349 353.
- Pnevmatikakis, E. A., Soudry, D., Gao, Y., Machado, T. A., Merel, J., Pfau, D., Reardon, T., Mu, Y., Lacefield, C., Yang, W., Ahrens, M., Bruno, R., Jessell, T. M., Peterka, D. S., Yuste, R. and Paninski, L. (2016) Simultaneous denoising, deconvolution, and demixing of calcium imaging data. *Neuron* 89(2), 285 299.
- Richardson, S. and Green, P. J. (1997) On Bayesian analysis of mixtures with an unknown number of components (with discussion). *Journal of the Royal Statistical Society: Series B* (*Statistical Methodology*) **59**(4), 731 792.
- Rodríguez, A., Dunson, D. B. and Gelfand, A. E. (2008) The nested dirichlet process. *Journal of the American Statistical Association* **103**(483), 1131 1154.
- Sethuraman, J. (1994) A constructive definition of Dirichlet priors. *Statistica Sinica* **4**(2), 639 650.
- Teh, Y. W., Jordan, M. I., Beal, M. J. and Blei, D. M. (2006) Hierarchical Dirichlet processes. *Journal of the American Statistical Association* **101**(476), 1566 1581.
- Vogelstein, J. T., Packer, A. M., Machado, T. A., Sippy, T., Babadi, B., Yuste, R. and Paninski, L. (2010) Fast nonnegative deconvolution for spike train inference from population calcium imaging. *Journal of Neurophysiology* **104**(6), 3691–3704.

Walker, S. G. (2007) Sampling the Dirichlet mixture model with slices. *Communications in Statistics - Simulation and Computation* **36**(1), 45–54.

Wei, X.-X., Zhou, D., Grosmark, A., Ajabi, Z., Sparks, F., Zhou, P., Brandon, M., Losonczy, A. and Paninski, L. (2019) A zero-inflated gamma model for post-deconvolved calcium imaging traces .