RESTING STATE FUNCTIONAL MRI ANALYSIS BY GRAPHICAL MODEL

by

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ABSTRACT

Functional magnetic resonance imaging (fMRI) measures the change of oxygen consumption level in the blood vessels of the human brain, hence indirectly detecting the neuronal activity. Resting-state fMRI (rs-fMRI) is used to identify the intrinsic functional patterns of the brain when there is no external stimulus. Accurate estimation of intrinsic activity is important for understanding the functional organization and dynamics of the brain, as well as differences in the functional networks of patients with mental disorders.

This dissertation aims to robustly estimate the functional connectivities and networks of the human brain using rs-fMRI data of multiple subjects. We use Markov random field (MRF), an undirected graphical model to represent the statistical dependency among the functional network variables. Graphical models describe multivariate probability distributions that can be factorized and represented by a graph. By defining the nodes and the edges along with their weights according to our assumptions, we build soft constraints into the graph structure as prior information. We explore various approximate optimization methods including variational Bayesian, graph cuts, and Markov chain Monte Carlo sampling (MCMC).

We develop the random field models to solve three related problems. In the first problem, the goal is to detect the pairwise connectivity between gray matter voxels in a rs-fMRI dataset of the single subject. We define a six-dimensional graph to represent our prior information that two voxels are more likely to be connected if their spatial neighbors are connected. The posterior mean of the connectivity variables are estimated by variational inference, also known as mean field theory in statistical physics. The proposed method proves to outperform the standard spatial smoothing and is able to detect finer patterns of brain activity. Our second work aims to identify multiple functional systems. We define a Potts model, a special case of MRF, on the network label variables, and define von Mises-Fisher distribution on the normalized fMRI signal. The inference is significantly more difficult than the binary classification in the previous problem. We use MCMC to draw samples from the posterior distribution of network labels. In the third application, we extend the graphical model to the multiple subject scenario. By building a graph including

the network labels of both a group map and the subject label maps, we define a hierarchical model that has richer structure than the flat single-subject model, and captures the shared patterns as well as the variation among the subjects. All three solutions are data-driven Bayesian methods, which estimate model parameters from the data. The experiments show that by the regularization of MRF, the functional network maps we estimate are more accurate and more consistent across multiple sessions.

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CHAPTER 1

INTRODUCTION

1.1 Problem statement

The human brain is organized into distributed functional modules that work independently for specific cognitive tasks, and also interact with each other. It has been shown that the spontaneous fluctuation of the blood oxygenation level dependent (BOLD) signal, as measured by functional magnetic resonance imaging (fMRI), is a valuable data source for delineating the functional network organization. More recently, the spontaneous activity of human brain has gained more attention because of its potential in helping understand the *baseline* patterns of cognitive activity as well as the cause of some cognitive diseases. Resting-state fMRI (rs-fMRI) is accordingly widely used for exploring such activities.

The analysis of rs-fMRI data is a challenging task, due to the scanner noise, physiological noise, head motion, and subject's random thoughts during data acquisition. Single subject's data are typically unreliable and inconsistent for the statistical inference of the whole population's intrinsic activity patterns. On the other hand, combining data from multiple subjects and jointly estimating the common functional networks is more robust. In group analysis of rs-fMRI data, one typically assumes that all subjects in the group share common functional connectivity patterns, These group networks can be estimated more accurately because the noise introduced in each subject is canceled by averaging. In practice, it is a major challenge to summarize the consistent patterns across subjects, as each subject's network patterns appear similar but have slight variation due to the anatomical and functional difference across subjects.

Recent years have seen substantial interest in estimating functional networks of individual subjects during group analysis. An accurate estimate of an individual's network is an important step towards understanding of brain-behavior relationships on a per-subject basis. The intersubject variation must be accounted for in order to obtain a good estimation of the networks of individual subjects as well as the group. Current methods [157, 39] either do not estimate individual functional network maps, or do not have an explicit statistical

model on the intersubject variations [27, 28]. Among the methods that do estimate subject functional networks, some have one or more of the drawbacks.

First, some methods map a common group functional network by concatenating the BOLD signal from all subjects. By doing this, one implicitly assumes that the voxels across subjects map to the same anatomical structure after coregistration, and share the same functional connectivity patterns. These assumptions are often violated due to the anatomical inhomogeneity between subjects, and also due to the imperfect alignment of the existing coregistration routine. In addition, the simple concatenation does not take into account the possible different variance across subjects. In particular, some participants may experience spontaneous, but active cognition during the scan even in the resting-state. These activities modulate each subject's functional network in a different way and to a different extent, and hence tamper the estimation of the group's functional networks. Such subject-specific confounding factors are less likely to be negligible by simple averaging compared to other sources of noise such as scanner noise, subject motion and coregistration.

Second, group analysis are often conducted in a one way procedure. In some scenarios [143, 38, 72, 70, 135, 110], each subject's functional network is estimated independently, and a group map is simply summarized by averaging the subjects' connectivity maps. The estimates of subject maps by these procedures do not use other subjects' information and are robust to noise. The group summary map extracted from these subject maps is hence suboptimal. In other scenarios [27], a group map is estimated first from the concatenated data, then is back-reconstructed to obtain the subject network maps. More recently, the subject network maps are estimated from the averaged group map using a dual regression approach [52, 11]. Such methods treat voxel intensity from all subjects the same way for group map estimation, ignoring that they may have subject specific variances. Both classes of approach do not iteratively refine the initial group or subject estimates, and the estimation of one subject's connectivity does not benefit from the information obtained from other subjects. Figure 1.1 gives an illustration of the various methods and their order of estimations.

Last, spatial smoothing is often used during preprocessing in order to address the issue of imperfect intersubject alignment. Although spatially blurring the time series increases the signal-to-noise ratio, the choice of the smoothing kernel size has a big impact on the estimated functional maps. Over-smoothing inevitably results in the loss of fine structures of the functional maps. In practice, the random field theory of statistical parametric mapping (SPM) [59] requires a smoothing kernel even larger than the anticipated region of interest.

One needs a model that uses the spatial dependency and the intersubject similarity of the rs-fMRI signals, without losing the finer details of the functional patterns.

A data-driven, unified probabilistic framework will help in solving the above issues. The model should integrate both group's and subjects' connectivity variables into this model. One can make inference from the posterior distribution of the variables in both subject and group levels given the observed BOLD signal.

In this dissertation we present a series of statistical methods for identification of the human brain's functional networks by using rs-fMRI data. All the methods aim to model the spatial dependency within a single subject in a principal way without the naive spatial smoothing, and model the intersubject similarity and variations for a more accurate group and subject network estimation. The main mathematical tools are Markov random fields (MRF) — an undirected graphical model, and Bayesian method. We use various methods including Markov chain Monte Carlo (MCMC) sampling and variational inference for solving the statistical inference problem in high dimensional space.

1.2 Dissertation statement

Here is the statement of this dissertation:

A multilevel Markov Random Field model improves the reliability of the functional network estimation in rs-fMRI group study by taking into account context information as a prior. The data-driven Bayesian model can jointly estimate both population and subjects' connectivity networks, as well as drawing inference on the uncertainty in the estimation, and on the variability across subjects.

The word *Context* has two meanings: 1) The functional patterns of the human brain are spatially coherent. Neighboring voxels have larger probability of being in the same functional network. 2) The network that a voxel belongs to in one subject is dependent on the networks of the same voxels in other subjects. The patterns of functional networks from the rs-fMRI study are to some extent shared by multiple subjects, while the variability across subjects must be taken into account.

By *reliability* we mean the decrease in the variance of the functional networks that we estimate with different subsets of all subjects. The reliable estimates will be closer to the true network in the simulation test, where we know the true answer.

To test our statement, we propose the following contributions:

• Full airwise connectivity with spatial coherence. We propose a method that estimates pairwise functional connectivity in the whole brain of a single subject,

without a priori knowledge of the seed region. The model needs to take into account the spatial context information, and learn the strength of the coherence from the data.

- Identify consistent, spatially coherent multiple functional networks. We propose a data-driven, generative model that can cluster the gray matter voxels of a single subject's brain into disjoint multiple functional networks, while respecting the spatial coherence of the voxels.
- Hierarchical model for group study. We propose a hierarchical model that can estimate functional networks from a group of subjects. The model will estimate an overall group's network map as well as individual subjects network maps at the same time. When clustering the voxels into different networks, spatial neighbors both within and across subjects will be used in a prior distribution of a Bayesian framework. The variability of each subject's connectivity due to noise and artifact will be reduced to the extent that is to be determined automatically from the data.
- Variability of resting-state functional network. Based on the hierarchical MRF model proposed above, we will draw inference on the variance and the confidence intervals of the functional network. we will test the variability of the network by using a subset of the data and perform bootstrap sampling. we also explore and visualize the modes of spatial variability of the functional network patterns.

1.3 Outline and contributions

The remainder of the dissertation is organized in the following chapters.

In Chapter 2, we will give a survey of the existing methods of modeling the brain's functional connectivities using rs-fMRI, as well as the statistical inference methods. we will describe the similarities and differences across these varied statistical approaches, evaluate their advantages and disadvantages, and relate them with our model.

Chapter 3 gives a general introduction of the mathematical tools that will be used in the following chapters. These tools include graph, graphical models and MRF, and various statistical inference methods including sampling, variational Bayesian, graph cuts and other approximation methods.

Chapter 4 will discover the first method that estimates a single subject's functional networks. This chapter is an application of the graphical model and MRF that we introduce in Chapter 3. In this particular case, the MRF is in a high dimensional space to model the prior distribution of the pairwise connectivity variables. Also, we use variational inference

to solve the problem. The inference is implemented on graphical processing unit (GPU) to speed up such a problem with N^2 complexity.

In Chapter 5, we apply the graphical model in different settings. Here the model is defined on the voxels in an original image domain instead of on the higher dimensional domain defined in previous chapter. Together with a mixture model and Bayesian method, we estimate all functional networks of a single subject dataset with spatial regularization.

Chapter 6 extends the concept of the graph to multiple subjects, and estimate the functional patterns with a group of rs-fMRI data. Both group and subject functional networks are estimated with higher accuracy, better consistency than standard methods. This chapter will also cover the validation of the hierarchical model. Besides the estimation accuracy, we focus on the consistency of the method across multiple sessions, and also under data perturbation.

Chapter 7 concludes the dissertation with a general discussion, and some future work that is worthy to explore.

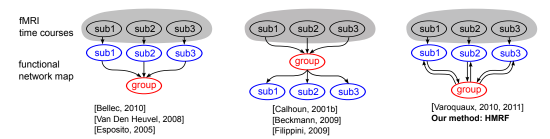


Figure 1.1: The order of estimation of various methods in group analysis.

CHAPTER 2

BACKGROUND AND RELATED WORKS

The human brain is organized into distributed functional modules that work independently but also interact with one another during cognitive activity. There appears two principles of how the brain function is organized: functional integration and functional specialization [59]. Functional specialization assumes the modules that process a particular cognitive activity can be localized to an anatomical region in the brain. In functional specialization, one studies the relationship of specific anatomical regions and cognitive activity. However, the brain regions or functional modules do not work alone. They interact with each other in a complex and dynamic way. Functional integration focuses on such interactions.

This chapter provides background information about the current research exploring brain's functional network using rs-fMRI. I will begin with the relationship between brain activity and fMRI, and give the definitions of functional connectivity and functional network. Then I will survey various classes of methods for estimating functional connectivity and networks.

2.1 Resting-state functional networks

The interactions among various functional modules of the brain can be represented by the connectivity among anatomical regions. Two sets of MRI techniques are widely used for mapping the *in vivo* connectivity of the human brain. Diffusion MRI, or more specifically, diffusion tensor imaging (DTI), as a structural imaging technique, detects the anisotropy of the neural axons in the brain's white matter by measuring the diffusion of water molecules. Such anisotropy is used for identifying the anatomical connectivities between two regions in the brain's white matter [87]. Functional MRI (fMRI) imaging, in contrast, serves to explore the functional links between two regions of interest in brain's gray matter. The functional connections are the actual flow of information through the anatomical links represented by the DTI. Researchers have demonstrated the correlation

between the anatomical and functional connections. Such correlation is an indication that functional connectivity is physically constrained and modulated by the structural connectivity [80]. However, functional and structural connectivity are not exactly equal, suggesting the dynamic characteristics of the functional patterns depend on the brain's state. The work in this dissertation is focused on functional connectivity.

Information is transferred in axons between neurons by the release of neurotransmitter molecules at synapses. The interactions between neurotransmitter and receptors consume energy. Because energy is produced by oxidative metabolism, increased synaptic activity will also increase local demand for delivery of oxygen. This, counter intuitively, increase the local blood flow, and increases the T_2^* image intensity [97, 140]. Even deoxygenated hemoglobin decreases during neural activity, MR signal increases. This is because more oxygen is supplied to the brain region than is consumed.

Haemoglobin have different magnetic property when it is bound to oxygen. This changes the local distortions of a magnetic field, which can be detected by MRI scanner. The relaxation times depends on the level of blood oxygenation, and the MRI signal depends on the relaxation time, and is called blood oxygenation level-dependent (BOLD) signal.

To understand the response of BOLD signal to a general stimulus signal, we need to know its response to a very short impulse signal, namely, a δ function. It is noted that the BOLD signal have about two seconds lags after the onset of stimulus. This initial dip is attributed to the transient increase of deoxygenated hemoglobin. After 4 to 6 seconds, the demand due to increase neuro activity results in an increased inflow of oxygenated blood, and it reaches the highest point. After the neuro activity stops, the BOLD goes below the baseline level. It takes about twenty seconds for the BOLD to go back to the baseline in this poststimulus undershot. The balloon model [25] is used to explain this extended period. The response function to a δ function is called the Hemodynamic response function (HRF). The response of a general boxcar function, would be the convolution of the boxcar function and the HRF.

Because the resolution of fMRI is much higher than the single cell or neuron, the BOLD signal reflects the energy demands of neuropopulation that fire together with a common functional purpose [84, 121]. Besides fMRI, other techniques are also used for mapping the functions of the brain. They differ in terms of both their temporal and spatial resolution. In general, electrophysiological methods, such as electroencephalography (EEG) or the associated magnetic version magnetoencephalography (MEG), record the neural events in real time; hence, these methods have relatively high temporal resolution. On the other

hand, fMRI and positron emission tomography (PET) detect the change of blood flow due to neuronal information processing, and have high spatial resolution (1-5 mm), but lower temporal resolution because of the delayed haemodynamic changes.

fMRI was initially discovered as a tool to map the brain's activity for subjects in specific cognitive experiments. One compares the BOLD signal at a specific region in the brain with the paradigm task signal. Because of the nature of the haemodynamics of the blood, the paradigm task signal is the convolution of the original paradigm function (for example, a boxcar function) with the haemodynamic response function (HRF). Later, researchers found that the BOLD signals can be used not only to detect the functional patterns in stimulus-guided experiments, but also to explore the co-activation of the brain when the subjects are not performing cognitive tasks [17]. In such experiments, the primary goal is to identify the functional connectivity between pairs of regions in the brain. Functional connectivity is defined as the temporal dependence of neuronal activity patterns of spatially remote (or anatomically separated) brain regions [61, 155, 60]. The temporal dependence is typically measured by the linear correlation across all time points. Although the temporal correlation within the BOLD signal of a single region or voxel violates the independence assumption of the time point samples, this temporal correlation is often safely ignored. Alternatively, one can also identify functional connectivity by transforming the signals into the frequency domain and using the coherence of two signals at a certain frequency band. The coherence as similarity measurement is equivalent to band pass filtering the original BOLD signal and computing the linear correlation [36, 35].

The pairwise correlation or coherence only measure the functional connectivity between regions as a local measurement. As the whole brain is organized as a complex system with many such pairwise interactions, it is of interest to find out those regions with similar patterns of neuronal activity. A functional network, or functional system (hereafter used interchangeably), is a collection of separate anatomical regions that have similar patterns of activity measured by BOLD signal. The regions within a functional system may have direct or indirect information flow among them. Together the system serves one or more cognitive tasks. An anatomical region may participate in different functional systems during different cognitive tasks.

The correlated fluctuation of multiple brain regions not only occurs in stimulus-evoked experiments, but also in experiments where the participants rest passively without any cognitive activity. Therefore, the resting-state fMRI (rs-fMRI) becomes a powerful tool for probing such intrinsic activity in a resting brain [124, 54]. The original name of the rs-fMRI

is not accurate, since the brain is not truly in a resting state even without any cognitive activity. The resting brain consumes about 20 percent of the energy of the whole body, but it occupies about only 5 percent of the body mass [56]. The study of the functional organization of the resting brain provides new insights into how functional connectivity relates to cognitive psychology and neurodegenerative diseases. Because the pathologic conditions appear to be reflected by the interactions within or between the functional systems, the rs-fMRI study also holds valuable diagnostic and prognostic information towards various neurological or psychiatric diseases including Alzheimer's diseases, depression, and schizophrenia [54, 70, 71], etc. The reason for this spontaneous activity is largely unknown, although some researchers reasonably postulate that it is a predictive intrinsic response to the unknown events in the outside environment [40].

2.2 Preprocessing fMRI data

Due to the noise and artifacts in fMRI data, multiple preprocessing steps are typically taken before the real analysis starts. The preprocessing steps include motion correction, slice timing correction, spatial and temporal filtering, registration between structural images and functional images, registration to the standard template, and removing physiological noise, etc.

Because of a subject's head movements during the scan, the volumes at various time points are usually not perfectly aligned. A rigid body registration is usually done between the volumes at different time points, with either the first volume or the middle volume as a reference. This is called motion correction. After correction, the same voxel coordinate is assumed to map to the same brain structure across all time points, although the correction is often not perfect. The motion correction parameters of each volume are often used as independent variables for a regression in order to remove the motion effect. The first few time points are usually discarded in case the scanner is not in a stationary state.

Slice timing correction is needed because each slice of fMRI volume is not scanned at the same time. Figure 2.3 shows the shifted time of scans for different slices and the method of interpolation. The shifted time will give suboptimal experiment results during the analysis, especially for the event-based experiments. One typically uses an interpolation step to obtain a volume in which all slices are at the same time point. Care must be taken for the ordering of the slices as the ordering is scanner dependent.

Because of the physiological process of the BOLD signal, most of the interesting information is concentrated at the low frequency range of the signal (0.01 - 0.1 Hz). The temporal band-pass filter is usually applied to remove the very low frequency (below 0.01).

Hz) and higher frequency (above 0.1 Hz). Also, due to the spatial process of fMRI data acquisition, neighboring voxels typically have similar signal patterns. The statistical parametric mapping method [59] makes use of this spatial dependency by applying a spatial Gaussian filter, and further enforces the smoothness of the signal on the spatial domain. This filtering increases the SNR, but inevitably discards finer patterns with size smaller than the smoothing kernel. Chapter 4 and Chapter 5 will discuss more principal methods that model this spatial dependency with no spatial smoothing, or very small, conservative smoothing.

In order to report the experiment results in a standard way such that others can understand, the structural images and functional images are both registered to a standard template, such as the MNI 152 template ¹. The functional images are first registered to structural images of the same subject by a rigid body transformation. The structural images are then transformed to the standard template by affine transformation. Because of the plasticity of the brain anatomy, the registration of the subject's structural image may need nonlinear transformation to register to the template [86]. After the structural images are registered to the template, the functional images will also be brought to the template space by the same transformation. The last step is the nuisance parameter regression. The nuisance parameters include the six motion correction parameters that are estimated in the previous motion correction step, and also the mean signal of the white matter and cerebrospinal fluid (CSF). The motion correction parameters are taken into account because even though the fMRI data are motion corrected at the first step of preprocessing, the motion may still have an impact on the estimation of the general linear model (GLM) of task-based fMRI, or the correlation for the rs-fMRI. The mean of white matter and CSF is regressed out as these average signals are assumed to consist of physiological noise. Since there is no information of the physiological signal such as heart beating and breathing, the mean of white matter and CSF is used as surrogates for such confounding signals.

With completion of the above preprocessing steps, the data are ready for analysis. The above preprocessing steps may vary depending on the specific experiments. For example, the slice timing shifting is not a severe issue for rs-fMRI analysis and can be skipped. Besides, the preprocessing steps may not have consistent impact on the fMRI processing pipeline. For example, Zhang [160] pointed out that slice timing correction and global intensity normalization have little consistent impact, but spatial smoothing, temporal detrending,

¹http://www.bic.mni.mcgill.ca/ServicesAtlases/ICBM152NLin2009

high-pass filtering and motion correction significantly improve the pipeline performance across all subjects.

2.3 Related methods

The temporal dependence of the spatial regions can be represented in multiple ways. Depending on whether one is interested in a specific region or a full brain's functional patterns, one can select the seed-based method or the full brain method. For the class of full brain methods, there are methods that define the region of interest (ROIs) and build a functional network by estimating the edges of the graph with nodes defined by the ROIs. Also, the parcellation defines an image segmentation problem where the regions with higher functional connectivities are grouped into a single cluster. In this section, I will give a short survey of the various methods, show their advantages and disadvantages, and the similarities and differences with our methods.

2.3.1 Seed-based methods

Depending on the specific experiment goal, functional networks may be represented in various ways. A straightforward yet statistically powerful method is to compute the linear correlation between $a\ priori$ given seed regions and all other regions in the brain [17, 21, 22]. The correlation values are typically Fisher transformed in order to meet the normal distribution assumption in the following hypothesis test. Those transformed correlation values with p value less than a certain threshold are regarded as the existence of the functional connectivity. All voxels or regions that are functionally connected to the seed regions belong to the same functional system. The seed-based methods are useful when a user asks a straightforward question and knows what functional system is of interest. The result is easy to interpret compared to other more complex methods. The advantage of seed-based methods is their simplicity and relatively ease of extention to multiple subjects. A user simply computes the average correlations across subjects for a given pair of regions. When users are interested in the connectivity to multiple regions, they can define more than one seed.

However, this method has a limitation: the user has to know the location of the seed in advance. The seed as *a priori* information is an advantage when it accurately represents the functional patterns of interests. However, a functional system cannot be identified if the seed region falls out of the system. Despite this limitation, researchers frequently use this method, sometimes with better visualization by dynamically moving the seed and showing the real-time functional system associated with the current seed region [157].

2.3.2 ICA and other decomposition methods

Because functional networks are in a large scale over the whole brain, and in a distributed manner, a multivariate analysis is a better way to explore the full brain's functional organization. A large class of multivariate methods use the signal-decomposition concept from the signal processing community and decompose the BOLD signal at each region or voxel into various independent components, each of which is part of one functional system. The signal and the weight coefficients of these independent signals represent how much of the current voxel belongs to certain functional network component. One widely accepted method in this class is the independent component analysis (ICA). ICA was originally introduced in the signal processing field to separate various sources of signals from samples of mixed signals and later was applied to rc-fMRI data [115, 9, 39]. The central limit theorem states that the sum of two independent signals is more like Gaussian, than any of the original signals. Therefore, maximizing the non-Gaussianity gives us the original independent components. Besides the maximization of non-Gaussianity, the independent components can also be estimated by minimization of mutual information, or by maximum likelihood [85].

There are two varieties of ICA method when applying to rs-fMRI dataset. Spatial ICA assumes all the voxel intensities at a certain time point as one mixed signal sample. Therefore, the mixed signals are independent across all spatial voxels. Alternatively, temporal ICA treats each BOLD time series at a voxel as a mixed signal; hence the source signals are independent across the time point [28]. Notice the functional map we are interested in is the rows of source signal matrix S for spatial ICA, and is the weight coefficients matrix \tilde{A} for temporal ICA (see Figure 2.4 for an illustration). Because of the large number of voxels compared to the number of time points, the spatial ICA is typically used for rs-fMRI analysis. Compared to seed-based methods, ICA is purely data-driven, and can identify networks over the whole brain, including the already well-known cognitive processing systems such as motor [17], visual [39], attention [55] executive control and salience network [136, 135].

However, because ICA needs to estimate both the independent components and mixing coefficients, it is a significantly more difficult task. Before applying ICA, the data usually needs a whitening step with principal component analysis, and is thus rotated such that it has unit variance in terms of the covariance matrix, which greatly simplifies the ICA problem. The estimated independent components are usually z transformed and thresholded for visualization purpose. Because the resting-state brain functional patterns are unpredictable, the output independent components need to be visually inspected in order to

identify physiologically meaningful components. The iterative optimization also introduces variability between multiple runs of ICA with different initial states. In addition, the dimensionality in the preprocessing step of principal component analysis (PCA) is typically chosen arbitrarily, adding more variation in the results. A recent test on the reliability of ICA [162] shows how the choice of the dimension and the number of components have an impact on the consistency of the results.

2.3.3 Segmentation-based methods

The functional networks estimated from ICA are represented by continuous numbers and have to be normalized to the z score and thresholded to give a binary map. The thresholding adds ambiguity to the consistency of the results. An alternative class of methods formulate the problem of identifying functional patterns as an image segmentation problem. The image segmentation problem can be also viewed as a data clustering problem in general data mining. However, since the data points are indeed voxels in the fMRI images, the spatial context information is also useful for the clustering. Therefore, we name this class of methods as image segmentation, to indicate the possible usage of spatial information. Once the fMRI images are segmented into various disjoint sets of regions, the voxels in the same regions have a higher correlation, and hence are believed to be functionally connected. Compared to ICA, the segmentation problem typically has a binary map for each functional network, i.e., the clusters, although some soft segmentation methods exist. The methods used for segmentation include hierarchical clustering [134, 32, 34, 12], partitioning clustering [12] and spectral clustering [143, 38]. The early works of fMRI image segmentation [34] are limited to a few slices due to the computation cost. Even within a few slices, the segmentation method is still able to identify the major functional components, and they shows that the results are robust to the confounds due to the subject motion and physiological noise. One interesting property of the hierarchical clustering technique is the potential of detecting the hierarchy of the brain's functional organization. Since the brain's functional patterns are widely believed to be also hierarchical [157], a computational method will be very useful if it can naturally identify the subclusters within a certain cluster. Notice that the hierarchical clustering method is different from the hierarchical MRF method in the following chapter. The former builds a hierarchy on the clusters, whereas our method builds the hierarchy on the group and subjects' functional network maps.

Among the segmentation methods, Mezer et al. [106] use the windowed Fourier transform and the spectrum below 0.2 Hz on the frequency domain, as well as the BOLD time series in the original image domain as features for the K-Means clustering. To test if the clusters

estimated by K-Means are significantly different, they used a repeated measure ANOVA test between each pair of clusters. The experiments in Mezer's work showed strong links between the functional patterns and the tissue architecture, suggesting the possibility that the rs-fMRI signal may include the contributions from physiological and artifact noise factors that cannot be easily separated. In Figure 2.1, we did a simple experiment for using spectral coherence as the similarity measure and use spectral methods for segmentation. From the figure, we can identify the visual area (the green) and the functional regions at temporal lobes (blue). Since we define each voxel as a node on the graph, the total number of data points is big, and the computation of the similarity matrix takes long time. The example here uses only $10\ z$ slices for illustration.

To decide the number of clusters in the functional network map is a difficult problem of model selection. Most of the methods use seven clusters, since the clustering into seven networks has a good match with the existing cognitive network configuration [157]. Other choices are possible if the goal is to identify the functional patterns at a finer scale. Another reason of choosing large number of clusters is to have a finer parcellation in the local scale. In a local parcellation, only the locally coherent functional regions are grouped into the same cluster. The remote functional regions may not be grouped into the same cluster eventhough they belong to the same functional network [106]. Such a local grouping approach is similar to the super-pixel approach widely used in the computer vision community [96, 1], where researchers group similar pixels into small regions as a preprocessing step for higher-level vision analysis. The grouping procedure is conservative in that only the spatially neighboring pixels are grouped. The super-pixels are used for the higher level image understanding problem in order to save computation time. One example of such functional network parcellation is Thiron [142], where the primary goal is to find spatially coherent clusters that are connected. Therefore, the spatially remote regions cannot be classified into the same cluster. The advantage of such parcellations is the small regions modeled as groups of voxels better represent true regions of activity in task-based fMRI data, and are less sensitive to the misregistration across subjects. The parcellation also reduces the multiple comparison problem that is typically found in the statistical parametric mapping method. Thiron et al. uses a spectral clustering approach with multidimensional scaling (MDS) representation of the dataset, and the C-Means method for clustering. The method is able to estimate a spatial coherent group clustering from the subject parcellations. Another example of finer parcellation is found in Craddock et al. [38]. The goal of their work is to evaluate the suitability of a few parcellation schemes on a group of subjects'

rs-fMRI data. The authors use the normalized cut method [138] to segment the brain into a large number of spatial and functional homogeneous regions. A graph is defined by adding edges only between spatial neighbors, with the weights on the edges defined by the linear correlation between BOLD signals. This definition of edges and weights means the functionally homogeneous voxels may be in different clusters if they are not spatially adjacent. Since the number of clusters in their experiments is relatively large (50 - 400), the estimated parcellation map will not be easily interpreted as functional networks, but will be used for further analysis. For group parcellation, Craddock et al. use second-level group clustering similar to the work of Van den Heuvel [143]. The estimated adjacency matrix estimated from subject parcellation was averaged across subjects, and the average matrix was used for a second-level normalized cut algorithm.

The segmentation of functional network is not necessarily limited on rs-fMRI data. It can also be used for the task-based fMRI. The work of Michel et al [107] is one example in this range. Michel et al. use a generalized hierarchical clustering based on the Ward algorithm to build a tree of hierarchical clusters. The leaves of the tree are individual voxels, and the root includes all voxels. The authors compute the optimal pruning of the tree by fitting the average signals of the leaves at each pruning depth to a general linear model. The clusters of optimal trees are able to predict the behavior variables. Varoquaux et al. from the same research group use the methods of Michel et al. [107] for solving an active region detection problem of task-based fMRI data [145]. Whereas the goal of Michel et al. is to identify the functional network patterns using the supervised clustering method with clinical variables, the goal of Varoquaux et al. [145] is to detect active regions when the observation is small and the number of predictors is large. By grouping spatially adjacent voxels with similar BOLD signals into clusters and using the clusters as the independent variables, varoquaux et al. solve the problem of large number of predictors, and are able to identify the functional patterns when there are more voxels than the number of observations.

Some of the methods we proposed in Chapters 5 and 6 belong to the class of segmentation. However, we defined the additional structures based on the spatial context information, and the intersubject coherence assumption, so the model is more complex than the above works, and therefore requires more advanced statistical inference methods.

2.3.4 Methods based on graph theory

Functional networks can also be represented by graph [23, 2, 139]. Our improved understanding of the complex systems and the increasing availability of large datasets of systems have led to the insight that these complex systems share similar behaviors that

can be represented by the same parameters [23]. To build an abstract graph from the fMRI data, one typically chooses some ROIs as the node of the graph, and estimates the existence or lack of edges between the nodes. The signal on each node is computed by averaging the BOLD signals of all the voxels within a certain radius of the ROI, usually a few millimeters (see Figure 2.2 for an illustration). The edge is estimated by computing the linear correlation or the frequency coherence between two ROIs. Once the functional network is represented by a graph, a rich class of methods in graph theory can be used to explore the global property of the graph. For example, the high degree of a node indicates its role as a hub in the network, and the distribution of the degrees is used to explore the vulnerability of the graph when some hubs are removed from the network. Various measures of the network topology are estimated from the graph to explore the similarities and differences between the functional network of the brain and other complex networks such as social networks and wireless networks. The graph property can also be used to explore how the functional system is configured to allow maximum flow of information with minimal wiring. For example, the *small-worldness* attribute is shared by many real-world networks including the functional networks of the human brain, and it is hypothesized that the small-worldness is chosen by evolution for high efficiency of information transfer between nodes at low connection cost [2, 23, 24].

The network estimation results depend on how we define the nodes of the graph. Nodes should be defined as brain regions with homogeneous functional and anatomical patterns. The comparison of functional and structural networks is meaningful only when they share the same definition of graph nodes [130]. Due to the limits on the available methods and the computation resources of estimating networks with continuous edge weights, typically one constructs a binary network by thresholding the similarity matrix. How to choose the optimal thresholding parameter is an open question. More importantly, there is no principal method for combing the networks from a group of subjects.

2.3.5 Group analysis

The analysis fMRI data is challenging, due to the scanner noise, physiological noise such as blood vessel and heart beat, and subject's head motion. Sometimes subjects have random thoughts even when specifically instructed not to think of anything during the data acquisition. The functional connectivity and network estimation is still not accurate and consistent even when various preprocessing techniques are used. On the other hand, because rs-fMRI experiments have much lower requirements for the patient subjects or control subjects compared to paradigm design experiments, it is possible to collect the

rs-fMRI data from a big group of subjects. The functional networks detected by either model-based methods such as seed-based correlation, or data-driven methods such as ICA, are highly reproducible across participants and scans. The more accurate and consistent functional networks can be estimated by using the statistical power of the group analysis. With the initiative of 1000 Functional Connectomes Project, a large cohort of data are available for researchers to explore the relationship of functional networks to the subjects' age, gender, etc.

Compared with single-subject analysis, the methods for multiple-subject functional network analysis are not yet established. The seed-based methods typically compute the correlation or regression coefficients between regions independently for each subject. The correlations or regression coefficients are treated as a statistic of each subject and go into a second-level analysis. The second level can be either fixed-effect analysis or a random-effect analysis with a standard hypothesis testing. The population-based z-score maps have to be corrected for multiple comparisons [56].

Group ICA is used as an extension of single-subject ICA in order to seek a set of independent components shared across multiple subjects [28]. In a typical group ICA study, all subjects are registered to a common atlas and are assumed to share a common spatial component map but have distinct time courses. The BOLD signals from all subjects are concatenated temporally, followed by a single-subject ICA. The subject component maps are then obtained by a back-reconstruction procedure.

Alternatively, a single-subject ICA is applied on each subject first, and a group summary is estimated from the estimates of subject functional patterns. Some methods that we have introduced such as the segmentation-based approach [12, 38] are in this range. If the ICA is used for estimating the spatial patterns at the subject level, it is not straightforward to conglomerate the spatial maps of multiple subjects and derive an optimal set of group maps. Such difficulty of matching the network label maps is because the component maps of subjects are not perfectly aligned with each other, and it is difficult to find the correspondence between subject A's component i and subject B's component j. To solve this problem, Esposito et al. [49] propose a self-organizing method for the group map estimation. A series of spatial component maps are estimated by the regular ICA method for each subject, and these maps are all put in a single pool. The similarity between each pair of component maps in this pool is defined and calculated. Then, Esposito et al. did a clustering on the pool of component maps. The cluster center of the maps is believed to be the group component map. Neither of the above approaches iteratively refines group (or

subject) maps once the subject (or group) maps are estimated.

Varoquaux et al. [146] give an extension of ICA method for group analysis. The authors define a generative model, in which the subject component maps are assumed to be generated from a group component map. In particular, the subject map is a weighted group map with an additive noise term, and the observed BOLD signal of each subject is a linear combination of the subject's components with additive noise. The estimation takes two steps. At the subject level, each subject's data are decomposed by principal component analysis (PCA) and an estimate of the subject component map is obtained. At group level, the authors use canonical correlation analysis (CCA) to find a common subspace among the estimated subject component maps. Such an approach is in the class of bottom-up methods that we introduced in Figure 1.1, since the group component map estimate is not used to iteratively refine the subject estimation. Besides, there seems to be a discrepancy between the group map definition and estimation. That is, the author [146] did not provide any justification for CCA giving an optimal estimate of the generate model defined on the group level.

Another interesting study by Varoquaux et al. [144] uses dictionary learning for the group analysis. Dictionary learning is in the class of linear signal decomposition, with ICA as a special case. In this work, the authors again defined a generative model, where the observed subject data matrix is the product of the loading matrix (i.e., the weighting coefficients) and a signal matrix (also called the code in the dictionary). The loading matrix is the subject-specific spatial map, and is generated from a population spatial map with an additive noise term. The objective function can be written as a nonhierarchical form, and is a function of the subject component map and group component map. The optimization is done by a coordinate descent approach. Overall, the model is similar to what we will propose in Chapter 6 in that a hierarchical map is defined and estimated. In addition, it is different from previous work [146] as the group and subject maps are estimated iteratively in the coordinate descent framework. So, the Varoquaux et al. method is indeed a counterpart to our hierarchical Markov random field model, but in the linear signal decomposition fields.

Despite the promising future of using large populations for studying the functional network and its relationship to the underlining structural patten, the group study also poses new challenges when putting all datasets together. The folding patterns of the cortical surface, and the structure and size of brain regions, are all different across individual subjects. Because of such structural variation, the intersubject registration is often not accurate enough to map the same anatomical structures to the same coordinates in standard

space. Such inaccurate registration may be due to the imperfect coregistration algorithm itself, or due to the difference of individual subjects' cortical structures. Even if the anatomical structures are aligned among multiple subjects, the functional patterns of each subject are not the same. Although numerous studies have shown the relationship of the functional connectivity is related to the structural connectivity [80, 78, 81], it is less well known how and to what extent this relationship is reflected by the variation of functional connectivity.

2.3.6 MRF and spatial constraints

Because of the univariate characteristic of the statistic parametric mapping (SPM) [59] method, the spatial context information is not taken into account in the model. That is, if one arbitrarily permutes the location of the voxels in the brain, the resultant activation map in a task-based experiment does not change, i.e., the original activated voxels are still activated. To take the spatial information into account, one typically forces the spatial homogeneity by applying a Gaussian filter on the spatial domain as a preprocessing step. The filter increases the signal-to-noise (SNR) ratio but inevitably has a blurring effect that sometimes results in the loss of finer structures of the functional patterns.

Markov random field (MRF), as a principal method of modeling spatial context, has been brought into image processing fields by Geman [65]. The first application of MRF to fMRI analysis was by Descombes et al. [43]. Although this is an early work of MRF's application, I give some details of their work because of its novelty. In their work, the authors propose to use MRF for two purposes. First, they used a spatiotemporal MRF to restore the noisy BOLD signal instead of a general temporal filter and spatial Gaussian filter. Second, they compared the restored BOLD signal with the stimulus haemodynamic response function for detecting the activation map, which is modeled by a spatial MRF. For the first goal of BOLD signal restoration, the authors defined a graph, whose nodes or voxels have 12 neighbors: 8 as spatial neighbors, and 4 as temporal neighbors. A hidden continuous variable was defined at each node of each time point to represent the true BOLD signal's intensity. To preserve the discontinuity, another line process MRF was defined on the dual lattice to model the existence or lack of edge between spatial neighbors. The statistical inference of this spatiotemporal MRF is solved by simulated annealing (SA). For activity detection, a second MRF was defined on the activation variables, and the MRF included only spatial neighbors. Although the work was applied on the fMRI data in paradigm design, the concept of spatial regularization and Bayesian method is also applicable to the rs-fMRI data.

Another early work of Hartvig et al. [77] aims at using the spatial context information for activation detection of task-based fMRI experiments. The authors' inductive bias is that the activation region should not be too big or to small. According to this assumption, they define a hidden activation variable taking value in $\{-1,0,1\}$ to represent negative-active, no-activation, and activation state. In addition, the estimated activation coefficients from the standard SPM are assumed to be Gaussian given the hidden variable is no-activation, and Gamma distribution given the hidden variable is activated. The prior distribution of a small set of neighboring voxels' hidden variables is defined to reflect the inductive bias, and the goal is to estimate the posterior of the hidden activation variable given the observed activation coefficients from SPM. The Hartvig et al. model is not MRF strictly speaking, although the model represents the authors' assumption of the spatial context.

A recent work of application of MRF to task-based fMRI data is by Ou et al. [118]. In their work, the authors define a MRF on the activation variables, and the MRF prior favors spatially coherent activated regions. The conditional probability of the observed data is defined by a general linear model (GLM) with the hidden activation variables as parameters. One important characteristic of the work of Ou is the parameter estimation, including both the GLM's parameter and the MRF's parameter. The authors define the MRF prior such that the samples drawn from the prior agree with the spatial properties of the true activation map in fMRI experiments. Therefore, they use the frequency counts to set the pairwise potentials with an additional parameter to control the overall sharpness of the joint component of the prior. This method is similar to the work of Boykov et al. [20] and Rother et al. [129], where a parameter is estimated from the data by counting over the whole image. The additional parameter is also estimated from the data in simulated experiments. The method of Ou is for single subjects.

Penny et al. [120] define a MRF on the regression coefficients of the general linear model on a task-based experiment. Because the regression coefficients are continuous, the authors model the prior distribution by the multivariate Gaussian distribution, with the precision matrix represented by a Laplacian operator. This Laplacian operator penalizes differences between the coefficients of neighboring voxels. In the generative model, the sample of this prior distribution can be generated by first draw independent samples from Gaussian, and then applying the Laplacian operator mapping.

Overall, the methods of functional network estimations have been ranged from the basic seed-based approach to more advanced multivariate, full brain analysis, and from singlesubject analysis, to a pooled summary of multiple subject analysis, to the more complex joint analysis of both population and individuals. In the following chapters, we will introduce our contribution to the methods by combining MRF and a hierarchical model. It should be noted that the more complex method will achieve better results only when the model fits the data well, and the statistical inference will successfully find the optimum or reasonably approximate the optimum solution.

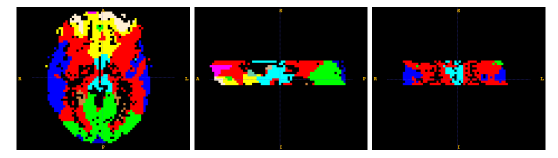


Figure 2.1: Segmentation map of a rs-fMRI volume. The spectral coherence between 0.01 to 0.1 Hz is used for similarity between pairs of voxels. A spectral clustering method [147] is used for dimension reduction followed by a K-Means clustering. We choose 12 clusters and 10 slices on z direction to save computation time.

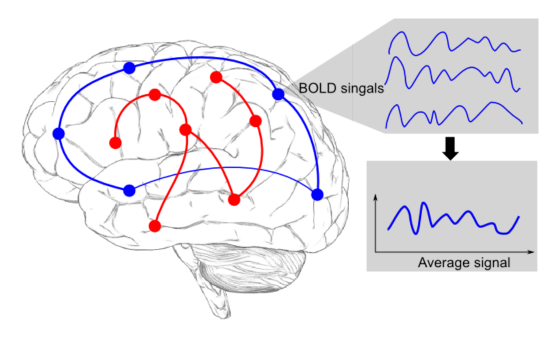


Figure 2.2: Using a graph to represent functional networks. A series of ROIs are chosen based on what questions asked. The signal at each ROI is computed by averaging the BOLD signals of all voxels within the sphere. The edge of the graph is estimated using the similarity of the signals between the ROIs.

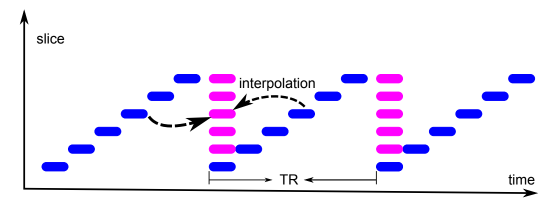


Figure 2.3: Slice timing correction. The data at temporally adjacent slices are resampled and interpolated to obtain a data point at the same time with the reference slices.

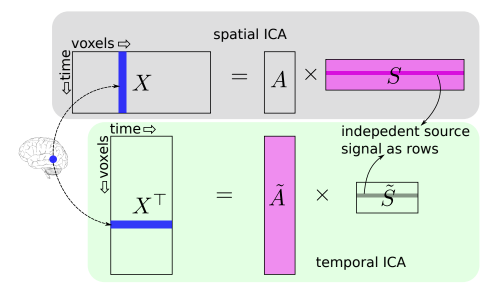


Figure 2.4: Spatial ICA versus temporal ICA for BOLD series of length T at N voxels. In spatial ICA, all the voxel intensity at a single time point is assumed to be a mixed signal of P independent signals, and there are T such mixed signals. The decomposition is in the form of $X = A \cdot S$, where A is the weight coefficient and each row of S is the independent source signal. We are interested in the S since each row is regarded as a functional component. In temporal ICA, the BOLD time series of each voxel is the mixed signals, and there are N such mixed signals. The decomposition is $\tilde{X} = \tilde{A} \cdot \tilde{S}$, with \tilde{A} the weights, and rows in \tilde{S} the independent signal. Here we are interested in the columns of \tilde{A} as they are regarded as representations of the functional networks.

CHAPTER 3

MATHEMATICAL TOOLS

In this chapter I will discuss some general graphical model concepts and inference methods. These mathematical tools will be used in the subsequent chapters, with various changes depending on the specific applications.

3.1 Graphical model

The main methodology we used in this work is statistical, and more precisely, the Bayesian method. Whether we define random variables on voxels of the fMRI image, on ROIs, or on pairwise connectivities between voxels or regions, the problems involve multivariate probability distribution. The variables in this collection interact in a complex way. Graphical models are tools for representing the dependency among the multivariate randomn variables, and the conditional independence among them. We begin the introduction of graphical models from the concept of the graph. A graph \mathcal{G} is defined by a collection of nodes \mathcal{V} and a collection of edges $\mathcal{E} \in \mathcal{V} \times \mathcal{V}$. For node $i, j \in \mathcal{V}$, if $(i, j) \in \mathcal{E}$, there is an edge between i and j. Otherwise, there is no edge between them. A graph can be either directed or undirected, depending on the statistical relationship of the random variables under consideration. The neighbors of a node i are the set of all nodes in \mathcal{G} having an edge to node i,

$$\mathcal{N}(i) = \{ j \in \mathcal{V} : (i, j) \in \mathcal{E} \}. \tag{3.1}$$

A directed graph is often used for modeling the causal relationship of variables. In this dissertation, we focus on the connectivity between pairs of variables without inference about causality, so we use undirected graphs to model the soft constraints between the variables that we are interested in. However, since we use a generative model, where the observed data are regarded to have been generated from hidden, unknown variables, the links between the hidden variables or the parameters and the observed data are directed. We use *chain graph* that includes both directed graphs and undirected graphs as special cases [94].

In statistics, the basic probability rules apply to either continuous or discrete variables, regardless how many dimensions the random variables have. Although the probabilistic inference and the learning can be addressed by applying the sum and the product rule of probability, it is advantageous to use a diagram for representing the probabilistic distributions. A probabilistic graphical model is a collection of random variables that factorize according to the structure of a graph. Graphical model is an intuitive way to visualize the structure of the probabilistic distribution. Because of the correspondence between the distribution and the graph, the conditional independence properties can be inferred by inspection of the graph [16].

To create a graph to represent an existing multivariate probabilistic distribution, we start by defining a graph and adding a node for each random variable in the distribution. If there is a conditional dependency between two variables, we add a link between the nodes associated with them. With this setting, the statistical dependency can be visually read from the graph. Furthermore, the conditional independence can also be read out from the graph.

3.2 Markov random field

A major class of undirected graphical model is the Markov random field (MRF). MRF is used extensively in this dissertation for modeling spatial constraints, and the constraints among the functional networks between subjects. Before introducing MRF, it is helpful to introduce a simplified one-dimensional version of MRF: Markov chains.

Definition 1. A Markov chain is a sequence of random variables x_1, x_2, x_3, \ldots with the Markov property that given the present state, the future and past states are conditionally independent.

$$P(x_{n+1}|x_1, x_2, \dots, x_n) = P(x_{n+1}|x_n)$$
(3.2)

The joint probability of the sequence X is given by

$$P(X) = P(x_1) \prod_{n=2}^{N} P(x_n | x_{n-1})$$
(3.3)

The joint distribution of a Markov chain can be represented by a linear directed graph in Figure 3.1. In this graphical model, the nodes are arranged in a one-dimensional space. The Markov property is equivalent to the conditional independence property, which states that nodes x_i and x_j are conditionally independent given other variables if there is no direct link between them. The directed graph can be used to represent physical processes such as

time series, where the dependency only happens on one direction as current events should not depend on future events. In other situations such as spatial statistics, an undirected linear graph may better represents bidirectional dependency.

When the nodes and their associated variables are defined in a multiple dimensional space, the bidirectional dependency of the variables defined on the undirected graph becomes a MRF. More specifically,

Definition 2. X is called a random field if $X = \{X_1, \ldots, X_N\}$ is a collection of random variables defined on a undirected graph $\mathcal{G} = (\mathcal{V}, \mathcal{E})$, where for each $s \in \mathcal{V}$, x_s takes a discrete value in $\mathcal{L} = \{1, \ldots, L\}$. A set of values of $X = \{x_1, \ldots, x_N\}$ is called a configuration of the field.

The set of edges in the undirected graph \mathcal{G} again represents the dependency between the variables associated with the nodes, without directional information on such dependency. The neighbor system $\mathcal{N}(s)$ is the set of nodes that are connected to node s by an edge. With the above definition of nodes and the neighbor system \mathcal{N} , the graph also gives the independence information between variables. The MRF is defined as:

Definition 3. X is said to be a Markov random field on the graph G with respect to a neighborhood system \mathcal{N} if for all $s \in \mathcal{V}$,

$$P(x_s|x_{-s}) = P(x_s|x_{\mathcal{N}(s)}). \tag{3.4}$$

The Markov property has three equivalent statements [132]. First, the node x_i and x_j are conditionally independent given all other variables if there is no edge between x_i and x_j . This is called the pairwise property. Second, given x_i 's neighbors $\mathcal{N}(s)$, x_i is independent of the remaining variables. This is the local property of MRF. Third, the set of nodes x_A and x_B are conditionally independent given set x_C , if C separates A and B. This is the global property. Usually one or the other properties are useful depending on the specific applications. Because of the lack of directions on the graph's edges, MRF is indeed a multidimensional Markov chain with isotropic statistical dependency between each node and its neighbors. Figure 3.2 gives an illustration of a MRF defined on a regular lattice, and a MRF defined on a general graph.

MRF is defined via the conditional independency property, which is a local property with regard to only a node and its neighbors on the graph. During the statistical inference of the marginal probability of certain variable x_s or a subset of variables X_A , where $A \in \mathcal{V}$, a global property will help infer the probability of X since we are interested in the joint

distribution of the variables. The Hammersley-Clifford theorem [31] builds the relationship between the local property $P(x_s|x_{\mathcal{N}(s)})$ and the global property P(X). Before introducing the theorem, we give the definition of the clique and Gibbs distribution (or Gibbs random field). A clique \mathcal{C} is a complete subgraph of \mathcal{G} , such that within the clique, each node in \mathcal{C} is linked to all other nodes. A maximal clique is a clique to which one cannot add a new node and still keep the subset a clique [92]. The clique is useful to rewrite the joint distribution P(X) in a factorized form. More formally, a set of random variables X is said to be a Gibbs random field (or is in Gibbs distribution) on the graph \mathcal{G} if and only if its probabilistic distribution takes the form of

$$P(X) = \frac{1}{Z} \exp\{-U(X)\}.$$

Here Z is a normalized constant to guarantee the function integrate to 1 and be a probabilistic density function. The exponential $U(Y) = \sum_{c \in \mathcal{C}} V_c(Y)$ is called the energy function. Each clique potential function V_c depends only on the variables in the corresponding clique c. The Hammersley-Clifford theorem states that Y is a MRF if and only if it distributes as a Gibbs distribution.

Unlike the joint probabilistic distributions represented by a directed graph, the clique potential functions in MRF do not have any probabilistic interpretation. One can convert a directed graph into an undirected graph and derive the clique potential from this conversion. A more direct way, however, is to define the clique potential function to reflect our constraints on the relationships between the variables. When the variable $x_s, \forall s \in \mathcal{V}$ takes values from $\mathcal{L} = \{0, 1\}$, and only pairwise neighbors are defined on a regular lattice, we obtain the *Ising* model [119]:

$$P(X) = \frac{1}{Z} \exp\{-U(X)\}, \qquad U(X) = \beta \sum_{(r,s)\in\mathcal{V}} \psi(x_r, x_s)$$
 (3.5)

$$\psi(x_r, x_s) = \begin{cases} 1 & x_r \neq x_s \\ 0 & x_r = x_s. \end{cases}$$

$$(3.6)$$

Because a realization of X with the same states between neighboring nodes has a lower energy according to the definition, such realization (also called configuration) has a higher probability and is therefore preferred. As the simplest MRF, the Ising model has all the important properties of a general MRF. The clique includes only two nodes and hence represents the pairwise relationship. When the variables have more than two possible states, we have a Potts model [122]. The Potts model will be extensively used in the following chapters when we apply MRF to the hidden labels of the brain functional networks and the number of networks is greater than two.

When x_s takes a value in a continuous domain, and is in a conditional Gaussian distribution given the remaining variables, the random field is called a Gaussian random field (GRF) [132]. GRF is an important model of spatial process, although we will not discuss it further.

3.3 Simulation of MRF

It is often important to draw samples from a multivariate distribution. A general usage of samples is the Monte Carlo integration. Consider the generic problem of evaluating the integral $\mathbb{E}_{f(x)}[h(x)] = \int_{\mathcal{X}} h(x)f(x)dx$, we can use a set of samples (x_1,\ldots,x_M) drawn from the density f(x) to approximate the above integral by the empirical average $\overline{h} = (1/M)\sum_{m=1}^{M} h(x_m)$. In imaging related problems, the samples are also used for model validation. By comparing the observed data with the samples drawn from the probability distribution assumed in our model, we can tell if our assumption of the distribution is valid. In our MRF model, thanks to the equivalence of the MRF and Gibbs distribution, we can simulate a MRF image by drawing samples from the corresponding Gibbs distribution.

3.3.1 Metropolis and Gibbs sampling

To draw a sample from a distribution in the form of $P(X) = (1/Z) \exp\{-U(X)\}$, one can use either Metropolis sampling [105] or Gibbs sampling [65]. Both methods are in the class of Markov chain Monte Carlo (MCMC) methods. In general, MCMC method draws samples from high-dimensional distributions by iteratively drawing a univariate sample given the other fixed variables, thus converting a multivariate sampling problem into a univariate one. The multivariate variable X with a single node changed at each step consists of a series of Monte Carlo samples, as illustrated in Figure 3.3. The algorithm of Metropolis sampling is shown in Algorithm 1.

It is noted that Algorithm 1 is slightly different from the general Metropolis sampling [105]. Here we compute the difference of the energy instead of the ratio of the density at X^m and candidate W. The difference of the energy is calculated because for MRF and Gibbs distribution, computing the ratio of two densities is equivalent to computing the difference of exponential terms, i.e., the energy function. Although both X^m and the candidate W are high-dimensional, they are different at only one node s. Therefore, we can sample x_s given all other variables are fixed, and construct W with the candidate w and the remaining variables. Now we have a univariate sampling problem that is significantly easier than the previous multivariate one. In practice, the proposal distribution can be a uniform distribution, and $\Delta E(W)$ can be computed just by looking at the cliques that involve x_s

Algorithm 1: Metropolis sampling algorithm for MRF.

since the energy of all other cliques does not change.

Figure 3.4 gives a simulated result from the Ising model in the form of (3.6). A binary image with resolution 256×256 is initialized with random states of 0 and 1. Each pixel is then updated in a raster scan order according to the procedures in Algorithm 1 with a uniform distribution as a proposal distribution. The order of the pixels for updating does not matter to the results as long as the sampling reaches stationary distribution of the Markov chain. We call it a scan once all pixels are visited just once, no matter if they are updated or not. Each of the subplots in Figure 3.4 has been scanned 1000 times to guarantee the sampling routine's convergence to stationary distribution. We show the simulated sample image with various values of β . In statistical physics a similar definition of Ising model has a parameter T, i.e., the temperature. The β in our definition is indeed the reciprocal of T. It has been shown [89] that the Ising model has a critical temperature with a corresponding β_c , such that the sampled image exhibits an unordered state with $\beta < \beta_c$, and exhibits an ordered state (either towards all zero or towards all one) with $\beta > \beta_c$.

The advantage of Metropolis sampling method is we do not need to sample from $P(x_s|x_{-s})$. Instead we sample from the proposal distribution, which is an easier problem than sampling from the original $P(x_s|x_{-s})$. As long as $\triangle E$ is easy to compute, the sampler will work. However, the convergence rate depends on the acceptance rate of the proposal distribution. For example, when the variables have more than two states, the same procedure in Algorithm 1 can be used to draw samples from the Potts model. Because of the greater number of states, the candidate label has a much larger probability of not being equal to its neighbors if we choose uniform distribution as a proposal. Therefore, the rejection rate will be higher than the two-class Ising model, and it may take more scans for the sampling of the Potts model to converge to the stationary distribution. Figure 3.5 shows the simulation of a 128×128 image from the Potts model with different values of β . A sample from a stationary Potts model distribution is a piecewise constant label map given $\beta > \beta_c$.

The Gibbs sampler is a special case of the Metropolis sampler in that the proposed candidates are always accepted. We use Gibbs sampling also in the multivariate problem and construct a Markov chain whose stationary distribution equals the target distribution P(X). As in Metropolis sampling, we use the Gibbs sampler to draws samples from $P(x_s|x_{-s})$, i.e., the univariate distribution of just one variable given the other variables are fixed. However, here the univariate distribution is a known distribution such that we can directly draw samples from it. This is different from Metropolis, where it may be difficult to draw samples from the univariate distribution conditioned on remaining variables, and we use

an easier proposal distribution as a surrogate. Once it is drawn, the sample is accepted with probability 1, and the sampler moves to the next variable. The procedure of Gibbs sampling is given in Algorithm 2. Compared to Metropolis sampling, the Gibbs sampler typically needs fewer iterations for convergence. However, that does not always mean less computation time compared to the Metropolis. If the direct sampling from $P(x_s|x_{-s})$ takes more time than sampling from the proposal distribution in the Metropolis sampler, the overall time may still be more than Metropolis sampling.

3.3.2 Swendsen-Wang sampling

Metropolis sampling and Gibbs sampling can be slow, especially when there are strong interactions between the neighboring nodes on the graph. When the sampler is not initialized correctly (i.e., the initial sample is far from the mode of the target distribution), the sampling may take an exponential number of steps to reach convergence [7]. The Swendsen-Wang algorithm [148] is proposed to address this issue. To understand the Swendsen-Wang (SW) algorithm, some background information is needed. There is a fundamental theorem [128] that underlies the slice sampler and also the SW algorithm. Assuming f is the pdf from which we want to draw samples, f(x) can be written as

$$f(x) = \int_0^{f(x)} 1du$$

f(x) can be seen as the marginal distribution of joint variables (x, u)

$$(x, u) \sim \mathcal{U}\{(x, u) : 0 < u < f(x)\},$$
 (3.7)

where \mathcal{U} is the uniform distribution, and u is usually named as the *auxiliary variable*. Thus, instead of drawing samples from f(x) directly (which might be difficult), we can draw samples (x, u) from their uniform joint distribution on the constrained set $\{(x, u) : 0 < u < f(x)\}$. Once we have the samples, we can discard u, and x will be in the original target distribution. This is the basic idea of a slice sampler.

In a slice sampler, we can generate a Markov chain with the stationary distribution equal to the joint uniform distribution of (3.7). We can generate x and u from their conditional distribution iteratively in a random-walk style: 1) generate u from $\mathcal{U}(\{u:u\leq f(x))\}$, and 2) given the new sample u, generate x from $\mathcal{U}(\{x:f(x)\leq u\})$. Robert and Casella [128] prove this Markov chain's stationary distribution is indeed (3.7).

When f(x) is a complex function, finding the set of x such that $f(x) \leq u$ can be difficult (step 2 in the above procedure), which can happen when the x is of large dimension (as in our fMRI study). The general slice sampler solves this problem by using multiple slices. In

Algorithm 2: Gibbs sampling for MRF.

Data: Definition of P(X)**Result**: Samples of P(X)

Start with any initial value X_0 with $P(X_0) > 0$;

while Not converged do

Given current state X^m , pick a node x_n and generate a new candidate w_n from $P(x_n|x_{-n})$;

Accept w_n with probability 1;

short, f(x) can be factorized into the products of $f_i(x)$, and each f_i is associated with an auxiliary variable u_i . In this way, the support of the conditional distribution p(x|u) in step 2 can easily be found. The SW sampler can be seen as one of such general slice samplers.

The settings of the SW algorithm are as follows: To sample from the Potts model in the form of (3.6) using the SW algorithm, we introduce a set of augmented binary random variables U as we do in the slice sampler. The variable u_{rs} corresponds to the bonds between spatially adjacent nodes x_r and x_s . For each u_{rs} , there are two states open or close denoted by $u_{rs} = 1$ or $u_{ij} = 0$. Conditioned on X, the u_{rs} are independent. Each u_{rs} is a uniform distribution on the interval $[0, a_{rs}]$, with $a_{rs} = \exp(-\beta \psi(x_r, x_s)) \leq 1$. So the conditional pdf of $u = \{u_{rs}\}$ given X is

$$f(U|X) = \prod_{(r,s)} \frac{\mathbf{1}_{(u_{rs} \le a_{rs})}}{a_{rs}} = \left(\prod_{(r,s)} \mathbf{1}_{u_{rs} \le a_{rs}}\right) \exp\left\{\beta \sum_{(r,s)} \psi(x_r, x_s)\right\}$$

The reason we define the distribution of U in this way is the joint distribution P(X, U) can be written simply as

$$P(X, U) = P(X) \cdot P(U|X) \propto \begin{cases} 1 & \text{if } u_{rs} \leq a_{rs}, \forall (r, s) \in \mathcal{V} \\ 0 & \text{otherwise.} \end{cases}$$

Therefore the P(X,Y) is uniformly distributed. More importantly, $P(X|U) \propto P(X,Y)$ is also uniformly distributed over the set $\mathcal{A} = \{X : u_{rs} \leq a_{rs}\}$. Now either $u_{rs} \in [0, e^{-\beta}]$ or $u_{rs} \in (e^{-\beta}, 1)$. If $u_{rs} \in [0, e^{-\beta}]$, it is impossible to tell if $x_r = x_s$ since such values of u can happen either $x_r = x_s$ or $x_r \neq x_s$. If, however, $u_{rs} \in (e^{-\beta}, 1)$, there must be $x_r = x_s$.

Therefore, the sites r and s for which $u_{rs} \in [e^{-\beta}, 1]$ can be gathered into clusters, and within each such cluster the x of all the nodes must be the same, and the value of x is uniformly distributed. The x_r and x_s values for those $u_{rs} \leq e^{-\beta}$ are not constrained to be the same and can be an arbitrary value. These values are also uniformly distributed. Therefore, we can generate samples of X given U, and generate samples of U given X. We can even further simplify the sampling by noting the exact value of u_{rs} is not required. We can simply record if $u_{rs} > e^{-\beta}$ by a binary variable v [131]. The variable v is in the Bernoulli distribution $\mathcal{B}er(1-e^{-\beta})$ such that $v_{rs}=1$ if $u_{rs}>e^{-\beta}$, and $v_{rs}=0$ otherwise. Then the SW algorithm iterates between the two steps:

- Given X, set $v_{rs} = 0$ if $x_i \neq x_j$. When $x_i = x_j$, set $u_{rs} = 1$ with probability $1 e^{-\beta}$, and set $v_{rs} = 0$ with probability $e^{-\beta}$. After this step, we have multiple connected components, each being a subset of the nodes on the graph.
- Given *U*, set all the nodes in a randomly chosen cluster (i.e., a connect component) with the same label. The label is drawn from a uniform distribution.

The stationary distribution of this Markov chain, like the slice sampler, is the joint distribution of U and X, which is again a uniform distribution [131]. It can be proved [150] that the marginal distribution P(X) is exactly (3.6). So the joint model is consistent with the original marginal distribution. If we sum out X and get the marginal distribution of the augmented variable P(U), we have a distribution called random cluster model [74]. After the sampling, we obtain samples of (X, U). We ignore the augmented variable U, and X will be the samples from the original target distribution.

The SW sampling is more efficient than Gibbs sampling, because at each step it changes the labels of the whole cluster, instead of only a single site. Even in low temperatures, the sampler flips the labels for larger clusters. Figure 3.6 gives a comparison of the samples of the Potts model in (3.6) drawn from Gibbs sampling and SW sampling. To show the difference between the two samplers, we choose a small burn-in period (100) and initialize the image with all-zero values. The all-zero initialization is far from the mode of the Potts model. Gibbs sampler has difficulty reaching the stationary distribution in a short burn-in period. On the other hand, the SW sampler converges during this short interval.

The mixing time of the sampling is polynomial in a regular lattice. Barbu et al. [7] discussed the convergence rate of SW algorithm on the Potts model. Huber [83] developed a new bounding chain algorithm to diagnose the convergence of Swendsen-Wang sampling. The number of steps to reach perfect sampling (which means convergence to stationary distribution) is in the order of $\mathcal{O}(\log |\mathcal{E}|)$, where $|\mathcal{E}|$ is the total number of edges. This running time applies when the temperature is far below or far above critical temperature. Cooper [33] shows the mixing time (or convergence time) is polynomial if the number of neighbors of each node does not increase with $|\mathcal{V}|$, the size of the nodes. The polynomial mixing time is good for the regular lattice where the number of adjacent nodes is a constant regardless of image size. Compared with the super-exponential rate of increase for the iteration number in standard Gibbs sampling, the SW algorithm is a big improvement for the convergence rate. These theoretical analysis are appropriate for cases without external fields, i.e., the data likelihood term.

3.4 Hidden Markov model

The main purpose of MRF in this dissertation is a prior distribution on the hidden variables to enforce the piecewise constant constraint for discrete variables, and the smoothness constraint for the continuous variables. In real-world applications, we are often provided with some noised data Y, and the goal is the inference of the true structures X behind

the observations. The true structures can be the true image pixels in an image denoising problem, or they can be the class labels in an image segmentation problem or a data clustering problem. Because the latent variables we are interested in are not observed, we call them hidden variables. The identification of hidden variables from the observations is often difficult, because multiple hidden variables can fit the data depending on the criteria. If we have some prior knowledge of the value of the hidden variables, such as whether they will be smooth or a piecewise constant on the image domain, such knowledge should be included in the estimation process. This prior knowledge or assumption is called *inductive bias* in machine learning. Inductive bias is the assumption of the learner to predict outputs that it has not encountered, given input data [109]. Although we are not in the training-testing framework here, the piecewise constant or continuity prior also applies as an assumption of the unseen hidden variables.

There are two classes of approaches of introducing the prior knowledge in the hidden variables. One is a Bayesian approach that is called the hidden Markov model (HMM). In this model, we define a MRF as an a priori distribution on the hidden variables X. Given X, we define another conditional probability P(Y|X) and assume Y is generated from the conditional distribution given X. This probability is also called the likelihood function of Y. Figure 3.7 gives an illustration of this model. Then the question to be answered is the posterior distribution of X given the data Y. According to the Bayesian rule,

$$P(X|Y) = \frac{P(X)P(Y|X)}{P(Y)} \propto P(X) \cdot P(Y|X)$$
(3.8)

The \propto is because we are not interested in P(Y) when looking at X as a variable, so P(Y) is a constant that can be ignored. Various methods exist for the inference in the form of (3.8), and we will discuss some of these methods in Section 3.5.

Another class of approaches to model both the observed data and hidden variables is conditional random field (CRF). In contrast to the HMM where P(X|Y) is decomposed into two separate parts P(X) and P(Y|X) by Bayesian method, CRF does not have such an explicit decomposition [93]. Instead, in CRF model, one assumes given the observed data Y, that X obeys the Markov property with respect to the graph \mathcal{G} . CRF directly defines a distribution on P(X|Y) such that the variables x_s at node s depend on other nodes. To put it another way, CRF's prior P(X) also depends on the observed data. This dependence is a violation of the Bayesian rule. However, the dependence does make sense in some situations. For example, the smoothness constraint should be relaxed if the observed data at two nodes are too different such that the underlying variables are impossible to be piecewise constant or smooth. In such cases, one cannot rewrite the P(X|Y) into the

product of P(X) and P(Y|X) up to a constant, and nor is it always necessary to do so. We give an example of the CRF model that is used in [20, 129]. Here the task is image segmentation with MRF defined on the hidden region labels. The clique potential function used in the MRF is defined as

$$V(X,Y) = \gamma \sum_{(m,n)\in\mathcal{E}} dis(m,n)^{-1} [x_m \neq x_n] \exp\{(y_m - y_n)^2\},$$
 (3.9)

where the [c] takes 1 if the conditional c is true, y_n and y_m are observed pixel intensities, and dis(m,n) is the distance between pixel m and n. We can see the clique potential, as part of the prior distribution's energy function, is also a function of the data Y.

3.5 Inference of graphical model and MRF

Given the definition of the graph and the observed data on some nodes, our goal is the statistical inference of the unknown variables. The graph inference addresses the issue of computing the posterior distribution or its expectation of the unknown variables given the observed data. The difficulty of the inference depends heavily on the structure of the graph. For example, a chain graph is the simplest graph structure, and the exact inference can be achieved by passing local messages on the chain. The time is linear in the number of nodes. Such methods can be generalized to trees without losing the linear computation time property [16, 113]. For an undirected graph, a tree is a graph that has no loops. For more general graphs, whether these existing inference methods work in a reasonable amount of time depends on the extent that the graph is like tree graph structures, measured by the tree width of the graph. The MRF mode defined in our work is different from a tree, so the exact inference is often intractable. However, we will look at some approximate inference methods that can find a good approximation to the optimal solution within reasonable computation time. These approximate methods will be applied to the specific problems in the following chapters, with some modifications.

To see why the exact inference is often not available on a general graph, we note for a graph with N nodes, and the variable x_s at each node s takes discrete values in $\{1, \ldots L\}$, the total number of possible realizations is L^n , an exponential function of the data points N. Because of the interactions among the variables, the inference of each variable cannot be factorized, therefore searching for an optimal solution in a big space will be difficult.

3.5.1 Iterated conditional modes

One method of finding the discrete random vectors X that maximizes the posterior P(X|Y) in the early year of MRF study is the iterated conditional Modes (ICM). Besag [15]

proposed this greedy strategy update each single node x_s that maximizes the conditional distribution $P(x_s|x_{-s},Y)$ given other nodes that are fixed. Algorithm 3 gives the procedures of the ICM algorithm. In practice, the results depend on the initial values of X and a typical choice of initialization is the maximum likelihood, i.e., an initialization of X that maximizes the likelihood function P(Y|X). The algorithm updates each data point in a prescheduled order until no more nodes are changed. The final result is a local optimal solution. The neighboring solutions in the search space would be the set of X that has only one node difference to the current solution. Therefore, the search region of ICM is small compared to the exponential large full space. In Section 3.5.5, we will see that a criterion to evaluate the performance of approximate algorithms is the size of the space within which the approximate solution is optimal. Compared to other modern methods such as the graph cuts method [20, 19], the neighboring space of the solution derived by ICM is small. Despite its limitations, ICM is widely used in practice due to its simplicity, and sometimes it achieves good results [161].

3.5.2 Sampling

In Section 3.3.1 we have shown that sampling techniques can be used to draw samples from complex distributions such as the Ising and Potts models. For the probabilistic inference from the posterior P(X|Y), we can again draw samples from this posterior by using Metropolis or Gibbs sampling. We want to do this for two reasons: First, since the ICM method tends to be stuck in the local minima if not initialized correctly, we can instead draw many samples from P(X|Y), and use Monte Carlo averaging to approximate the random functions we are interested in, such as the posterior mean. With a good design of samplers, the samplers can jump out of the local minima and the averaging of the samples is a good approximation of the posterior mean. Second, with the samples available, we are not only able to perform a point estimation, but also can estimate the confidence of the point estimates, i.e., the variance of such estimates. The set of samples has all the information of the posterior P(Y|X) as long as the number of samples is big enough and the samples are indeed from the target distribution.

The sampling procedure from the posterior P(X|Y) is similar to that of the prior P(X) for both Gibbs and Metropolis, except that now the local distribution that we draw univariate sample x_s from is also conditioned on the observed data y_s . For example, if we define P(X) as an Ising model, and the likelihood function is Gaussian, i.e., $P(y_s|x_s) \sim \mathcal{N}(\mu(x), \sigma(x)^2)$, the conditional probability of x_s will be

Algorithm 3: Iterated conditional modes (ICM) for finding approximate posterior of discrete random vector given data

```
Data: Definition of P(X|Y)

Result: A realization of X that maximize P(X|Y)

Start with a realization X_0 = \operatorname{argmax}_X P(Y|X);

while Not converged do

foreach s \in \mathcal{V} do

x_s \leftarrow \operatorname{argmax}_{x_s} P(x_s|x_{\mathcal{N}(s)}, y_s);
```

$$\log P(x_s|x_{\mathcal{N}(s)}, y_s) = -\beta \sum_{r \in \mathcal{N}(s)} \psi(x_r, x_s) - \frac{(y_s - \mu(x_s))^2}{2\sigma^2(x_s)} - \sigma(x_s).$$
 (3.10)

With a slight modification based on Algorithm 2, the Gibbs sampling routine is given in Algorithm 4.

The original SW sampling applies only to Ising and Potts models. The generalized SW sampling algorithm proposed by Barbu and Song-Chun Zhu [7] can be applied when there is data likelihood and we seek sampling from the posterior distribution of P(X|Y). In addition, the generalized SW algorithm makes use of the observed data when sampling the cluster labels from the proposal distribution, and this specialized proposal function makes the convergence faster than the standard SW algorithm. The last strength of the generalized SW sampler is to adaptively increase or decrease the number of labels.

There are two significant changes from standard SW to generalized SW. First, the probability of turning on the edges (bonds) at step 1 is changed from $q_0 = 1 - e^{-\beta}$ to $q_e = -g(h_i, h_j)$, where h is the observed data (or features). $g(h_i, h_j)$ would take a larger value when the observed data at i and j are similar. The similarity is represented by the KL divergence [7], but can be defined differently in other applications. Second, the sampling of labels in step 2 can have an acceptance rate smaller than one, instead of the 100% acceptance in the original SW algorithm. The acceptance probability to move to new labels also depends on posterior probability given the observed data, as shown in theorem 2 in the work of Barbu et al. [7]. The third version SWC-3 of the generalized SW replaces the Metropolis-Hasting sampling in step 2 with a Gibbs sampler and achieves the acceptance rate of 1. The Gibbs sampler draws labels from the posterior probability given the data.

Relating the generalized SW sampling to the fMRI application, there are two issues to address in order to use SW sampling. First we need to define a function $g(h_i, h_j)$ to replace the KL divergence in eq (12) of Barbu and Song-Chun Zhu [7]. The function will be plugged into the acceptance probability when sampling the augmented variable U (edge variables). One straightforward solution is to use the correlation between the BOLD signal of two voxels or two ROIs. An edge will be open with larger probability if two voxels connecting the edge have a higher correlation. More work needs to be done to find the relationship between the acceptance probability q_e and the posterior probability p(X|Y).

3.5.3 Simulated annealing

Depending on whether we are interested in a point estimation or a full Bayesian analysis, the statistical inference of the problem of P(X|Y) aims either at the full posterior distri-

Algorithm 4: Gibbs sampling for the posterior distribution.

Data: Definition of P(X|Y)**Result**: Samples of P(X|Y)

Initialize X by maximum likelihood estimates: $X_0 = \operatorname{argmax} P(Y|X)$;

while Not converged do

Pick a node x_n ;

Draw sample w from $P(x_n|x_{-n},Y)$. Construct a new candidate vector W with

the existing X^m and the new w; Accept W with probability 1; bution, or a maximum a posterior (MAP) estimation. When the latter is of the interest, one makes use of the sampling technique, together with the simulated annealing method to find the mode of the posterior distribution.

Simulated annealing (SA) optimization is a method originally introduced in statistical mechanics and later used by Kirkpatrick [90] for optimization. The goal of finding the mode of P(X|Y) is indeed a combinatorial optimization problem that cannot be solved in polynomial time. Given the definition of a probability distribution function $P(x) = (1/Z) \exp(-E(X))$, we can introduce a new temperature parameter T and construct a new distribution $P(x) = (1/Z) \exp(-E(X)/T)$. When T is high, all the possible states of the variables have similar probabilities, and the sampling will be in a near-random state. When T is low, only the most probable samples will happen. The annealing process is similar to the fact that material solidifies at lower temperature [65].

The SA algorithm with Metropolis or Gibbs sampling is fundamentally different from the iterative method such as ICM. In the iterative gradient descent method, one iteratively moves each variable in the system towards the descent direction of the gradient. Furthermore, the system may get stuck in a local minimum. Figure 3.8 gives an illustration of the difference of the coordinate descent and SA method.

3.5.4 Variational inference

Another major class of approximation methods is variational inference. Here the goal is to find the *best* posterior distribution from a subset of all possible distributions. Although the original variational methods address the issue of finding a derivative with respect to a function, we can use this concept to find the approximate solutions of the P(X|Y). Instead of optimizing the objective functional over the whole space of possible P(X|Y), we can search in the restricted set of the posterior function. For our specific problem with MRF as the prior P(X), a typical restriction is that P(X|Y) must be able to be factorized into the form

$$Q(X) = \prod_{s=1}^{S} q_s(x_s),$$

where each x_s is a disjoint subgroup or a single variable in the original random vector X. The task now is to look for a best Q(X) within the subset with the above factorized form. Next, we need to define an objective function of X. The marginal likelihood function of Y, or equivalently the $\log P(Y)$ can be written as the sum of two terms [16]:

$$\log P(X) = LB(Q) + KL(q||p) \tag{3.11}$$

$$LB(Q) = \sum_{X} Q(X) \log \left\{ \frac{P(X,Y)}{Q(X)} \right\}$$
 (3.12)

$$KL(Q||P) = -\sum_{X} Q(X) \log \left\{ \frac{P(X|Y)}{Q(X)} \right\}. \tag{3.13}$$

where P is the true posterior distribution P(X|Y) we look for. Because the second term $\mathrm{KL}(Q||P)$ is the KL divergence between distribution Q and true posterior distribution P, it is always greater than or equal to zero. The first term $\mathrm{LB}(Q)$ is essentially the lower bound of P(X). Therefore, in order to maximize P(X), we instead maximize its lower bound. If we search Q in the full possible space, we will end up with Q = P(X|Y) and the KL divergence will be zero. In practice, since it is intractable to search the full space, we search an approximate solution within a subspace. Because of the factorization of Q(X), we optimize the KL divergence with respect to each factor $Q_s(x_s)$ in turn. Bishop [16] and Murphy [113] have shown that within this restricted search space, the optimal $Q_s(x_s)$ has the following property

$$\log Q_s(x_s) = \mathbb{E}_{r \neq s}[\log P(X, Y)] + \text{const}$$
(3.14)

This property means we can compute the optimal factor $Q_s(x_s)$ by computing the log of joint distributions of all hidden variables and observed variables, and then take expectations with respect to all the hidden variables excluding x_s . The terms will be absorbed into the constant term unless they are functions of x_s . Because x_s is a discrete variable in our problem, we can compute $Q_s(x_s)$ by taking the exponential of both sides of (3.14), repeat this for all possible values of x_s , and then normalize such that $Q_s(x_s)$ sums to 1.

To apply the variational inference to our model P(X|Y) with X a MRF, we first assume P(X) is a MRF in the form of

$$P(X) = \frac{1}{Z} \exp \left\{ \beta \sum_{(r,s) \in \mathcal{V}} \langle x_r, x_s \rangle \right\}.$$
 (3.15)

Here, for convenience, we rewrite x as a vector of length L, with L the number of possible discrete values. Each element of x, denoted by x_k , is a binary indicator variable. The angle bracket computes the dot product of two vectors. We see that although this definition of MRF takes a different form to (3.6), it is also a valid MRF that prefers smoothness within the neighbors on the graph. For example, if x_r and x_s are equal, their dot product will be 1, and hence has greater probability. We further define $P(y_s|x_s)$ a Gaussian distribution for the convenience. It can be any other distribution in other applications and does not change our derivation of the variational methods. Then we follow the assumption of variational

inference and assume the posterior has to be in the form of (3.5.4). By using the property of (3.14), we can write the log of the posterior at node s as

$$\log Q_s^*(x_s) = \mathbb{E}_{r \neq s}[\log P(X, Y)] + \text{const}$$
(3.16)

$$= \mathbb{E}_{r \neq s} \left[\beta \sum_{(r,s) \in \mathcal{V}} \langle x_r, x_s \rangle - \log Z\right] + \mathbb{E}_{r \neq s} \left[\log P(Y|X)\right] + \text{const}$$
 (3.17)

$$= \mathbb{E}_{r \in \mathcal{N}(s)} \left[\beta \sum_{r \in \mathcal{N}(s)} \langle x_r, x_s \rangle \right] + \mathbb{E}_{r \in \mathcal{N}(s)} \left[\log P(y_s | x_s) \right] + \text{const}$$
 (3.18)

$$= \beta \sum_{r \in \mathcal{N}(s)} \langle \overline{x}_r, x_s \rangle + \log P(y_s | x_s) + \text{const.}$$
(3.19)

In the above derivation, $\overline{x}_r = \mathbb{E}_{x_r}[x_r]$. We note in (3.17), because of the MRF property, current node x_s is conditionally independent of the remaining nodes given its neighbors $x_r, \forall r \in \mathcal{N}(s)$. Therefore the expectation can be simplified to those nodes neighboring x_s . Furthermore, $\log Z$ and $\log P(y_r|x_r), \forall r \neq s$ are not functions of x_s , so they can be absorbed into the constant term. From (3.18) to (3.19), the expectation goes inside of the dot product because of the linearity of expectation, and the expectation of $\log P(y_s|x_s)$ with respect to $r \in \mathcal{N}(s)$ is just itself since it is not functions of x_r . Finally, we got a simplified form, and it is just the original clique potential functions with neighbors replaced by their expectations, plus a log-likelihood term. This updating is called *mean field* theory in statistical physics [159], and here we derive it by the variational method, with the only assumption that the target posterior P(X|Y) must be factorized.

Because the solution includes the expectation of other nodes that are also unknown, this is not a closed-form solution. We will adopt an iterative approach here. With an appropriate initialization of x, we update each factor x_s in a scheduled order, given the expected value of its neighboring nodes fixed. A crux of the mean field theory for MRF with number of labels $|\mathcal{L}| > 2$ is, the kth element x_k of the variable x's is a binary variable, therefore the posterior probability of x_k is just equal to its expectation. To see that, by the definition of expectation, $\mathbb{E}_{x_k}[x_k] = 1 \cdot P(x_k = 1|\cdot) + 0 \cdot P(x_k = 0|\cdot) = P(x_k = 1|\cdot)$, where $P(x_k|\cdot)$ means the posterior of x_k given the neighboring nodes. Therefore, the posterior estimated at x_s is used as \overline{x}_s when updating its neighbors x_r . The cycling continues until convergence where no more node change its expectation value.

One of the advantage of using variational inference approximation to interpret the standard mean field theory is we can build a full Bayesian model that integrated MRF and other prior knowledge on the parameters, and still use variation inference to solve it. This is because variational inference treats the parameters the same way as the hidden

variable once we assume a distribution on the parameters. So both the hidden variable and parameters are just one factor $Q_s(x_s)$ in the variational approximation.

3.5.5 Graph cut optimization

If the goal is the mode of the posterior distribution P(X|Y), graph cut segmentation is an alternative algorithm that can achieve global optimum in a polynomial time when there are only two possible states for each node. The earliest connection of graph cut and the combinatorial optimization of computer vision problem is found by Greig et al. [73], and later re-introduced to multiple labels segmentation by Boykov et al. [19]. The function that is being optimized by this class of methods is of the form

$$E(X) = \sum_{(r,s)\in\mathcal{E}} U_{rs}(x_r, x_s) + \sum_{s\in\mathcal{V}} D_s(x_s).$$
 (3.20)

The function E(X) has two terms, a pairwise smoothness term, where U is the clique potential function of the pairwise nodes, and a data term, where D_s represents how the label x_s match the observed data y_s (not shown since it is a given constant). We can verify that this objective function is indeed the negative log likelihood of P(X|Y) in our model (3.6), so a maximum a posteriori estimation is equivalent to minimizing the E(X) in (3.20). Greig [73] pointed out that we can construct a graph with all the variables defined on the nodes, and pairwise constraints defined on the edges. We further define two additional nodes, a source node s and a sink node t. There are edges between each regular node and s, t, and the weights of the edges are defined according to the data term D_s . Between regular nodes x_r and x_s , an edge is added if they are neighbors in the original image domain, and the weights of the edges are defined by the clique potential functions of the MRF. Given the above settings, finding the minimal value of E(X) is equivalent to finding a cut, i.e., a subset of edges that has minimal total weights, because the sum of the weights is equal to E(x) up to an additive constant. According to Ford and Fulkerson [53], the minimum of E(X) is the maximum flow through the graph from the source node s to the sink node t subject to the edge capacities (weights), and there is efficient algorithm for solving this problem. Figure 3.9 shows the segmentation of images with MRF prior to using graph cut algorithm.

Boykov et al. [19] generalize the graph cut algorithm to multiclass segmentation. Optimizing the general form of (3.20) is a combinatorial optimization and a NP-hard problem. Boykov et al. still use a greedy algorithm to search the locally optimal configuration. The difference to the standard greedy algorithm is the search space is much larger. Note in standard greedy algorithm such as ICM, the algorithm only searches the optimal label

configuration with one voxel distance of the current configuration. That is, only one voxel move at a time to find the better solution. Boykov et al. define two types of moves: α expansion and $\alpha\beta$ swap, such that a large number of voxels change labels simultaneously within a single move. The local minium solution under such movement is much closer to the global minimum. Actually the α expansion move is a strong one such that the local optimal labeling with respect to this moves is within a known factor of the global minimum.

The α expansion and $\alpha\beta$ swap moves both have an exponential number of possible moves of voxels. The authors convert the problem of finding optimal solution within one α expansion or one $\alpha\beta$ swap moves, to the problem of a binary-label graph cut problem. Because graph cut is able to find the optimal solution efficiently by max-flow algorithm, the α expansion move can also be solved efficiently.

The graph cut algorithm and its extension is used not only in HMM but also in CRF. For HMM, the log of data likelihood includes part of the data term in (3.20), together with the unitary prior term in MRF. For CRF, the prior also includes the data, but can also be used as the smoothness term in (3.20). Therefore, the definition of the prior term is transparent to graph cut algorithm, as long as the objective function has the form of (3.20).

To show an example of using graph cut for segmentation of binary images, we generate a true label map of resolution 200×200 using the Ising model in (3.6) with $\beta = 0.7$, and scan 100 times on each pixel. The Gaussian noise of zero mean and $\sigma^2 = 9$ is added to the label map in order to obtain a noisy observed image. We then use the graph cut to segment the image, given the correct β and σ^2 . Figure 3.10 shows the generated true label map, the noisy image, and the recovered label map by graph cut. From the histogram of the observed image's intensity, it is difficult to find an optimal threshold to separate two classes as the two Gaussian components are heavily overlapped. By using the spatial soft constraints modeled by the MRF prior, and the graph cut for global optimal solution, we can recover most of the label map. Some finer structures are lost, though, as can be seen from the circled region in the true label map in Figure 3.10. This is believed to be one disadvantage of graph cut, i.e., the global criteria of energy minimization often is achieved at the price of losing local structure. Because of that disadvantage, the label map estimated by graph cut mostly has blob-like patterns, even the original image does not have such patterns. This makes it difficult to apply graph cut for segmentation of thin structures such as blood vessel and trees. Also because of the preference of blob-like shapes, if we use the the estimated label map for parameter estimation, the estimated parameter will tend to be larger than the true value. This is the reason we did not choose graph cut as the optimization method in our expectation maximization framework introduced in Section 3.6.2.

3.6 Parameter estimation

Functional MRI data include multiple sessions of one subject, and multiple subjects in one site, and even the data from multiple sites. The heterogeneous nature of the data acquisition process means the model parameters are probably different across sites and even subjects. A data-driven model does not need the user to give the model parameters. Instead, the estimation process includes both the hidden variable inference and parameter estimation. In this section, we will address an easier question: Given the observed data and also the values of the hidden variables, we seed to estimate the parameters in our hidden Markov model.

More specifically, we will study the parameters in the model P(X|Y) with the prior defined by (3.6) and the likelihood P(Y|X) defined in any appropriate form. One possible model is that X is modeled by a MRF, and each y_s is in a univariate Gaussian given x_s , as shown by a graphical model in Figure 3.11. There are various criteria and methods to estimate the parameters in such a model, and in this work we give one method that we use extensively in the following chapter, i.e., the maximum likelihood (ML) estimation. For ML estimation, we aim to find a set of parameters θ that maximize the likelihood (or equivalently, log-likelihood) of the form $P(X) \cdot P(Y|X)$. Furthermore, the set of parameters θ may include the set of parameters θ_P in the prior distribution P(X), and the set of parameters θ_L in the conditional distribution P(Y|X). Because the factorization of P(X) and P(Y|X), the optimal parameters should be

$$\theta_P^* = \operatorname*{argmax}_{\theta_P} \log P(X; \theta_p),$$

$$\theta_L^* = \operatorname*{argmax}_{\theta_L} \log P(Y|X; \theta_L).$$

Depending on the specific form of the conditional probability P(Y|X), the estimation of θ_L could be either closed form, or through iterative refinement. For example, the estimation of μ and σ^2 in the model of Figure 3.11 is in closed form given X and Y. Here we will focus on the estimation of θ_P . Since the logarithm is monatomic function, we just need to optimize $\log P(X;\theta_L) = -U(X;\theta_L) - \log(Z(\theta_L))$. The normalization constant Z is also a function of θ_L . The evaluation of Z is intractable given the combinatorial nature of X. Therefore, one has to resort to approximate the likelihood function. One simple method is the pseudo-likelihood [14]. The likelihood function is approximated by the product of the distribution at each node. For the Ising model in (3.6), the approximation is

$$\widetilde{P}(X;\beta) = \prod_{s \in \mathcal{V}} P(x_s | x_{\mathcal{N}(s)}) = \prod_{s \in \mathcal{V}} \frac{1}{Z_s} \exp \left\{ -\beta \sum_{(r,s) \in \mathcal{V}} \psi(x_r, x_s) \right\}.$$

Because Z_s is the summation over a univariate x_s , it is easily computed. The pseudo-likelihood is the sum of the conditional probability of each variable assuming other variables are given. This approximation is, in spirit, similar to the variational inference approximation in Section 3.5.4, as both use factorized univariate distributions to approximate the original multivariate distribution. However, they serve different goals. In variational approximation, the factorization defines a restricted space in which the approximate solution is found. Here the approximation is used to convert an intractable function to one that can be easily evaluated. Once the likelihood is represented by pseudo-likelihood, the optimization can be solved by the standard gradient descent method. We will defer the estimation in specific applications to the following chapters.

Other estimation methods include coding method and least squares estimation. In coding method, the variables are split into K disjoint subset. With the subset, the variables are independent with each other given all other subsets. The number K depends on the neighborhood definition of the original graph structure. For 8-neighborhood system of a two-dimensional image, the pixels can be split into four groups, as shown in Figure 3.12. For a single group, the parameter θ can be estimated by optimizing the joint likelihood of the variables $\prod_{s \in \mathcal{V}_k} P(x_s|x_{-s};\theta_k)$, where \mathcal{V}_k is the set of nodes for kth group. The joint likelihood does not have the normalization constant Z thanks to the independence of the variables and is therefore tractable. Once θ_k is estimated from group k, a final estimate of θ is computed by averaging θ_k .

3.6.1 Expectation maximization

Often we face the problem that both the hidden variables and the parameters need estimation. In such a situation, the expectation maximization (EM) is the standard approach. EM was proposed for maximum likelihood parameter estimation when the likelihood function is too difficult to evaluate or take derivatives [42]. By introducing a hidden variable x_s for each data point y_s , the EM algorithm takes two steps to estimate the parameters: 1) Given the current parameter values θ^{old} , estimate the posterior distribution of the hidden variables $P(X|Y;\theta^{\text{old}})$. 2) Given the posterior $P(X|Y;\theta^{\text{old}})$ estimated from previous step, optimize parameters by maximizing the so called Q function

$$\theta = \operatorname*{argmax}_{\theta} Q(\theta) = \operatorname*{argmax}_{\theta} \mathbb{E}_{P(X|Y)} \log P(X, Y; \theta). \tag{3.21}$$

The expectation is required in (3.21) because the hidden variables are random. We need to marginalize the hidden variables when estimating the parameters. It has been shown that the expectation step maximizes a lower bound of the likelihood function $P(Y;\theta)$, and the maximization step maximizes the actual $P(Y;\theta)$ [16]. In some situations (as in our work), the hidden variables are actually the variable we are interested in, as well as the parameters, and the EM framework is still a valid method to simultaneously estimate both. When the parameters are also defined as random variables, the inference can be done by variation inference that we discussed in Section 3.5.4. In such a case, there is no difference between the hidden variables and the parameters as both are random variables.

The EM is significantly more difficult when the prior of X is a MRF. There are mainly two difficulties. First, the expectation of the $\log P(X,Y;\theta)$ is difficult to evaluate. This is in contrast to the standard Gaussian mixture model (or other mixture model), where the hidden variables at each data point are independent, and the expectation with respect to P(X|Y) can be factorized into the expectation with each individual variable. Second, the normalization constant Z (also called partitioned function) in P(X) again is intractable to evaluate, and that makes $\log P(X,Y;\theta)$ also difficult to evaluate. We will give two approximate solutions for these issues.

3.6.2 Monte Carlo EM

We replace the expectation step in EM algorithm with a sampling step. Given the current state of the parameters and the observed data, we draw samples from the posterior $P(X|Y;\theta)$, and get a series of samples $X^m, m = 1, \dots, M$, where M is the number of samples. Then we can use Monte Carlo averaging to approximate

$$\mathbb{E}_{P(X|Y)}[\log P(X,Y;\theta)] \approx \frac{1}{M} \sum_{m=1}^{M} \log P(X^m,Y;\theta), \qquad X^m \sim P(X|Y;\theta^{\text{old}}). \tag{3.22}$$

This method is indeed the Monte Carlo EM (MCEM) that was first introduced by Wei and Tanner [149]. The general idea of the MCEM is to modify the EM algorithm where the expectation in the E-step is computed numerically through Monte Carlo simulations. There are other variants of this class of methods such as stochastic EM [50], where we only draw one sample of X from the posterior $P(X|Y;\theta^{old})$ and use that in place of the expectation $\mathbb{E}_{P(X|Y;\theta)}$. While the stochastic EM asymptotically converges to the local maximum of the likelihood, we prefer the Monte Carlo EM in our work. This is because the average of $(1/M) \sum_{m=1}^{M} \log P(X,Y;\theta)$ is a better approximation of the target function $\mathbb{E}_{P(X|Y)} \log P(X,Y;\theta)$ when the number of M is large.

The numerical approximation of the expectation by sample average also introduces another source of variance, and the variance of the averaged log likelihood function depends on the number of samples M. Large M will reduce the variance, but inevitably increase computation time. To reduce the computation time on sampling, Levine and Casella proposed a method that uses important sampling instead of draw samples of X at each E-step [95]. The authors only draw samples X^m from $P(X|Y;\theta_0)$, once with an initial set of parameters θ_0 . The log likelihood function is approximated by the weighted averaging of the initial samples

$$\mathbb{E}_{P(X|Y}[\log P(X,Y;\theta)] \approx \frac{1}{\sum_{m=1}^{M} w_m} \sum_{m=1}^{M} w_m \log P(X^m,Y;\theta).$$

The original sample X^m is reused with weights w_m . At each E-step, no new samples of X are generated. Instead, the w is updated such that if the newly estimated θ increases the likelihood of a sample X^m , the w_m is accordingly increased to reflet the importance of X^m . The update of w takes less time compared to generating new samples of X. Therefore, the total computation time is less than standard MCEM.

Caffo [26] solves the convergence problem form a different view. Since the goal of the EM is to iteratively maximize the expectation of the joint likelihood, i.e., the Q function of (3.21) that is approximated by the MC sample averages of (3.22), we can use the approximated Q function as a criteria of the sample quality. For ascent-based EM, a lower bound is calculated for

$$B(\theta, \theta^{old}) = Q(\theta) - Q(\theta^{old}), \tag{3.23}$$

where θ^{old} is the parameters at previous EM iteration. If the $B(\theta)$ is positive, the new parameter is accepted and the algorithm continues. Otherwise, the θ is rejected. However, the generated samples at this iteration are kept. We then generated another MC sample, append it to the existing set of samples and obtain a new parameter estimate by using the new MC sample set. The process is repeated until $B(\theta, \theta^{old})$ reaches positive. This algorithm is different from the regular convergence of the MCMC sampling. Here, whether the MC samples are from the target distribution is not a important factor. Instead, the samples are believed good as long as the approximated Q function is maximized. Therefore, we can start the EM with a small number of samples, since the early stage of EM can often easily increase the Q function. With more EM iteration, the Q tends to converge, and we increase the sample size to guarantee the convergence of the Q. Caffo has proved that when the lower bound is positive, there is sufficient evidence to conclude that the new parameter

 θ increases the likelihood. When B is negative, the estimate of Q is deemed swamped with MC error and a large sample size is required to estimate a more accurate Q.

3.6.3 Convergence of MCEM

If the staring point of the Markov chain is poorly chosen, the burn-in period will increase dramatically. The rule of thumb is choosing the starting sample close to the center of the distribution, i.e., the mode of the probability density function (pdf). The proposal distribution of the Metropolis-Hasting sampling also has a big impact on the steps needed to reach stationary distribution. For example, the random walker, a special case of Metropolis-Hasting sampler, has a symmetric proposal distribution (either uniform, or Normal distribution) with a tunable variance parameter. Increasing this variance parameter will have larger movement, which is good to explore the whole support space, but at the risk of low acceptance rate and high correlation between samples. If the variance is too small, there is higher probability of accepting the candidates, but less opportunity to explore all modes of the target distribution, and the samples are also highly correlated. In such a case, the chain will converge too slowly.

For the convergence test of the sampling from univariate distribution, perhaps the single most popular approach is that of Gelman [64]. To use their method, we need to run multiple parallel MCMC chains with different starting points. These chains must be overdispersed initially with respect to the posterior. Each chain has length 2N and the first half of points are discarded. If we use φ_{mn} to represent the statistics of chain m at time n. The Gelman-Rubin method computes the between- and within-sequence variances of the statistics φ

$$B = \frac{N}{M-1} \sum_{m=1}^{M} (\overline{\varphi}_m - \overline{\varphi})^2,$$

$$\overline{\varphi}_m = \frac{1}{N} \sum_{n=1}^{N} \varphi_{mn}, \quad \overline{\varphi} = \frac{1}{M} \sum_{m=1}^{M} \overline{\varphi}_m,$$

$$s_m^2 = \frac{1}{N-1} \sum_{n=1}^{N} (\varphi_{mn} - \overline{\varphi}_m)^2,$$

$$W = \frac{1}{M} \sum_{m=1}^{M} s_m^2.$$

We can estimate the marginal posterior variance of the statistics φ by a weighted average of W and B

$$\widehat{Var}(\varphi|data) = \frac{N-1}{N}W + \frac{1}{N}B. \tag{3.24}$$

This estimator overestimates the marginal variance when the staring points of the chain are overdispersed, but is unbiased when $n \to \inf$. On the other hand, the within-variance W will under-estimate the variance of φ and converge to $Var(\varphi)$ when $n \to \inf$. So we can compare the value of (3.24) with W. If they are very different, that means the chain is not converged yet. The Gelman-Rubin algorithm uses the estimated potential scale reduction

$$\widehat{R} = \sqrt{\frac{\widehat{Var}(\varphi|data)}{W}},$$

which declines to 1 as $n \to \inf$.

One issue with this method is getting overdispersed starting points, one needs to have some knowledge of the pdf of interest, for example, the modes and shape of high density regions. If multiple chains all start from a single mode of the density function, they may take long steps (if ever possible) to explore other modes. In such cases, multiple chains do not help much compared to a single long chain, and the Gelman-Rubin method can not verify the convergence to the stationary distribution.

The Gelman-Rubin method is difficult to apply to sampling of high dimensional random vectors, because saving multiple independent chains will require large memory, and sampling these chains also has high computation cost.

The second test for convergence is a nonparametric test. It is applied to the single chain. Assume $\theta^{(t)}$ is the statistics derived from the chain of $1, \ldots, t$. When the chain reaches stationary, $\theta^{(t_1)}$ and $\theta^{(t_2)}$ have the same marginal distribution for any t_1 and t_2 . Given a MCMC chain $\theta^{(1)}, \cdots, \theta^{(T)}$, we can compare the empirical distributions of two half chain $(\theta_1^{(1)}, \ldots, \theta_1^{(T/2)})$ and $(\theta_2^{(T/2)}, \ldots, \theta_2^{(T)})$. The Kolmogorov-Smirnov statistics are defined as the supremum of the absolute value on the difference of two empirical distribution functions

$$K = \sup_{\eta} |F_1(\eta) - F_2(\eta)| = \frac{1}{M} \sup_{\eta} \left| \sum_{m=1}^{M} \mathbf{1}_1(\eta) - \sum_{m=1}^{M} \mathbf{1}_2(\eta) \right|,$$

where F_1 and F_2 is θ 's empirical distribution for two half chains, and $\mathbf{1}$ is an indicator function. It is noted that because of the correlation between adjacent samples in MCMC, the half chain θ is sampled in a batch mode, i.e., θ_1^m and θ_1^{m+1} are separated by a interval to make a quasi-independent chain.

Under the stationary assumption, the limiting distribution of $\sqrt{M}K$ has the cumulative distribution functions (cdf) [128]

$$R(x) = 1 - \sum_{k=1}^{\infty} (-1)^{k-1} \exp\{-2k^2x^2\}.$$

Now we can construct a hypothesis, with the null hypothesis that the two chains are from the same distribution (i.e., the MCMC chain reaches stationary). The null hypothesis is rejected if $\sqrt{M}K > K_{\alpha}$, where K_{α} is computed from $\Pr(x < K_{\alpha}) = 1 - \alpha$.

It is not straightforward to generalize the Kolmogorov-Smirnov test into higher dimension, especially for the Ising model with dimension as the number of image voxels. This is because the maximum difference between two joint cdf is not generally the same as the maximum difference of any of the complementary distribution functions. One solution is to compare the cdfs of the two samples with all possible orderings, and take the largest of the set of resulting K-S statistics. In d dimensions, there are 2^{d-1} such orderings, which is intractable for Ising model.

To test the convergence of a Gibbs sampler on a simple MRF such as an Ising model, we can instead compute the upper bound of the number of samplings. One method is to use the coupled sampled paths to study the convergence property [88]. Two coupled processes have the same transition kernel but different starting point. The coupled paths reach the same state after a certain number of iterations. The iteration is defined as the sweep of all the data points in the mode. By examining the distribution of the iterations need for coupling, convergence properties of the sampling can be established. Johnson [88] uses a 64×64 regular lattice and assumes an Ising model on the binary variables on the lattice. He tried to look for the relationship between the number of required sampling iterations and the Ising parameter β . Figure 3.13 shows that when β is small, the required number of iterations is also small. However, because the growth of the iteration number is super-exponential, a large value of β will need much more iterations to converge. When $\beta = 0.9$, the 95th and 99th quantile of the iterations distribution reached 1 million.

Gibbs [67] also gave a upper bound of the iterations of Gibbs sampling on a onedimensional Ising model. His upper bounds, however, are also a function of the square of data points, and of a tolerance ε , on the variation distance. Similar to [88], the upper bound increases fast with β . For $\varepsilon = 0.01$, $\beta = 0.5$ gives an upper bound of $128N^2$, while $\beta = 1.5$ gives $6162N^2$, where N is number of points in the lattice. Gibbs [67] noted that there is no phase transition in this one-dimensional model. For higher dimensional Ising model, his upper bounds only apply to small β , and convergence is slow when $\beta > \beta_0$, where β_0 is the reciprocal of the critical temperature. For a higher dimension of Ising model such that the number of neighbors of each data point increases, the convergence upper bounds increase even faster. Actually, the upper bound is a function of n, the number of neighbors of each node, and will increase when n increases. As a side note, Barbu et al. [7] talked about the convergence rate of Swendsen-Wang algorithm on a Potts model. Huber [83] developed a new bounding chain algorithm that can diagnose the convergence of Swendsen-Wang sampling. The number of steps to reach perfect sampling (which means convergence to stationary distribution) is in the order of $\mathcal{O}(\log |\mathcal{E}|)$, where \mathcal{E} is the set of edges. This running time applies when the temperature is far below or far above critical temperature. Cooper [33] showed that the mixing time (or convergence time) is polynomial if the number of neighbors of each node does not increase with $|\mathcal{V}|$, the size of the nodes. This is good for regular lattice where the number of adjacent nodes is fixed, regardless of image size. Compared with the super-exponential rate of increase for the iteration number in standard Gibbs sampling, Swendsen-Wang algorithm is a big improvement for convergence rate. One thing that needs to be noted is that these theoretical analysis are for the cases without external fields (data likelihood term).

3.6.4 Variational inference with EM a.k.a mean field theory

Because the variational inference is an approximate method to compute the posterior distribution of the hidden variables, it can naturally be used in the EM methods for approximating $\mathbb{E}_{P(X|Y;\theta^{old})}\log P(X,Y;\theta)$. However, depending on which parameter we want to estimate from $\mathbb{E}_{P(X|Y;\theta^{old})}\log P(X,Y;\theta)$, it may or may not be possible to use variational methods for parameter estimation. In the remaining parts of this section, we choose the definition of equation (3.15) for the convenience of variation inference approximation. Suppose the conditional distribution of P(Y|X) is Gaussian with unknown mean μ and Σ . In this case, we can estimate parameters in P(Y|X), i.e., the μ and Σ by the variational methods. Suppose we have already computed the posterior of the indicator variable x_s by using equation (3.19) and the mean field update converged, and we use $\gamma_{sk} = P(x_{sk} = 1|Y) = \mathbb{E}_{P(X|Y)}x_{sk}$, the estimates of μ and Σ take the following form

$$\hat{\mu}_k = \frac{1}{N_k} \sum_s \gamma_{sk} \cdot y_s$$

$$\hat{\Sigma}_k = \frac{1}{N_k} \sum_s \gamma_{sk} (y_s - \hat{\mu}_k) (y_s - \hat{\mu}_k)^{\top}.$$

This is essentially the same update equation for standard EM algorithm on the Gaussian mixture model. Such equivalence is necessary since the standard EM can be interpreted by the variational methods. On the other hand, the variational inference method is not able to estimate some parameters. For example, if we want to estimate the β parameter in the MRF prior, we will try to compute the $\mathbb{E}_{P(X|Y)} \log P(X,Y)$ by the following two steps.

First, we observe that the partition function Z in (3.15) is still difficult to compute due to the combinatorial nature of X, we use the pseudo-likelihood to approximate it

$$P(X) \approx \widetilde{P}(X) = \prod_{s \in \mathcal{V}} \frac{1}{Z_s} \exp\left\{\beta \sum_{r \in \mathcal{N}(s)} \langle x_s, x_r \rangle\right\}$$

$$\log \widetilde{P}(X) = \beta \sum_{r \in \mathcal{N}(s)} \langle x_s, x_r \rangle - \log Z_s$$

$$= \beta \sum_{r \in \mathcal{N}(s)} \langle x_s, x_r \rangle - \log \sum_{x_{s,k} = 1, x_{s,-k} = 0} \exp\left\{\beta \sum_{r \in \mathcal{N}(s)} \langle x_s, x_r \rangle\right\}. \tag{3.25}$$

The $\widetilde{P}(X)$ is the pseudo likelihood of X. Then we observe that first term $\beta \sum_{r \in \mathcal{N}(s)} \langle x_s, x_r \rangle$ is a linear function of x_s and x_r , so the expectation operator directly goes into the term. That is, $\mathbb{E}[\beta \sum_{r \in \mathcal{N}(s)} \langle x_s, x_r \rangle] = \beta \sum_{r \in \mathcal{N}(s)} \langle \bar{x}_s, \bar{x}_r \rangle$, where $\bar{x}_s = \mathbb{E}[x_s], \bar{x}_r = \mathbb{E}[x_r]$. However, the log of partition function $\log \sum_{x_{s,k}=1,x_{s,-k}=0} \exp \left\{\beta \sum_{r \in \mathcal{N}(s)} \langle x_s, x_r \rangle\right\}$ is a nonlinear function of x_r , so we cannot just use $\mathbb{E}[x_r]$ to replace the x_r in this term in order to compute the term's expectation. Therefore, in such case it is not possible to estimate β with the variational inference method.

There is an alternative approximation by using the traditional mean field theory for estimating the β parameter in the MRF prior. Zhang [159] proposed to use the mean field approximation in place of the expectation

$$\mathbb{E}_{P(X|Y;\theta)}[P(X,Y)] \approx \prod_{s \in \mathcal{V}} \mathbb{E}_{P(x_s|y_s)}[P(x_s|x_{-s}) \cdot P(y_s|x_s)]. \tag{3.26}$$

Note that (3.26) is not exactly the same with the Q function of EM. In the Q function, we take the logarithm first, then take the expectation of the log likelihood, while (3.26) takes the expectation first. The ML estimation of β is hence done by maximizing (3.26) with respect to β , or equivalently, the log likelihood of (3.26). Note because of the factorization, the normalization constant of $P(x_s|x_{-s})$ is tractable to compute.

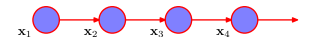


Figure 3.1: A graph model that represents the Markov chain.

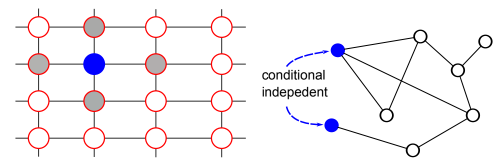


Figure 3.2: Two graphical models represented by graphs. A graphical model representing a MRF can be either a regular grid or an general graph. For the regular grid, The node in blue color is conditional independent of the white node given it's adjacent neighbors, colored gray. For the general graph example, the two nodes in blue color are conditional independent given the remaining nodes.

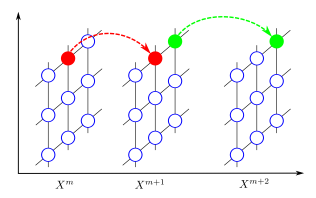


Figure 3.3: A simulation of MRF. When a new candidate w is accepted to replace current x_s , we get a new set of variables X^{m+1} that differs from the current variable X at only s. The set of variable X^m and W^{m+1} is a sample of a Markov chain, since X^{m+1} depends only on the previous X^m . Upon convergence, X will be a sample from the target distribution P(X).

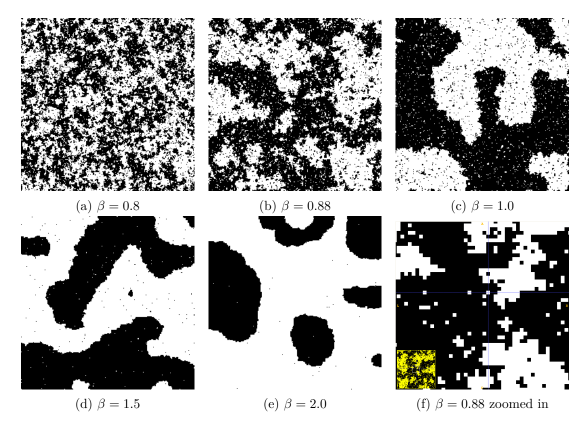


Figure 3.4: Simulating Ising model with various values of β . For each simulation, the image is initialized with random states, and then scanned 1000 times. Notice when β is small, the image is less spatially coherent. When β is large, the image has more spatial coherent regions.

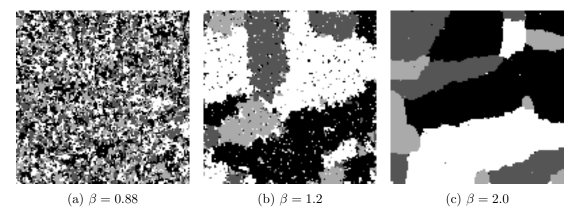


Figure 3.5: Simulating Potts model of four states with various values of β . For all simulations, the image was initialized with random states, and then was scanned 1000 times.

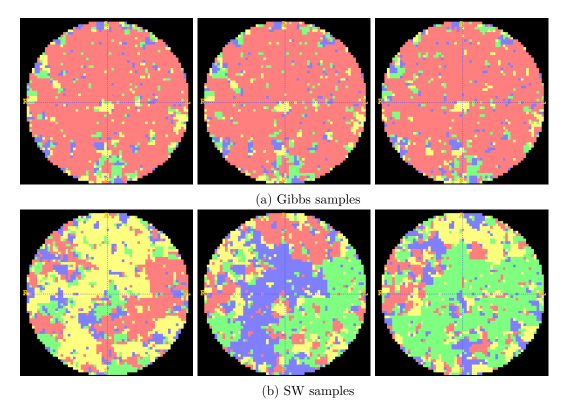


Figure 3.6: Consecutive samples of Potts model with $\beta=1.1$ using SW and Gibbs sampling. Both samplers initialize the sample image with all-zero values, have 100 burn-in sampling and then save three consecutive samples. Note for the SW samples, multiple voxel labels have been changed between the consecutive sample images. Such multiple updates speed up convergence. For Gibbs, the three sample images are similar due to the strong interactions (relatively large β) between the neighboring nodes.

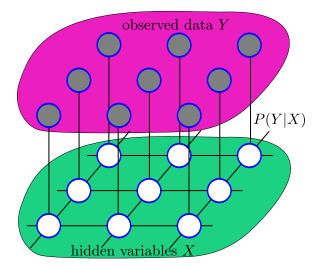


Figure 3.7: A graphical representation of the hidden Markov model(HMM). X is defined on a regular lattice graph and is given a MRF prior to represent our knowledge of the smoothness or piecewise constant. Y is the observed data that is generated from the likelihood function given the hidden X.

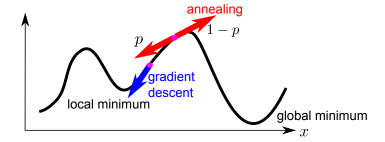


Figure 3.8: Simulated annealing samples one variable at a time. Unike coordinate descent that always moves in the gradient descent direction (blue color arrow), the SA algorithm updates the variable based on a certain probability, which depends on the difference of the function value of two configurations (red arrow).

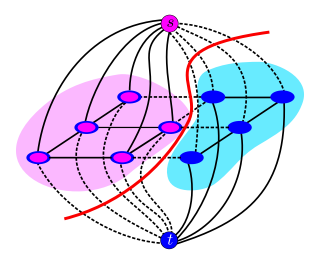


Figure 3.9: Graph cut segmentation. Each voxel is defined as a node on a graph. Neighboring voxels have edges between them with weights given by MRF. A source node s and a sink node t are added. All nodes have links to both sources and sink nodes with weights depend on the likelihood function (data term). Graph cut algorithms find a cut, i.e., a set of edges whose overall weights are minimized. In the figure, edges with solid lines are kept, and edges with dashed lines are removed after the cut. Red think links are the cut. Node is assigned to source or sink label if they are connected to either of them.

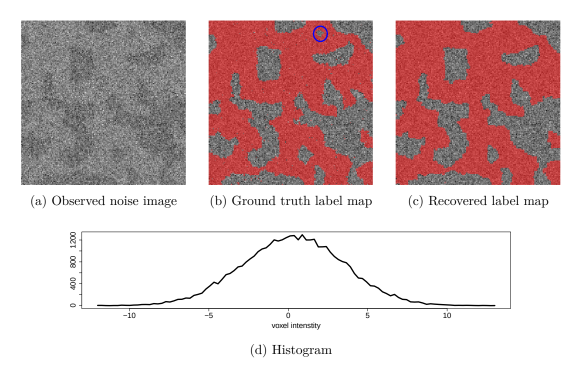


Figure 3.10: Recovering noise image by graph cut. Top row from left to right: a) observed noised image, b) ground truth label map, c) recovered label map. Bottom d) histogram of the observed image intensity. Note the region in blue circle of the true map is misclassified.

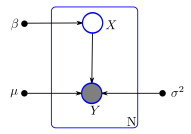


Figure 3.11: A hidden Markov model with X in MRF, and each y_s is independent Gaussian given x_s . The parameters are black dots, the hidden variables are circles, and the observed data are grayed circles. The MRF structure on X is not shown in this diagram. Instead a box is on X and Y to represent that there are N such nodes.

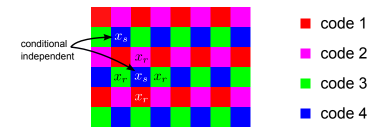


Figure 3.12: Coding scheme for parameter estimation. For four-neighbors system of two-dimensional image, the voxels are separated into four groups. The voxels in the same group are conditionally independent given other groups.

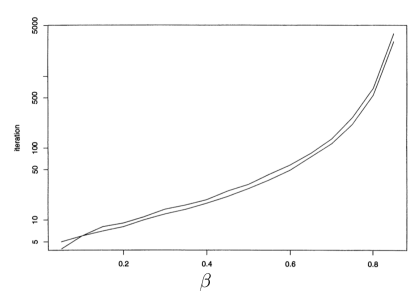


Figure 3.13: Percentile of coupling iterations for Ising model of size 64×64 . Top curve shows the 99% and bottom shows the 95% percentile from the distribution of the iterations needed for coupling, as a function of β parameter. The percentiles are estimated using 1000 repetitions of Gibbs sampling initialized with all-white and all-black value. (adapted from Johnson [88].

CHAPTER 4

FULL PAIRWISE CONNECTIVITY WITH SPATIAL COHERENCE

In this chapter, we present a new method for spatial regularization of functional connectivity maps based on MRF priors. The high level of noise in fMRI leads to errors in functional connectivity detection algorithms. A common approach to mitigate the effects of noise is to apply spatial Gaussian smoothing, which can lead to blurring of regions beyond their actual boundaries and the loss of small connectivity regions. Recent work has suggested MRF as an alternative spatial regularization in the detection of fMRI activation in task-based paradigms. We propose to apply MRF priors to the computation of functional connectivity in resting-state fMRI. Our Markov priors are in the space of pairwise voxel connections, rather than in the original image space, resulting in a MRF whose dimension is twice that of the original image. The high dimensionality of the MRF estimation problem leads to computational challenges. We present an efficient, highly parallelized algorithm on the graphics processing unit (GPU). We validate our approach on a synthetically generated example as well as real data from a resting state fMRI study.

4.1 Motivation

To show the noise level and the impact of spatial smoothing on the functional connectivity estimation, we take a seed voxel at the MNI coordinate (42, 16, 25) and compute the correlation between this seed region and every other voxel's BOLD signal in the same volume of the spatially smoothed rs-fMRI volume, and show the correlation map at Figure 4.1. The seed signal is computed by averaging the BOLD signal of voxels within 5 mm distance from the seed voxel. Following standard seed-based functional connectivity methods, we average the BOLD signal of the voxels within r=5 mm from the seed voxel, and use the averaged signal as a surrogate of the BOLD signal at this coordinate. We then compute the sample correlation between this average signal and every other voxel's BOLD signal in the full volume. For display purposes, we only show the correlation map of one slice where the

seed region is located. Figure 4.1a shows the first time point of the original BOLD signal. As expected, it is impossible to see any meaningful patterns from this image, since the functional connectivity is not represented by a voxel's intensity at a single time point, but the average effect across all time points. Figure 4.1b shows the linear correlation map with the seed regions, with the red dot giving the location of the seed region. We can see the symmetric location of the seed region on the other hemisphere shows higher intensity of correlation, suggesting these voxels are more positively correlated with the seed. In Figure 4.1c, we show the correlation map thresholded at 0.3. Since the seed region belongs to the default model network (DMN), an important component in the intrinsic functional system, we expect to see other spatially remote regions in the same network have higher correlation in 4.1c, We observe one region on the same location of the opposite hemisphere, and also one region at the prefrontal lobe remains after thresholding. However, overall the spatial patterns are hard to identify, and there are also some false-positive detections. One reason for the difficulty in detecting the connectivities is the large amount of noise in the data, and the spatial smoothing in the preprocessing step. We aim to address this problem in this chapter.

In both task-based and rs-fMRI the impact of imaging noise can be reduced by taking advantage of the spatial correlations between neighboring voxels in the image. A common approach used for instance in statistical parametric mapping (SPM)[155] is to apply a spatial Gaussian filter to smooth the signal prior to statistical analysis. However, this can lead to overly blurred results, where effects with small spatial extent can be lost and detected regions may extend beyond their actual boundaries. An alternative approach to spatial regularization that has been proposed for task activation paradigms is to use a Markov random field (MRF) prior [117, 43, 44, 152, 37], which models the conditional dependence of the signals in neighboring voxels.

In this chapter we propose to use MRF models in rs-fMRI to leverage spatial correlations in functional connectivity maps. Unlike previous MRF-based approaches, which use the neighborhood structure defined by the original image voxel grid, the neighborhoods in functional connectivity must take into account the possible relationships between spatially distant voxels. Therefore, we define the neighborhood graph on the set of all voxel pairs. This results in a Markov structure on a grid with twice the dimensions of the original image data, i.e., the pairwise connectivities for three-dimensional images results in a six-dimensional MRF. The neighborhood structure is defined so that two voxels are more likely to be connected if they are connected to each other's spatial neighbors.

We combine the Markov prior on functional connectivity maps with a likelihood model of the time series correlations in a posterior estimation problem. Furthermore, we solve for the unknown parameters of the MRF and likelihood using the EM algorithm that we discussed in Section 3.6.1. In the estimation step the posterior random field is sampled using Gibbs sampling and estimated using mean field theory, also known as variational inference.

In the next section we describe our MRF model of functional connectivity maps. In Section 4.2 we give the details of the algorithm to estimate the functional connectivity probabilities, including implementation details for the GPU solver. Finally, in Section 4.4 we demonstrate the advantages of our approach on a synthetically generated data set as well as on real rs-fMRI data.

4.2 Methods

Our framework for functional connectivity is a Bayesian approach in which we estimate the posterior distribution of the connectivity between voxels, conditioned on the fMRI data. Let $X = \{x_{ij}\}$ denote the functional connectivity map, i.e., a map denoting whether each pair of voxels i, j is connected, and let Y denote the original fMRI data, or some measurement derived from the fMRI. In this work we take Y to be the map of correlations between pairs voxel time series. The posterior distribution is then proportionally given by

$$P(X \mid Y) \propto P(X) \cdot P(Y \mid X). \tag{4.1}$$

In this work we model P(X), the prior on the connectivity map, using a MRF, and the likelihood P(Y | X) using Gaussian models of the Fisher transformed correlation values. We now give details for both of these pieces of the model.

4.2.1 Markov prior

Conventional image analysis applications of MRFs [99] define the set of sites of the random field as the image voxels, with the neighborhood structure given by a regular lattice. Because we are studying the pairwise connectivity between voxels, we need to define a MRF in the higher-dimensional space of voxel location pairs. Thus, if $\Omega \subset \mathbb{Z}^d$ is a d-dimensional image domain, then the sites for our connectivity MRF form the set $\mathcal{S} = \Omega \times \Omega$. Let $i, j \in \Omega$ be voxel locations, and let $\mathcal{N}_i, \mathcal{N}_j$ denote the set of neighbors of voxel i and j, respectively, in the original image lattice. Then the set of neighbors for the site $(i, j) \in \mathcal{S}$ is given by $\mathcal{N}_{ij} = (\{i\} \times \mathcal{N}_j) \cup (\mathcal{N}_i \times \{j\})$. In other words, two sites are neighbors if they share one coordinate and their other coordinates are neighbors in the original image lattice. This neighborhood structure will give us the property in the MRF that two voxels i, j in

the image are more likely to be connected if i is connected to j's neighbors or vice-versa. Equipped with this graph structure, S is a regular 2d-dimensional lattice. With the node set X and neighboring system N, we define a graph G that we call the *connectivity graph*. Figure 4.2 gives an illustration of how this connectivity graph is defined.

We next define a multivariate random variable $X = \{x_{ij}\}$ on the set \mathcal{S} , where each x_{ij} is a binary random variable that denotes the connectivity $(x_{ij} = 1)$ or lack of connectivity $(x_{ij} = -1)$ between voxel i and j. If $A \subset \mathcal{S}$, let X_A denote the set of all x_{ij} with $(i, j) \in A$, and let X_{-ij} denote the collection of all variables in X excluding x_{ij} . For X to be a MRF it must satisfy

$$P(x_{ij} \mid X_{-ij}) = p(x_{ij} \mid x_{\mathcal{N}_{ij}}).$$

According to the Hammersley and Clifford theorem [13], X is a Markov random field if and only if it is also a Gibbs distribution, defined as

$$P(X) = \frac{1}{Z} \exp\left(-U(X)\right),\tag{4.2}$$

where U is the energy function $U(X) = \sum_{c \in \mathcal{C}} V_c$, with potentials V_c defined for each clique c in the clique set \mathcal{C} . The partition function $Z = \sum_X \exp(-U(X))$ is a normalizing constant, where the summation is over all possible configurations of X. We use a particular form of MRF—the Ising model—a commonly used model for MRFs with binary states. In this model the energy function is given by

$$U(X) = -\beta \sum_{(ij,mn)} x_{ij} x_{mn}, \tag{4.3}$$

where the summation is over all edges (ij, mn) on the graph \mathcal{G} , i.e., all adjacent voxel pairs (i,j), (m,n) in the connectivity graph. When $\beta > 0$, this definition favors similarity of neighbors. An alternative definition is to make the MRF prior also contain the data term, such as the strength of the connection between neighboring nodes also depends on the data. Where the observed correlation value of two nodes are different, the strength of the MRF prior will be decreased. This is the conditional random field we have discussed in Section 3.4.

4.2.2 Likelihood model

We now define the likelihood model, P(Y|X), which connects the observed data Y to our MRF. Because we are interested in the functional connectivity between pairs of voxels, we compute the correlation between the time courses of each pair of voxels, and get a correlation matrix $Y = \{y_{ij}\}$. Just as in the case of the random field X, the correlation

matrix Y is also defined on the 2d-dimensional lattice S. Linear correlation is not the only choice for Y. We can use any statistic, as long as it indicates the affinity between two voxel time series. Another possibility could be frequency domain measures, such as the coherence [112], although the definition of likelihood function is not straightforward for such similarity measurement. On the other hand, simple linear correlation has the advantage that after the Fisher transformation, the correlation statistics have a well defined distribution. If the original variables y_i and y_j 's intensities across all time points are Gaussian, then the sample correlation estimated from the intensities are also Gaussian after Fisher transformation. Therefore, the conditional distribution of $P(y_{ij}|x_{ij})$ is a well-defined Gaussian distribution.

Before defining the full data likelihood, we start with a definition of the emission function at a single site $s_{ij} \in \mathcal{S}$. This is defined as the conditional likelihood, $P(y_{ij} | x_{ij})$, and is interpreted as the probability of the observed correlation, given the state of the connectivity between voxels i and j. We model the emission function as a Gaussian distribution with unknown mean and variance on the Fisher transformed correlation y_{ij} , that is,

$$P(y_{ij} \mid x_{ij} = k) = \frac{1}{\sqrt{2\pi}\sigma_k} \exp\left(-\frac{(F(y_{ij}) - \mu_k)^2}{2\sigma_k^2}\right),$$
(4.4)

where F denotes the Fisher transform. Notice that each correlation y_{ij} on site s_{ij} only depends on the latent variable x_{ij} on the same site, and does not depend on neighbors of x_{ij} . Alternative definitions exist. For example, the observed correlation data y_{ij} can also depend on the neighbors of x_{ij} , similar to the work of Geman et al. [65]. This is equivalent to adding more edges on the graphical model, and therefore the posterior inference will be more complicated. In this work, we use a simple model that has a one-to-one correspondence between x_{ij} and y_{ij} . Given the above setting, the full likelihood is given by

$$P(Y | X) = \prod_{s_{ij} \in S} P(y_{ij} | x_{ij}).$$
 (4.5)

4.3 Estimation via expectation maximization

Having defined the data likelihood and MRF prior in the previous section, we are now ready to describe the maximization of the posterior given by (4.1). For this we need to determine the model parameters, β in (4.3) and (μ_k , σ_k^2) in (4.4). Rather than arbitrarily setting these parameters, we estimate them using the EM algorithm. Exact computation of the full posterior (4.1) is intractable, due to the combinatorial number of terms in the

partition function Z. Therefore, we instead maximize the approximate posterior given by the pseudo-likelihood function [99, 13],

$$PL(X,Y) = \prod_{ij} P(x_{ij}|x_{N_{ij}})P(y_{ij}|x_{ij}).$$
(4.6)

In the E-step, the parameters are held fixed and we compute the posterior probability for each x_{ij} , and sample x_{ij} from the posterior distribution using Gibbs sampling. We then compute the expected value of each connectivity node by mean field theory. After we compute the posterior of current point x_{ij} , we update x_{ij} with its expected value $\langle x_{ij} \rangle$. The equivalence of posterior probability and the expectation of binary random variable is discussed in Section 3.6.4.

In the M-step, the complete data $\{\langle X \rangle, Y\}$ are available, and the parameters can be estimated by maximizing the joint pseudo-likelihood given by (4.6) using Newton's method. After several iterations of this EM algorithm, we get parameters as our MAP estimates. During parameter estimation, the joint likelihood can be factorized into two separate terms of the prior and the conditional likelihood, and the parameter β can be estimated by maximizing the MRF prior (equivalently the pseudo-likelihood) by Newton's method. The estimation of μ and σ^2 can be done in closed from just like the standard Gaussian mixture model, as shown in Section 3.6.

4.3.1 GPU implementation

The whole algorithm involves updating a high dimensional connectivity matrix X iteratively, and hence it has high computation cost. We designed a parallel Markov random field updating strategy on a graphics processing unit (GPU). The algorithm takes only a few minutes compared with more than 1 hour on the CPU counterpart.

To fit the algorithm into GPU's architecture, we use some custom strategies. First, because GPU only support three-dimensional array, we need to reshape X and Y defined originally on a higher dimensional graph by linear indexing their original subscripts. This is especially difficult for brain fMRI data because the gray matter voxels reside in an irregular three-dimensional lattice. Specific data structures are used for mapping between original voxel subscripts and their linear index i and j. Second, to update each site of the MRF in parallel, we have to make sure a site is not updated simultaneously with its neighbors, otherwise the field tends to be stuck in a checkerboard-like local maximum, as indicated by Figure 4.3. Our strategy is to divide all the sites of the field into several subgroups, such that a site is not in the same subgroup with its neighbors. We then can update the

subgroup sequentially, while the data in subgroups are updated simultaneously. The whole procedure is summarized in Algorithm 5.

It is noted this updating scheme is closely related to the coding method of parameter estimation of MRF that we discussed in Section 3.6. Both approaches split the variables into multiple subgroups. They differ in that for coding methods, the separation is used to get independent variables within the subgroup, while here the voxels in subgroups are independent so we can update them in parallel.

4.4 Results

In this section, we give the experimental results for both simulated data and real fMRI data. We compare the results of various methods on simulated data with the ground truth to show the proposed method's performance. In the real data experiment, we compare the results with the functional connectivity map of other literatures.

4.4.1 Synthetic data

We first construct a synthetic data set consisting of a 100×1 one-dimensional image, with each pixel a 300-point time course signal. The one-dimensional image will guarantee the connectivity map be in two-dimensional space and can be visualized easily. The time course was constructed with a baseline DC signal of 800, plus additive Gaussian noise of zero mean and variance 50. We then added a sine wave signal of frequency 0.2 and amplitude 20 to two distant regions of the image. The goal is to detect the connectivity between these two distant regions. The true connectivity value will be one between those pixels containing the sine wave signal, otherwise it is defined as -1 for lack of connectivity between signal and noise, and between noise time series. The true connectivity map is shown in Figure 4.4a.

To compare our MRF model with conventional Gaussian blurring of the correlation map, we applied both approaches to the synthetic data (Figure 4.4). On the correlation map in the top row, we see smoothing does remove noise and results in a correlation map that looks more like the true connectivity map. However, it also creates several errors, most noticeably false positives around the diagonal (Figure 4.4e). This is because without prior knowledge of the scale of the patterns we are interested in, the choice of smoothing kernel size is arbitrary. Therefore, a large kernel size will enforce the signals of many voxels to look similar, and the following correlation map will have more false-positive detections. Figure 4.4f shows the proposed MRF method better detects the true connectivity regions while removing most false positives.

Algorithm 5: MAP estimation by EM

Require: Sample correlation matrix Y.

Init posterior matrix by maximizing conditional likelihood $P(y_{ij}|x_{ij})$.

while
$$\Delta\{\beta,\mu,\sigma^2\} > \varepsilon$$
 do

E step:

- (a) Based on the current parameters, compute the posterior by (4.6).
- (b) Repeatedly Do Gibbs Sampling until the field stabilizes.
- (c) Based on current value of x_{ij} , iteratively compute the mean field for all nodes in S until the field is stable.

M step:

(d) With complete data $\{X, Y\}$, estimate β , μ and σ^2 by maximizing (4.6).

end while

return posterior matrix X.

4.4.2 Resting-state fMRI

Next we tested our method on real data from healthy control subjects in a resting-state fMRI study. BOLD EPI images (TR= 2.0 s, TE = 28 ms, GRAPPA acceleration factor = 2, 40 slices at 3 mm slice thickness, 64 x 64 matrix, 240 volumes) were acquired on a Siemens 3 Tesla Trio scanner with 12-channel head coil during the resting state, eyes open. The data was motion corrected by SPM software and registered to a T2 structural image. We used a gray matter mask from an SPM tissue segmentation so that only gray matter voxels are counted in the connectivity analysis. We do *not* spatially smooth the data, in order to see the benefit of replacing spatial smoothing with our MRF method. Before computing the correlations, the time series at all voxels are linearly detrended by least squares regression.

Figure 4.5 compares the real data results using no spatial regularization, Gaussian smoothing, and the proposed MRF model. Though the posterior connectivity of the MRF is computed between every pair of voxels within a slice, for visualization purposes, only the posterior of the connectivity between one voxel and the slice is shown. We chose to visualize the connectivity to a voxel in the posterior cingulate cortex (PCC) because this is known to be involved in the default mode network [125], with connections to the medial prefrontal cortex (MPFC). The results show that Gaussian smoothing is able to remove noise, but is unable to find a clear connection between the PCC and the MPFC. Our proposed MRF model (4.5c and 4.5f) is able to remove spurious connections, and also clearly shows a connection to the MPFC.

To show the strongest connections found by each method, Figure 4.6 shows the thresholded connectivity maps overlaid on T2 structural image. Images in the first two columns are thresholded such that the top 5% voxel correlations are shown. For the MRF in 4.6c and 4.6f, the MAP estimate is shown.

4.5 Discussion

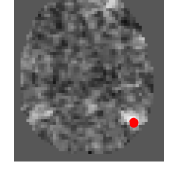
Functional connectivity is a key step of understanding the human brain's functional networks, since it contains all the information of how each functional region interacts with other regions. There are multiple ways of modeling functional systems. One is defining some regions of interest (ROI) as the nodes on the graph, and estimate the connectivities among these ROIs. The ROI definition has no standard to follow. One can either define them according to the anatomical structure, or by a parcellation on the functional data. The other way of representing functional networks is to cluster the full brain volume (indeed the gray matter voxels) into small regions. Regions in the same cluster are highly connected even if they are spatially remote. Our method outputs a posterior matrix that includes

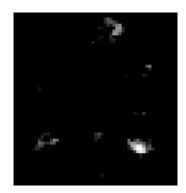
all the connectivity information between each pair of voxels. One potential usage of the full connectivity matrix is exploring the connectivity given a single seed coordinate. That is, given the seed, one can explore the posterior connectivity between the seed voxel and the full brain volume in real time. Such full connectivity will be dynamically shown with the changing seed's location. Such a dynamic, and potentially user-interactive functional connectivity exploratory concept has been partly realized by Yeo et al. [157], though they did not use MRF to model the spatial constraints of the connectivity variables. Instead, Yeo et al. use a large group of subjects and obtained a highly consistent population functional connectivity network. The network can be visualized given a seed region and also the change of the network patterns when the seed changes.

On the optimization side, the MCEM algorithm, and the wider class of sampling methods are potentially good candidates for the application of GPU. Although theoretically the computation on GPU needs independence between the data points, the order of voxels being sampled have no effect on the quality of the samples once the sampling reaches stationary distribution. Therefore, even at each scan, the order, or the schedule of the data points change in an undetermined way within the GPU cores, we can still get samples in the target distribution give enough burn-in time. Alternative optimization methods other than MCEM exist, such as graph cut. However, in our experiments with graph cuts, we found the local finer patterns are sometimes ignored by graph cut. The estimated hidden variables are accordingly mostly blob-like patterns with small boundary-area ratio. So graph cut optimization is more suitable for the recognition of blob-like patterns. Such preference does not necessarily reflect the true structure in our fMRI data. As a result, the estimation of the parameters β in (4.3) is also biased towards a larger value than the true value, since the true X is not as smooth as the estimated one. Therefore, we choose to use MCEM in our EM procedure for unbiased estimation of parameters.

One weakness of the pairwise connectivity estimation routine is lack of visualization for all the functional networks as spatial maps. Only when the input image is one-dimensional, such as the synthetic data example we used in Section 4.4.1, can the connectivity map be visualized on a two-dimensional plane. In general three-dimensional volume image as input, the connectivity matrix is in a six-dimensional space and cannot be shown in a straightforward way. Once the posterior connectivity matrix is computed, we still need to give a seed region in order to map all the functional systems that are connected to the seed. We will discuss methods of estimating multiple functional networks and show them in a single volume image in the next chapter.







(a) One slice of a resting-state fMRI dataset at z=25.

(b) Correlation map between a seed and the current slice.

(c) The correlation map thresholded at 0.3.

Figure 4.1: An example of correlation map with a seed in the default model network on the rs-fMRI data set.

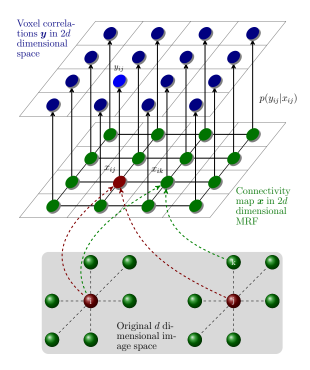


Figure 4.2: MRF prior of the connectivity variables. Each node of the graph represents a pairwise connectivity variable between voxel i and j. An edge is added between two nodes x_{ij} and x_{ik} if k is the neighbor of voxel j. The graph where the MRF is defined is twice the dimensions of the original image domain, i.e., six dimensions. Given the hidden variable X, the observed sample correlation values are assumed to be generated from a Gaussian distribution with unknown parameter $\mathcal{N}(y_{ij}|x_{ij};\mu,\sigma^2)$.

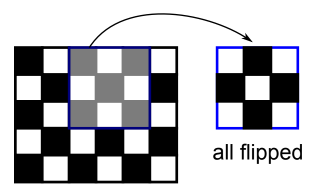


Figure 4.3: Ideally the update of each voxel is independent of other voxels in order to be used on the GPU. In our Gibbs sampling, although the sampling of each voxel depends on its neighbors, the order of the voxels being updated does not matter. Upon convergence, the image will be a sample of the target Gibbs distribution. However, numerically, the sampling tends to be stuck in this local minimum of checkerboard image. At the current state, each voxel has a neighbor with a different state, and the sampling flips the color of all voxels in the next stage.

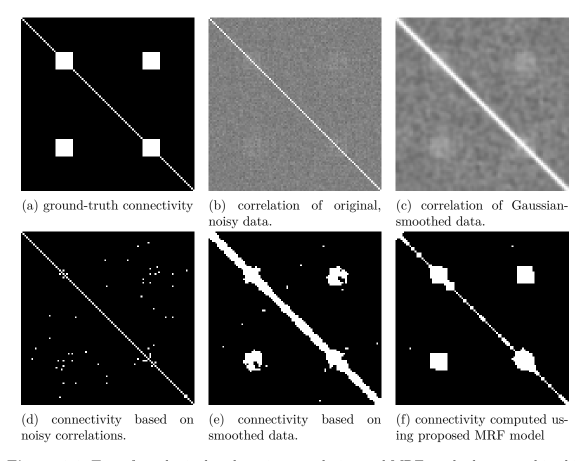


Figure 4.4: Test of synthetic data by using correlation and MRF method proposed in this work. (a) Ground truth connectivity map. (b) Connectivity based on smoothed data. (c) Correlation of Gaussian-smoothed data. (d) Connectivity based on noisy correlations. (e) Connectivity based on smoothed data. (f) Connectivity computed using proposed MRF model.

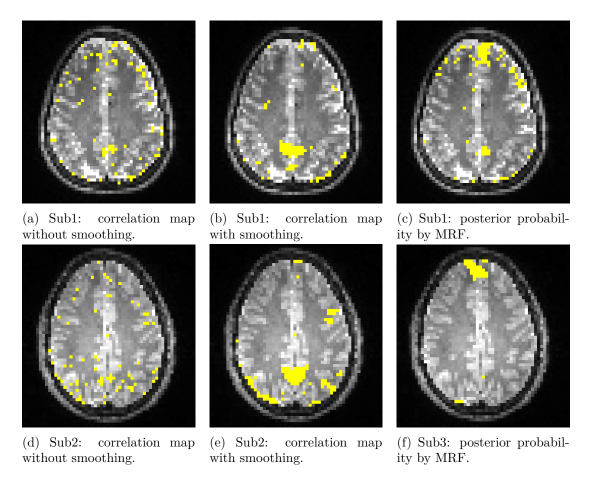


Figure 4.5: Threshold correlation mappind Posterior Connectivity map between seed voxel and the current slice, overlaid to T2 image. (a) Subject 1: correlation without smoothing. (b) Subject 1: correlation with smoothing. (c) Subject 1: posterior estimated from MRF. (d) Subject 2: correlation without smoothing. (e) Correlation without smoothing. (f) Posterior estimated from MRF.

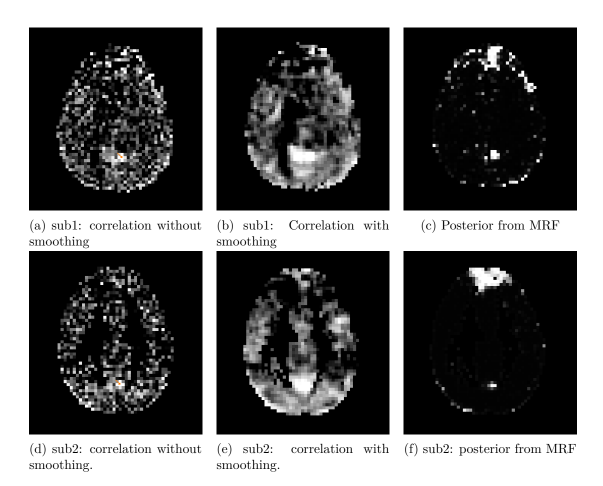


Figure 4.6: Correlation map and posterior connectivity map between seed voxel and slice containing the seed. (a) Subject 1: correlation without smoothing. (b) Subject 1: correlation with smoothing. (c) Subject 1: posterior estimated from MRF. (d) Subject 2: correlation without smoothing. (e) Correlation without smoothing. (f) Posterior estimated from MRF.

CHAPTER 5

CONSISTENT AND SPATIALLY COHERENT FUNCTIONAL NETWORKS

The pairwise functional connectivity estimation method in Chapter 4 estimates the connections between each pair of voxels, and outputs a symmetric matrix with element (i,j), the connectivity of voxel i and j. However, in order to explore functional networks, we still need a seed. Moreover, only those regions connected to the selected seed region can be identified. If a seed is incorrectly chosen, it may fall outside of the functional networks that we are interested in, the resulting network map will not be informative. In this chapter, we propose a new data-driven method to partition the brain's gray matter into disjoint partitions of functional networks. Unlike the previous chapter, the proposed algorithm does not require specification of a seed, and there is no ad hoc thresholding or parameter selection. The algorithm identifies all of the functional network maps in a single run. To achieve the above properties, we define the model in the original image space and cluster the voxels with higher connectivity into the same class. This is indeed an image segmentation problem, or an unsupervised clustering problem in data mining. The proposed approach is, in spirit, similar to the functional parcellation work that has been proposed by Yeo et al. [157].

We make a natural assumption that functionally homogeneous regions should be spatially coherent. Our method incorporates spatial information through a MRF prior on voxel labels, which models the tendency of spatially-nearby voxels to be within the same functional network. Here we use a MRF to model the network label's spatial soft constraint, such that the network component maps are spatially coherent with a piecewise constant labeling.

The BOLD time course at each voxel is first normalized to zero mean and unit norm, which results in data lying on a high-dimensional unit sphere. We then model the normalized time-series data as a mixture of von Mises-Fisher (vMF) distributions [6]. Each component of the mixture model corresponds to the distribution of time series from one functional network. Solving for the parameters in this combinatorial model is intractable, and we

therefore use a MCEM algorithm, which approximates the expectation step using Monte Carlo integration. The stochastic property of MCEM makes it possible to explore a large solution space, therefore it performs better than a standard mode approximation method such as iterated conditional modes (ICM). Finally, we demonstrate on real fMRI data that our method is able to identify visual, motor, salience, and default mode networks with considerable consistency between subjects.

The vMF distribution was first introduced in the machine learning community by Banerjee et al. [6] to address the parametric clustering problem on the dataset of documents. In such data, the similarity of two data samples is better represented by the inner product of normalized feature vectors. By normalization, the feature vector has zero mean and unit variance, which is equivalent to being projected onto a high-dimensional sphere. Original usage of the inner product as distance metric is by Dhillon and Modha's spkmeans algorithm [45], where the cosine similarity is used for clustering. When the features have high dimensions, the spkmeans is superior to standard K-Means algorithm in terms of speed and classification accuracy. The extension of spkmeans to vMF [6] is similar to the extension of Gaussian mixture model to the standard K-Means, except that vMF has a simplified definition of variance (or, precision) such that the distribution is isotropic. The vMF distribution is indeed a generalization of the lower dimensional von Mises distribution, which is discussed by Mardia [103] in the context of directional statistics.

5.1 Hidden Markov models of functional networks

We use a Bayesian statistical framework to identify functional networks of the gray matter in fMRI data. We formulate a generative model, which first generates a spatial configuration of functional networks in the brain, followed by an fMRI time series for each voxel based on its network membership. We employ an MRF prior to model network configurations, represented by the hidden network label variables. Given a label, we assume that the fMRI time series, normalized to zero mean and unit norm, are drawn from a von Mises-Fisher distribution.

Let \mathcal{V} be the set of indices for all gray-matter voxels. We assume that the number of networks L is a known free parameter. Let $\mathcal{L} = \{1, 2, \dots, L\}$ be the set of labels, one for each network. We denote a label map for functionally-connected networks as a vector $X = (x_1, \dots, x_N), x_s \in \mathcal{L}$. Let $\mathcal{X} = \mathcal{L}^N$ be the set of all possible X's configurations.

5.1.1 A Markov prior model

Functional networks should consist of few, reasonably-sized, possibly distant regions. We model such networks X using a special case of MRF model that we discussed in Chapter 3, i.e., the Potts [99]:

$$P(X) = \frac{1}{Z} \exp \left\{ -\beta \sum_{(r,s)\in\mathcal{E}} \psi(x_r, x_s) \right\},\,$$

where the function ψ takes 1 if its argument is not equal and 0 otherwise; (r, s) is the set of voxel pairs that are spatial neighbors on the graph; $\beta > 0$ is a model parameter controlling the strength of label-map smoothness; Z is a normalization constant that is the sum of P(X) over all possible configuration of X. The Markov-Gibbs equivalence [99] implies that the conditional distribution of x_s at site s is:

$$P(x_s|X_{-s}) = P(x_s|X_{\mathcal{N}_s}) = \frac{\exp\left\{-\beta \sum_{r \in \mathcal{N}_s} \psi(x_s, x_r)\right\}}{\sum_{l \in \mathcal{L}} \exp\left\{-\beta \sum_{r \in \mathcal{N}_s} \psi(x_r, l)\right\}},$$
(5.1)

where X_{-s} is the collection of all variables in X excluding site s. The neighborhood is the usual six adjacent voxels, which does not overly smooth across boundaries. Previous works [149, 43] have demonstrated the advantages of MRFs over Gaussian smoothing in preserving segment boundaries.

5.1.2 Likelihood model

We observe that in order to make the analysis robust to shifts or scalings of the data, one typically normalizes the time series at each voxel to zero mean and unit length. This results in the data being projected onto a high-dimensional unit sphere. After normalization, the sample correlation between two time series is equal to their inner product, or equivalently, the cosine of the geodesic distance between these two points on the sphere. Thus, we reformulate the problem of finding clusters of voxels with high correlations to the problem of finding clusters with small within-cluster distances on the sphere. Figure 5.1 shows this equivalence.

In the previous chapter, the observed data are the linear correlations between a prior of voxels BOLD signal. Since the correlation is a scalar, the emission function, i.e., the conditional probability of P(Y|X) can be easily modeled by the Gaussian distribution once the correlation has been Fisher-transformed. In the model of this chapter, the observed data are the BOLD signal itself, so we need a multivariate distribution to model its conditional probability. We use the notation $Y = \{(y_1, \ldots, y_N) | y_s \in S^{p-1}\}$ to denote the set of normalized time series. Observe that given $X \in \mathcal{X}$, the random vectors $y_s, \forall s \in \mathcal{V}$ are

conditional independent. Thus, the likelihood $\log P(Y|X) = \sum_{s \in \mathcal{V}} \log P(Y_s|x_s)$. We model the emission function $P(y_s|x_s)$ using the von Mises-Fisher (vMF) distribution

$$f(y_s; \mu_l, \kappa_l | x_s = l) = C_p(\kappa_l) \exp(\kappa_l \mu_l^\top y_s), \quad y_s \in S^{p-1}, \quad l \in \mathcal{L}.$$
 (5.2)

For the cluster labeled l, μ_l is the mean direction, $\kappa_l \geq 0$ is the concentration parameter, and the normalization constant $C_p(\kappa)$ is given by

$$C_p(\kappa) = \frac{\kappa^{\frac{p}{2}-1}}{((2\pi)^{\frac{p}{2}}I_{\frac{p}{2}-1}(\kappa))},$$

where I_{ν} denotes the modified Bessel function of the first kind with order ν . The larger the κ , the greater is the density concentrated around the mean direction. When $\kappa = 0$, $f(y_s; \mu, \kappa)$ reduces to the uniform distribution on S^{p-1} . If $\kappa \to \inf$, y_s will be in a point density. Since (5.2) depends on y only by $\mu^{\top}y$, the vMF distribution is unimodal and rotationally symmetric around μ .

In the Bayesian framework, we also define distributions on parameters. We assume that $\forall l \in \mathcal{L}, \kappa_l \sim \mathcal{N}(\mu_\kappa, \sigma_\kappa^2)$ with hyperparameters μ_κ and σ_κ^2 that can be set empirically. This prior enforces constraints that the clusters should not have extremely high or low concentration parameters. We empirically tune the hyperparameters μ_κ and σ_κ^2 and have found the results to be robust to specific choices of the hyperparameters.

5.1.3 Monte Carlo EM

The model of the functional network variables can be illustrated by Figure 5.2. Given the definition of the prior and likelihood function, our goal is the statistical inference of the posterior probability of X given the observed data y. Because of both the hidden variables X and the model parameters μ, κ, β are unknown, we use EM method to estimate them in an iterative way. To estimate the model parameters and the hidden labels, we use the MCEM [149] algorithm. The standard EM algorithm maximizes the expectation of the log-likelihood of joint PDF of Y and the hidden variable X with respect to the posterior probability P(X|Y), i.e., $\mathbb{E}_{P(X|Y)}[\log P(X,Y;\theta)]$. The combinatorial number of configurations for X makes this expectation intractable. Thus, we use Monte Carlo simulation to approximate this expectation as

$$\widetilde{Q}(\boldsymbol{\theta}; X, Y) \approx \frac{1}{M} \sum_{m=1}^{M} \log P(X^m; \beta) + \log P(Y|X^m; \boldsymbol{\theta}_L),$$
 (5.3)

where X^m is a sample from P(X|Y), $\boldsymbol{\theta}_L = \{\mu_l, \kappa_l : l \in \mathcal{L}\}$ is the parameter vector of the likelihood, and $\boldsymbol{\theta} = \{\beta, \boldsymbol{\theta}_L\}$ is the full parameter vector of the model. Computing the MRF

prior in (5.3) is still intractable due to the normalization constant, and we instead use a pseudo-likelihood approximation [99], which gives

$$\widetilde{Q} \approx \frac{1}{M} \sum_{m=1}^{M} \sum_{s \in \mathcal{V}} \log P(x_s | x_{\mathcal{N}_s}; \beta) + \frac{1}{M} \sum_{m=1}^{M} \sum_{s \in \mathcal{V}} \log P(y_s | x_s; \boldsymbol{\theta}_L) = \widetilde{Q}_P + \widetilde{Q}_L.$$

We use \tilde{Q}_P to denote the log-pseudo-likelihood of the prior distribution, and use \tilde{Q}_L to denote the log-likelihood distribution. Now the prior term \tilde{Q}_P is in a tractable form for evaluation, and we can estimate β by optimization of \tilde{Q}_P with a Newton-Raphson method, initialized by $\beta = 0$. Because of the separation of the parameters in \tilde{Q}_P and \tilde{Q}_L , we can estimate the parameter θ_L by optimizing Q_L . There is an approximated closed form solution for this estimation, and we will discuss it in Section 5.1.5.

5.1.4 Sampling from the posterior

Given the observed data Y and parameter value $\theta = \{\beta, \theta_L\}$, we sample from the posterior distribution $P(X|Y;\theta)$ using Metropolis sampling. We define the posterior energy, which is to be minimized, as the negative log of the posterior $P(x_s|Y_s)$. Thus, Bayesian rule implies:

$$U(x_s = l|X) = \beta \sum_{r \in \mathcal{N}_s} \psi(x_s, x_r) - \log C_p(\kappa_l) - \kappa_l \mu_l^{\top} y_s + \text{const},$$
 (5.4)

which is the sum of the prior energy, the conditional energy, and a parameter-independent quantity. Then, given a current configuration X^m , Metropolis sampling generates a new candidate label map \boldsymbol{w} as follows: (i) Draw a new label l' at site s with uniform distribution; W has value l' at site s, with other sites remaining the same as X^m ; (ii) compute the change of energy $\Delta U(W) = U(W|Y) - U(X^m|Y) = U(x_s = l'|Y) - U(x_s = l|Y)$; (iii) accept candidate W as X^{m+1} with probability min $(1, \exp\{-\Delta U(W)\})$; (iv) after a sufficiently long burn-in period, generate a sample of size M from the posterior distribution P(X|Y). This is indeed the Metropolis sampling algorithm (Algorithm 1) that we have introduced in Chapter 3.

5.1.5 Parameter estimation

In order to estimate μ and κ of the vMF distribution, we need to maximize \widetilde{Q}_L with the constraint $||\mu_l|| = 1$ and $\kappa > 0$. For one sample label map, the maximum likelihood maximization of the mean vector μ_l for label l is [6, 46]

$$\mu_l = \underset{\mu_l}{\operatorname{argmax}} \prod_{s \in \mathcal{V}_l} P(y_s; \mu_l, \kappa_l | X)$$
$$= \underset{\mu_l}{\operatorname{argmax}} \sum_{s \in \mathcal{V}_l} \log P(y_s; \mu_l, \kappa_l | X)$$

$$= \underset{\mu_l}{\operatorname{argmax}} \sum_{s \in \mathcal{V}_l} N \log C_p(\kappa_l) + \kappa_l \mu_l^{\top} R_l, \tag{5.5}$$

where $\mathcal{V}_l = \{s \in \mathcal{V} : x_s = l\}$ is the set of data points in cluster l, and $R_l = \sum_{s \in \mathcal{V}_l} y_s$. We maximize (5.5) with the constraints $\mu_l^{\top} \mu$ by introducing a Lagrangian multiplier and obtain

$$\hat{\mu}_l = R/\|R_l\|. \tag{5.6}$$

In MCEM, instead of having one label map, we have M maps sampled from P(X|Y). The M sample maps are not independent since they were obtained in a consecutive time point. But for the purpose of parameter estimation, we can safely assume the independence and pool them into same dataset, where we extract the subset of label l to estimate μ_l . We can estimate μ as

$$R_l = \sum_{m=1}^{M} \sum_{s \in \mathcal{V}_l} y_s, \qquad \hat{\mu}_l = \frac{R_l}{||R_l||},$$
 (5.7)

We have no *a priori* knowledge for μ_l , so a maximum likelihood estimation in (5.7) is the best we can do. For κ_l we maximize the posterior distribution $P(\kappa_l|Y,X^1,\ldots,X^M)$. Since \widetilde{Q}_P is not dependent on κ , we maximize $\widetilde{Q}_L(\kappa_l) + \log P(\kappa_l;\mu_\kappa,\sigma_\kappa^2)$ and get [6]

$$A_p(\hat{\kappa}_l) + \frac{\hat{\kappa}_l - \mu_{\kappa}}{N_l \sigma_{\kappa}^2} = R_l, \tag{5.8}$$

where $A_p(\hat{\kappa}_l) = I_{\frac{p}{2}}(\hat{\kappa}_l)/I_{\frac{p}{2}-1}(\hat{\kappa}_l)$ and $N_l = |\mathcal{V}_l|$ is the number of data points in cluster l. Because (5.8) contains the ratio of two modified Bessel functions, an analytic solution is unavailable and we have to resort to a numerical solution. We use Newton's method for solving $g(\hat{\kappa}_l) = A_p(\hat{\kappa}_l) - (\hat{\kappa}_l - \mu_{\kappa})/(N_l \sigma_{\kappa}^2) - R_l = 0$. The choice of initial value for Newton's algorithm depends on the strength of the prior on κ_l (i.e., the σ_{κ} value). For a noninformative prior, $\hat{\kappa}_l = (pR_l - R^3)/(1 - R^2)$ is a good a good initial value [6]. For a strong prior, a reasonable initial value is the current value of κ_l .

To estimate β , we again rely on Newton's method to find the solution numerically. The derivatives $\partial \tilde{Q}_P/\partial \beta$ and $\partial^2 \tilde{Q}_P/\partial \beta^2$ for the pseudo-likelihood approximation of the MRF prior are easily computed. With all the settings above, we give the steps of the E step and M step iterations in Algorithm 6.

Given the methods for sampling and parameter estimation, we estimated the hidden MRF model by iteratively using (i) MCEM to learn model parameters and (ii) using ICM to compute optimal network labels. In the expectation (E) step, we draw samples from the posterior P(X|Y), given current estimates for parameters $\boldsymbol{\theta}$. In the maximization (M) step, we use these samples to update estimates for the parameters $\boldsymbol{\theta}$.

Algorithm 6: MCEM-ICM Algorithm for Hidden-MRF Model Estimation

```
input : Preprocessed 4D fMRI data; number of clusters
output: Labeled functional network map
Initialization: Run k-means clustering a few times and choose \mathbf{z} with the smallest
sum-of-square errors; estimate \theta_L and set \beta to a small value;
while MCEM not converged do
    E step: Given current \theta, for m \leftarrow 1 to M do
       foreach s \in \mathcal{V} do Draw sample x_s^m from P(x_s|Y_s) using (5.4);
   M step: Given (X^1, \ldots, X^M), estimate \beta and \theta_L;
   Estimate labels with ICM using the current estimates for \beta and \boldsymbol{\theta}_L;
```

5.2 Experiment results

In this section, we give preliminary test results on simulated fMRI dataset, in order to show its accuracy under large noise. We then apply our MCEM method on the *in vivo* data.

5.2.1 Synthetic data

We first simulate low-dimensional time series (two-dimensional 64×64 image domain; three time points, for visualization on sphere S^2) to compare the (i) proposed method using MCEM with (ii) the mode-approximation approach that replaces the E step in EM with a mode approximation. We simulate a label map by sampling from a MRF having $\beta = 2$ and number of labels L=4. Given the label map, we simulate vMF samples on the sphere S^2 . The method we used to simulate samples from vMF distribution is from Dhillon [46] and Wood [151]. Figure 5.3 gives the simulated data and the estimation by the two methods. We note the simulated network label map is piecewise constant, with occasional small regions scattered between large patches. This is similar to the estimated functional network maps from the real data. Although there are only three time points, it is difficult to visualize the dynamic change of the voxel intensity over time, so we choose to show only the volume at the first time point on the second image of Figure 5.3. Both the ICM and MCEM model are initialized with a K-Means clustering. We observe that ICM easily converges to a local minimum of the energy function within a few iterations, and its estimated labels are less accurate compared with our MCEM approach. The MCEM solution is close to the ground truth, while the mode-approximation solution is stuck in a local maximum.

5.2.2 Real rs-fMRI

We evaluated the proposed method on real data obtained from healthy control subjects, in a resting-state fMRI study. BOLD EPI images (TR = 2.0 s, TE = 28 ms, 40 slices at 3 mm slice thickness, 64 x 64 matrix, 240 volumes) were acquired on a Siemens 3 Tesla Trio scanner. The data was preprocessed in SPM, including motion correction, registration to T2 and T1 structural MR images, spatial smoothing by a Gaussian filter, and masked to include only the gray-matter voxels. We used the conn software to regress out signals from the ventricles and white matter, which have a high degree of physiological artifacts. A bandpass filter was used to remove frequency components below 0.01 Hz and above 0.1 Hz. We then projected the data onto the unit sphere by subtracting the mean of each time series and dividing by the magnitude of the resulting time series. We then applied the proposed method to estimate the functional network labels with the number of clusters set to L=8.

Figure 5.4 shows the optimal label maps, produced by the proposed method for 3 of all 16 subjects in the dataset. We note that among the eight clusters, one cluster, with the largest κ value and largest number of voxels, corresponds to background regions with the weakest connectivity and is not shown in the figure. Among the clusters shown, we can identify the visual, motor, dorsal attention, executive control, salience, and default mode networks (DMN) [125]. Four networks: the visual, motor, executive control, and DMN, were robustly found across all subjects. More variability was found in the dorsal attention network (notice that it is much larger in subject 3) and salience network (notice that it is missing in subject 2). We found that changing the number of clusters, although leading to different label maps, preserves the four robust networks. For instance, we also ran the analysis with the number of clusters set to 4 or 6 (results not shown) and were able to recover the same four robust networks.

The next experiment compares our results with ICA. A standard ICA toolbox (GIFT; mialab.mrn.org) was applied on the same preprocessed data of each subject independently, which we call "Individual ICA". We also applied standard Group ICA, using all data from the 16 subjects simultaneously. In both ICA experiments the number of components are set to 16. The component maps are converted to z score and thresholded at 1. For each method we computed an overlap map for each functional network by adding the corresponding binary label maps of all 16 subjects. The results in Figure 5.5 show our method can detect the motor, attention, and visual network with accuracy comparable with Group ICA. Besides, our method also detects DMN with posterior cingulate cortex (PCC) and medial prefrontal cortex (MPFC), while Group ICA split the DMN into two components, one with the MPFC and another with the PCC (not shown).

To see the consistency of the label map between subjects for all three methods, we look at each method's overlapped label map and count the number of voxels whose value are greater than 8. Table 5.1 shows that our method exhibits better consistency than both Individual and Group ICA.

5.3 Discussion

We proposed a segmentation algorithm and applied it on single subject rs-fMRI data. The output of the algorithm is a label map, in which voxels of the same functional network are assigned the same labels. The label of each voxel has a spatial dependency on its neighbors, and we use a MRF to model the interdependency, and accordingly define a prior distribution of the label map. The posterior inference is an intractable problem due

to the complex dependency of multiple variables. We use MCEM to draw samples from the posterior distribution and use the samples for parameter estimation. The Metropolis sampling uses a uniform distribution as a proposal distribution. Because there are L possible labels in this uniform distribution, there is a higher chance that the uniform does not propose the best label value during the first few scans. When L is large, we accordingly need more scans for the proposal distribution to catch the best label. Therefore, the burn-in times are longer than the binary case in Chapter 4.

Table 5.1: The number of voxels with value greater than 8 in the overlapped label map.

	DMN	Motor	Attention	Visual
MCEM	5043	7003	3731	5844
Individual ICA	114	167	228	134
Group ICA	3075	5314	3901	3509

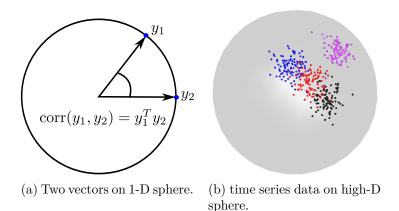


Figure 5.1: Data points with Von Mises-Fisher distribution.

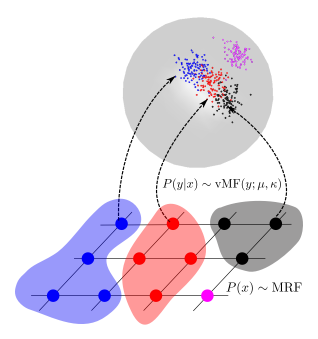


Figure 5.2: A generative model of the functional network. The network variable X is a multivariate variable defined on a MRF. Given X, Y is seen as being generated from a vMF distribution whose parameter μ and κ are functions of X.

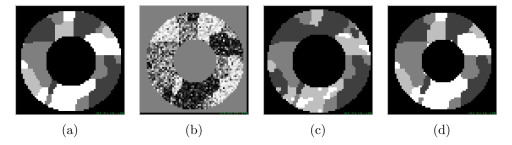


Figure 5.3: Synthetic example. (a) True labels, (b) First time point of observed time series, (c) Time series plot on sphere, (d) label map estimated by mode-approximation, and label map estimated by MCEM.

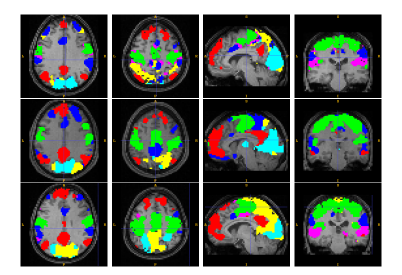


Figure 5.4: Functional networks detected by the proposed method for 3 subjects overlaid on their T1 images. The clusters are the visual (cyan), motor (green), executive control (blue), salience (magenta), dorsal attention (yellow), and default mode (red) networks.

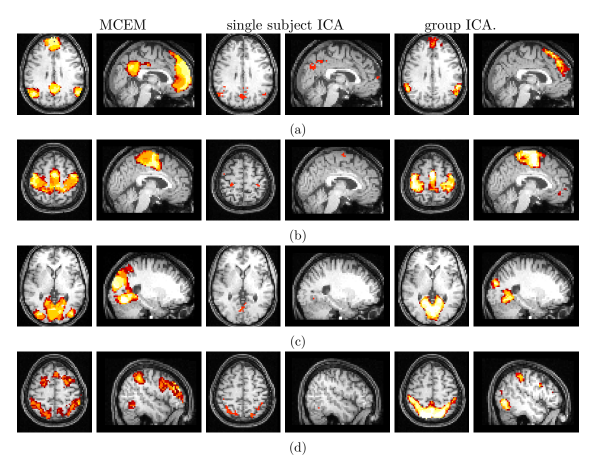


Figure 5.5: Comparison of the overlap of the label maps estimated by our MCEM approach, group ICA and single subject ICA on 16 subjects. Color map ranges from 8 (red) 16 (yellow). (a) MDN, (b) motor, (c) visual, (d) attentive.

CHAPTER 6

HIERARCHICAL MODEL FOR GROUP ANALYSIS

In this chapter, we will apply the graphical model and MRF to one of the core problems in functional network analysis: estimation of networks from a group of subjects. To study the brain's intrinsic activity with rs-fMRI data, one either models the data of a single subject or a group of subjects. The BOLD signals of a single subject are often contaminated with the noise of various sources, and the results are typically unreliable for the inference of the whole population. On the other hand, combining data from multiple subjects and jointly estimating the common functional networks is more robust. In group analysis of rs-fMRI one assumes that all subjects in the group share common functional connectivity patterns and that the group networks can be estimated more accurately because the noise from each subject is canceled by averaging. In practice, it is a major challenge to summarize the consistent patterns across subjects, as each subject's network structure appears similar but has slight variations.

In this chapter we propose a Bayesian hierarchical model for estimating the functional networks by using the rs-fMRI data from a group of subjects. The hierarchy comes from an additional level of group map defined on top of the conventional subject functional network maps. The group effect goes into the subject network label's probabilistic distribution as a parameter. Both group and subject networks are jointly estimated in an iterative way. We give a natural interpretation of the regularization with a Bayesian perspective. Once the group's network map is known, it can help the individual subject's estimation as a prior probability. Because the group map combines the information from all subject maps, this prior distribution is equivalent to using other subjects' data for the current subject's inference. Besides, a subject's network estimates also help to iteratively refine the group map estimates. We model the intersubject variability by balancing between a common map for all subjects (no variability, maximal shared information) and a separate map for each subject (no shared information, maximal variability). We achieve the optimal balance in the

Bayesian sense by computing the subject network label's posterior density. This posterior density combines the prior information from the group map and the data likelihood from the subject-specific BOLD signal. We further model the within-subject spatial coherence by a Markov random field (MRF). In the remaining part of the paper, we refer to our model a hierarchical Markov random field (HMRF).

A classical occurrence of hierarchical modeling in fMRI is the inclusion of random effects in a general linear model (GLM) [10], which is later extended to a full Bayesian framework [153]. The multilevel model has richer structures and can capture the structures in multiple-group, multiple-session data, and distinguish between the influence of the fixed effect and that of the random factors. In our model, the hierarchy is defined on a latent variable mixture representation.

A Markov random field is a multivariate distribution defined on an undirected graph to represent the soft constraints between the variables. In fMRI analysis, it is a principal regularization method of obtaining a spatially coherent solution. Depending on the context, previous works have defined MRF on different variables. They have been used for the regularization priors on the coefficients of the general linear model (GLM) [120], on the parameters of a spatiotemporal auto-regression model [154], and on the hidden activation variables in task-based experiments [77]. In this article, we define MRF on the latent network label variables of hidden Markov model (HMM), to represent our prior knowledge of the spatial coherence of the network patterns within a subject. There is a key difference between our model and conventional HMMs, though. We generalize the conventional concept of spatial regularization by defining a joint graph that includes the network variables of both the group and subject levels. In our model, the neighbors of each node on the graph include the corresponding nodes at another level, as well as the spatially adjacent voxels in the same level. The new graph introduces our additional assumption that one subject's functional networks should share similar patterns with another subject's, implicitly represented by the group. With this definition, we map all the variables in a hierarchical model on to a single graph, and formulate a problem conceptually appealing and feasible in practice.

The exact inference of MRF is a combinatorial optimization of discrete variables, hence it is computationally infeasible except in special cases [73, 114]. The iterated conditional mode (ICM) is typically used to obtain a local optimum of the mode of the posterior density [15]. In this work we are interested in the posterior variance of the network label variables as well as the mode, and we use MCEM sampling algorithm for the inference of

both group and subject label maps. MCEM is data-driven in that the model parameters are estimated together with the network label variables. The only parameter that needs special treatment is the link strength between the group and subjects. MCEM integrates the Markov chain Monte Carlo sampling in the expectation-maximization loop. The price to pay is the longer computation time than in other approximate inference methods such as variational Bayes.

We show our HMRF model is able to recover both group and subject functional networks in simulated group fMRI data. While HMRF's group estimates are comparable or more accurate than the two other methods under comparison, we are especially interested in the higher accuracy of the individual subjects' estimates. We further show the strength of the model by a real multiple-session dataset, where we achieve significantly higher intersession consistency by using our joint-estimation model. The method also proves to be more stable under the data perturbation in a bootstrap experiment. This paper is based on our earlier work [101], and we extend previous work to redefine the model in an integrated graphical model context. The new simulated data experiments explore the performance of the algorithm under various levels of spatial smoothing. In the real data experiments, we added a new intersession consistency test and the algorithm stability test with bootstrapping. We also improved the parameter estimation by using the Bayesian posterior predictive distribution of the test subjects in a cross-validation framework.

In the remainder of this chapter, we define the model in Section 6.2, and give the approximate inference procedure in Section 6.3. We compare the accuracy and consistency of our method, together with other methods by testing on synthetic and real data experiments in Section 6.4 and Section 6.4.2 and discuss the model and algorithm performance in Section 6.5.

6.1 Related works

ICA is a powerful tool for identifying functional networks by using rs-fMRI of a single subject [28]. ICA is used to recover the statistically independent functional components without a priori knowledge of the regions of interest. See Figure 2.4 for how spatial and temporal ICA methods are defined. Group ICA is used as an extension of single-subject ICA in order to seek a set of independent components shared across all subjects [28]. In a typical group ICA study, all subjects are registered to a common atlas and assumed to share a common spatial component map but have distinct time courses. The BOLD signals from all subjects are concatenated temporally, followed by a single-subject ICA analysis. The subject

component maps are then obtained by a back-reconstruction procedure. Alternatively, single-subject ICA is applied on each subject first, and a self-organized clustering algorithm applies to all subjects' components such that similar components are assigned into one cluster. The group ICA components are represented by the centers of the clusters [49]. Neither of the above approaches iteratively refine group (or subject) maps once the subject (or group) maps are estimated.

ICA as a signal decomposition method obtains overlapped spatial components and needs ad-hoc thresholding. Such ambiguity makes interpreting the results difficult. On the other hand, functional networks estimation can also be defined as an image segmentation problem. The region-of-interest (ROI), or even the whole brain voxels can be partitioned into disjoint spatial patches. The patches with the same network labels, even when spatially remote from each other, belong to the same functional networks. To extend the segmentation method to a group of subjects, segmentations are performed first on individual subjects. The connectivity maps are averaged to obtain a group affinity matrix. A second level segmentation is performed on this affinity matrix [12, 143]. Again, subject network maps are not refined from the estimates of the group network.

It is worth noting that Ng et al. [114] also use MRF for group study. The spatial neighborhood is extended to cross-subject voxels, in order to address the imperfect anatomical alignment and functional correspondence. Our model is different from Ng's group MRF model in that 1) a group level is defined in our model, whereas in Ng's work, a combined structure including all subjects is defined without a group level. In such a flat model, a voxel directly uses the information of the corresponding voxels of other subjects. Instead, we expect a second level enriches the model and better decomposes the fixed and random effects in the subject network map. 2) Ng et al. define the MRF prior on the GLM coefficients in task-based experiments so the posterior inference is a two-class problem (active versus inactive), and an exact solution can be obtained by a graph-cuts algorithm. Our model applies to the network labels in a rs-fMRI study, and hence is a multiclass segmentation problem that is significantly more difficult to solve. 3) The unary potential function in the model of Ng is defined via the posterior probability of the label variable given the GLM's coefficients, and there is no explicit splitting between the MRF prior and the likelihood. This modeling method is essentially a conditional random field [93]. In our model, the unary potential in the MRF prior is defined as the links between group and subject level, and thus is separated from the data likelihood, so our model is a variant of the hidden Markov model. Thanks to the Bayesian perspective of our hierarchical model, the trade-off between prior and likelihood is accounted for automatically. 4)The MRF model of Ng et al. is built on certain regions of the brain, while ours is built on the whole brain's gray matter voxels.

Another class of methods identifies the functional spatial patterns by decomposing the BOLD signal, which can be seen as generalizations of ICA [146, 144] methods. The authors of both works introduce generative models that include a population level and subject level. Each subject's mixing weight coefficients are regarded as being generated from population level latent factors, and the population and subject level's mixing matrices are solved jointly as a convex optimization problem [144]. With regard to the joint estimation of both levels of the hierarchy, we can see such methods as a counterpart of our model in the class of signal decomposition methods.

6.2 Hierarchical MRF For modeling group fMRI

We begin by defining each subject's network label map as a Markov random field (MRF) with the neighborhood structure given by a regular lattice. The statistical dependency between adjacent voxels acts as a prior model favoring spatial coherence of estimated functional regions. To generalize the MRF to a hierarchical setting, an additional group label map is defined in addition to all subject label maps. The group label map has the same number of voxels and the same Markov structure as the individuals' maps, again to encourage spatial coherence of the functional regions in the group level. In addition, each voxel in the group map is connected to the corresponding voxel in each subject map. These connections model the relationships between the group and the individuals. The subjects' functional network labels are regarded as generated from the group labels, and the rs-fMRI time courses are regarded to be generated from a mixture of high-dimensional distributions given the subject network labels. All voxels of subjects and group label map are jointly connected into a single MRF. The functional network estimation is the inverse problem of the above data generation process, as the labels are inferred from their posterior distribution given the data. See Figure 6.1 for an illustration.

More specifically, we follow the notation in Chapter 3 and define an undirected graph $\mathcal{G} = (\mathcal{V}, \mathcal{E})$. The set of vertices $\mathcal{V} = (\mathcal{V}_G, \mathcal{V}_1, \dots, \mathcal{V}_J)$ is the union of the gray matter voxels \mathcal{V}_j for all J subjects as well as those in the group volume \mathcal{V}_G . An edge $(s, t) \in \mathcal{E}$ is defined in one of three types: (1) $s \in \mathcal{V}_G$, $t \in \mathcal{V}_j$ and s, t have the same physical coordinates, (2) $s, t \in \mathcal{V}_G$, and s, t are spatial neighbors, or (3) $s, t \in \mathcal{V}_j$, and s, t are spatial neighbors. In our model we use a 26-neighbor system in a three-dimensional volume image (a voxel at the boundary of the gray matter may have ; 26 neighbors). We will refer to the first type of

link as between-level links, and the second and third types of links as within-subject links. On each node $s \in \mathcal{V}$, a discrete random variable $x_s \in \mathcal{L} = \{1, \dots, L\}$ is defined to represent the functional network label. We use -s for the set of nodes excluding node s, and $\mathcal{N}(s)$ for the set of neighboring nodes of s. Last we define clique c as a complete subgraph of c, such that every pair of nodes in c has a link between them, and define c the set of all cliques in c.

6.2.1 MRF prior

MRF is a principal regularization method for modeling spatial context information. In a Bayesian setting, we use it as a prior distribution of the network label variables $X = \{x_s \in \mathcal{L} | s \in \mathcal{V}\}$. Formally, with the definition of the graph \mathcal{G} and neighbor system $\mathcal{N}(s), \forall s \in \mathcal{V}$ above, X is said to be a MRF on \mathcal{G} if $P(x_s|x_{-s}) = P(x_s|x_{\mathcal{N}(s)})$, i.e., a variable is conditional independent of the variables on the remaining nodes of the graph given its neighbors [99]. This local conditional independence property is difficult to apply to the inference of the joint distribution. Thanks to the equivalence of MRF and Gibbs fields [13], one can transform the local property into a global property. A Gibbs random field or Gibbs distribution takes the form of $P(X) = (1/Z) \exp\{-U(X)\}$, where Z is a normalization constant called partition function in order to guarantee the function integrates to 1, and $U(X) = \sum_{c \in \mathcal{C}} V_c(X_c)$ is called the energy function. Each clique potential function V_c only depends on the variables in the corresponding clique c. The Hammersley-clifford theorem [31] states that X is a MRF if and only if it obeys a Gibbs distribution. In this specific problem, the energy function takes the following form:

$$U(X) = \sum_{s,r \in \mathcal{V}_G} \beta \psi(x_s, x_r) + \sum_{j=1}^J \left(\sum_{s \in \mathcal{V}_G, \tilde{s} \in \mathcal{V}_j} \alpha \psi(x_s, x_{\tilde{s}}) + \sum_{s,r \in \mathcal{V}_j} \beta \psi(x_s, y_r) \right). \tag{6.1}$$

The binary function ψ takes zero if the two inputs are equal and takes 1 otherwise. Parameters α and β determine the strength of the links. The pair of voxels (s,r) is spatially adjacent within the subject volume or the group volume (the type two and type three links), and (s,\tilde{s}) is a pair of neighboring voxels at a different level in the hierarchy, but sharing the same physical coordinates (type one link).

This regularization encodes two physiologically meaningful a priori assumptions on the functional networks under investigation: (1) The networks are spatially coherent within the single subject map and within the group map. This spatial coherency is modeled by the β potential term. (2) The subject's intrinsic functional activity must share similar patterns, regardless of the possible confounding of the noise and subject-specific effect.

This between-subject constraint is modeled by the α potential term. The proposed energy function represents both priors without introducing blurring artifacts. As for the inference, although appearing different in the image domain, the three types of links are no different when looking from the abstract graph layer, and can be treated equivalently in the inference procedure. Our MRF prior is essentially a Potts model with different weights defined on three types of edges [122]. However, we extend the Potts model such that the cliques in a graph include both within-subject links and between-level links, so the model favors not only spatial coherence but also the intersubject coherence. Figure 6.2 gives an alternative view of the same graphical model that are illustrated in Figure 6.1.

6.2.2 Likelihood model

In the generative model, for any individual subject, the observed time course at each voxel is assumed to be generated from a distribution conditioned on the network label at that voxel. In fMRI analysis the BOLD signal is typically normalized to be zero mean and unit norm, so the analysis is invariant of shifting or scalings of the data [68]. The normalization results in the data being projected onto a high-dimensional unit sphere, and the sample correlation between the two time series is equal to their inner product. The rs-fMRI segmentation aims at a clustering such that within-cluster voxels have a high correlation, and between-cluster voxels have a low correlation. The equivalence of the correlation and inner product makes it possible to reformulate the original problem into a new one. Now we can find a clustering where voxels with a larger inner product are put into one cluster. The new problem can be modeled and solved using a mixture of the von Mises-Fisher (vMF) distribution.

We use $Y = \{(y_1, \ldots, y_N) \mid y_s \in S^{p-1}\}$ to denote the set of normalized time series on the p-sphere, where p is the number of time points in the original BOLD signal, and N is the total number of gray matter voxels of all subjects. Given X, the random vectors y_s are conditionally independent, hence $\log P(Y|X) = \sum_j \sum_{s \in \mathcal{V}_j} \log P(y_s|x_s)$. The likelihood function $P(y_s|x_s)$ is naturally modeled by a vMF distribution

$$f(y_s|x_s = l; \mu_l, \kappa_l) = C_p(\kappa_l) \exp\left(\kappa_l \mu_l^{\mathsf{T}} y_s\right), \qquad y_s \in S^{p-1}, \quad l \in \mathcal{L},$$
 (6.2)

where for the network cluster label l, μ_l is the mean time course, $\kappa_l \geq 0$ is the concentration parameter, and C_p is the normalization constant. The larger the κ_l , the greater the density concentrated around the mean. Since eq. (6.2) depends on x only through $\mu^{\mathsf{T}}x$, the vMF distribution is unimodal and rotationally symmetric around μ .

6.3 Bayesian inference

The exact inference of P(X|Y) is computationally intractable due to the pairwise interaction of MRF prior distribution. Various approximate solutions exist for such types of undirected graphical model inference problems, including Gibbs and Metropolis sampling, expectation-propagation, and some variation inference methods such as mean field approximation and message-passing methods. In this work, we choose Gibbs sampling because of its simple formulation and straightforward implementation in a multiprocessor system. In addition, compared to a point estimate such as $maximum\ a\ posteriori\ (MAP)$ framework, the samples of the label map can be used to approximate the full posterior density, and to help understand the confidence of the point estimates such as posterior mean or modes.

6.3.1 Gibbs sampling

The Gibbs sampler, as a special case of the Metropolis-Hastings sampler, solves a multivariate sampling problem using iterative univariate sampling. When all the random variables but one are fixed, the transition probabilities depend only on the local conditional distributions. The resultant equilibrium density of the Markov chain is exactly the target density P(X|Y). In general MCMC sampling, the variables are visited either at random, or according to a predefined order. As a way of incorporating domain-specific information in the design of our Gibbs sampler, we schedule the sampling order also in a multilevel fashion. At the image level, we draw the mth sample of the group label map X_G^m given all the previous subject label maps $\{X_j^{m-1}, j=1\dots J\}$. Next, we draw a sample of subject j's label map X_j^m given the current group map sample X_G^m (Figure 6.3). At the voxel level, we sample and update x_s given the rest of the nodes are fixed (Figure 6.4). We call it a scan when each $x_s, \forall s \in \mathcal{V}$ is updated once. The conditional distribution used to generate samples at the group and subject voxels can be derived from equation (6.1) and are given as

$$P(x_s|x_{-s}, Y) = \frac{1}{Z_s} \exp\left\{-U_p(x_s|x_{\mathcal{N}(s)}, y_s)\right\} \quad \text{where,}$$

$$\forall s \in \mathcal{V}_G, \quad U_p = \alpha \sum_{j=1}^J \psi(x_s, x_{\tilde{s}}^j) + \beta \sum_{r \in \mathcal{N}(s)} \psi(x_s, x_r), \qquad (6.3)$$

$$\forall s \in \mathcal{V}_j, \quad U_p = \alpha \psi(x_s, x_{\tilde{s}}) + \beta \sum_{r \in \mathcal{N}(s)} \psi(x_s, x_r) - \kappa_l \mu_l^{\mathsf{T}} y_s - \log C_p, \qquad (6.4)$$

where -s is the set of all the nodes excluding node s, Z_s is the partition function of x_s , U_p is the posterior energy, and $\mathcal{N}(s)$ is the set of neighbors of s. The $x_{\tilde{s}}^j$ in (6.3) is the network label of subject j's voxel with the same physical coordinates with s, and the $x_{\tilde{s}}$

in (6.4) is the label of the group map's voxel with the same physical coordinates as s. Note the evaluation of Z_s is easy since it is in a univariate distribution and is the sum of only L terms. Because of the dependency on previous samples, the sequence of label map samples $\{X^m, m=1\ldots, M\}$ is indeed a Markov chain; hence our method falls into Markov chain Monte Carlo (MCMC) sampling. After a sufficient burn-in period, a series of samples $\{X^m, m=1, \cdots, M\}$ is saved. The samples have all the information of P(X|Y) and can be used for approximating the expectation $\mathbb{E}_{P(X|Y)}[\log P(X,Y;\theta)]$ as well as estimating the posterior variance.

6.3.2 Parameter estimation

The parameters $\{\beta,\kappa,\mu\}$ in our model are data-dependent, and manual assignment can easily result in over-fitting. For example, β 's optimal value depends on the number of neighbors of a voxel and also on the number of subjects in the group. In this data-driven model, we propose to estimate the parameters θ from the data using an expectation maximization (EM) algorithm, with the network labels X as the hidden variable. However, the high-dimensionality and dependency between spatially adjacent voxels in MRF make it infeasible to obtain a closed form solution of the expectation of $\log P(X,Y;\theta)$ with respect to P(X|Y). Here we propose to approximate the expectation using Monte Carlo EM (MCEM) algorithm. The set of samples, (X^1,\cdots,X^M) generated from density P(X|Y) is used to approximate the expectation by the empirical average $(1/M)\sum_{m=1}^M \log P(Y,X^m;\theta)$. Furthermore, in order to evaluate $\log P(Y,X^m;\theta) = \log P(X^m;\theta) + \log P(Y|X^m;\theta)$ as a function of θ , we face the difficulty of evaluating the partition function Z in $P(X^m)$. In practice the likelihood function $P(X;\theta)$ is approximated by pseudo-likelihood [13], which is defined as the product of the conditional likelihoods $P(x_s|x_{-s};\theta), \forall s \in \mathcal{V}$. Therefore the label map's log-likelihood can be written as

$$\log P(X;\theta) \approx \sum_{s \in \mathcal{V}} -U(x_s | x_{-s}; \theta) - \log Z_s, \tag{6.5}$$

$$\text{where} \quad Z_s = \sum_{l=1}^L \exp\{-U(x_s = l | x_{-s})\}, \tag{6.6}$$

$$\forall s \in \mathcal{V}_G, \quad U(x_s | x_{-s}) = \alpha \sum_{j=1}^J \psi(x_s, x_{\tilde{s}}^j) + \beta \sum_{r \in \mathcal{N}(s)} \psi(x_s, x_r);$$

$$\forall s \in \mathcal{V}_j, \quad U(x_s | x_{-s}) = \alpha \psi(x_s, x_{\tilde{s}}) + \beta \sum_{r \in \mathcal{N}(s)} \psi(x_s, x_r);$$

where $x_{\tilde{s}}^{j}$ and $x_{\tilde{s}}$ have the same definition as in equation (6.3) and (6.4). With the pseudo-likelihood approximation, there is no need to compute the original Z. Instead we compute

 Z_s for each voxel s, just like what we do in the Gibbs sampling

6.3.3 HMRF algorithm using MCEM

With all the preparation above, parameter estimation can be done by maximizing $(1/M)\sum_{m=1}^{M}\log P(Y,X^m)$. More specifically, β exists only in the prior, and can be estimated by maximizing $\frac{1}{M}\sum_{m=1}^{M}\log p(X^m)$ with the Newton-Raphson method. Since $\{\mu,\kappa\}$ exist only in the data likelihood, the normalization constant Z in the prior is not a problem, hence $\{\mu,\kappa\}$ are estimated by maximizing $(1/M)\sum_{m=1}^{M}\log P(Y|X^m)$. The α parameter is treated differently and will be discussed in Section 6.3.4. In order for MCMC sampling to converge quickly to the posterior, we need a reasonably good initial network label map. Here the K-Means clustering on a concatenated group dataset is used for the initial maps of both the group and subjects. After the EM iteration converges, we save M Monte Carlo samples as output. The Monte Carlo samples have all the information of the posterior distribution of network labels, and will be used in postprocessing for inference. Putting this all together, the HMRF method to estimate the group and individual label maps is given in Algorithm 7.

6.3.4 Estimating α parameter by cross-validation

The parameter α in our model represents the strength of the links between the group and subject network label maps. The parameter implicitly represents the extent to which the functional patterns are shared among the subjects. Unfortunately, this parameter cannot be estimated in a MCEM framework by a Newton-Raphson method, as such a direct optimization will result in a collapsed solution. A solution of $\alpha = 0$ would minimize the energy associated with the between-level links, and the group map \mathcal{V}_G would degenerate into a constant label map because such a map would minimize the energy associated with the links within the group map. We instead use the posterior predictive distribution [62] of a test subject's BOLD signal Y_t , defined as

$$P(Y_t|Y;\alpha,\theta_t) = \int P(Y_t|X_t;\theta_t)P(X_t|Y;\alpha) \,dX_t, \tag{6.7}$$

where $\theta_t = \{\mu_t, \kappa_t, \beta_t\}$ is the parameter set of the test subject. With a leave-one-out procedure, the same as that in the standard cross-validation, we pick one subject as the test subject Y_t , and the remaining J-1 subjects as the training data. We then compute the average $P(Y_t|Y;\alpha,\theta_t)$ across all test subjects given a list of prespecified α values, and choose α with the highest average predictive distribution.

Algorithm 7: HMRF: Monte Carlo EM algorithm for network label inference and parameter estimation

```
Data: Normalized rs-fMRI, initial group label map Result: MC samples of label maps \{X^m, m = 1, \dots, M\}, parameters \{\beta, \mu, \sigma\} while \mathbb{E}_{P(Y|X)}[\log P(Y, X; \theta)] not converged do

repeat

foreach s \in \mathcal{V}_G do Draw sample of x_s from P(x_s|x_{-s}, y_s; \theta) using (6.3);

foreach j = 1 \dots J do

foreach s \in \mathcal{V}_J do Draw sample of x_s from P(x_s|x_{-s}, y_s; \theta) using (6.4);

Save sample X^m after B burn-ins;

until B + M times;

foreach l = 1 \cdots L do

Estimate \{\mu_l, \kappa_l\} by maximizing (1/M) \sum_{m=1}^M \log P(Y|X^m; \theta);

Estimate \beta by maximizing (3.25);
```

The test subject's predictive distribution in (6.7) for a chosen α can be evaluated through a Monte-Carlo approximation

$$P(Y_t|Y;\alpha;\theta_t) \approx \frac{1}{M} \sum_m P(Y_t|X_t^m;\alpha,\theta_t), \quad X_t^m \sim P(X_t|Y;\alpha).$$
 (6.8)

One economical way of generating sample $\{X_t^m, m = 1..., M\}$ can be done within the MCEM loop of Algorithm 7. After the current group map is generated in E step, one sample X_t^m can be generated from $P(X_t|Y;\alpha,\theta)$. The corresponding posterior energy function at voxel s is $U_p(x_s|x_{N(s)}) = \alpha \psi(x_s,x_{\bar{s}}) + \beta \sum_{r \in N(s)} \psi(x_s,x_r)$. This energy is the same as equation (6.4), except that there is no time series data term $\kappa_l \mu_l^\mathsf{T} y_s - \log C_p$ since the test subject data Y_t are not given in this distribution. For one sample map X_t^m , the test subject parameter set θ_t is obtained by optimizing $P(Y_t|X_t^m)$. As a simple reasoning of why we can use the equation (6.7) for estimating α , when α is too small, most of the X_t^m will depend less on the group map X_G and tend to be random clusterings, which will have low data likelihoods in (6.8). When α is too big, X_t^m will be almost the same as X_G , again resulting in a suboptimal value for (6.8). Only with an appropriate α , could X_t^m sufficiently explore the sampling space including the regions where the predictive distribution is maximized. In practice, we evaluate eq. (6.8) for a fixed set of α values, and choose α with the largest predictive density value $P(X_t|Y;\alpha)$.

6.4 Experiments on simulated data

Given the lack of ground truth of the functional network of the *in vivo* rs-fMRI data, we begin the experiments with a simulated dataset. We focus primarily on the estimation accuracy on the simulated dataset, and on the estimation consistency on the *in vivo* data.

We compare our method with two other clustering methods – K-Means and normalized-cuts (N-Cuts) – as well as two degenerated versions of the HMRF algorithm: HMRF-A and HMRF-B. The K-Means algorithm, as a simple and fast clustering method, is applied to the paradigm fMRI study in Baumgartner et al. [8], and is later used by Bellec et al. [12] for bootstrap analysis of the rs-fMRI group study. In our experiment, the distance metric of K-Means is defined as $1-x_s^{\rm T}x_r$. To estimate an individual subject's network, we apply K-Means on each subject's BOLD signal 20 times, and choose the segmentation map with the minimal ratio of the sum of the intercluster distance and the sum of the intracluster distance. For the group study, we construct a group dataset by concatenating all subjects' time courses and run K-Means 20 times also on this group's dataset to estimate a group network label map. The initial cluster centers for both subject and group clustering are chosen randomly while at the same time maximizing the between-center distance [5].

N-Cuts formulates the fMRI image segmentation as a graph partitioning problem. A global criterion is used to find a subset of edges to remove from a full-connected graph, and the voxels are partitioned into multiple disjoint sets [138]. N-Cuts is used by Heuvel et al. [143] and Craddock et al. [38] for the group rs-fMRI study. Following Heuvel et al. [143], we also apply N-Cuts in two stages. First, N-Cuts is run on each subject's affinity matrix, as computed from the pairwise correlation between time courses. A second N-Cuts is applied on a group affinity matrix, computed by summing all subjects' binarized affinity matrices derived from their segmentation maps. We use the same toolbox Ncutclustering_9 [138] as in Heuvel et al. [143], as well as the same parameter setting.

Both HMRF-A and HMRF-B, as simplified versions of HMRF, serve to check whether a reduced model would be able to achieve the same or better performance compared to the proposed full model. Both models are the same as HMRF except $\alpha=0$ for HMRF-A, and $\beta=0$ for HMRF-B. The model HMRF-B indeed amounts to defining a MRF on each single subject and estimating each subject's networks independent of other subjects. Such a strategy is equivalent to the hidden Markov model we proposed in Liu et al. [100].

For HMRF, we skip the first 500 burn-in samples before saving 100 samples of the label map at each EM iteration. The convergence testing of MCMC sampling, especially in high-dimensional space is an open question and there is no widely accepted method to address this issue. We empirically choose the number of burn-in and MC samples by observing that the posterior probability estimated from samples has no significant change. The β parameter is estimated by the M step, as well as the μ and κ for each vMF component. As an optional postprocessing step, the discrete label map is obtained by running a iterated conditional mode [15] algorithm based on the last MCMC sample map.

Before a discussion of synthetic data generation, we briefly discuss how to measure the data quality of rs-fMRI. The separability of a data set for the purpose of clustering depends on both the within-cluster variance and between-cluster variance. In this specific rs-fMRI dataset, the signal-to-noise ratio (SNR) is represented by the ratio of the average between-cluster distance (defined as $1 - \mu_i^{\mathsf{T}} \mu_j$, where μ_i and μ_j are the cluster's mean time series), and the average within-cluster variance (defined by $1/\kappa$).

We generated synthetic rs-fMRI data in two steps. First, a group network map with five network labels is generated by drawing samples from a Potts model with $\beta = 2.0$ and 500 scans. Given the group map, a subject map is generated according to equation (6.1) with $\alpha = 0.5$ and $\beta = 2.0$. The subject map generation procedure is repeated 25 times to obtain a group of 25 subjects. To simulate the BOLD signal given the network label map, we

first generate mean time courses μ_l , $l=\{1,\ldots,5\}$ from a first-order auto-regressive process $x_t=\varphi x_{t-1}+\varepsilon$, with $\varphi=0.8$ and $\varepsilon=0.1$. The sample correlations between the mean time series are in the range of (-0.15,0.3). Then, we add independent Gaussian white noise on each cluster's mean time course. The variance of the white noise is chosen such that the simulated BOLD signals have SNR=24, which is close or slightly lower than that of the real rs-fMRI data used in our experiments. Once the time series are generated, they are spatially smoothed with a Gaussian filter. Because the size of the smoothing filter may have interactions with our HMRF model and hence have an impact on the estimation accuracy, we spatially smoothed the generated BOLD signals with three levels of scale: FWHM = 0, FWHM = 1.88 mm, and FWHM = 4.7 mm. Furthermore, the synthetic data are generated randomly, so the experimental results from the data may also vary. To take account of the variability of the results, we repeated the above data generation process 100 times. For each generated data set, we run the five methods on the BOLD signals preprocessed by three levels of Gaussian filters, respectively, and compare the Monte Carlo average of the estimated label maps with the ground truth.

6.4.1 Synthetic data results

Among the 100 Monte Carlo runs of the data generation and estimation procedure, we choose one dataset smoothed at FWHM = 1.88 mm. The corresponding estimates are shown in Figure 6.5. We use the Rand index [126] to measure the similarity between simulated ground truth subject maps and the true group map. Rand index (RI) ranges in [0, 1], and takes 1 if the two maps under comparison are the same. The RI value for this particular simulated dataset is 0.88 (similar values for other generated datasets), which we find is empirically close to the real data. From the figure, all methods appear to estimate the group map well (except HMRF-B, which does not allow a group map estimate), but perform differently on the subjects. K-Means tries to identify the finer details of the individual subject's spatial patterns but fails due to the high noise level. N-Cuts and HMRF-A can detect the large patterns but lose some detail; HMRF-B does estimate the smooth subject map thanks to the within-subject smoothness links but the maps do not match the ground truth well. Finally, the HMRF is able to recover subjects' network maps with good matching to the ground truth.

To quantitatively evaluate the accuracy of the segmentation map from various methods, we calculate the RI values between the true map and the estimated map. The boxplot in Figure 6.6 shows the RI across all Monte Carlo runs and subjects. In all three settings of smoothing kernel size, HMRF achieves higher accuracies compared to other methods.

In addition, for individual subjects' estimation, our model performs best at a moderate smoothing size of FWHM = 1.88 mm, which is smaller than the typical 5-8 mm smoothing size. This is because the HMRF model benefits from the reduced noise variance resulting from the moderate smoothing, but avoids losing finer details due to excessive smoothing. In practice, this means when applying HMRF, the BOLD signal should be smoothed by a small-kernel Gaussian filter, and we choose FWHM = 1.5 mm in the following real data experiments. We also note that the K-Means optimal smoothing kernel size is larger than that of HMRF, because it lacks the spatial coherence regularization and hence needs more smoothing in preprocessing stage. Last, we found that the two reduced models HMRF-A and HMRF-B do not perform as well as the full model, indicating that the hierarchy in the full model is indeed necessary. For all possible smoothing sizes, HMRF's estimation accuracy is comparable or moderately better than the other four methods on the group label map, and significantly higher on subject maps.

6.4.2 Real data experiments

In this work we test our methods on the publicly available NYU test-retest (TRT) dataset that has been used previously [137, 162]. While the original goal of the above works was to verify the voxel-wise intra- and intersession TRT reliability, our goal is to verify whether the methods under consideration are able to estimate consistent functional network maps across sessions, given the fair amount of intersession consistency in the data set [39, 30, 104, 57]. We present two experiments with the NYU-TRT datasets. The first experiment aims at demonstrating the intersession consistency of the estimated subject network maps, and the second one evaluates how the algorithms behave under the perturbation of the data by using bootstrap sampling. We compare three methods, HMRF, K-Means, and N-Cuts, in both experiments. The other two methods, HMRF-A and HMRF-B, are not taken into account in this section since they are a simplified version of HMRF and have been shown to be suboptimal compared to the full model.

6.4.3 Preprocessing

Twenty-six healthy control participants (11 males, mean age 20.5 ± 4.8 years) were scanned three times. The participants had no history of psychiatric or neurological illness. BOLD EPI images (TR = 2 s, TE = 25 ms, flip angle = 90, 39 slices at 3 mm slice thickness, 64×64 matrix, field of view = 192 mm, 197 volumes) were acquired on a Siemens Allegra 3.0 Tesla scanner. Scans 2 and 3 were conducted in a single session, 45 minutes apart, and were 5-16 months after the first scan. The subjects were asked to relax and remain still with

their eyes open during the scan. A high resolution T1-weighted image was also obtained (MPRAGE with TR = 2.5 s, TE = 4.35 ms, TI = 900 ms, flip angle = 8° , 176 slices, FOV = 256 mm).

The fMRI data was preprocessed using the scripts of the 1000 functional connectomes projects, as well as FMRIB's FSL toolset and the Medical College of Wisconsin's AFNI tool. The volumes are motion corrected by aligning to the mean volume with a six-parameter rigid body transformation. The BOLD signals were bandpass filtered to 0.01 to 0.1 Hz, and nuisance variables were regressed out including white matter, CSF mean time courses and six motion parameters. The signal is then filtered by a FWHM = 1.5 mm Gaussian filter for spatial smoothness. The small kernel size of spatial smoothing guarantees that noise is canceled by averaging neighboring voxels while keeping the finer functional network patterns (see the simulated test and Figure 6.6). The functional images are first registered to the corresponding T1 images, and both functional and T1 images are registered to MNI152 (Montreal Neurological Institute) space with a 12-parameter affine transformation. Finally, after masking out white matter and CSF voxels, we have 39,080 gray matter voxels remaining in each subject. We construct a joint graph with over one million nodes including all subjects and the group map.

6.4.4 Choosing parameters

In this work we do not address the problem of how many functional networks exist in the human brain. Instead, we use existing reports [157] and choose seven functional networks for segmentation throughout the real data experiments. With this setting, we expect to identify the following typical functional networks: visual and primary motor [39], attention [55], default mode network (DMN) [72], saliency, and executive control system [136], regardless of the segmentation methods used. The K-Means is repeated 20 times with random initialization [5] for segmentation of both the subject and group maps. For N-Cuts, we threshold each subject's correlation matrix at 0.4 before applying N-Cuts on a single subject. After the individual segmentation, we average all subjects' binary segmentation matrices, and threshold the averaged matrix at 0.3. The result represents the group correlation matrix. Both cut-off thresholds are suggested by Heuvel et al. [143]. Our implementation is different with from Heuvel et al. [143] only in that we partition the subject map into seven clusters instead of 20. This is because we need to compare the subject maps estimated by N-Cuts with those estimated by the HMRF method at the same number of networks. We also run N-Cuts with 20 clusters on subject maps to compare with our seven-cluster configuration (results now shown), and find the group level segmentation has not been impacted by our lack of over-segmentation at the subject level. For HMRF, we initialize both the group and subject label maps with the group label map estimated from K-Means. The sampling routine (E-step of MCEM algorithm) skips 500 burn-in samples before saving 100 MC samples. The parameters $\{\beta, \mu, \kappa\}$ are estimated from the data. With α estimated from the posterior predictive distribution (see Section 6.3.4), we found the similarity between estimated group and subject maps measured is around 0.85 measured by RI value.

6.4.5 Intersession consistency

Since the TRT dataset and the general rs-fMRI data have been shown to share consistent functional networks across all sessions [39, 30, 104, 57], we verify the consistency of the HMRF algorithm by applying it to all three sessions of data. A method is said to be consistent if it is able to derive similar network estimates across sessions. We compare three pairs of sessions' consistency: S1 vs S2, S1 vs S3 and S2 vs S3. For each subject in each pair of sessions, we compute the consistency score between this subject's network map estimates in two sessions. The similarity is again represented by the RI values. We expect the proposed HMRF algorithm has higher average similarity compared with other methods. The consistency scores of all subjects are summarized in a boxplot as in Figure 6.7. For comparison, the same boxplots are also drawn for K-Means and N-Cuts. From the figure, the subject network label maps estimated from HMRF have significant higher intersession consistency scores compared to the other two methods. This indicates that our algorithm is able to capture the common functional patterns across sessions. In addition, both K-Means and HMRF have higher intersession consistency scores between session two and session three, compared to the other two intersession comparisons. This is consistent with the fact that sessions two and three have a smaller interval (45 minutes apart), compared to session one and two (5-16 months). K-Means has slightly better between-session consistency than N-Cuts, probably because we have run K-Means multiple times and have chosen the best solutions.

The RI values in Figure 6.7 only give a single number of similarity between two network label maps, rather than a voxel-wise consistency map. To visualize the consistency at the voxel level, we first match session two and session three's segmentation maps to session one's by permuting the cluster labels (this is not needed for the between-session RI, which is invariant to label permutation). Then we define a variance map as follows: the variance at certain voxels takes the value zero if the estimates of all three sessions have the same labels. The variance takes 1 if two of the three sessions have the same labels, and takes 2 if none of the estimates are the same. We then average the variance map across all subjects

and obtain a mean variance map. This map shows how the algorithm performs in term of consistency at the voxel level across all subjects. The results are shown in Figure 6.8. Image visualization is done by using nipy, a python package for neuroimaging data. We note that although K-Means and N-Cuts have low variance at the visual cortex, they have larger variance in most voxels of dorsal attention and the DMN. These findings confirm the different level of consistency between the functional networks, as has been shown in the original work of Zuo et al. [162]. Overall, the HMRF method's estimates have the lowest level of variance and hence the highest level of consistency.

6.4.6 Bootstrapping

In these experiments we aim to evaluate the performance of the three algorithms with bootstrapping. In the bootstrapping method, one covers the whole distribution of the estimator with the independent samples drawn from the original dataset with replacements, and estimates the stability of an algorithm [48]. An approximate solution of an algorithm is stable if the solution is not highly sensitive to the input data. It is unstable if a slight change in the data can cause the predicted values to change significantly. In this experiment, the bootstrapped samples can be seen as a small perturbation of the input data and will be used to test the algorithm stability.

There are various approaches for resampling the available data. One may resample the subjects from the original dataset [39]. Here for each voxel of each subject in session one, we sample with replacement from the 197 time points of preprocessed data, and obtain a bootstrap sample volume with the same BOLD signal length and number of subjects with the original dataset. The sampling is similar to the circular block bootstrap in Bellec et al. [12], except that we do not model the temporal correlation between time points. Since all methods under comparison here do not model temporal correlation, the shuffling of the time points has no effect on the segmentation results. After repeating the sampling 100 times, we obtain a set of 100 bootstrap samples, each of which includes all subjects' time series data. Then, all three segmentation methods are applied on each of the bootstrap datasets. We estimate group and subject level maps from each bootstrap dataset by using the three methods. All the estimated label maps are postprocessed by a label permutation routine to guarantee that the same networks have the same labels.

Figure 6.9 shows seven average group-level functional network maps across all bootstrap sampled data. For each network, we extract a binary map with voxel intensity taking 1 in that network and 0 outside. This binary map is then averaged over all bootstrap samples. We also show the variance of this binary label map over all samples in Figure

6.10. Small variance indicates more stability under bootstrap sampling. All three methods have moderate to high stability across bootstrap samples. For visual, motor, and DMN networks, K-Means and N-Cuts have reasonably high stability, although some voxels at the boundary of the network regions are labeled differently across bootstrap samples. For the attention, salience and executive control networks estimated by K-Means and N-Cuts, the ambiguity not only happens on the boundary of the network regions, but also on some bigger regions inside the networks. For example, in some bootstrap runs, K-Means incorrectly assigns the posterior cingulate cortex (PCC) to the attentive network (see the red regions in dorsal attentive in Figure 6.9), whereas PCC has been shown to be part of the DMN [71]. K-Means also miss part of the primary motor network. K-Means even in a few runs merges the brain stem into the DMN. For N-Cuts, the dorsal attentive, salience, and executive control networks have larger variance under this data perturbation. Compared to the other two methods, HMRF has the smallest variance, and hence the highest stability in all seven networks including the brain stem. A small number of voxels in motor and DMN still shows unstable assignments.

To demonstrate the stability of the estimates on each of the subject functional networks, we first pick 3 subjects from the 25 subjects in the dataset. For each subject, we show the average network patterns over all bootstrap samples. See Figure 6.11. We show only six physiologically meaningful networks, excluding the one corresponding to the brain stem. For each network, one representative slice is shown. From the figure, all three subjects' mean network maps have lower stability compared to their corresponding group networks. Certain subjects' networks are significantly less stable than other subjects, due to the various degree of perturbation by the random sampling even using the same bootstrapping procedure. Among the six networks, attentive networks exhibit the most dramatic change under bootstrap sampling. Some voxels of salience and executive control networks are absorbed into attentive networks. This misclassification happens most on subject 2, and also happens a moderate amount on subjects 1 and 3. Compared to the other two methods, HMRF is able to estimate reasonably stable functional networks even with data resampling. Attentive networks and executive control networks tend to change more than other networks, but still less than K-Means and N-Cuts.

Another way to show the stability of the subject label maps is the variance map. Since we are interested in comparing among three methods the variance of the networks across all subjects, we show the variance not for each single subject, but an average variance over all subjects. See Figure 6.12. Because of the averaging over all subjects, the variance is

more spread over the voxels. Again, HMRF shows significantly smaller variance than the other two methods, indicating its subject map estimates are more stable under bootstrap sampling.

6.4.7 Between-level links estimation

We also run the cross-validation and use the posterior predictive distribution in equation (6.7) for estimating the optimal α parameter. Figure 6.13 gives a plot of the average predictive density with alpha ranging in [0.15, 0.5], with interval 0.05. We found that with too small α , the model has low prediction values on the test data, and too large α values improve the prediction but are still not the optimal. The best α value is around 0.3 to 0.35.

6.5 Discussion

We proposed a new hierarchical model for identifying functional networks of the human brain from a group of rs-fMRI dataset. The model assumes a group functional network map is shared among all subjects in the population, and individual subjects' functional patterns are generated as variations from this group level network. If we see the functional network pattern as a clustering of the fMRI data, we actually assume the subject maps are samples from an unknown distribution of the clusterings, with its mean given by the group map. We reformulate the distribution of clusterings as a distribution of network labels, where a subject's labels at each voxel are seen as generated from the group's network labels. While the intersubject statistical dependency is defined by the links between group and subject labels, the spatial coherence of the functional networks is guaranteed by the within-subject MRF. All the network label variables at both levels with their links, and the parameters, are defined in an integrated graph, and the general techniques of graphical models can be used for (approximate) inference. This multilevel view is typically used in general statistical analysis when the individuals are grouped into units, and the variance of the variables is decomposed into the group-specific and subject-specific terms. We borrow this idea and apply it in a clustering problem where the intensities of voxels at each time point are grouped into units (subjects), and the vMF's κ parameter represents the individual subject's variance. The α parameter is equivalent to the pooling factor in the standard hierarchical linear model [63], and controls the degree to which the estimates of subject label maps are pooled together.

We use the MCMC sampling for the inference because of its good approximation of the posterior density. An alternative class of methods is variational inference, including mean field approximation and expectation propagation. Both variational methods and MCMC

are the approximation of the true posterior distribution. The former approximate the target distribution by the product of factorized distributions, and the latter achieve the approximation by Monte Carlo averaging. Both classes of methods depend on initial conditions. However, with Gibbs sampling, we obtain a full posterior density estimate of the network variables, while variational methods usually have point estimates. Besides, the derivation of the conditional expectation used for updating of variational methods would be cumbersome in our multilevel model. On the other hand, the Gibbs sampling is straightforward as the conditional probability is easy to compute in our Bayesian setting. Therefore, we choose Gibbs sampling due to its simplicity, as well as the fact that the application does not require real time computation. An additional critical property of the MCMC sampling is that its convergence does not depend on the dimension of the variables [128]; thus we can achieve reasonable compute time even in this million-dimensional problem. The whole Monte Carlo expectation maximization procedure uses 45-50 cores on a multiprocessor machine, and takes about 2 hours for a group of 25 subjects.

As a practical guide for applying HMRF, the introduction of within-subject MRF is not meant to replace the spatial smoothing in the preprocessing steps. This is one step further from what we found in our previous work [101], where no spatial smoothing is conducted when the HMRF model is used. In the simulated experiments, we found a moderate spatial smoothing, plus our HMRF model can achieve the best estimation accuracy. The best accuracy of the combined model is because moderate smoothing does help to decrease the noise, without overly impacting the signal at finer spatial scales. The MRF regularization further favors spatial coherence and intersubject coherence.

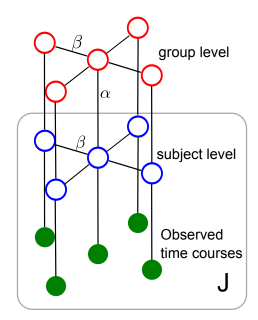


Figure 6.1: We define a MRF on a graph that includes the voxels of all subject maps as well as the group map. The set of edges includes the between-level links with weight α , and within-subject links with weight β . The square box on the subject level and time courses repeats J times the nodes in the square, representing all the subjects. Only the central voxels connection is shown for the between-level links, whereas in practice the links exist on all other voxels. The BOLD signal variables are shaded, meaning they are set to the observed value.

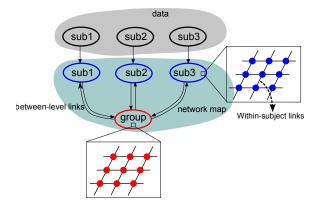


Figure 6.2: An alternative representation of the graphical model of the HMRF. A regular MRF is defined on the network variables within subject, and within group label maps. Then between-level links are added between the group voxel and each subject voxel at the same anatomical location. The added edges, together with the original edges, consist of a new graph which integrates two levels of variables.

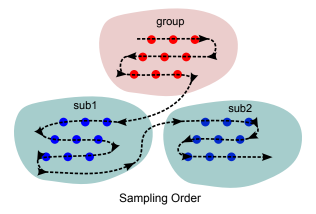


Figure 6.3: Gibbs sampling schedule on a high level view. The sampling scan of all voxels in the group before updating each subject. This schedule repeats until convergence.

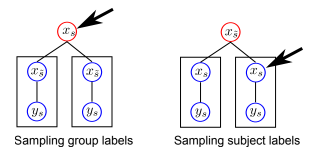


Figure 6.4: Gibbs sampling iterates between group and subjects. On the voxel-level, the sampler draws samples of one voxel given its neighbors that includes both with-subject and between-level neighbors.

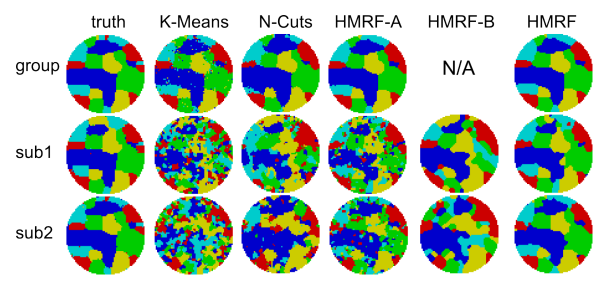


Figure 6.5: The estimated group and subject functional network label maps from various methods, as well as the ground truth maps. Only two are shown among the 25 subjects.

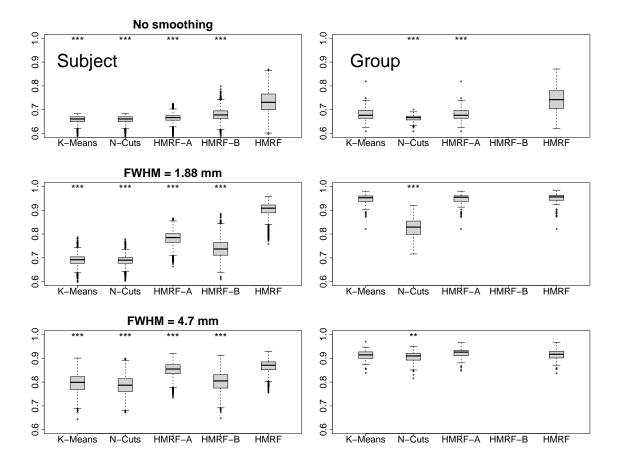


Figure 6.6: Box-and-whiskers plots of the estimation accuracies of all methods for three levels of spatial smoothing. The accuracies of subject labels are across all subjects and MC samples. The group map accuracies are across all MC samples. The upper and lower "hinges" correspond to the 25th and 75th percentiles. The asterisk on top of each box indicates the p-value of the standard two-tailed T test between HMRF and the corresponding method. No asterisk: significant p > 0.05; *: significant at p < 0.01; **: signi

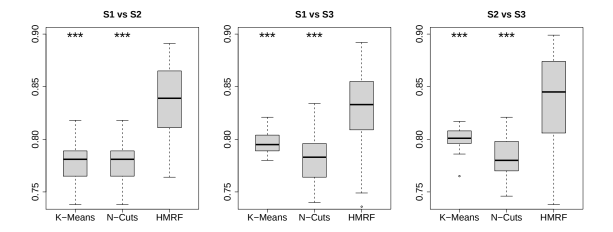


Figure 6.7: Box-and-whiskers plots of the RI value between each pair of sessions over the all subjects' label map. The bottom and top of the boxes are the 25th and 75th percentile, and the whiskers extend to the whole range of the data except the outliers.

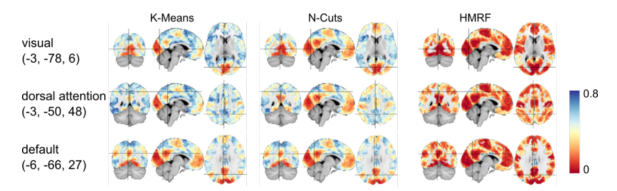


Figure 6.8: The intersession variance maps for three segmentation methods. The variance maps are obtained for each subject, averaged across subjects, and finally normalized to [0, 1]. A few voxels with intensity above 0.8 are rendered the same as those with intensity 0.8. This single map covers all seven functional networks, and we selectively show the slices corresponding to the three major networks. The image left is the subject's left, and we use the same convention in the following figures.

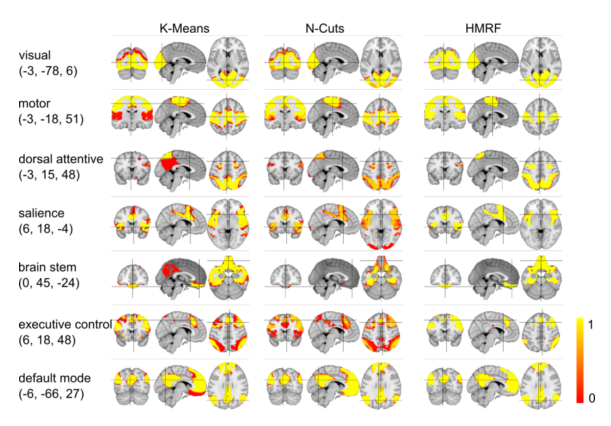


Figure 6.9: The group level's mean functional networks estimated from all bootstrapped data by three segmentation methods. The binary map of each network is averaged over all bootstrap samples. The average intensity ranges from 0 to 1.

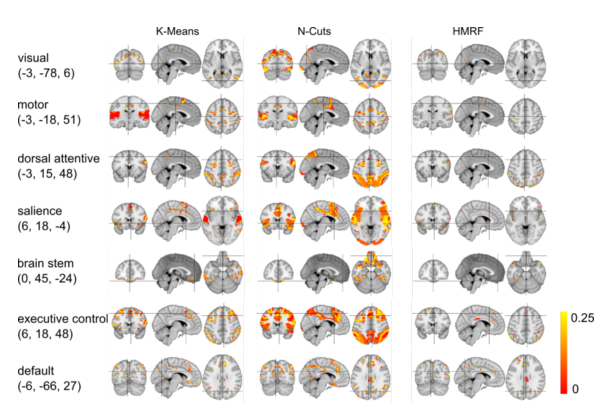


Figure 6.10: The group variance map estimated from all bootstrap data by the three segmentation methods. The variance ranges from 0 to 0.25.

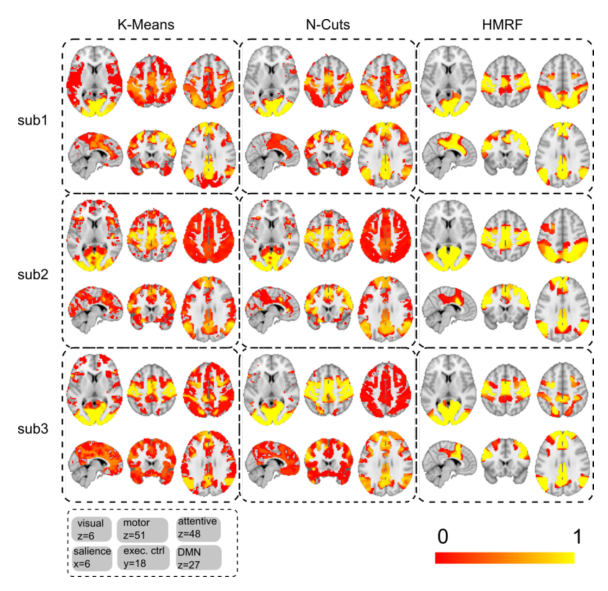


Figure 6.11: The three subjects' average network label maps estimated from all bootstrap samples. One representative slice is shown for each of the seven networks for each subject (row) and each method (column), excluding brain stem component. The average values range from 0 to 1.

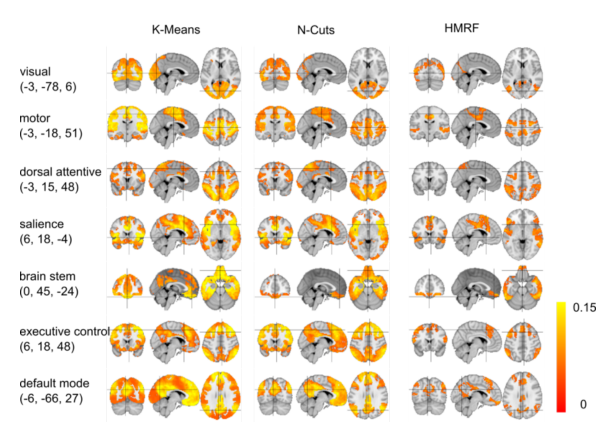


Figure 6.12: The subjects' variance maps estimated from all bootstrap samples. The maps are averaged across all subjects, and their values range from 0 to 0.25. The color map is in [0, 0.15] since most of the variance values fall into this range.

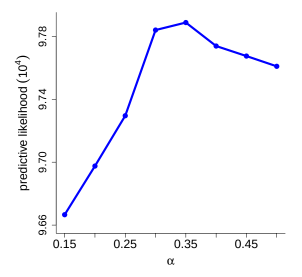


Figure 6.13: Estimation of parameter α with the average predictive distributions using the leave-one-out cross-validation. We use the data from only the first session of the NYU-TRT dataset but find similar patterns in the other two sessions. α are sampled between 0.15 and 0.5, with interval 0.05.

CHAPTER 7

GENERAL DISCUSSION

In this chapter we will give a summary of the dissertation work, with some discussions on the models we have used. We will also discuss the possible extension of our models in the future, and other general promising approaches of rs-fMRI study.

7.1 Summary of dissertation work

The analysis of functional networks using rs-fMRI is a difficult problem, due to various sources of noise, the artifact introduced by the preprocessing, and the variations between the subjects in the same group. In this dissertation we proposed a hypothesis that a hierarchical MRF model is able to describe the within-subject spatial coherence and the between-subject similarity, with higher estimation accuracy on simulated data and higher consistency on real data. The three applications of our models proved our hypothesis. More specifically:

- We proposed a high-dimensional MRF model to depict our prior information on the pairwise connectivity variables of a single subject rs-fMRI dataset. The MRF enforced similar values of the adjacent variables, where the adjacency is defined by the MRF structure. We are able to identify finer functional patterns without resorting to spatial smoothing.
- We also presented a *maximum a posteriori* framework for estimating multiple functional networks of a single subject, by using MRF in original image space, and a vMF distribution as likelihood function. Again, with the proposed method, we identified functional networks that are spatially coherent, and more consistent across subjects compared to other standard methods.
- Last, we extended the MRF model to a hierarchical setting, where both the withinsubject spatial coherence and between-subject similarity are modeled by a single graphical model. By sampling from the posterior distribution of the network variables,

we are able to show that the network maps are more accurate for synthetic data, and more consistent for real data compared to the flat model of group analysis.

Here we give a brief comparison of the three methods that we discussed in Chapters 4, 5 and 6. The pairwise connectivity estimation method in Chapter 4 is a lower level information extraction compared to other methods. The algorithm aims to estimate the functional connectivities between voxel pairs, without knowledge of the functional system. Because the connectivity variables we are interested in only have two states, connected or not connected, we deal with a unsupervised binary classification problem. We define a model similar to the standard Gaussian mixture (GMM) model, and estimate the posterior of the hidden variables. Our model is different from the regular GMM in that we define a prior distribution on the hidden variables, the connectivity. The prior distribution is indeed a MRF, or equivalently, Gibbs distribution, defined on a high-dimensional graph to represent our knowledge of the spatial soft constraints. The inference, as a result of this additional prior model, is significantly more difficult than GMM, since the inference of each pairwise connectivity cannot be factorized like standard GMM due to the statistical dependency on other variables. We use variational inference to approximate the posterior distribution of the connectivity variables and solved the intractable optimization problem.

The second method we discussed in Chapter 5 is in a higher level than the pairwise connectivity estimation in Chapter 4. The algorithm outputs a three-dimensional label map, and the labeling represents the functional systems. Compared to Chapter 4, here we are able to detect all functional systems without defining a seed region, and show them in a single spatial map. The algorithm is aware of the functional system, as each cluster represents one system. Although the ability to identify multiple systems is a strength compared to the previous pairwise connectivity estimation, it is hard to state the multiple network detection method is superior to the pairwise connectivity estimation method, as the two methods are at different levels. The lower-level pairwise connectivity estimated can be used in other models as input. For example, it can be used to define the ROIs for the graph-based analysis. Or, the prior distribution of the connectivity variables can be modified to include an additional parameter for multisubject analysis. The current prior distribution of the connectivity variables are defined by a MRF including only smoothing constraints. In the case of multiple subjects, we can assume the same pairwise connectivity variable for all subjects are random variables from a population distribution. By redefining the six-dimensional graph and including other subjects, and a group layer, we have a hierarchical model for the pairwise connectivity variables, much like what we have done in the HMRF model in Chapter 6.

Both the algorithms in Chapters 4 and 5 deal with single subject. By contrast, the hierarchical MRF model in Chapter 6 explores the functional patterns from the rs-fMRI of a group of subjects. It turns out if we integrate our knowledge of the within-subject smoothness and the between-subject similarity into the graph structure, we can convert the group inference problem into a standard inference of the posterior distribution. So we defined an abstract layer of the graph including all the gray matter voxels in the subject volumes, and a *virtual* volume of the group. Below this layer is the image voxels of the subjects and groups volume. Above the layer, we have a standard graph and we are able to apply a standard optimization algorithm on it. The single-subject MCEM method we proposed in Chapter 5 is a special case of our extended HMRF model. We have shown that the extended model is able to enforce the similarity of the functional patterns across subjects, as well as estimating a group map. Because a single subject can borrow statistical power from other subjects via the group level, we can estimate a reasonable map even from noisy single-subject data.

A classical occurrence of hierarchical models is the inclusion of the random-effects in the linear model [127]. For a simple example, suppose we randomly choose M schools from all the schools in the country, and randomly choose N students from each school and record their scores on an exam. We use y_{ij} to denote the score of the student n from school m. The score can be modeled by a linear model with random-effects $y_{nm} = \beta + \theta_m + w_{nm}$, where β is the average score of the population, θ_m is the school-specific random effect that represents the difference between the population's average score β and school m's score, and w_{nm} is the individual student's deviation from the school's average score. Both θ_m and w_{nm} are random variables since the schools and the students are randomly chosen. The above linear model can be decomposed into a hierarchical model as

$$y_{nm} = u_m + w_{nm}$$
$$u_m = \beta + \theta_m.$$

Here u_m is the average score for the school m, and is further decomposed into the fixed effects β and random effects θ_m . In our multilevel model for the rs-fMRI group analysis, the BOLD time series correspond to the student score y, the subject network label maps correspond to the average score u of the school, and the group label map corresponds to the population's average score β . vMF distribution denotes the random effects of the BOLD signals, hence is similar to the w in the above example. Because the parameters (called hidden variables in our model) are discrete values, there is no direct counterpart of θ in our

model, but the MRF and its equivalent Gibbs distribution represent the random properties of the group and subject label maps.

There are reasons of this decomposition and the formulation by Bayesian rules [127]. First, the two levels of models represent the prior knowledge of the metapopulation. The hierarchical models naturally appeared in the metaanalysis. In the case of fMRI study, the results of the analysis on other subjects can be used as a prior of current subjects. Second, the hierarchical model can separate the priors into two components. One component corresponds to the soft constraints applied on the subject level parameters (subject network label map), and the other component represents the uncertainty of such constraints, i.e., the distribution of the group label map, as well as the unknown weight parameters between the subject and the group. Also, the decomposition can often simplify the posterior inference of the hidden variable, at least on the concept level.

7.2 HMM versus CRF

The MRF, as a extension of the one-dimensional hidden Markov model (HMM), is a model that does not depend on the observed data. The conditional random field (CRF), on the contrary, has a regularization energy function that depends on the data. One of the reasons that the prior energy should be dependent on the data is that the smoothness assumption is often violated at the discontinuities, i.e., the edges in the image, be it natural scene images or fMRI images. When the BOLD signal at certain voxel is significantly different from one of its spatial neighbor's, we have reason not want to borrow information from this neighbor. Intuitively this makes sense, although in such a definition, the separation of the prior P(X) and the conditional probability P(Y|X) would not be possible, since P(X) also involves the data Y. The CRF is called adaptive smoothing in some literature [98].

Historically, the existence or lack of links between spatially adjacent nodes on a regular lattice is modeled by a line process by Geman [65]. The inference is iterated between estimating the MRF with image pixels at nodes, and estimating line process. At last, when both the MRF and the line process converge, we just discard the line process estimation but keep the estimation of the nodes in MRF. The estimated nodes will be from the marginal distribution according to the original MRF. However, even the links are modeled as a line process, the hidden variable X is still in a distribution that does not include Y. So fundamentally such model is the same with regular MRF, rather than a CRF. It is the optimization methods that differ.

CRF is a reasonable model in some situations, when the disparity of the neighboring voxels represents the discontinuity of the images. In some cases, if a voxel is very noisy, such

that its signal looks significantly different than its neighbors, while in fact the difference is due to large noise, the CRF may assume the discontinuity at this edge, and does not define, or defines a weak interaction. Therefore, the noisy voxel may not be correctly classified. In such situations, the data Y is essentially used twice, once in the prior distribution P(X), and the other in the likelihood function P(Y). Because we increase the weight of the data term, the CRF will fail to estimate the correct hidden variables when the data are noisy. In practice, in a multimodality image segmentation problem, we used CRF and found some noisy voxels were not labeled correctly because the CRF removed the edge between the noisy voxels and their neighbors due to the large intensity difference. One possible solution of this incorrect edge determination is to use a larger neighborhood when computing the discontinuity, thus decreasing the influence of the single noisy pixel [141].

In our model, we choose to use HMM instead of CRF with the belief that a simple Bayesian model will conceptually fit our problem and have good generalization power. It is worth applying the CRF model to our hierarchical MRF framework. Because of its Bayesian concept, HMM is easier to understand than the CRF. However, in some situations if the CRF outperforms HMM and Bayesian-based models, it will be an open question if we are willing to give up a little performance in order to obtain a simple model that we can understand.

7.3 Convergence rate of MCMC

One of the main disadvantages of the MCMC algorithm, or the general Monte Carlo sampling algorithm, is the slow convergence and long computation time. Although, the computation time is not a critical issue in the fMRI study, this is an interesting and actively evolving topic that is worth discussion. With the high-level data parallel frameworks such as MapReduce, the processing of large-scale data is greatly simplified. These data processing frameworks do not support the core data mining and machine learning algorithms. Many datasets have variables that are interdependent with each other, and a graphical model is a good abstract representation for these multivariate data. The statistical inference often involves computing the posterior probability of the variables given the observation. When the prior distribution and likelihood function cannot be defined as a conjugate distribution, one alternative method will be sampling from the posterior and use Monte Carlo averaging for the inference. In this parallel framework, it is natural to run the sampling on the parallel machine for faster convergence. The state-of-the-art method implemented in the GraphLab, the parallel graph abstract library [102], uses the graph coloring technique to divide the

nodes in the original graph into K subsets. Within the subsets, the nodes have the same color, and are updated in parallel. Between different colors of subsets, the nodes are updated in sequential. Such strategy is exactly the same with our implementation in Section 4.3, where we also divide the nodes into four subsets to avoid the checkerboard local minium.

Without multiple-core machines, a common method of accelerating the mixing of the Gibbs sampler is to update the variables in a batch mode, i.e., in a block. The Swendsen-Wang algorithm [148] and its extension [7] are in this category. The key to the block update is to find those variables that are strongly coupled with each other and can be updated together. A recent development along this line is by Hamze and Freitas [76]. In their work, the authors divide the nodes into two subsets such that each subset of nodes is a tree. A tree, as a special case of general graph, does not have a loop of edges, and it is therefore significantly easier to compute the posterior distribution. By using the Rao-Blackwell transformation, they give an algorithm that combines analytical and sampling steps. Given a sample of one tree, the algorithm uses belief propagation [156] to compute the exact distribution of the other tree conditioned on the sample. This block update scheme is proved to have better convergence property than stand Gibbs sampling. The algorithm of Gonzalez et al. [69] also uses a similar concept.

7.4 Future works

Graphical model and its undirected variants of MRF are powerful tools for modeling multivariate random variables and their uncertainty. Here we give possible extensions of our computational models to dynamical settings, to specific mental health disorders and to the integrated spatiotemporal models.

7.4.1 Dynamics of the functional network

In conventional brain connectivity analysis one assumes that the connections between regions are static and do not change with time. The functional connectivity may not be in a stationary state. Not like structural connectivity measured by DTI, the functional connections depend on the the cognitive activity of the human brain, and may change over time. For example, the most active regions during resting-state seem to be the regions that shows the greatest deactivation during external cognitive challenges. Besides, functional links can persist even without direct anatomical connection.

Zalesky et al. [158] tested the variation of the fMRI time series by splitting the time series into two parts, and used paired T test to test any difference between the two halves with respect to the various statistical measures, including variance, maximum and minimum

amplitude. They did not identify a significant difference. This suggests that there is no long-term variation on the time series, but this is not sufficient to suggest that the functional network is static.

One interpretation of the intrinsic activity, especially the dynamics of the patterns is an inner state of exploration where the brain generates predictions about the best possible network patterns that would be optimal for an impending future external event [40]. If that is the case, it will be beneficial to study the dynamics of the functional network for better understanding how the brain predicts external events and prepares for them. There have been works that model the complex network's dynamics with time delay, mostly notably the models proposed by Honey et al. [81], Deco et al. [41], Ferguson et al. [51] and Ghosh et al. [66]. In Honey et al. [81], the structure connectivity matrix is first computed, and neural dynamical potential signal is simulated. The neural potential has a high sampling rate. The functional connectivity is computed from this simulated time course signal at different temporal scales. BOLD signal is also simulated from the neural potential signal by the balloon model. The simulated BOLD signal has coarser spatial resolution compared the neural potential signal, i.e., BOLD is simulated per region. The authors concluded that functional connectivity estimated from whole neural potential signals are more consistent with structural network, while the network estimated from a segment of neural signals have more transient network patterns. In Ghosh et al. [66], the low level neural signal is also simulated similar to Honey et al. [81], and BOLD signal is computed from these simulated signals. The difference from the work of Honey et al. is that the authors added time delay between the regions, and the delay is proportional to the distance between the regions. For the study of fMRI, we do not have to assume a model for neural potential signal and simulate it. An direct way is to define a dynamic model for BOLD signal. To model the dynamics of the network, we can either explicitly model the time delay due to the transmission of the neural signal in the fibers, or we can choose not model this time delay. Honey et al. [81] do not model the time delay, which is reasonable due to the low temporal resolution of BOLD signal. The time delay, compared to the TR of BOLD signal, can be ignored.

These models mainly explore the dynamic patterns of a single subject, with the anatomical constraints applied on the possible functional structure. Although the dynamics may differ across subjects, it would intriguing to study a group of subjects' network dynamics and see if some patterns are shared among them. It would be a reasonable assumption that the parameters of the dynamic model are similar across subjects. For instance, if we assume a ROI swith between mental states with certain probability, the probability as a parameter

on the Markov chain, might be shared across subjects with some variation. Along this line of thought, we can even model the common parameters across subjects in a hierarchical way. To model the dynamics of the functional networks, some new techniques about network dynamics [18, 79, 91, 4, 111] are required besides our hierarchical model.

7.4.2 Functional connectivity in clinical study

The intrinsic activity, as the baseline signal for task-based analysis not only helps understanding subject's functional patterns for specific tasks, but also in studying mental disorders such as autism spectrum disorders (ASDs), attention deficit hyperactivity disorder (ADHD) and Alzheimer's [22, 82, 108, 3, 116]. For instance, in a review of an ADHD study [29], the author found the default model network has less anticorrelation for ADHD patients. Since the strong anticorrelation is believed to be related to better behavioral performance, such findings help the understanding and treatment of ADHD patients.

One particular interesting application of functional connectivity is the classification of ASDs and typical developing group. The classification of autism and control groups can be cast into a regression problem by defining the pairwise functional connectivity as independent variables and the autism diagnostic scores as the dependent variables. By this definition, we aim to use rs-fMRI data to predict autism patients. The correct prediction of autism patients is not the only goal. By solving the linear regression problem, it becomes possible to pinpoint one or more functional connectivity variables that are strongly correlated to autism clinical scores, and even the anatomical regions that are involved in these important connectivity variables. Because of connectivity variables are pairwise, the number of such variables is large. For a set of ROIs defined by Power et al. [123] with 264 nodes, the number of pairwise connectivity variables will be $264 \times 263/2 = 34,716$. For the denser ROIs used By Anderson et al. [3], there are 7,266 seed regions, and the number of pairwise connectivity variables will be 26, 397, 378. On the contrary, the number of observations (subjects) for such types of studies is much smaller. Therefore, we face the typical curse of dimensionality problem. Rich literature in the machine learning community are available for addressing such problems. An early and simple solution is feature selection, i.e., selecting only a subset of the independent variables and discarding others. The regression is performed between those chosen variables and the dependent variables.

To select k features with good predictability from n features is an intractable optimization problem due to the combinatorial number of choices. One can use univariate methods with a hypothesis test to tell if a feature can significantly separate apart the training sample set, or use more advanced methods taking into account the interactions of

the features [75, 133]. More modern methods integrates feature and variable selection and classification in a single framework, such as the L1 regularization methods, least absolute shrinkage and selection operator (LASSO) [58].

With the autism brain imaging data exchange (ABIDE) repository open to the public [47], thousands of rs-fMRI datasets of both autism patients and control groups from various sites are available for analysis. This data sharing initiative is an opportunity to explore the relationship between the functional connectivity and the ASDs in a large data cohort. However, due to the different source of fMRI data, a simple pooling of all the subjects in a one layer model may not work well, as has been shown by Nielsen et al. [116]. The authors use 964 subjects from 16 separate international sites and use the pairwise connectivity variables of all the subjects to predict the autism variable, with the age variable taken into account. Nielson et al. show the classification accuracy significantly outperformed chance but is much lower than their previous results on single site [3]. One possible reason is the heterogeneity of the datasets due to the different sites, scanner parameters and practitioners. We can extend our HMRF model to this problem by defining a hierarchical regression and classification problem. By the hierarchy, we assume the regression coefficients are also random variables, instead of fixed unknown parameters. The coefficients are from a prior distribution with hyper-parameters. With the hierarchical model, the regression is balanced between the estimation purely from the data, and the estimation of other sites. For example, if for one site, the training data has good quality and the variance of the regression coefficients are small, the posterior distribution of the coefficients of subjects in this particular site will be closer to the value that fits the data, i.e., a estimate from standard regression. If the data of one site is highly overlapped and nonseparable, the variance of the regression coefficients will be larger, so the posterior mean of the coefficients will be closer to the population's coefficients, which is estimated from other sites. In this way, each site has its own parameter to model the data quality, and we may achieve an overall higher rate of classification, and find more reliable correspondence between the function regions and Autism disorder.

7.4.3 Spatiotemporal modeling

One interesting model that can be used for this spatio-temporal analysis is the Gaussian process (GP). GP originates from linear regression problem $\boldsymbol{y} = \boldsymbol{w}^{\top} \boldsymbol{\phi}(\boldsymbol{x})$ with arbitrary basis function $\boldsymbol{\phi}(\boldsymbol{x})$. When the weight parameters \boldsymbol{w} are unknown fixed constants, the estimation of \boldsymbol{w} can be solved be least square estimation, or equivalently, maximum likelihood estimation when the data are Gaussian distributed. We can define the \boldsymbol{w} also as

random variables, and accordingly the response variable y will also be random. When \boldsymbol{w} is of Gaussian distribution, the joint probability $P(\boldsymbol{y},\boldsymbol{w})$, the conditional probability P(w|y), and the predictive distribution of y' for a new data point x' are also Gaussian, and therefore the statistical inference of y and y' can be done in a closed form. The standard linear regression with prior distribution defined on w is solved in parameter space, i.e., by searching the optimal parameter w in a space that can maximize the posterior distribution of w. GP adopts a function view in that we define a distribution on the function y(x). The variance of the functional value $y(x_n)$ and $y(x_m)$ can be represented by the inner product of the basis function $\phi(x_n)$ and $\phi(x_m)$, which can be defined directly by a kernel function of x_n and x_m . Most of the kernel function is defined such as the predicted value y' depends more on data points x that are closer to x', although the estimation of y' uses the y value of all the x in the training dataset. We can, however, make use of the Markov property and assume that y' only depends on the nearby y. In one-dimensional case, the auto-regressive (AR) model is a special case where the variable y only depends on the limited number of values of y in the past. In such a model, the covariance matrix of the function value y is dense, but the inverse covariance matrix, i.e., the precision matrix, is sparse, since it represents the conditional independence.

The extension to the spatiotemporal model has two possible options. We can either start from a time series model for each spatial data point, such as auto-regressive (AR), moving average (MA), or ARMA models, and assume the model parameters depend on their spatial neighbors. Or, we can assume a spatial model such as MRF, or Gaussian random field, and allow the parameters to change over time instead of being fixed. Such a model would be helpful for the analysis of fMRI data. For now, the spatial and temporal properties of BOLD signal of fMRI data are typically addressed separately. The temporal model is either represented by a frequency filter with the assumption of the interested frequency band, or by using a explicit model such as AR process. The spatial dependency is either enforced by a spatial Gaussian filter as a preprocessing step, or modeled by MRF such as the HMRF in our work. A unified spatiotemporal model will help the denoising, estimation, interpolation and prediction of the fMRI data.

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