APPLICATIONS OF MACHINE LEARNING TO LOCATION DATA

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Abstract

Positioning devices are generating location data at an unprecedented pace. Coupled with the right software, these data may enable a virtually unlimited number of valuable services. However, to build such software, there is a need for sophisticated algorithms that can extract the relevant information from location data. In this thesis, we use machine learning to develop such algorithms for three fundamental location-based problems.

First, we introduce a new graphical model for tracking radio-tagged animals and learning their movement patterns. The model provides a principled way to combine radio telemetry data with an arbitrary set of spatial features. We apply our model to real datasets and show that it outperforms the most popular radio telemetry software package used in ecology, produces accurate location estimates, and yields an interpretable model of animal movement.

Second, we develop a novel collaborative ranking framework called Collaborative Local Ranking (CLR), which is designed to solve a ranking problem that occurs frequently in the real-world but has not received enough attention in the scientific community. In this setting, users provide their affinity for items via local preferences among a subset of items instead of global preferences across all items. We justify CLR with a bound on its generalization error and derive an alternating minimization algorithm with runtime guarantees. We apply CLR to a venue recommendation task and demonstrate it outperforms state-of-the-art collaborative ranking methods on real datasets.

Third, we design two Bayesian probabilistic graphical models that predict users’ future geographic coordinates based on sparse observations of their past geographic coordinates. Our models intelligently share information across users to infer their locations at any future weekhour, determine the number of significant places and the spatial characteristics of these places, and compute the conditional distributions that describe how users spend their time at these places. We apply our models to real location datasets and demonstrate that, despite the sparsity, they provide accurate representations of users’ places and outperform existing methods in estimating users’ future locations.
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To my brother, Mert,
and my parents, Sait and Zerrin,
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Introduction

During the last few years, there has been a surge in the amount of location data that has been generated by positioning devices, which are devices that determine their users’ location. There are three reasons for this surge. First, recent advances in hardware have allowed positioning devices to be smaller and use less power [59], making it easier to embed them into mobile phones. Second, the US government’s “Enhanced 911” requirement has mandated US wireless carriers to provide accurate location estimates for emergency calls, accelerating the adoption of positioning technologies by mobile phone manufacturers [23]. Third, widespread adoption of positioning hardware has catalyzed the development of location-aware software, which are software that collect location data to perform their tasks. The culmination of these factors generated an unprecedented amount of location data in a few short years.

The number of valuable services that can be provided with such large amounts of location data is virtually unlimited. For example, with the right software, location data can prevent Alzheimer patients from getting lost, automatically deliver traffic alerts or targeted advertisements based on users’ future location, or predict the venues that users will visit and make any necessary reservations. However, collecting large amounts of location data is not sufficient to deliver such services; instead, we need algorithms that can convert raw location data into a more refined form these services require. Depending on the problem, this might mean predicting a user’s future location based on her past locations, or it might mean extracting a user’s preferences over venues she did not yet visit based on venues she did visit. These algorithms are not trivial to design, since a user’s location data is generated by complex processes, and these processes are not necessarily amenable to simple rules.
In this thesis, we use machine learning to design and analyze novel algorithms that leverage location data. As a field that is at the intersection of computer science, statistics, and applied mathematics, machine learning is the study of algorithms that improve their performance based on experience. In the general machine learning setting, one provides the algorithm with a training dataset, the algorithm processes it to extract a relevant hypothesis, and this hypothesis is set aside to be applied to future data. This workflow is preferable to the alternative where researchers extract the hypotheses themselves, because designing such hypotheses manually requires domain-specific knowledge and the hypotheses extracted might not easily transfer to new datasets. In contrast, machine learning methods can more easily adapt to new datasets as long as the assumptions made during the derivations of the algorithms are still valid.

When designing new algorithms, machine learning researchers focus on optimizing their efficiency and generalization ability. Algorithms are efficient when they are guaranteed to compute a desired solution in a reasonable amount of time, which is an important measure when the size of the training data is large. Algorithms generalize well when the hypotheses they extract from their training data also extend to data that will be processed in the future, such as generalizing from a user’s past venue visits to infer her preferences over venues she has not visited yet. In this thesis, we apply these theoretical concepts to each problem we study, in addition to evaluating our algorithms on synthetic and real location datasets.

In Chapter 1, we assist biologists on a fundamental problem in ecology by developing algorithms that model how animals move through their environment. For the past few years, biologists have been studying the plant and animal life that thrives on the Barro Colorado Island, which is a tropical island in the middle of the Panama canal. Some of these animals have been tagged with radio transmitters that allow researchers to track their location, but due to the interference caused by the rainforest canopy, the resulting signal is extremely noisy. Biologists have also logged the species and the location of the trees on the island, providing researchers with valuable features that describe the surrounding geography. In this project, we combine radio telemetry data and geographical data in a principled way using a state space model (SSM) to provide accurate estimates of the animals’ locations. Compared to previous work, our SSM comprises a richer movement model that in-
corporates spatial features, generalizing Gaussian random walk models that are associated with Kalman filters, and it yields an interpretable model that enunciates the relationship between the animal and its environment. In order to handle the challenges of incorporating a very large and feature-rich representation of movement, we derive an efficient stochastic gradient descent algorithm for learning the parameters of the model. We demonstrate the algorithm’s effectiveness by comparing it to the expectation-maximization (EM) algorithm, both via asymptotic analysis and synthetic experiments. Finally, we apply our model to real datasets and demonstrate that it outperforms the most popular radio telemetry software package used in ecology.

In Chapter 2, we turn our attention to collaborative filtering, and develop a new framework for ranking items. In typical collaborative filtering, a learning algorithm is provided with users’ ratings over items they have rated, and in return, the algorithm is expected to predict users’ ratings over items they have not yet rated. In contrast to supervised learning algorithms, collaborative filtering algorithms do not require explicit features to be prepared for the users or the items [28], making it easy to apply them to new domains. Furthermore, collaborative filtering algorithms achieve state-of-the-art performance on recommendation tasks, and consequently, are the method of choice for companies such as Amazon [34] and Netflix [5]. Despite their success, existing collaborative filtering methods are not designed for a common type of recommendation problem, one where users only provide local preferences instead of global preferences. This type of feedback exists in many real-world applications, two of which are movie recommendation and venue recommendation. For example, when users choose to watch a movie after reading the reviews of a set of candidate movies, they are implicitly preferring the chosen movie over the reviewed movies, as opposed to preferring the chosen movie over all other movies. Similarly, when users choose to visit a venue after exploring nearby venues through a mobile application, they are providing local preferences among nearby venues instead of providing preferences across all venues.

We call this problem collaborative local ranking and provide a formal characterization of it. We prove a Rademacher-based bound on the generalization error using a hypothesis class of low-rank matrices. We then transform the empirical loss into a tractable form, derive a simple alternating minimization for it, and prove that
each minimization step converges to the correct solution in time independent of the size of the training data. Finally, we demonstrate the performance of our algorithm on a venue recommendation task and show that it outperforms state-of-the-art collaborative ranking methods.

In Chapter 3, we develop probabilistic graphical models for location prediction, where the goal is to predict users’ future geographic coordinates based on their past geographic coordinates. Location prediction is a problem [8, 20] that underlies virtually all location-based tasks, and in fact, De Domenico et al. [12] recently won Nokia’s Mobile Data Challenge by devising a location prediction algorithm that predicts where the user is going to be during the next 24 hours. Despite being a problem of extreme interest, past work on location prediction has focused exclusively on dense datasets, where location data has been collected with high frequency. This is a valid assumption for certain datasets, such as those collected by cellular carriers; however, most location-aware applications can collect location data only when the user is actively using the application, either due to privacy restrictions or to save the device’s battery. For example, a reminder application might only record the user’s location when running in the foreground, and even though it collects very few location data points from the user during this period, it would nevertheless be expected to remind the user whenever she arrives at the target destination. Similarly, an application which delivers customized news based on the user’s location might not record any data when running in the background, but it would nevertheless be expected to deliver news alerts continuously based on the user’s estimated location. In fact, as users expect applications to be more intelligent and as location-aware applications collect location data more judiciously, there will be a growing need for location prediction algorithms that can leverage sparse datasets.

In order to estimate users’ locations from sparse data, we design two novel Bayesian probabilistic graphical models. Our models intelligently share information across users to infer their locations at any future weekhour even when the user has not been previously observed on that day or at that hour. In addition to estimating users’ locations, our models infer the number of significant places for each user, infer the spatial characteristics of these places, and compute the conditional distributions that describe how users spend their time during each weekhour. We derive a fast collapsed Gibbs sampler for estimating model parameters and demon-
strate that our models not only infer significant places accurately, but also achieve state-of-the-art predictive performance in estimating users’ future locations.

The research described in Chapter 1 was published as Kapicioglu et al. [27], and it is joint work with Robert Schapire, Martin Wikelski, and Tamara Broderick. The researches described in Chapters 2 and 3 are currently under submission, and they are joint work with David Rosenberg, Robert Schapire, and Tony Jebara.
Chapter 1

Learning Animal Movement Models

1.1 Background

Animals move through their environments in complex ways. Understanding the processes that govern animal movement is a fundamental problem in ecology and has important ramifications in areas such as home-range and territorial dynamics, habitat use and conservation, biological invasions and biological control [25]. Ecologists rely heavily on collecting and analyzing animal movement data to deepen their understanding of these processes.

In recent years, various technological advances, such as radio telemetry systems and the Global Positioning System (GPS), have created new avenues for collecting data from animals [57]. In radio telemetry, animals are tagged with a tiny radio transmitter. At regular time intervals, fixed-location towers in the environment record the signal from the transmitter and use this information to infer the direction of the animal with respect to the tower. In contrast to GPS, radio telemetry systems can use smaller transmitters, can collect data more frequently, are simpler to implement, and can be used under rainforest canopies. An implementation of such a system in Barro Colorado Island, called the Automated Radio Telemetry System Initiative (ARTS), provides researchers access to hundreds of thousands of directional data measurements collected from a variety of animals [10]. However, even though this method has led to a proliferation of directional data measurements,
it has been difficult to harness the full potential of these datasets because: 1) directional measurements obtained through radio telemetry are notoriously noisy; 2) telemetry databases may contain very large amounts of data; and 3) directional measurements are not in a form that is easily interpretable. What is needed are computational tools that would efficiently and accurately convert these large and noisy datasets into a form that would enhance ecological research.

Previously, various sequential probabilistic graphical models have been proposed to solve this problem. Anderson-Sprecher and Lodelter [1, 2] used an iterated extended Kalman filter-smoother to estimate animal locations. Jonsen et al. [26, 25] represented animal movements as correlated random walks and used Bayesian techniques to infer posterior model parameters and animal locations. Ovaskainen [42] used a diffusion approach to model movement in heterogeneous landscapes. Morales et al. [38] fitted multiple random walks to animal movement paths and modeled switching probabilities between them as a function of landscape variates. Patterson et al. [43] and Schick et al. [48] further review past probabilistic graphical models and their applications to ecology and animal movement.

In this work, we propose a new state space model (SSM) approach. In contrast to previous work, our SSM includes a richer and more general animal movement model. It can utilize arbitrary geographical features, such as the population densities of various tree species or the location of water sources, to represent geographically-dependent animal movement models that cannot be represented by previous approaches at this generality. While our animal movement model also generalizes Gaussian random walks, which are models associated with Kalman filters, combining and incorporating spatial features, especially geographical ones, yields an interpretable model that enunciates the relationship between the animal and its environment. Furthermore, a richer model leads to increased accuracy in estimating animal locations from telemetry data.

In order to handle the challenges of incorporating a very large and feature-rich latent state space, we present an efficient stochastic gradient algorithm for learning the parameters of the model. We demonstrate the algorithm’s effectiveness in training our model by comparing it to the expectation-maximization (EM) algorithm both via asymptotic analysis and synthetic experiments. We note that, in contrast to sequential Monte Carlo methods such as particle filters, our inference
algorithms allow us to train parameters and infer past estimates based on all observations.

Finally, we apply our model to real datasets and demonstrate that it outperforms the most popular radio telemetry software package used in ecology. In conclusion, we show that our methods aid us in producing interpretable results and accurate animal location estimates from large and noisy telemetry datasets.

This chapter is organized as follows. In Section 1.2, we specify the SSM and formalize its parameter and state estimation problems. In Section 1.3, we detail the EM, stochastic gradient descent, and Viterbi algorithms for solving the estimation problems and analyze their time-complexity. We report the results of our experiments on synthetic and real datasets in Section 1.4 and conclude in Section 1.5.

The research described in this chapter was published as [27], and it is joint work with Robert Schapire, Martin Wikelski, and Tamara Broderick.

1.2 A State Space Model

Our telemetry datasets comprise a long sequence of directional measurements that are observed by a small number of fixed-location towers at regular time intervals. Note that all directional measurements in a given telemetry dataset are received from the same animal. We model the data as if it were generated by an SSM. In this section, we describe the SSM by detailing its latent state space as well as its start, transition and observation models. Then, we formalize the parameter and state estimation problems, and argue how solving the estimation problems yields an interpretable model of animal movement and accurate animal location estimates.

Intuitively, the latent state space is the space of all possible animal locations. In order to use the SSM machinery, we discretize a continuous latent state space of animal locations into a finite and discrete space of coordinates $Q \subset \mathbb{R}^2$. In practice, $Q$ is constructed by partitioning a larger state space into finitely many equally sized grid cells and assigning each grid cell midpoint as a coordinate in $Q$.

The start model generates the first state of the latent animal trajectory. For simplicity’s sake, we use the uniform distribution as the start model. Let the latent random state at time step $t \in \{1, \ldots, T\}$ be denoted by $x_t \in Q$. Then, the start model
The transition model, a Gibbs distribution, generates the rest of the latent animal location data. Gibbs distributions, also known as the conditional exponential model, have emerged as popular models in machine learning due to their practical success and their theoretical elegance. These distributions are defined using features, which in our case are functions that encode information about the spatial properties of the environment. For example, a feature may encode the distance between two coordinates in the latent state space, or it may encode the minimum distance to a certain tree species. In a slightly different but related setting, it has been shown that the maximum likelihood estimate of a Gibbs distribution is equivalent to the maximum entropy distribution with various constraints imposed by the features [6]. In our case, we adopt the Gibbs distribution as the transition model because it provides a means to incorporate spatial features without making any independence assumptions about them, remains resilient to the extension of feature space by irrelevant features, and generalizes discrete versions of simpler random walk models which have previously been used to model animal movement. More formally, let \( f_k : Q \times Q \to \mathbb{R} \) denote the \( k \)th feature and \( \lambda_k \in \mathbb{R} \) denote the corresponding weight parameter, where \( k \in (1, \ldots, K) \). We use \( \lambda \) as a vector representation of feature weights. Then the transition model is

\[
p(x_{t+1}|x_t; \lambda) = \frac{\exp \left( \sum_{k=1}^{K} \lambda_k f_k(x_t, x_{t+1}) \right)}{\sum_{x \in Q} \exp \left( \sum_{k=1}^{K} \lambda_k f_k(x_t, x) \right)}.
\]

The observation model generates the directional measurements observed by the towers. We model the behavior of towers as a von Mises distribution, a circular analogue of the normal distribution [36]. A von Mises distribution is parameterized by \( \mu \) and \( \kappa \), roughly the analogs of the mean and the variance of a normal distribution, respectively. Intuitively, a radial bearing is sampled from a von Mises distribution, and it is added as noise to the true bearing that points in the direction of the animal. More formally, fix a radial bearing system shared by all towers (i.e., bearing
 Let \( y_{t,n} \in [-\pi, \pi) \) denote the random variable, a radial bearing, observed by tower \( n \in \{1, \ldots, N\} \) at time step \( t \). This is a noisy observation that is supposed to point in the direction of the animal. Let \( z_n \in \mathbb{R}^2 \) denote the coordinate of tower \( n \) and define \( h(x, z_n) \in [-\pi, \pi) \) to be the true radial bearing of the vector that points from tower \( n \) towards location \( x \). Let \( \mu_n \in \mathbb{R} \) and \( \kappa_n \geq 0 \) denote the parameters of the von Mises distribution for tower \( n \). Let \( I_0 \) denote the modified Bessel function of the first kind with order 0, which simply acts as the normalization constant for the von Mises distribution. Finally, let \( \theta, \sum, \mu, \kappa \) be the vector representations of the corresponding parameters and random variables. Then, the observation model is

\[
p ( y_t | x_t; \mu, \kappa ) = \prod_{n=1}^{N} p ( y_{t,n} | x_t; \mu_n, \kappa_n ) \\
= \prod_{n=1}^{N} \exp \left( \kappa \cos \left( \theta_{t,n} - h(x, z_n) - \mu_n \right) \right) / 2\pi I_0(\kappa) .
\]

(1.3)

The factorization occurs because of the conditional independence assumptions that hold between the observations.

The start, transition, and observation models can be used to compute all the marginal and conditional probability distributions of the SSM. Let \( y \) denote the vector of all observations and let \( \theta = (\lambda, \mu, \kappa) \) denote the vector of model parameters. As usual, the joint probability distribution of the SSM is

\[
p ( x, y; \theta ) = p ( x, y; \lambda, \mu, \kappa ) \\
= p ( x_1 ) \prod_{t=1}^{T-1} p ( x_{t+1} | x_t; \lambda ) \prod_{t=1}^{T} p ( y_t | x_t; \mu, \kappa ) .
\]

(1.4)

Parameter estimation is the problem of estimating the model parameters (i.e. \( \theta \)). We choose maximum likelihood estimation (MLE) as the method of fitting model parameters to data. We define the problem of parameter estimation as

\[
\hat{\theta} = (\hat{\lambda}, \hat{\mu}, \hat{\kappa}) = \arg\max_{\lambda \in \mathbb{R}^K, \mu \in \mathbb{R}^N, \kappa \geq 0} \log p ( y; \lambda, \mu, \kappa ) .
\]

(1.5)

State estimation, on the other hand, is the problem of estimating values of unobserved random variables. We choose to seek the hidden state sequence that max-
imizes the probability conditioned on the observations and the MLE parameter estimates. More formally, we define the problem of state estimation as

\[ \hat{x} = \arg\max_{x \in Q^T} p(x|\hat{\theta}). \]  

(1.6)

Our intention is to use the parameter estimates of the Gibbs distribution as an interpretable model of animal movement, and the state estimates of the latent random variables as animal location estimates. We note that an analysis of Gibbs weights might not conclusively explain the processes governing animal movement, but we hope that it will be a first step in further exploration of these processes.

### 1.3 Algorithms

In this section, we detail the algorithms for solving the estimation problems and analyze their time-complexity. Later in the chapter, we will see that for the type of datasets we have, where both the cardinality of the latent state space and the number of time steps is large, but the cardinality of the feature space is small, the stochastic gradient algorithm is asymptotically superior to EM. The analysis in this section will allow us to compare EM and the stochastic gradient algorithms.

#### 1.3.1 Expectation Maximization

**Algorithm 1.1 EM(\(\theta^0\), maxTime)**

1: \(i = 0\).
2: for all until elapsed time exceeds maxTime do
3: \(\forall t \in \{1, \ldots, T\}, \text{compute } \log p(x_t|y; \theta^i)\).
4: \(\forall t \in \{1, \ldots, T-1\}, \text{compute } \log p(x_t, x_{t+1}|y; \theta^i)\).
5: \(\forall n \in \{1, \ldots, N\}, \text{compute } (\mu_{n}^{i+1}, \kappa_{n}^{i+1}) \leftarrow \arg\max_{\mu_n, \kappa_n} \sum_{t=1}^{T} E_{x_t|y; \theta^i} \left[ \log p(y_t, x_t|\mu_n, \kappa_n) \right]\).
6: \(\lambda^{i+1} \leftarrow \arg\max_{\lambda} \sum_{t=1}^{T} E_{x_t|y; \theta^i} \left[ \log p(x_{t+1}|x_t, \lambda) \right]\).
7: \(i \leftarrow i + 1\).
8: return \(\theta^i\).

In Algorithm 1.1, we present the adaptation of EM to our setting. Here, maxTime denotes the maximum allowed running time and \(\theta^0 = (\lambda^0, \mu^0, \kappa^0)\) denotes
the initial parameter settings. Lines 3 and 4 describe what is traditionally called the E-step and involve computations of various conditional log-probabilities. These log-probabilities can be computed using algorithms such as the forward-backward algorithm. Similarly, Lines 5 and 6 describe what is traditionally called the M-step and involve maximization of expected complete log-likelihoods. The maximization problem in Line 5, after substitution of Equation (1.3) and some trigonometric manipulations, can be rewritten as

$$\arg\max_{\theta} \sum_{t=1}^{T} \kappa_n \cos \mu_n \mathbb{E}_{x_t \mid y_t; \theta} \left[ \cos \left( y_{t,n} - h(x_t, z_n) \right) \right]$$

$$+ \sum_{t=1}^{T} \kappa_n \sin \mu_n \mathbb{E}_{x_t \mid y_t; \theta} \left[ \sin \left( y_{t,n} - h(x_t, z_n) \right) \right] - T \log 2\pi I_0(\kappa_n).$$

In this form, it is equivalent to the problem of finding MLE estimates of von Mises parameters from i.i.d samples, and it can be solved in closed form using techniques from directional statistics [36]. The maximization problem in Line 6 is convex and unconstrained, and it can be solved using numerical optimization algorithms such as BFGS [35, 40].

EM is an iterative algorithm and it is hard to predict beforehand how many iterations are necessary until convergence. Here, we analyze the time-complexity of each iteration for a direct implementation of Algorithm 1.1. When it is clear from context, we write $Q$ for $|Q|$. We also remind the reader that $Q$ denotes the latent state space, $K$ denotes the number of features, and $T$ denotes the length of the animal trajectory. Almost all EM computations rely on the transition matrix, which can be computed in $O(Q^2K)$. The E-step, Lines 3 and 4, consists of computation of conditional log probabilities, which takes $O(Q^2T)$. The M-step, Lines 5 and 6, consists of both iterative and non-iterative optimization algorithms for estimating Gibbs and von Mises parameters, respectively. Optimization of von Mises parameters can be computed in $O(QT)$, whereas optimization of Gibbs parameters is itself conducted iteratively using BFGS. However, in practice, the number of BFGS iterations is small. For each BFGS iteration, it takes $O(Q^2T)$ to evaluate the objective function and $O(Q^2K)$ to evaluate its gradient. Finally, we note that we suppressed the dependence on $N$, the number of towers, since it is small. Overall, each EM iteration takes $O(Q^2(K + T))$. 

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For solving the problem of state estimation, Equation (1.6), the Viterbi algorithm may be used. Viterbi is not iterative and it only needs to be executed once. Once the transition matrix is computed in $O(Q^2K)$, the most likely path is computed in $O(Q^2T)$, yielding, like EM, a total running time of $O(Q^2(K+T))$.

An important factor in determining both computational efficiency and statistical accuracy is the resolution of the grid used to construct $Q$, the latent state space. As the diagonal distance between corners of each grid cell in $Q$ gets shorter, the latent state space representation becomes more accurate\(^1\) linearly, whereas the cardinality of the latent state space increases quadratically. Thus, an increase in statistical accuracy at a linear rate is offset by an increase in computational complexity at a \textit{quartic} rate, since EM and Viterbi themselves have time-complexity quadratic in the cardinality of the latent state space. This trade-off leads to a rapid increase in the running time of these algorithms in exchange for small gains in statistical accuracy.

\subsection*{1.3.2 Stochastic Gradient}

In this subsection, we propose a stochastic gradient alternative to EM, which under certain conditions, is asymptotically superior to EM. The algorithm is an optimization method that can be used when one has access to a noisy approximation of the gradient of the objective function [51]. Younes [58] used it to estimate parameters of partially observed Markov Random Fields. Delyon et al. [13] proved convergence results for a family of stochastic approximations to EM, including the stochastic gradient algorithm of the form depicted in Algorithm 1.2. In the rest of the subsection, we present our implementation of the stochastic gradient algorithm, analyze its time-complexity and compare it to that of EM, and detail how our implementation differs from other implementations.

The algorithm is presented formally as Algorithm 1.2. Here, $\theta^0$ denotes the initial parameter settings, maxTime denotes the maximum allowed running time, and numBurn denotes the number of Markov Chain Monte Carlo (MCMC) iterations executed before a sample obtained through Gibbs sampling is accepted. The

\footnote{One measure of accuracy is the diagonal distance between the corners of a grid cell. In the worst-case scenario, a larger grid cell would yield a higher absolute error between the location estimate and the true location even when the true location is estimated by the closest cell midpoint.}
Algorithm 1.2 SG($\theta^0$, numBurn, maxTime)

1: $i \leftarrow 0.$
2: for all until elapsed time exceeds maxTime do
3: Sample $x^i \sim p(.|y; \theta^i)$ using randomized Gibbs sampling with a burn-in period of numBurn.
4: $\theta^{i+1} \leftarrow \theta^i + \gamma_i \nabla_{x^i} L_x(\theta^i)$, where $\gamma_i$ is chosen to satisfy Wolfe conditions with respect to $L_x(\theta^i)$.
5: $i \leftarrow i + 1.$
6: return $\theta^i$.

The log-likelihood of the complete data, denoted $L$, is defined as

$$L_x(\theta) = L_x(\lambda, \mu, \kappa) = \log p(x, y; \lambda, \mu, \kappa).$$  \hfill (1.7)

In Line 3, latent random variables are sampled from the conditional distribution using Gibbs sampling, and in line 4, model parameters are updated in the direction of the gradient. We note that the algorithm is a stochastic gradient algorithm because the gradient of the log-likelihood of the complete data, Equation (1.7), is a noisy approximation of the gradient of the objective function, Equation (1.5).

The main difference between our implementation of the stochastic gradient algorithm and previous implementations is the choice of the learning rate $\gamma$. Common implementations of the stochastic gradient algorithm use a decreasing learning rate sequence that guarantees the convergence of the algorithm to a stationary point [13]. However, in practice, the choice of the constant factor associated with such sequences significantly influences the speed of convergence and it is a difficult task to identify the optimal factor. We tune the learning rate automatically by choosing one that satisfies the Wolfe conditions. These are line search conditions that guarantee the convergence of gradient updates to an optimum in deterministic convex optimization settings [40], but in our case, we use them in a stochastic optimization setting. In practice, parameters that satisfy these conditions can be found by providing the objective function and its gradient to line search methods.

Even though we lack a proof of convergence, we have observed informally that our criterion for selecting the learning rates ensures better performance than manually choosing learning rates that satisfy the customary assumptions. Furthermore, our approach provides an automated way to set the learning rate based on the dataset.
We conclude the subsection with an analysis of the time complexity of our implementation of Algorithm 1.2. Line 3 takes $O(Q(B + VK))$, where $B$ is the burn-in period and $V$ is the number of unique states visited during sampling. In order to achieve this complexity, we compute the transition probabilities only for the sampled states, and use memoization to store and recall them. Line 4 is conducted iteratively, due to the line search that involves finding the right learning rate, but number of iterations is very small in practice. Computing the objective function takes $O(QVK + T)$, and computing the gradient takes $O(K(QV + T))$.

In conclusion, the choice between EM and the stochastic gradient algorithm depends on the relative size of the latent state space, the feature space, and the number of time steps; however, in our problem, where both the cardinality of the latent state space and the number of time steps is large, but the cardinality of the feature space is small, the stochastic gradient algorithm is asymptotically superior.

An asymptotic comparison of each iteration does not suffice to determine which algorithm to use; thus in the next subsection, we perform empirical tests that compare these algorithms with respect to a variety of performance measures. Also, we note that both algorithms may be optimized such that only state transitions to a small neighborhood are taken into account in the computations. This would reduce time-complexity from $O(Q^2)$ to $O(QR)$, where $R$ is the cardinality of the largest neighborhood. Our implementation of all discussed algorithms take advantage of this optimization.

### 1.4 Experiments

In this section, we demonstrate our model on both synthetic and real datasets. First, we generate synthetic datasets to evaluate our model in a setting where true feature weights are known. We also use the synthetic datasets to compare the performance of the EM and the stochastic gradient algorithms. Then we use the real datasets to compare our model with LOAS, the most ubiquitous radio telemetry software package used in ecology.
1.4.1 Synthetic Datasets

In order to simulate a real-world application as closely as possible, we created a
virtual island that has approximately the same dimensions and the same tower
locations as the Barro Colorado Island. We partitioned the virtual island into 3654
grid cells (latent states), where each grid cell is 100 \times 100 meters square. Then
we created 10 different animal movement models (transition models). Each animal
movement model consisted of 5 features whose function values were generated uni-
formly at random from \([0, 1]\), and each feature’s weight was generated uniformly at
random from \([-10, 10]\). For each animal movement model, we generated an animal
path of length 1000. In order to be as realistic as possible, we constrained the ani-
mal to move at most 500 meters at each time step and normalized the probabilities
of the animal movement model accordingly. We also used the same constraint dur-
ing the parameter estimation. As for the von Mises parameters of the towers, we
set each \(\mu\) to 0 and \(\kappa\) to 15, which approximates a normal distribution with a stan-
dard deviation of 15 degrees. For each of the 10 animal paths, we generated the
corresponding noisy bearings, and together with the corresponding features (but
without the feature weights), provided them as input to the parameter estimation
algorithms.

In the first batch of experiments, we compared the performance of the stochastic
gradient algorithm and the EM algorithm. For both algorithms, we set their initial
Gibbs weights to 0, which corresponds to a uniform transition model, and their ini-
tial \(\kappa\) parameters to 50. For the stochastic gradient algorithm, we used a burn-in
period of 100,000. We executed the algorithms on each dataset for 10 hours. To
evaluate their performance, we measured the arithmetic mean error of the loca-
tion estimates, the Euclidean distance between the learned weights and the true
weights, and the log-likelihood of the observed bearings. We used the Euclidean
distance between the weights as a measure of the interpretability of the model,
where the closer weights were considered more interpretable.

As suggested by the asymptotic analysis, the stochastic gradient algorithm ex-
ecuted many more iterations than EM, and outperformed EM with respect to both
the arithmetic mean error of the location estimates and the Euclidean distance be-
tween the learned weights and the true weights. The stochastic gradient algorithm
also attained a higher log-likelihood than EM. We display the results, averaged
over the 10 datasets, in Figure 1.1.

In the second batch of experiments, we compared the performance of our animal movement model with a discrete version of the animal movement model used by Anderson-Sprecher [1]. By doing so, we both wanted to demonstrate how our model can generalize previous animal movement models and we wanted to observe whether having a richer model leads to an improvement in the accuracy of location estimates. The animal movement model used by Anderson-Sprecher is an isotropic bivariate Gaussian random walk which was trained using an extended Kalman filter-smoother. His model is defined as

\[
p(x_{t+1}|x_t; \sigma^2) = \frac{1}{2\pi\sigma^2} \exp\left(-\frac{\|x_{t+1} - x_t\|^2}{2\sigma^2}\right).
\]  

(1.8)

We can represent the same model approximately as a Gibbs distribution using the single distance-based feature \( f_{\text{dist}} = -\frac{1}{2} \|x_{t+1} - x_t\|^2 \). Similar to the first batch of experiments, we executed two copies of the stochastic gradient algorithm on the same datasets; one copy used the features that generated the datasets and the other copy used the single feature that represents the bivariate Gaussian random walk.
The richer model outperformed the Gaussian random walk model with respect to the accuracy of the location estimates. We display the results of these experiments, averaged over 10 datasets, in Figure 1.2.

1.4.2 Real Datasets

We applied our model to the radio telemetry data collected from two sloths, named Chispa and Wendi, that live on the Barro Colorado Island. These sloths were one of the few animals on the island whose true locations were labeled every few days by human researchers via GPS devices. Thus, we were able to use the radio telemetry data to train our algorithms and the GPS data to test them.

The radio telemetry data was collected every 4 minutes for 10 days, yielding approximately 3600 time points. Both datasets had considerable noise in the bearing measurements. In order to apply our model, we discretized the island into 1200 grid cells, each of size 50 × 50 meters square. As features, we used both the distance-based feature that encodes a bivariate random walk, Equation (1.8), and tree-based features that encode the change in the population density of various tree species across grid cells. In particular, the model included 18 features: 1 distance-based feature and 17 tree-based features. Each of the 17 tree-based features corresponded
to a different tree species and they were normalized to have values in $[-1, 1]$. We initialized the Gibbs weights to 0, the $\mu$ parameters to 0, and the $\kappa$ parameters to 10. We executed 10 different instances of the stochastic gradient algorithm, each using a different random seed, and averaged the results. As a baseline comparison, we used LOAS, which is the most popular radio telemetry software package used in ecology [31].

In Figure 1.3, we display the distance error obtained by SSM and LOAS on the two datasets. For both datasets, SSM outperformed LOAS. In Figure 1.4, we show how the feature weights evolved while training on the Wendi dataset. Our model successfully learned a diminishing movement variance for the sloth, represented in the figure as the growing weight associated with the distance-based feature. It also identified a small portion of the tree species that the sloth seems to have a preference for. In Figure 1.5, we display the true locations, SSM estimates, and the LOAS estimates obtained on the Wendi and Chispa datasets. SSM estimates are clustered much more closely to the true locations than the LOAS estimates.

We also generalized our movement model features to be temporally-dependent. In particular, we partitioned each day into different time zones, and for each such zone, we allowed the animals to move according to a different movement model.
Figure 1.4: The evolution of the transition weights for the Wendi dataset. We only plotted the weights that exceeded the 0.5 threshold. "D" and "T" denote the distance-based and tree-based features, respectively.

The changes in the daily activity of the sloths have already been studied by field biologists; so for our experiments, we set the time zones, "day" and "night", based on biologists' feedback.

In our last experiment, we were interested in evaluating whether feature weights converge to meaningful values, whether they are interpretable, and how different types of features interact with each other. For this purpose, we created an artificial tree type, where for each of the very few locations GPS data was available for Wendi, we placed a virtual tree. We defined this tree-feature formally as $f_{\text{tree}} = -\frac{||\text{tree}_{\text{dist}}(x_{t-1}) - \text{tree}_{\text{dist}}(x_t)||^2}{2}$, where $\text{tree}_{\text{dist}}(x)$ is the Euclidean distance between $x$ and the closest tree of that type. By defining the tree-based feature this way, we set it to the "same" unit as the distance-based feature associated with Equation (1.8), allowing the feature weights to be numerically comparable. After normalizing all feature values to be within $[-1, 1]$, the weights converged to 0.704 for distance-based feature at night and 0.316 during day; −1.279 for tree-based feature at night and −1.312 during day. As expected, interpreting distance-based weights indicate that Wendi moves more during day than night and that the positive magnitude of these weights indicate that the animal is more likely to stay in place in succeeding time steps. Also, tree-based weights correctly indicate that
Figure 1.5: Plots of the true locations, SSM estimates, and LOAS estimates on the two animal datasets. SSM estimates are based on the last stochastic gradient iteration. Top plot displays the results for the Wendi dataset and the bottom plot displays the results for the Chispa dataset.
Wendi has a strong preference for certain neighborhoods on the map (i.e. the true locations), and that the strength of these preferences are indifferent to the time zone.

1.5 Discussion

In this chapter, we presented an SSM approach to locate radio-tagged animals and learn their movement patterns. We presented a model that incorporates both geographical and non-geographical spatial features to improve animal location estimates and provide researchers an interpretable model that enunciates the relationship between the animal and its environment. We showed that the model generalizes discrete versions of random walk models and demonstrated empirically that a richer model improves animal location estimates. We also provided a fast parameter estimation algorithm, the stochastic gradient algorithm, and demonstrated its effectiveness against EM both asymptotically and empirically. Finally, we applied our model to real datasets. Our model outperformed LOAS, the most popular radio telemetry software package in ecology, with respect to the accuracy of the location estimates. Our model also hypothesized that the sloth has a relatively strong preference for certain types of trees. We are currently working on applying our model to other telemetry datasets collected at the Barro Colorado Island.
Chapter 2

Collaborative Ranking of Local Preferences

2.1 Background

Since the early days of the Netflix Prize competition, matrix factorization (MF) \([30]\) has become a popular method for modeling users’ preferences over a set of items. MF achieves state-of-the-art performance on very large-scale datasets, such as the Netflix dataset \([4]\), which comprises more than one hundred million ratings. MF also does not require user and item features \([29]\), making it particularly useful for practitioners, as they can easily apply it to new domains without designing domain-specific features.

Most MF methods optimize root-mean-square error (RMSE) \([53][46][47]\), largely because the winner of the Netflix Prize was determined by it. However, for many applications of collaborative filtering, RMSE is inappropriate because actual performance is based on predicting a rank rather than a rating. For example, when choosing which sci-fi movie to watch, a user might wonder how Star Trek ranks against Serenity, and predicting 3.8 for Star Trek and 4.5 for Serenity is relevant only in so far as helping the user infer a relative ranking.

As a remedy, researchers have devised various algorithms for collaborative ranking. Weimer et al. \([54]\) proposed CofiRank, which is an MF method that optimizes ranking measures, such as normalized discounted cumulative gain (NDCG). Rendle et al. \([44]\) analyzed the collaborative ranking problem from a Bayesian perspective.
and provided a meta-optimization criterion called Bayesian Personalized Ranking (BPR), which when coupled with MF, reduces into a differentiable version of CofiRank’s objective. Balakrishnan and Chopra [3] used a two-stage model to rank; the first stage learns the latent factors via Probabilistic Matrix Factorization (PMF) [46], and the second stage processes these factors as features using supervised regression and ranking algorithms.

Existing collaborative ranking algorithms, including CofiRank and BPR, require users’ preferences to be totally ordered, but there are many tasks where this is too restrictive. For example, for a movie recommendation task, when a user browses a small list of sci-fi titles and eventually chooses one, she prefers the chosen movie over the remaining sci-fi movies in that list, but she does not necessarily prefer the chosen movie over all other movies. Similarly, for a venue recommendation task, when a user considers nearby restaurants and settles on one, she demonstrates a local preference among a small set of nearby restaurants, rather than a global preference between the chosen restaurant and all other restaurants.

We are interested in an algorithm that not only learns from local preferences encountered during training, but also is evaluated according to the local preferences it predicts during deployment. For example, for a venue recommendation task, the user would query the recommendation system when she is at a particular neighborhood, and the system would return a local, not global, ranking over nearby venues. Thus, a successful recommendation system does not just use local preferences it observes during training to piece together a global ranking; what the user ultimately cares about are local rankings themselves.

To tackle these issues, we introduce a matrix factorization framework called Collaborative Local Ranking (CLR). In Section 2.2, we formally set up the problem, and in Section 2.3, we prove a bound on its generalization error using low-rank matrices as hypotheses. In Section 2.4, we describe the CLR objective and explore its relationship with other learning methods. In Section 2.5, we derive our algorithm for training CLR, and analyze its correctness and running time. We present empirical results on a venue recommendation dataset in Section 2.6 and conclude in Section 2.7.

The research described in this chapter is currently under submission and it is joint work with David Rosenberg, Robert Schapire, and Tony Jebara.
2.2 Problem Formulation

In this section, we formally set up the problem by describing our assumptions about the data, the hypothesis space, and the loss function. We assume that data is generated from a sample space $\mathcal{X}$, and each data point consists of a user, a candidate set of items, a local time, and a single item that the user prefers over the remaining candidates (i.e. the label). More formally, let $\mathcal{U} = \{1, \ldots, m\}$ be the set of users, let $\mathcal{V} = \{1, \ldots, n\}$ be the set of items, and let $\mathcal{T} = \{1, \ldots, T\}$ indicate the local time. Then, the sample space is defined as

$$\mathcal{X} = \{(u, C, i, t) \mid u \in \mathcal{U}, C \subseteq \mathcal{V}, i \in C, t \in \mathcal{T}\}. \quad (2.1)$$

Given a training dataset, the learning algorithm’s goal is to choose a hypothesis $g: \mathbb{Z}^3 \rightarrow \mathbb{R}$, which would map a user $u$, item $i$, and time index $t$ to a scalar value. Once the training is complete and the hypothesis is deployed, the recommendation system may be queried with a user $u$, candidate set $C$, and local time. In return, the hypothesis assigns a scalar value to each candidate item and induces a ranking over the candidate set, and the higher the value of an item, the higher its rank. For example, in case of venue recommendation, the user sends the system her geographical coordinates, the system identifies the nearby venues, forms the candidate set, applies the hypothesis, and ranks the candidate venues.

The performance of the recommendation system depends on how high it ranks the item that the user will ultimately choose. More formally, let $P[\cdot]$ denote probability, let $C_i$ be the set $C$ excluding element $i$, and let $c \overset{U}{\sim} C$ mean that $c$ is sampled uniformly from $C$. Then, the local ranking loss associated with hypothesis $g$ is

$$L_g(u, C, i, t) = \frac{P}{c \overset{U}{\sim} C_i} [g(u, i, t) - g(u, c, t) \leq 0]. \quad (2.2)$$

Intuitively, if the highest scalar value is assigned to the correct item $i$, there is no loss. Otherwise, the loss is proportional to the number of items in the candidate set that are ranked higher than the correct item.

To simplify the discussion, for most of the paper, we will assume that the sample space, hypotheses, and loss functions all exclude references to local time. We will revisit this issue in Section 2.4.
2.3 A Bound on the Generalization Error

Our ultimate goal is to devise algorithms that minimize the generalization error of collaborative ranking. In this section, we focus specifically on deriving a bound on the generalization error, which in turn will influence our algorithmic design.

We assume that the hypothesis class is based on the set of low-rank matrices. Given a low-rank matrix $M$, let $g_M \in \mathcal{F}$ be the associated hypothesis, where $g_M(u, i) = M_{u,i}$. Throughout the paper, we abuse notation and use $g_M$ and $M$ interchangeably. We assume that data is generated with respect to $\mathcal{D}$, which is an unknown probability distribution over the sample space $\mathcal{X}$, and we let $E$ denote expectation. Then, the generalization error of hypothesis $M$ is $E_{(u,C,i) \sim \mathcal{D}} L_M(u, C, i)$, which is the quantity we bound below.

We will derive the generalization bound in two steps. In the first step, we will bound the empirical Rademacher complexity of our loss class, defined below, with respect to samples that contain exactly 2 candidates, and in the second step, we will prove the generalization bound with a reduction to the previous step.

**Lemma 2.1.** Let $m$ be the number of users and let $n$ be the number of items. Define $\mathcal{L}_r = \{ L_M | M \in \mathbb{R}^{m \times n} \text{ has rank at most } r \}$ as the class of loss functions associated with low-rank matrices. Assume that $S_2 \subseteq \mathcal{X}$ is a set of $d$ samples, where each sample contains exactly 2 candidate items; i.e. if $(u, C, i) \in S_2$, then $|C| = 2$. Let $R_{S_2}(\mathcal{L}_r)$ denote the Rademacher complexity of $\mathcal{L}_r$ with respect to $S_2$. Then,

\[
R_{S_2}(\mathcal{L}_r) \leq \sqrt{\frac{2r(m + n) \ln \left( \frac{16emn^2}{d} \right)}{d}}.
\]

**Proof:** Because each sample in $S_2$ contains exactly 2 candidates, any hypothesis $L_M \in \mathcal{L}_r$ applied to a sample in $S_2$ outputs either 0 or 1. Thus, the set of dichotomies that are realized by $\mathcal{L}_r$ on $S_2$, called $\Pi_{\mathcal{L}_r}(S_2)$, is well-defined. Using Equation (6) from Boucheron et al. [7], we know that $R_{S_2}(\mathcal{L}_r) \leq \sqrt{\frac{2\ln |\Pi_{\mathcal{L}_r}(S_2)|}{d}}$. Let $\mathcal{X}_2 \subseteq \mathcal{X}$ be the set of all samples that contain exactly 2 candidates, $|\Pi_{\mathcal{L}_r}(S_2)| \leq |\Pi_{\mathcal{L}_r}(\mathcal{X}_2)|$, so it suffices to bound $|\Pi_{\mathcal{L}_r}(\mathcal{X}_2)|$.

We bound $|\Pi_{\mathcal{L}_r}(\mathcal{X}_2)|$ by counting the sign configurations of polynomials using proof techniques that are influenced by Srebro et al. [52]. Let $(u, \{i, j\}, i) \in \mathcal{X}_2$ be
a sample and let \( M \) be a hypothesis matrix. Because \( M \) has rank at most \( r \), it can be written as \( M = UV^T \), where \( U \in \mathbb{R}^{m \times r} \) and \( V \in \mathbb{R}^{n \times r} \). Let \([\cdot]\) denote an indicator function that is 1 if and only if its argument is true. Then, the loss on the sample can also be rewritten as

\[
L_M(u, \{i, j\}, i) = [M_{u,i} - M_{u,j} \leq 0] = [(UV^T)_{u,i} - (UV^T)_{u,j} \leq 0] = \left\{ \sum_{a=1}^{r} U_{u,a} (V_{i,a} - V_{j,a}) \leq 0 \right\}.
\]

Since cardinality of \( \mathcal{X}_2 \) is at most \( 2m(n) \leq mn^2 \), putting it all together, it follows that \( |\Pi_{\mathcal{X}_r} (\mathcal{X}_2)| \) is bounded by the number of sign configurations of \( mn^2 \) polynomials, each of degree at most 2, over \( r (m + n) \) variables. Applying Corollary 3 from Srebro et al. [52], we obtain

\[
|\Pi_{\mathcal{X}_r} (\mathcal{X}_2)| \leq \left( \frac{16emn^2}{r(m+n)} \right)^{r(m+n)}.
\]

Taking logarithms and making basic substitutions yield the desired result.

We proceed to proving the more general result via a reduction to Lemma 2.1.

**Theorem 2.2.** Let \( m \) be the number of users and let \( n \) be the number of items. Assume that \( S \) consists of \( d \) independently and identically distributed samples chosen from \( \mathcal{X} \) with respect to a probability distribution \( \mathcal{D} \). Let \( L_M \) be the loss function associated with a matrix \( M \), as defined in Equation 2.2. Then, with probability at least \( 1 - \delta \), for any matrix \( M \in \mathbb{R}^{m \times n} \) with rank at most \( r \),

\[
\mathbb{E}_{(u, C, i) \sim \mathcal{D}} L_M(u, C, i) \leq \mathbb{E}_{(u, C, i) \sim \mathcal{S}} L_M(u, C, i) + 2\sqrt{\frac{2r(m+n)\ln \left( \frac{16emn}{r} \right)}{d}} + \sqrt{\frac{2\ln \left( \frac{2}{\delta} \right)}{d}}. \quad (2.3)
\]

**Proof.** We will manipulate the definition of Rademacher complexity [7] in order to
use the bound given in Lemma 2.1:

\[
R_S(\mathcal{L}_r) \triangleq \mathbb{E}_{\sigma} \left[ \sup_{L_M \in \mathcal{L}_r} \left( \frac{1}{d} \sum_{a=1}^{d} \sigma_a L_M(u_a, C_a, i_a) \right) \right] \quad \text{(definition of } R_S(\mathcal{L}_r))
\]

\[
= \mathbb{E}_{\sigma} \left[ \sup_{L_M \in \mathcal{L}_r} \left( \frac{1}{d} \sum_{a=1}^{d} \sigma_a \mathbb{E}_{i_a \in (C_a \setminus \{i_a\})} L_M(u_a, \{i_a, j_a\}, i_a) \right) \right] \quad \text{(definition of } L_M)
\]

\[
\leq \mathbb{E}_{\sigma} \left[ \sup_{L_M \in \mathcal{L}_r} \left( \mathbb{E}_{j_1, \ldots, j_d} \left( \frac{1}{d} \sum_{a=1}^{d} \sigma_a L_M(u_a, \{i_a, j_a\}, i_a) \right) \right) \right] \quad \text{(linearity of expectation)}
\]

\[
= \mathbb{E}_{j_1, \ldots, j_d} \left[ \mathbb{E}_{\sigma} \left( \sup_{L_M \in \mathcal{L}_r} \left( \frac{1}{d} \sum_{a=1}^{d} \sigma_a L_M(u_a, \{i_a, j_a\}, i_a) \right) \right) \right] \quad \text{(reversing expectation order)}
\]

\[
= \mathbb{E}_{j_1, \ldots, j_d} \left[ R_{S_2}(\mathcal{L}_r) \right] \quad \text{(definition of } R_{S_2}(\mathcal{L}_r))
\]

\[
\leq \sqrt{\frac{2r(m + n) \ln \left( \frac{16emn^2}{r(m+n)} \right)}{d}} \quad \text{(by Lemma 2.1)}
\]

Plugging the bound to Theorem 3.2 in Boucheron et al. [7] proves the theorem. □

Srebro et al. [52] used covering numbers to prove a generalization error bound for collaborative binary classification, which is matrix factorization framework where the matrix entries are binary and the loss function is 0-1. Even though their learning setting is different from ours, we can still compare our bound in Theorem 2.2 with their bound, and it can be seen that they match up to logarithmic factors. Thus, collaborative local ranking maintains the same generalization error as collaborative binary classification in exchange for a small increase in sample size.

We can improve the tightness of our generalization error bound by restricting the sample space \( \mathcal{X} \). In our proof of Lemma 2.1, we loosely bounded the cardinality of the sample space with \( |\mathcal{X}_2| \leq mn^2 \), but many collaborative ranking tasks have additional structure which lead to better bounds. For example, in case of venue recommendation, \( \mathcal{X}_2 \) is restricted because two venues are only assigned to the same candidate set if they are near each other. If we assume that every venue has at most \( b \) nearby venues, then \( |\mathcal{X}_2| \leq mnb \), and the \( n \) in the logarithm of Equation 2.3 gets replaced by a small constant. Thus, by making additional assumptions about
the structure of the sample space, the generalization error bound in Theorem 2.2 becomes asymptotically equivalent to the bound provided in [52], even though the latter bound was derived for the simpler collaborative classification setting.

2.4 Collaborative Local Ranking

In this section, we describe CLR by formulating its objective function, which we justify using the generalization error bound derived above. We also make connections between CLR and other learning methods, such as maximum-margin matrix factorization (MMMF) [45] and ranking support vector machine (ranking SVM) [24].

Given the generalization error bound in Theorem 2.2, a reasonable objective is to minimize the empirical local ranking loss, which is the first term on the right hand side of Equation 2.3. However, this function is discontinuous and difficult to minimize with respect to $M$, so we propose minimizing a more tractable upper bound instead. Let $h(x) = \max(0, 1 - x)$ be the hinge function, let $M$ be the hypothesis matrix with rank at most $r$, and let $M = UV^T$, where $U \in \mathbb{R}^{m \times r}$ and $V \in \mathbb{R}^{n \times r}$. Then, we can bound the empirical local ranking loss as

$$
\mathbb{E}_{(u,C,i) \in S} L_M(u, C, i) = \frac{1}{|S|} \sum_{(u,C,i) \in S} L_M(u, C, i)
\leq \frac{1}{|S|} \sum_{(u,C,i) \in S} \left| C_i \right| \left( \frac{1}{|C_i|} \sum_{e \in C_i} h\left( (UV^T)_{u,i} - (UV^T)_{u,c} \right) \right). \quad (2.4)
$$

The resulting upper bound is not necessarily convex with respect to $U$ and $V$ jointly, but it is convex with respect to $U$ if $V$ is fixed, and vice versa.

We use trace norm regularization, in addition to the low-rank constraint, to further restrict the hypothesis class. Let $\| \cdot \|_T$ denote the trace norm and let $\| \cdot \|_F$ denote
the Frobenius norm, then we use the equality $\|M\|_T = \min_{U, V, M = U V^T} \frac{1}{2} (\|U\|_F^2 + \|V\|_F^2)$
given in Lemma 6 of Mazumder et al. [37] to state the objective in its final form.

**Definition 2.3** (CLR Objective). Let $S$ be the training data. Let $U \in \mathbb{R}^{m \times r}$ and $V \in \mathbb{R}^{n \times r}$ be the factor matrices, and let $\lambda > 0$ be the regularization parameter. The CLR objective is

$$f_{CLR}(S; U, V) = \frac{\lambda}{2} (\|U\|_F^2 + \|V\|_F^2) + \frac{1}{|S|} \sum_{(u, C, i) \in S} \frac{1}{|C_i|} \sum_{c \in C_i} h \left( (U V^T)_{u,i} - (U V^T)_{u,c} \right).$$

If $\lambda = 0$, then the CLR objective is equivalent to minimizing the bound in Equation 2.4. If $\lambda \neq 0$, then it can be shown that the equivalence still holds, but under the constraint that matrices have bounded trace norm in addition to having bounded rank. Even though minimizing Equation 2.4 is more directly justified by our generalization bound, we use the CLR objective instead, which is more general, to allow additional regularization. Note that the resulting hypothesis still satisfies the assumptions of Theorem 2.2. Rank-truncated trace norms have been used as regularizers in other collaborative learning settings, such as by Foygel et al. [18] and by Rennie and Srebro [45], and have been demonstrated to work well.

We note that the CLR and the ranking SVM [24] objectives are closely related. If $V$ is fixed and we only need to minimize $U$, then each row of $V$ acts as a feature vector for the corresponding item, each row of $U$ acts as a separate linear predictor, and the CLR objective decomposes into solving simultaneous ranking SVM problems. In particular, let $S_u = \{(a, C, i) \in S \mid a = u\}$ be the examples that correspond to user $u$, let $U_u$ denote row $u$ of $U$, and let $f^{rSVM}$ denote the objective function of ranking SVM, then

$$f^{CLR}(S; U, V) = \frac{\lambda}{2} (\|U\|_F^2 + \|V\|_F^2) + \frac{1}{|S|} \sum_{(u, C, i) \in S} \frac{1}{|C_i|} \sum_{c \in C_i} h \left( (U V^T)_{u,i} - (U V^T)_{u,c} \right).$$

$$= \sum_{u=1}^{m} \left[ \frac{\lambda}{2} (\|U_u\|_F^2 + \|V\|_F^2) + \frac{1}{|S_u|} \sum_{(u, C, i) \in S_u} \frac{1}{|C_i|} \sum_{c \in C_i} h \left( (U V^T)_{u,i} - (U V^T)_{u,c} \right) \right]$$

$$= \sum_{u=1}^{m} f^{rSVM}(S_u; U_u, V).$$

This correspondence is not surprising, since in the ranking SVM setting, items have features, the training data consists of binary orderings between items, and the
ranking SVM objective itself is a convex upper bound on the number of misordered pairs. Additionally, just like optimizing the CLR objective decomposes into solving simultaneous ranking SVM objectives, optimizing the MMMF [45] objective also decomposes into solving simultaneous binary SVM objectives, and this observation influences how we design our algorithms in the next section.

Lastly, we extend the CLR objective to incorporate features. In the context of venue recommendation, such features might be obtained from an external database and indicate the venue type or the query time stamp. We assume an extended sample space, and given a sample \((u, C, i, t) \in \mathcal{X}\), we let \(t\) denote the query time stamp.

**Definition 2.4 (CLR.F Objective).** Let \(S\) be the training data. Given a sample \((u, C, i, t) \in S\), assume that \(F_{u,i,t} \in \mathbb{R}^q\) denotes the corresponding feature vector. Let \(U \in \mathbb{R}^{m \times r}\) and \(V \in \mathbb{R}^{n \times r}\) be the factor matrices, let \(w \in \mathbb{R}^q\) be the feature coefficients, and let \(\lambda, \gamma > 0\) be the regularization parameters. The CLR.F objective is

\[
\begin{align*}
    f_{\text{CLR.F}}(S; U, V, w) = & \frac{\lambda}{2} \left( \|U\|_F^2 + \|V\|_F^2 \right) + \frac{\gamma}{2} \|w\|^2 \\
    & + \frac{1}{|S|} \sum_{(u, C, i, t) \in S} \frac{1}{|C_i|} \sum_{c \in C_i} h\left( (UV^T)_{u,i} - (UV^T)_{u,c} \right) + w^T (F_{u,i,t} - F_{u,c,t}).
\end{align*}
\]

### 2.5 Algorithms

#### 2.5.1 Derivation

In this subsection, we derive and describe our algorithm for optimizing the CLR objective. As we noted before, the CLR objective is not necessarily jointly convex in \(U\) and \(V\), but it is convex in \(U\) when \(V\) is fixed and vice versa. We minimize the CLR objective using alternating minimization, where we sequentially alternate between solving the convex subproblems, and we solve each such subproblem using projected stochastic subgradient descent. The pseudocode is depicted in Algorithm 2.1.

Now, we derive the projected stochastic subgradient descent algorithm for minimizing \(V\) while keeping \(U\) fixed. At each iteration, the algorithm approximates the objective function based on an example selected at random, updates the weight
Algorithm 2.1 Alternating minimization for optimizing the CLR objective.

Require: Training data \( S \subseteq X \), regularization parameter \( \lambda > 0 \), rank constraint \( r \), number of iterations \( T \).

1: \( U_1 \leftarrow \) Sample matrix uniformly at random from \( \left[-\frac{1}{\sqrt{\lambda r}}, \frac{1}{\sqrt{\lambda r}}\right]^{m \times r} \).
2: \( V_1 \leftarrow \) Sample matrix uniformly at random from \( \left[-\frac{1}{\sqrt{\lambda r}}, \frac{1}{\sqrt{\lambda r}}\right]^{n \times r} \).
3: for all \( t \) from 1 to \( T - 1 \) do
   4: \( U_{t+1} \leftarrow \arg\min_U f^{CLR}(S; U_t, V_t) \)
   5: \( V_{t+1} \leftarrow \arg\min_V f^{CLR}(S; U_{t+1}, V) \)
4: return \( U_T, V_T \).

We introduce various matrix notation to help us define the approximate subgradients. Given a matrix \( M \), let \( M_{k,} \) denote row \( k \) of \( M \). Define the matrix \( \hat{M}_{p, q, z} \), for \( p \neq q \), as

\[
\hat{M}_{s,} = \begin{cases} 
M_{z,} & \text{for } s = p, \\
-M_{z,} & \text{for } s = q, \\
0 & \text{otherwise,}
\end{cases}
\]  

(2.5)

and define the matrix \( \check{M}_{s,} \) as

\[
\check{M}_{s,} = \begin{cases} 
M_{p,} - M_{q,} & \text{for } s = z, \\
0 & \text{otherwise.}
\end{cases}
\]

(2.6)

Then, the subgradient of the approximate objective function with respect to \( V \) is

\[
\nabla_V f^{CLR}((u, C, i); U, V) = \lambda V - \frac{1}{|C_i|} \sum_{c \in C_i} [(UV^T)_{u,i} - (UV^T)_{u,c}] \quad \text{for } i \neq c, u.
\]

(2.7)
Setting $\eta_t = \frac{1}{\lambda t}$ as the learning rate at iteration $t$, the approximate subgradient update becomes $V_{t+1} = V_t - \eta_t \nabla f_{\text{CLR}}((u, C, i); U, V)$. After the update, the weights are projected onto a ball with radius $\frac{1}{\sqrt{\lambda}}$. The pseudocode for optimizing both convex subproblems is depicted in Algorithms 2.2 and 2.3. We prove the correctness of the algorithms and bound their running time in the next subsection.

**Algorithm 2.2** Projected stochastic subgradient descent for optimizing $U$.

**Require:** Factors $V \in \mathbb{R}^{n \times r}$, training data $S$, regularization parameter $\lambda$, rank constraint $r$, number of iterations $T$.

1: $U_1 \leftarrow 0^{m \times r}$
2: for all $t$ from 1 to $T - 1$ do
3:     Choose $(u, C, i) \in S$ uniformly at random.
4:     $\eta_t \leftarrow \frac{1}{\lambda t}$
5:     $C^+ \leftarrow \{ c \in C \mid (U_t V^T)_{u,i} - (U_t V^T)_{u,c} < 1 \}$
6:     $U_{t+1} \leftarrow (1 - \eta_t \lambda) U_t + \frac{\eta_t}{|C|} \sum_{c \in C^+} \hat{V}^{i,c,u}$
7:     $U_{t+1} \leftarrow \min \left\{ 1, \frac{1}{\sqrt{\lambda \|U_{t+1}\|_F}} \right\} U_{t+1}$
8: return $U_T$.

**Algorithm 2.3** Projected stochastic subgradient descent for optimizing $V$.

**Require:** Factors $U \in \mathbb{R}^{m \times r}$, training data $S$, regularization parameter $\lambda$, rank constraint $r$, number of iterations $T$.

1: $V_1 \leftarrow 0^{n \times r}$
2: for all $t$ from 1 to $T - 1$ do
3:     Choose $(u, C, i) \in S$ uniformly at random.
4:     $\eta_t \leftarrow \frac{1}{\lambda t}$
5:     $C^+ \leftarrow \{ c \in C \mid (U V^T)_{u,i} - (U V^T)_{u,c} < 1 \}$
6:     $V_{t+1} \leftarrow (1 - \eta_t \lambda) V_t + \frac{\eta_t}{|C|} \sum_{c \in C^+} \hat{U}^{i,c,u}$
7:     $V_{t+1} \leftarrow \min \left\{ 1, \frac{1}{\sqrt{\lambda \|V_{t+1}\|_F}} \right\} V_{t+1}$
8: return $V_T$.

We conclude the subsection by commenting on the implementation details. A naive implementation of Algorithm 2.2 would execute each iteration in time $\Omega(bmnkr)$, where $b$ denotes the size of the largest candidate set. One can reduce the running time of each iteration considerably by normalizing the matrix efficiently. We represent $U$ as a triplet $(W, a, v)$, where $a$ is a scalar, $U = a W$, and $\|U\|_F = v$. It can be
verified that, using the new representation, a single iteration in both Algorithms 2.2 and 2.3 can be executed in time $O(br)$.

### 2.5.2 Analysis

In this subsection, we analyze the running time and correctness of our algorithms. In particular, we prove that projected stochastic subgradient descent converges to the correct solution in time independent of the size of the training data. We do not bound the total number of alternating minimization steps a priori; nevertheless, we demonstrate that our methods are especially suitable for large datasets since they solve each individual minimization problem efficiently.

The convex subproblems we analyze have the general form

$$\min_{X \in \mathcal{D}} f(X; \ell) = \min_{X \in \mathcal{D}} \frac{\lambda}{2} \|X\|^2_F + \frac{1}{|S|} \sum_{(u, C, i) \in S} \ell(X; (u, C, i)).$$  \tag{2.8}

One can obtain the individual subproblems by specifying the domain $\mathcal{D}$ and the loss function $\ell$. For example, in case of Algorithm 2.2, the corresponding minimization problem is specified by

$$\min_{X \in \mathbb{R}^{m \times r}} f(X; \ell_V) \text{ where } \ell_V(X; (u, C, i)) = \frac{1}{|C^i|} \sum_{c \in C^i} h\left(\left(XV^T\right)_{u, i} - \left(XV^T\right)_{u, c}\right),$$  \tag{2.9}

and in case of Algorithm 2.3, it is specified by

$$\min_{X \in \mathbb{R}^{n \times r}} f(X; \ell_U) \text{ where } \ell_U(X; (u, C, i)) = \frac{1}{|C^i|} \sum_{c \in C^i} h\left(\left(UX^T\right)_{u, i} - \left(UX^T\right)_{u, c}\right).$$  \tag{2.10}

Let $U^* = \text{argmin}_U f(U; \ell_V)$ and $V^* = \text{argmin}_V f(V; \ell_U)$ denote the solution matrices of Equations 2.9 and 2.10, respectively. Also, given a general convex loss $\ell$ and domain $\mathcal{D}$, let $\tilde{X} \in \mathcal{D}$ be an $\epsilon$-accurate solution for the corresponding minimization problem if $f(\tilde{X}; \ell) \leq \min_{X \in \mathcal{D}} f(X; \ell) + \epsilon$.

In the remainder of this subsection, we show that Algorithms 2.2 and 2.3 are adaptations of the Pegasos [49] algorithm to the CLR setting. Then, we prove certain properties that are prerequisites for obtaining Pegasos’s performance guarantees. In particular, we show that the approximate subgradients computed by Algo-
gorithms 2.2 and 2.3 are bounded and the loss functions associated with Equations 2.9 and 2.10 are convex. In the end, we plug these properties into a theorem proved by Shwartz et al. [49] to show that our algorithms reach an \(\epsilon\)-accurate solution with respect to their corresponding minimization problems in \(\tilde{O}\left(\frac{1}{\epsilon^2}\right)\) iterations.

Aside from having different loss functions, another difference between our collaborative ranking setting and the supervised learning setting of Shalev-Shwartz et al. [49] is the treatment of features. In the supervised learning setting, features are constant, but in our setting, the factor matrices \(U\) and \(V\) act like features, and their values vary from one step of alternating minimization to the next. For example, when optimizing \(U\), the factor matrix \(V\) acts like a feature matrix, and vice versa, and both matrices change in value during subsequent steps of alternating minimization. As a result, feature values that are treated as constants in the supervised setting cannot be treated as constants in the collaborative setting. We treat these issues in our proofs as well.

**Lemma 2.5.** \(\|U^*\| \leq \frac{1}{\sqrt{\lambda}}\) and \(\|V^*\| \leq \frac{1}{\sqrt{\lambda}}\).

**Proof.** One can obtain the bounds on the norms of the optimal solutions by examining the dual form of the optimization problems and applying the strong duality theorem. Equations 2.9 and 2.10 can both be represented as

\[
\min_{v \in \mathbb{D}} \frac{1}{2} \|v\|^2 + \sum_{k=1}^{K} e_k h(f_k(v)),
\]

where \(e_k = \frac{1}{\lambda |\mathcal{S}| |C_k|}\) is a constant, \(h\) is the hinge function, \(\mathbb{D}\) is a Euclidean space, and \(f_k\) is a linear function. We rewrite Equation 2.11 as a constrained optimization problem

\[
\min_{v \in \mathbb{D}, \xi \in \mathbb{R}^K} \frac{1}{2} \|v\|^2 + \sum_{k=1}^{K} e_k \xi_k \quad \text{subject to} \quad \xi_k \geq 1 - f_k(v), \quad k = 1, \ldots K, \\
\xi_k \geq 0, \quad k = 1, \ldots K.
\]
The Lagrangian of this problem is

\[ L(v, \xi, \alpha, \beta) = \frac{1}{2} \|v\|^2 + \sum_{k=1}^{K} e_k \xi_k + \sum_{k=1}^{K} \alpha_k (1 - f_k(v) - \xi_k) - \sum_{k=1}^{K} \beta_k \xi_k \]

\[ = \frac{1}{2} \|v\|^2 + \sum_{k=1}^{K} \xi_k \left(e_k - \alpha_k - \beta_k\right) + \sum_{k=1}^{K} \alpha_k (1 - f_k(v)), \]

and its dual function is

\[ g(\alpha, \beta) = \inf_{v, \xi} L(v, \xi, \alpha, \beta). \]

Since \( L(v, \xi, \alpha, \beta) \) is convex and differentiable with respect to \( v \) and \( \xi \), the necessary and sufficient conditions for minimizing \( v \) and \( \xi \) are

\[ \nabla_v L = 0 \quad \Leftrightarrow \quad v = \sum_{k=1}^{K} \alpha_k \nabla_v f_k(v), \]

\[ \nabla_\xi L = 0 \quad \Leftrightarrow \quad e = \alpha + \beta. \quad (2.13) \]

We plug these conditions back into the dual function and obtain

\[ g(\alpha, \beta) = \inf_{v, \xi} L(v, \xi, \alpha, \beta) \]

\[ = \frac{1}{2} \left\| \sum_{k=1}^{K} \alpha_k \nabla_v f_k(v) \right\|^2 + \sum_{k=1}^{K} \alpha_k \left(1 - f_k \left(\sum_{k=1}^{K} \alpha_k \nabla_v f_k(v)\right)\right) \]

\[ = \frac{1}{2} \left\| \sum_{k=1}^{K} \alpha_k \nabla_v f_k(v) \right\|^2 + \sum_{k=1}^{K} \alpha_k - \sum_{k=1}^{K} \alpha_k f_k \left(\sum_{k=1}^{K} \alpha_k \nabla_v f_k(v)\right). \quad (2.14) \]

Since \( f_k \) is a linear function, we let \( f_k(v) = \bar{k} \cdot v \), where \( \bar{k} \) is a constant vector, and
\[ \nabla_v f_k(v) = \tilde{k}. \] Then,

\[
\left\| \sum_{k=1}^{K} \alpha_k \nabla_v f_k(v) \right\|^2 = \left\| \sum_{k=1}^{K} \alpha_k \tilde{k} \right\|^2 = \left( \sum_{k=1}^{K} \alpha_k \tilde{k} \right) \cdot \left( \sum_{k=1}^{K} \alpha_k \tilde{k} \right) = \sum_{k=1}^{K} \alpha_k \tilde{k} \cdot \left( \sum_{k=1}^{K} \alpha_k \tilde{k} \right) = \sum_{k=1}^{K} \alpha_k f_k \left( \sum_{k=1}^{K} \alpha_k \tilde{k} \right)
\]

\[
= \sum_{k=1}^{K} \alpha_k f_k \left( \sum_{k=1}^{K} \alpha_k \nabla_v f_k(v) \right). \tag{2.15}
\]

Simplifying Equation 2.14 using Equation 2.15 yields

\[
g(\alpha, \beta) = -\frac{1}{2} \left\| \sum_{k=1}^{K} \alpha_k \nabla_v f_k(v) \right\|^2 + \sum_{k=1}^{K} \alpha_k
\]

\[
= -\frac{1}{2} \left\| \sum_{k=1}^{K} \alpha_k \tilde{k} \right\|^2 + \sum_{k=1}^{K} \alpha_k. \tag{2.16}
\]

Finally, we combine Equations 2.13 and 2.16, and obtain the dual form of Equation 2.12,

\[
\max_{\alpha} -\frac{1}{2} \left\| \sum_{k=1}^{K} \alpha_k \tilde{k} \right\|^2 + \sum_{k=1}^{K} \alpha_k \tag{2.17}
\]

subject to \(0 \leq \alpha_k \leq e_k, \quad k = 1, \ldots K.\)

The primal problem is convex, its constraints are linear, and the domain of its objective is open; thus, Slater’s condition holds and strong duality is obtained. Furthermore, the primal problem has differentiable objective and constraint functions, which implies that \((v^*, \xi^*)\) is primal optimal and \((\alpha^*, \beta^*)\) is dual optimal if and only if these points satisfy the Karush-Kuhn-Tucker (KKT) conditions. It follows that

\[
v^* = \sum_{k=1}^{K} \alpha^*_k \tilde{k}. \tag{2.18}
\]
Note that we defined \( e_k = \frac{1}{|S[|C_k|]} \), where \( \sum_{k=1}^K e_k = \frac{1}{\lambda} \), and the constraints of the dual problem imply \( 0 \leq \alpha_k \leq e_k \); thus, \( \sum_{k=1}^K \alpha_k^* \leq \frac{1}{\lambda} \). Because of strong duality, there is no duality gap, and the primal and dual objectives are equal at the optimum,

\[
\frac{1}{2} \| v^* \|^2 + \sum_{k=1}^K e_k \xi_k^* = -\frac{1}{2} \left( \sum_{k=1}^K \alpha_k^* \right)^2 + \sum_{k=1}^K \alpha_k^*
\]

(by Equation 2.18)

\[
\leq -\frac{1}{2} \| v^* \|^2 + \frac{1}{\lambda}
\]

\[
\Rightarrow \| v^* \|^2 \leq \frac{1}{\lambda}.
\]

This proves the lemma.

Given the bounds in Lemma 2.5, it can be verified that Algorithms 2.2 and 2.3 are adaptations of the Pegasos [49] algorithm for optimizing Equations 2.9 and 2.10, respectively. It still remains to show that Pegasos’s performance guarantees hold in our case.

**Lemma 2.6.** In Algorithms 2.2 and 2.3, the approximate subgradients have norm at most \( \sqrt{\lambda} + 2\sqrt{\frac{1}{\lambda}} \).

**Proof.** The approximate subgradient for Algorithm 2.3 is depicted in Equation 2.7. Due to the projection step, \( \| V \|_F \leq \frac{1}{\sqrt{\lambda}} \), and it follows that \( \| \lambda V \|_F \leq \sqrt{\lambda} \). The term \( \hat{U}^{i,c,u} \) is constructed using Equation 2.5, and it can be verified that \( \| \hat{U}^{i,c,u} \|_F \leq \sqrt{2} \| U \|_F \leq \sqrt{\frac{2}{\lambda}} \). Using triangle inequality, one can bound Equation 2.7 with \( \sqrt{\lambda} + \sqrt{\frac{2}{\lambda}} \). A similar argument can be made for the approximate subgradient of Algorithm 2.2, yielding the slightly higher upper bound given in the lemma statement.

We combine the lemmas to obtain the correctness and running time guarantees for our algorithms.

**Lemma 2.7.** Let \( \lambda \leq \frac{1}{4} \), let \( T \) be the total number of iterations of Algorithm 2.2, and let \( U_t \) denote the parameter computed by the algorithm at iteration \( t \). Let \( \hat{U} = \)
\( \frac{1}{T} \sum_{t=1}^{T} U_t \) denote the average of the parameters produced by the algorithm. Then, with probability at least \( 1 - \delta \),

\[
f(\bar{U}; \ell_V) \leq f(U^*; \ell_V) + \frac{21 \left( \sqrt{\lambda} + 2 \sqrt{\frac{1}{\lambda}} \right)^2 \ln \left( \frac{T}{\delta} \right)}{\lambda T}.
\]

The analogous result holds for Algorithm 2.3 as well.

**Proof.** First, for each loss function \( \ell_V \) and \( \ell_U \), variables are linearly combined, composed with the convex hinge function, and then averaged. All these operations preserve convexity, hence both loss functions are convex. Second, we have argued above that Algorithms 2.2 and 2.3 are adaptations of the Pegasos [49] algorithm for optimizing Equations 2.9 and 2.10, respectively. Third, in Lemma 2.6, we proved a bound on the approximate subgradients of both algorithms. Plugging these three results into Corollary 2 in Shwartz et al. [49] yields the statement of the theorem.

The theorem below gives a bound in terms of individual parameters rather than average parameters.

**Theorem 2.8.** Assume that the conditions and the bound in Lemma 2.7 hold. Let \( t \) be an iteration index selected uniformly at random from \( \{1, \ldots, T\} \). Then, with probability at least \( \frac{1}{2} \),

\[
f(U_t; \ell_V) \leq f(U^*; \ell_V) + \frac{42 \left( \sqrt{\lambda} + 2 \sqrt{\frac{1}{\lambda}} \right)^2 \ln \left( \frac{T}{\delta} \right)}{\lambda T}.
\]

The analogous result holds for Algorithm 2.3 as well.

**Proof.** The result follows directly from combining Lemma 2.7 with Lemma 3 in Shwartz et al. [49].

Thus, with high probability, our algorithms reach an \( \epsilon \)-accurate solution in \( \tilde{O} \left( \frac{1}{t^2c} \right) \) iterations. Since we argued in Subsection 2.5.1 that the running time of each stochastic update is \( O(br) \), it follows that a complete run of projected stochastic subgradient descent takes \( \tilde{O} \left( \frac{br}{t^2c} \right) \) time, and the running time is independent of the size of the training data.
2.6 Experiments

In this section, we provide an empirical analysis of the CLR framework by applying it to a venue recommendation task. We assess CLR's generalization and running time performance by comparing it against CofiRank, an algorithm which is representative of the state-of-the-art in collaborative ranking.

We created our dataset by collecting publicly available “check-ins” via the Twitter and Foursquare APIs. A check-in is a virtual announcement where a user shares her whereabouts with other people on her social network. A check-in contains a user ID, a venue ID (e.g. “Shake Shack”), and a local time; optionally, it may also contain a text (e.g. “Trying the shackburger!!”), photos, and other metadata. We collected public check-ins that occurred in New York City during the 9 month period ranging from January 2011 to September 2011. We filtered our dataset to only include users who have checked in at least 5 times and venues which have been checked into at least 5 times, yielding a total of 13750 users, 11700 venues, and 269597 check-ins. We note that all the check-ins we collected were shared with the entire Internet, so our dataset does not contain any private information.

At a high level, our goal is to build a venue recommendation application for mobile platforms, where the user specifies a geographical region she is interested in exploring, and the application returns a personalized ranking over the venues in that region. If such an application already existed, it would have been an ideal fit for the CLR setting for the following reasons: 1) the regions explored by the users are generated by an unknown probability distribution; 2) when checking into a venue, the user implicitly prefers it over the remaining venues in the explored region; 3) the application’s goal is to provide a relevant ranking over the venues by minimizing the local ranking loss. Even though the Foursquare dataset we collected is not guaranteed to be generated under similar assumptions, we use it as a proxy regardless, and assume that when a user checks into a venue, she is implicitly preferring that venue over the remaining venues in a specified radius. We note that similar assumptions about implicit preferences have been made by Joachims [24] in the supervised learning setting, where the author introduced ranking SVMs to rerank search engine results using clickthrough logs.

We partition the check-ins chronologically, and place the earliest 60% of check-
ins to train, the subsequent 20% to validation, and final 20% to test sets. Each check-in corresponds to a tuple \((u, i)\), where \(u\) is the user and \(i\) is the checked-in venue. In order to apply CLR, we augment each check-in with a candidate set \(C\), which consists of venues within a specified distance to \(i\), and form the CLR sample \((u, C, i)\). We train a variety of CLR and CofiRank models on a combined train and validation set, determine the best performing model parameters using the validation set, and report the performance of the chosen parameters on the test set. The model parameters include the rank \(r \in \{2, 5, 10, 20\}\), CofiRank regularization parameter \(\lambda \in \{10, 100, 1000, 10000\}\), and CLR regularization parameter \(\lambda \in \{0.01, 0.001, 0.0001, 0.00001, 0.000001\}\). For testing, we form queries with a user \(u\) and a candidate set \(C\), hide the label \(i\), and measure the algorithms’ performances accordingly. We vary the radius that determines the size of the candidate set \(C\) from 50m to 300m, in 50m increments.

Figure 2.1 displays the local ranking loss of various algorithms on the held-out test set. We compare CLR against both CofiRank and a baseline algorithm called “Popular”, which simply outputs the most frequently checked in venue from the candidate set. We execute CofiRank with three different loss functions: ordinal, NDCG, and squared [55]. CofiRank performs best when coupled with the squared loss, where it effectively mimics MMMF [45]; however, the “Popular” baseline outperforms CofiRank. Our algorithm, CLR, outperforms all methods and achieves the lowest local ranking loss. We note that, unlike competing methods, CLR does not suffer an additional loss when the radius and size of its candidate set increases.

Figure 2.2 displays the recall@\(k\) performance on the held-out test set, for \(k = \{1, 5, 10\}\). The order of algorithms with respect to recall performance is exactly the same as their order with respect to local ranking loss. For recall@1, the “Popular” baseline outperforms all versions of CofiRank, but as the radius of candidate sets increase beyond 150m, the performance of CofiRank-squared matches that of “Popular”. In contrast, CLR outperforms them all. For recall@5 and recall@10, performance of CLR and “Popular” coincide up to 100m radius, but CLR outperforms competing methods for higher radii.

We also empirically analyze the running time of CofiRank and CLR. For both algorithms, for each radius, we choose the parameter settings which perform the best on the validation set, and record their respective training time. We plot the
Figure 2.1: The local ranking loss of various algorithms on the held-out test set. CofiRank performs best when coupled with squared loss, “Popular” performs better than CofiRank, and CLR outperforms them all. The error bars correspond to standard error and are barely noticeable.

results using log-scale in Figure 2.3. Even though the running time of CLR has a linear dependence on the size of the candidate sets; nevertheless, CLR trains orders of magnitude faster than CofiRank, as suggested by our theoretical analysis in Subsection 2.5.2.

2.7 Discussion

In this chapter, we formulated a new collaborative ranking framework, called Collaborative Local Ranking, which allows us to formulate a wide variety of real-world ranking tasks in a collaborative setting. We justified CLR with a bound on its generalization error. We also derived a simple alternating minimization algorithm and showed that each minimization step can be efficiently computed in time independent of the size of the training data. We applied CLR to a venue recommendation task and demonstrated that it outperforms state-of-the-art collaborative ranking methods, such as CofiRank, both in terms of generalization performance and running time.
Figure 2.2: Recall@k of various algorithms on the held-out test set, for \( k = \{1, 5, 10\} \). CLR outperforms all competing methods. The error bars correspond to standard error and are barely noticeable.
Figure 2.3: Training time of CLR and CofiRank in log-scale. We report the results for CofiRank-squared, which is the best performing CofiRank variant. CLR trains orders of magnitude faster than CofiRank, even when we increase the size of the radius and candidate sets.
Chapter 3

Collaborative Place Models

3.1 Background

During the last three years, positioning devices that measure and record our locations have become ubiquitous. The most common positioning device, the smartphone, is projected to be used by a billion people in the near future [11]. This surge in positioning devices has increased the availability of location data, and provided scientists with new research opportunities, such as building location-based recommendation systems [22, 60, 61], analyzing human mobility patterns [8, 20, 50], and modeling the spread of diseases [16].

For many location-based tasks, a fundamental problem is predicting users’ future locations. For example, a navigation application that has access to traffic conditions can warn the user about when to depart, without requiring any input from the user, as long as the application can accurately predict the arrival time and location of user’s next destination. Similarly, a restaurant application can provide a list of recommended venues, even reserve them while space is still available, by modeling the probable locations the user might visit in the evening. Predicting future locations based on past ones is a difficult problem, and to solve it, one needs efficient algorithms that identify the structures and routines hidden in users’ past locations.

Complicating the problem further, in many real-world applications, location datasets are sparse. Due to privacy considerations [56] and high energy consumption of positioning hardware [41], most mobile phone users only allow applications
to log their location when the application is active, but not when it is running in the background. Consequently, compared to previous datasets used in location prediction research, many real-world datasets have much less information available for each user, as depicted in Figure 3.1. Even when users share very little with their applications, they nevertheless expect these applications to infer as much as possible; thus, there is a need for location prediction algorithms that are designed with sparsity in mind.

There have been various studies on location prediction. Gao et al. [19] designed a Markov model that takes temporal context into account in order to predict the user’s location in the immediate future. Cho et al. [9] proposed a two-state mixture of Gaussians that leverages the social relationships between users, but they limited their model to only represent “home” and “work”. De Domenico et al. [12] presented a location prediction algorithm that won the Nokia Mobile Data Challenge (MDC), where the dataset consisted of tens of thousands of GPS observations, collected every few minutes and over the span of a year. Their algorithm exploited the high density of the location dataset, as well as the social relationships between users, by comparing all historical trajectories that spanned a day and making a forecast based on the similarities between such trajectories.

In addition to location prediction, there have also been studies on algorithms that detect significant places and routines in location data. Eagle and Pentland [14] applied eigendecomposition to the Reality Mining dataset, where all locations were already labeled as “home” or “work”, and extracted users’ daily routines. Farrahi and Gatica-Perez [17] used the same dataset, but extracted the routines using Latent Dirichlet Allocation (LDA) instead of eigendecomposition. Liao et al. [32, 33] proposed a hierarchical conditional random field to identify activities and significant places from the users’ GPS traces, and since their algorithm was supervised, it required locations to be manually labeled for training.

In this chapter, we propose two unsupervised Bayesian probabilistic models for location prediction that are designed specifically for sparse location datasets. Both models assume that each user is characterized by a number of place clusters, whose spatial characteristics, such as their means and covariances, are determined probabilistically from the data. Similar to a mixed-membership model [15], we allow the place clusters to be global to each user, but we restrict the user’s preferences over
Figure 3.1: Histograms of number of observations per user for a dense and a sparse dataset. The top plot is based on a random sample of 1395 mobile carrier users and is representative of dense datasets used in previous work on location prediction. The bottom plot is based on a random sample of 5007 smartphone application users and is the sparse dataset used in this chapter. On average, the dense dataset has 21 times more observations per user than the sparse dataset.
the clusters to be local to each weekhour. For example, our models may learn that a user partitions her time between four places, two of which are in West Village and midtown Manhattan, determine the spatial characteristics of these places, infer that on Thursday at 7pm the user has a high likelihood of visiting West Village and a low likelihood of visiting midtown Manhattan, and achieve these inferences in an unsupervised manner from the observed geographic coordinates alone.

A source of difficulty is that, due to sparsity, each user has many weekhours for which no location has been observed before. Since our goal is to construct a week-long profile of users’ whereabouts, our models need to infer the place distributions for the missing weekhours, which they accomplish by sharing information across users. It is the details of how users share this information that sets the two models apart.

Our first model, called Constrained Place Model (CPM), assumes that for each weekhour all users have the same place distribution, even though the spatial characteristics of these places remain unique to each user. For example, users Rob and Tony may both have a work cluster, with different means and covariances, but for any given weekhour, Rob and Tony’s probabilities of being at work would be the same. By assuming that the place distributions are the same for all users, CPM is able to estimate place distributions for all weekhours, even when data is sparse.

Since constraining all users to have the same place distribution may be too restrictive, we propose a second model called Smooth Place Model (SPM). This model assumes that, for each user, each weekhour’s place distribution is a convex combination of component place distributions, and it is the coefficients of these convex combinations that are shared across all users. As a result, SPM allows users to have different place distributions for each weekhour, different number of places for each user, as well as different spatial characteristics for each place. Despite its flexibility, SPM accurately constructs place distributions for hours and days where the user has not been observed before, in addition to providing state-of-the-art performance in sparse location prediction.

The chapter proceeds as follows. In Section 3.2, we provide a formal description of our models. In Section 3.3, we derive the inference algorithms. In Section 3.4, we provide details of our experimental setup and apply our algorithms to a sparse smartphone application dataset and a dense mobile carrier dataset. We conclude
in Section 3.5.

The research described in this chapter is currently under submission and it is joint work with David Rosenberg, Robert Schapire, and Tony Jebara.

### 3.2 Collaborative Place Models

In this section, we provide a formal description of our models, CPM and SPM, and describe their generative process. Both models comprise a spatial component, which represents the inferred place clusters, and a temporal component, which represents the inferred place distributions for each weekhour. Both CPM and SPM have the same spatial components, but they have different temporal components. The models are depicted in Figure 3.2 using the graphical model representation.

#### 3.2.1 Constrained Place Model (CPM)

In this subsection, we describe the temporal and spatial components of CPM, and discuss the intuition behind some of the technical decisions we have made about the model. The temporal component of CPM constrains all users to have the same place distribution for a given weekhour. Let $w$ represent an hour in a week, ranging from 1 to $W$, and let $k$ represent a place index, ranging from 1 to $K$. In case of CPM, all users have exactly $K$ places, but we will relax this requirement for SPM. Let $\theta_w \in \mathbb{R}^K$ denote a Dirichlet random variable that represents, intuitively, the users’ global place preferences at weekhour $w$. These preferences are used to generate $z_{u,w,n}$, the latent place assignment for user $u$ and observation $n$ at weekhour $w$. The place assignment is in turn used to sample the observed coordinates $\ell_{u,w,n}$ from the corresponding place cluster.

The spatial component of CPM models the $K$ place clusters, whose means and covariances are unique to each user. Intuitively, we expect each place cluster to correspond to locations such as “home”, “work”, and “gym”. Given a user $u$ and a place index $k$, each place cluster is characterized by a bivariate normal distribution, with mean $\phi_u^k$ and covariance $\Sigma_u^k$. The observed coordinates $\ell$ are considered to be noisy observations sampled from these place clusters.

Our models use conjugate priors because of the computational advantages they
Figure 3.2: Graphical model representations of CPM (top) and SPM (bottom). CPM assumes that all users share the same place distribution. In contrast, SPM assumes that all users share the same coefficients over the component place distributions, whose convex combination yields the final place distribution.
provide in Bayesian inference. Since \( \theta_w \) is the parameter of a categorical distribution, we set its prior to a symmetric Dirichlet with concentration parameter \( \alpha > 0 \) and let \( \text{Dirichlet}_K(\alpha) \) denote the \( K \)-dimensional distribution. In practice, we set \( \alpha \) to 1 and make all parameters in Dirichlet’s support equally probable, a reasonable assumption given the lack of prior information.

We set the normal distribution’s prior to the normal-inverse-Wishart (NIW) distribution. Among its parameters, \( \Lambda \in \mathbb{R}^{2 \times 2} \) is a positive definite scale matrix, \( \nu > 1 \) indicates the degrees of freedom, and together they define the distribution over the covariance matrix. The prior mean of the mean, \( \mu_u \), is customized for each user \( u \) and is computed as the mean of the user’s historical locations. The \( p_k^u \) parameter is a function of the user and place, and it ensures that the covariance of the mean matches the empirical covariance of the user’s past location data.

In location datasets, a user sometimes logs locations that are one-offs and are not representative of the user’s regular location profile. To ensure that such outliers do not affect how the model infers the user’s regular place clusters, we designate place \( K \) as a special outlier place and set its covariance’s prior mean, as determined by \( \Lambda_K \), to be very large. As for the covariance of regular clusters, we set the remaining scale matrices, \( \Lambda_{-K} \), such that each coordinate of the covariance’s prior mean has a standard deviation of 250 meters. This is a reasonable size for a place cluster, as it is large enough to encapsulate both the potential inaccuracies of location hardware and the inherent noise in the users’ locations, but small enough to ensure that each place cluster corresponds to a single intuitive place, such as “home” or “work”.

Let \( \mathcal{N} \) denote the normal distribution and \( \mathcal{IW} \) denote the inverse-Wishart distribution. The generative process of CPM is described in more formal terms below. Further technical details about the distributions we use in our model can be found in the appendix.

1. For each weekhour \( w \), draw a distribution over places \( \theta_w \sim \text{Dirichlet}_K(\alpha) \).

2. For each user \( u \) and each place \( k \),
   
   (a) Draw a place covariance \( \Sigma_u^k \sim \mathcal{IW}(\Lambda_k, \nu) \).

   (b) Draw a place mean \( \phi_u^k \sim \mathcal{N}(\mu_u, \Sigma_u^k / p_k^u) \).

3. For each user \( u \), weekhour \( w \), and observation index \( n \),
(a) Draw a place assignment \( z_{u,w,n} \sim \text{Categorical}(\theta_w) \).

(b) Draw a location \( \ell_{u,w,n} \sim \mathcal{N}(\Phi_{u}^{z_{u,w,n}}, \Sigma_{u}^{z_{u,w,n}}) \).

### 3.2.2 Smooth Place Model (SPM)

In this subsection, we turn our attention to SPM, and specify its spatial and temporal components. SPM’s spatial component is essentially identical to that of CPM, except SPM does not restrict all users to have the same number of places, whereas CPM does. In contrast, SPM’s temporal component is quite different from that of CPM, and that is what sets the two models apart.

SPM’s temporal component assumes that, for each user \( u \) and weekhour \( w \), the corresponding place distribution is a convex combination of \( F \) component place distributions. The component place distributions are unique to each user, but the coefficients of the convex combination are shared across all users. This way, SPM is able to infer each user’s place distribution for weekhours where the user has not been observed before, as well as predict each user’s future geographic coordinates accurately despite sparsity. Furthermore, by not restricting each user to have the same number of places or each weekhour to have the same distribution over places, SPM allows a more realistic and flexible representation of the users’ location profiles.

For each user \( u \) and component index \( f \), let \( \theta_{u}^{f} \in \mathbb{R}^{K_{u}} \) be a Dirichlet random variable that represents the corresponding component place distribution, where \( K_{u} \) denotes the number of places for each user \( u \). Let \( \gamma_{w} \in \mathbb{R}^{F} \) denote a Dirichlet random variable that represents the component weights, which when combined with the component place distributions, results in the user’s final place distribution for weekhour \( w \). The variable \( \gamma_{w} \) generates \( y_{u,w,n} \), the component assignment for user \( u \) and observation \( n \) at weekhour \( w \), and the component assignment generates the place assignment \( z_{u,w,n} \) from the component place distribution \( \theta_{u}^{y_{u,w,n}} \). The place assignment is in turn used to sample the observed coordinates \( \ell_{u,w,n} \) from the corresponding place cluster.

We abuse notation and let \( \text{Dirichlet}_{K}(\cdot) \) denote a symmetric Dirichlet if its parameter is a scalar and a general Dirichlet if its parameter is a vector. The generative process of SPM is described in more formal terms below. Further technical
details about the distributions we use in our model can be found in the appendix.

1. For each weekhour $w$, draw a distribution over components $\gamma^w \sim \text{Dirichlet}_F(\beta^w)$.

2. For each user $u$ and component $f$, draw a component place distribution $\theta^f_u \sim \text{Dirichlet}_{K_u}(\alpha)$.

3. For each user $u$ and place $k$,
   
   (a) Draw a place covariance $\Sigma^k_u \sim IW(\Lambda_k, v)$.

   (b) Draw a place mean $\phi^k_u \sim \mathcal{N}(\mu_u, \Sigma^k_u p_k)$.

4. For each user $u$, weekhour $w$, and observation index $n$,

   (a) Draw a component assignment $y^u, w, n \sim \text{Categorical}(\gamma^w)$.

   (b) Draw a place assignment $z^u, w, n \sim \text{Categorical}(\theta^y_u, w, n)$.

   (c) Draw a location $\ell^u, w, n \sim \mathcal{N}(\phi^z_u, w, n, \Sigma^z_u, w, n)$.

### 3.3 Inference

In this section, we derive our inference algorithm, and we present our derivation in multiple steps. First, we use a strategy popularized by Griffiths and Steyvers [21], and derive a collapsed Gibbs sampler to sample from the posterior distribution of the categorical random variables conditioned on the observed geographic coordinates. Second, we derive the conditional likelihood of the posterior samples, which we use to determine the sampler’s convergence. Third, we derive formulas for approximating the posterior expectations of the non-categorical random variables conditioned on the posterior samples. Finally, in the last step, we combine all the previous derivations to construct a simple algorithm for efficient posterior inference. In later sections, we will also show how these posteriors are used to compute users’ significant places, to analyze the way users partition their time across these places, and to predict the users’ future locations.
3.3.1 Collapsed Gibbs Sampler

Since the derivations for CPM and SPM are fairly similar, we avoid duplication and only present the derivation for SPM. In Lemmas 3.1 and 3.2, we derive the collapsed Gibbs sampler for variables \( z \) and \( y \), respectively.

Given a vector \( x \) and an index \( k \), let \( x_{-k} \) indicate all the entries of the vector excluding the one at index \( k \). For Lemmas 3.1 and 3.2, assume \( i = (u, w, n) \) denotes the index of the variable that will be sampled.

**Lemma 3.1.** The unnormalized probability of \( z_i \) conditioned on the observed location data and remaining categorical variables is

\[
p(z_i = k \mid y_i = f, z_{-i}, y_{-i}, \ell) \propto t_{\nu_k}^{-1} \left( \ell_i \mid \tilde{\nu}_k^u, \tilde{\Lambda}_u^k \right) \left( \tilde{\nu}_k^u + 1 \right) \left( \alpha + \tilde{m}^{k,f}_{u} \right)
\]

The parameters \( \tilde{\nu}_k^u, \tilde{\Lambda}_u^k, \) and \( \tilde{\nu}_k^u \) are defined in the proof. \( t \) denotes the bivariate t-distribution and \( \tilde{m}^{k,f}_{u} \) denotes counts, both of which are defined in the appendix.

**Proof.** We decompose the probability into two components using Bayes’ theorem:

\[
p(z_i = k \mid y_i = f, z_{-i}, y_{-i}, \ell) = \frac{p(\ell_i \mid z_i = k, y_i = f, z_{-i}, y_{-i}, \ell_{-i})}{p(\ell_i \mid y_i = f, z_{-i}, y_{-i}, \ell_{-i})} \times \frac{p(z_i = k \mid y_i = f, z_{-i}, y_{-i}, \ell_{-i})}{p(\ell_i \mid y_i = f, z_{-i}, y_{-i}, \ell_{-i})}
\]

\[
= \frac{p(\ell_i \mid z_i = k, z_{-i}, \ell_{-i})}{p(\ell_i \mid y_i = f, z_{-i}, y_{-i}, \ell_{-i})} \times \frac{p(z_i = k \mid y_i = f, z_{-i}, y_{-i})}{p(\ell_i \mid y_i = f, z_{-i}, y_{-i}, \ell_{-i})}
\]

\[
\propto p(\ell_i \mid z_i = k, z_{-i}, \ell_{-i}) \times p(z_i = k \mid y_i = f, z_{-i}, y_{-i})
\]

(3.1)

(3.2)
In the first part of the derivation, we operate on (3.1). We augment it with \( \phi \) and \( \Sigma \):

\[
p(\ell_i \mid z_i = k, z_{-i}, \ell_{-i}) = \int \int p(\ell_i \mid z_i = k, z_{-i}, \phi_u^k, \Sigma_u^k) \\
\times p(\phi_u^k, \Sigma_u^k \mid z_i = k, z_{-i}, \ell_{-i}) \, d\phi_u^k \, d\Sigma_u^k
\]

\[
= \int \int p(\ell_i \mid z_i = k, \phi_u^k, \Sigma_u^k) \\
\times p(\phi_u^k, \Sigma_u^k \mid z_i = k, z_{-i}, \ell_{-i}) \, d\phi_u^k \, d\Sigma_u^k
\]

\[
= \int \int \mathcal{N}(\ell_i \mid \phi_u^k, \Sigma_u^k) \\
\times p(\phi_u^k, \Sigma_u^k \mid z_i = k, z_{-i}, \ell_{-i}) \, d\phi_u^k \, d\Sigma_u^k. \quad (3.3)
\]

We convert (3.4) into a more tractable form. Let \( \tilde{M}_u^k \) be a set of indices, which we define in the appendix, and let \( \ell_{\tilde{M}_u^k} \) denote the subset of observations whose indices are in \( \tilde{M}_u^k \). In the derivation below, we treat all variables other than \( \phi_u^k \) and \( \Sigma_u^k \) as a constant:

\[
p\left(\phi_u^k, \Sigma_u^k \mid z_i = k, z_{-i}, \ell_{-i}\right) = p\left(\phi_u^k, \Sigma_u^k \mid z_i = k, z_{-i}, \ell_{\tilde{M}_u^k}\right)
\]

\[
= p\left(\phi_u^k, \Sigma_u^k \mid z_{-i}, \ell_{\tilde{M}_u^k}\right)
\]

\[
= p\left(\ell_{\tilde{M}_u^k} \mid \phi_u^k, \Sigma_u^k, z_{-i}\right) \\
\times p\left(\phi_u^k, \Sigma_u^k \mid z_{-i}\right)
\]

\[
= p\left(\ell_{\tilde{M}_u^k} \mid \phi_u^k, \Sigma_u^k, z_{-i}\right) \frac{p\left(\phi_u^k, \Sigma_u^k \mid z_{-i}\right)}{p\left(\ell_{\tilde{M}_u^k} \mid z_{-i}\right)}
\]

\[
\propto p\left(\ell_{\tilde{M}_u^k} \mid \phi_u^k, \Sigma_u^k, z_{-i}\right) p\left(\phi_u^k, \Sigma_u^k\right)
\]

\[
= \left(\prod_{j \in \tilde{M}_u^k} p\left(\ell_j \mid \phi_u^k, \Sigma_u^k, z_{-i}\right)\right) p\left(\phi_u^k, \Sigma_u^k\right)
\]

\[
= \left(\prod_{j \in \tilde{M}_u^k} p\left(\ell_j \mid \phi_u^k, \Sigma_u^k, z_j\right)\right) p\left(\phi_u^k, \Sigma_u^k\right)
\]

\[
= \left(\prod_{j \in \tilde{M}_u^k} \mathcal{N}\left(\ell_j \mid \phi_u^k, \Sigma_u^k\right)\right) \mathcal{N}\left(\phi_u^k \mid \mu_u^k, \Sigma_u^k \frac{1}{p_k^u}\right)
\]

\[
\times IW\left(\Sigma_u^k \mid \Lambda_k, v\right).
\]

Since the normal-inverse-Wishart distribution is the conjugate prior of the multi-
variate normal distribution, the posterior is also a normal-inverse-Wishart distribution,

\[ p(\phi_u^k, \Sigma_u^k | z_i = k, z_{-i}, \ell_{-i}) = \mathcal{N}(\phi_u^k | \tilde{\mu}_k^u, \Sigma_u^k | \tilde{\nu}_k^u, \tilde{\beta}_k^u) \mathcal{IW}(\Sigma_u^k | \tilde{\Lambda}_u^k, \tilde{\nu}_k^u), \]  

(3.5)

whose parameters are defined as

\[
\begin{align*}
\tilde{\mu}_k^u &= p_u^k + m_u^k, \\
\tilde{\nu}_k^u &= \nu + m_u^k, \\
\tilde{\ell}_k^u &= \frac{1}{m_u^k, j \in M_u^k} \sum_j \ell_j, \\
\tilde{\lambda}_k^u &= \frac{p_u^k \mu_u + m_u^k \tilde{\ell}_k^u}{\tilde{\mu}_k^u}, \\
\tilde{S}_u^k &= \sum_{j \in M_u^k} (\ell_j - \tilde{\ell}_k^u)(\ell_j - \tilde{\ell}_k^u)^T, \\
\tilde{\Lambda}_u^k &= \Lambda_k + \tilde{S}_u^k + \frac{p_u^k m_u^k}{p_u^k + m_u^k} \left( \tilde{\ell}_k^u - \mu_u \right) \left( \tilde{\ell}_k^u - \mu_u \right)^T.
\end{align*}
\]

The posterior parameters depicted above are derived based on the conjugacy properties of Gaussian distributions, as described in Murphy [39]. We rewrite (3.1) by combining (3.3), (3.4), and (3.5) to obtain

\[
p(\ell_i | z_i = k, z_{-i}, \ell_{-i}) = \int \int \mathcal{N}(\ell_i | \phi_u^k, \Sigma_u^k) \\
\times p(\phi_u^k, \Sigma_u^k | z_i = k, z_{-i}, \ell_{-i}) \, d\phi_u^k \, d\Sigma_u^k \\
= \int \int \mathcal{N}(\ell_i | \phi_u^k, \Sigma_u^k) \mathcal{N}(\phi_u^k | \tilde{\mu}_k^u, \Sigma_u^k | \tilde{\nu}_k^u, \tilde{\beta}_k^u) \\
\times \mathcal{IW}(\Sigma_u^k | \tilde{\Lambda}_u^k, \tilde{\nu}_k^u) \, d\phi_u^k \, d\Sigma_u^k \\
= t_{\tilde{\nu}_k^u - 1} \left( \ell_i | \tilde{\mu}_k^u, \tilde{\Lambda}_u^k (\tilde{\beta}_k^u + 1) \right), \quad (3.6)
\]

where \( t \) is the bivariate \( t \)-distribution. (3.6) is derived by applying Equation 258 from Murphy [39].

Now, we move onto the second part of the derivation. We operate on (3.2) and
augment it with $\theta$:

$$ p\left(z_i = k \mid y_i = f, z_{-i}, y_{-i}\right) = \int p\left(z_i = k \mid y_i = f, z_{-i}, y_{-i}, \theta_u^f\right) p\left(\theta_u^f \mid y_i = f, z_{-i}, y_{-i}\right) \, d\theta_u^f $$

$$ = \int p\left(z_i = k \mid y_i = f, \theta_u^f\right) \times p\left(\theta_u^f \mid y_i = f, y_{-i}, z_{-i}\right) \, d\theta_u^f. \quad (3.7) $$

We convert (3.8) into a more tractable form. As before, let $\tilde{M}^f_u$ be a set of indices, which we define in the appendix, and let $z_{\tilde{M}^f_u}$ denote the subset of place assignments whose indices are in $\tilde{M}^f_u$. In the derivation below, we treat all variables other than $\theta_u^f$ as a constant:

$$ p\left(\theta_u^f \mid y_i = f, y_{-i}, z_{-i}\right) = p\left(\theta_u^f \mid y_i = f, y_{-i}, z_{\tilde{M}^f_u}\right) $$

$$ = \frac{p\left(z_{\tilde{M}^f_u} \mid y_i = f, y_{-i}, \theta_u^f\right) p\left(y_i = f, y_{-i}, \theta_u^f\right)}{p\left(y_i = f, y_{-i}, z_{\tilde{M}^f_u}\right)} $$

$$ \propto \prod_{j \in \tilde{M}^f_u} p\left(z_j \mid y_i = f, y_{-i}, \theta_u^f\right) p\left(\theta_u^f\right) $$

$$ = \prod_{j \in \tilde{M}^f_u} \text{Categorical}\left(z_j \mid \theta_u^f\right) \text{Dirichlet}_K\left(\theta_u^f \mid \alpha\right) $$

$$ \implies p\left(\theta_u^f \mid y_i = f, y_{-i}, z_{-i}\right) = \text{Dirichlet}_K\left(\theta_u^f \mid \alpha + \tilde{m}^{1,f}_u, \ldots, \alpha + \tilde{m}^{K_u,f}_u\right), \quad (3.9) $$

where the last step follows because Dirichlet distribution is the conjugate prior of the categorical distribution. We rewrite (3.2) by combining (3.7), (3.8), and (3.9):

$$ p\left(z_i = k \mid y_i = f, z_{-i}, y_{-i}\right) = \int p\left(z_i = k \mid y_i = f, \theta_u^f\right) p\left(\theta_u^f \mid y_i = f, y_{-i}, z_{-i}\right) \, d\theta_u^f $$

$$ = \int \theta_u^{f,k} \text{Dirichlet}_K\left(\theta_u^f \mid \alpha + \tilde{m}^{1,f}_u, \ldots, \alpha + \tilde{m}^{K_u,f}_u\right) \, d\theta_u^f $$

$$ = \frac{\alpha + \tilde{m}^{k,f}_u}{K_u \alpha + \tilde{m}^{f}_u}. \quad (3.10) $$
The last step follows because it is the expected value of the Dirichlet distribution.

Finally, we combine (3.1), (3.2), (3.6), and (3.10) to obtain the unnormalized probability distribution:

\[
p(z_i = k \mid y_i = f, z_{-i}, y_{-i}, \ell) \propto p(\ell_i \mid z_i = k, z_{-i}, \ell_{-i}) \\
\times p(z_i = k \mid y_i = f, z_{-i}, y_{-i}) \\
= t_{v_k - 1} \left( \ell_i \mid \tilde{\mu}_k^u, \tilde{\nu}_k^u \right) \frac{\alpha + \tilde{m}_k^{f, u}}{K_u \alpha + \tilde{m}_k^{f, u}}.
\]

\[\text{(3.11)}\]

**Lemma 3.2.** The unnormalized probability of \( y_i \) conditioned on the observed location data and remaining categorical variables is

\[
p(y_i = f \mid z_i = k, y_{-i}, z_{-i}, \ell) \propto \frac{\alpha + \tilde{m}_k^{f, u}}{K_u \alpha + \tilde{m}_k^{f, u}} \left( \beta_{w,f} + \tilde{m}_w^{f} \right),
\]

where the counts \( \tilde{m}_k^{f, u}, \tilde{m}_w^{f} \) are defined in the appendix.

**Proof:** We decompose the probability into two components using Bayes' theorem:

\[
p(y_i = f \mid z_i = k, y_{-i}, z_{-i}, \ell) = p(y_i = f \mid z_i = k, y_{-i}, z_{-i}) \\
= p(z_i = k \mid y_i = f, z_{-i}, y_{-i}, \ell) \\
\times p(y_i = f \mid z_{-i}, y_{-i}) \\
= p(z_i = k \mid y_i = f, z_{-i}, y_{-i}) \\
\times p(y_i = f \mid z_{-i}, y_{-i}) \tag{3.11}
\]

Since (3.11) is equal to (3.2), we rewrite it using (3.10)

\[
p(z_i = k \mid y_i = f, z_{-i}, y_{-i}) = \frac{\alpha + \tilde{m}_k^{f, u}}{K_u \alpha + \tilde{m}_k^{f, u}} \tag{3.13}
\]
We operate on (3.12) and augment it with $\gamma$:

$$
\begin{align*}
\mathbb{P}(y_i = f | z_{-i}, y_{-i}) &= \mathbb{P}(y_i = f | y_{-i}) \\
&= \int \mathbb{P}(y_i = f | y_{-i}, y_w) \mathbb{P}(y_w | y_{-i}) \, dy_w \\
&= \int \mathbb{P}(y_i = f | y_w) \mathbb{P}(y_w | y_{-i}) \, dy_w \\
&= \int \gamma_{w,f} \, dy_w \\
&\quad \times \mathbb{P}(y_w | y_{-i}) \, dy_w. 
\end{align*}
$$

(3.14)

(3.15)

We convert (3.15) into a more tractable form. As before, let $\bar{M}_{i,:}$ be a set of indices, which we define in the appendix, and let $y_{\bar{M}_{i,:}}$ denote the subset of component assignments whose indices are in $\bar{M}_{i,:}$. In the derivation below, we treat all variables other than $y_w$ as a constant,

$$
\begin{align*}
\mathbb{P}(y_w | y_{-i}) &= \mathbb{P}(y_w | y_{\bar{M}_{i,:}}) \\
&= \mathbb{P}(y_{\bar{M}_{i,:}} | y_w) \frac{\mathbb{P}(y_w)}{\mathbb{P}(y_{\bar{M}_{i,:}})} \\
&\propto \mathbb{P}(y_{\bar{M}_{i,:}} | y_w) \mathbb{P}(y_w) \\
&= \prod_{j \in \bar{M}_{i,:}} \mathbb{P}(y_j | y_w) \mathbb{P}(y_w) \\
&= \prod_{j \in \bar{M}_{i,:}} \text{Categorical}(y_j | y_w) \text{Dirichlet}_F(y_w | \beta_w) \\
\Rightarrow \mathbb{P}(y_w | y_{-i}) &= \text{Dirichlet}_F\left(y_w | \beta_{w,1} + \tilde{m}_{i,1}, \ldots, \beta_{w,F} + \tilde{m}_{i,F}\right),
\end{align*}
$$

(3.16)

where the last step follows because Dirichlet distribution is the conjugate prior of the categorical distribution. We rewrite (3.12) by combining (3.14), (3.15), and...
(3.16):

\[
p(y_i = f \mid z_{-i}, y_{-i}) = \int y_{w,f} p(y_w \mid y_{-i}) \, dy_w
\]

\[=
\int y_{w,f} \text{Dirichlet}(y_w \mid \beta_{w,1} + \tilde{m}_{:,w,1}, \ldots, \beta_{w,F} + \tilde{m}_{:,w,F}) \, dy_w
\]

\[=
\frac{\beta_{w,f} + \tilde{m}_{:,w,f}}{\sum_f (\beta_{w,f} + \tilde{m}_{:,w,f})}. \tag{3.17}
\]

Finally, we combine (3.11), (3.12), (3.13), and (3.17) to obtain the unnormalized probability distribution:

\[
p(y_i = f \mid z_i = k, y_{-i}, z_{-i}, \ell) \propto p(z_i = k \mid y_i = f, z_{-i}, y_{-i})
\times p(y_i = f \mid z_{-i}, y_{-i})
\]

\[= \frac{\alpha + \tilde{m}_{:,w,f}^{k,f}}{K_u \alpha + \tilde{m}_{:,w,f}} \frac{\beta_{w,f} + \tilde{m}_{:,w,f}}{\sum_f (\beta_{w,f} + \tilde{m}_{:,w,f})}.
\]

\[\square
\]

3.3.2 Likelihoods

In this subsection, we derive the conditional likelihoods of the posterior samples conditioned on the observed geographical coordinates. Later in the chapter, when we present the algorithm for posterior inference, we will use these conditional likelihoods to determine the algorithm’s convergence.

We present the derivations in multiple lemmas and combine them in a theorem at the end of the subsection. Let \( \Gamma \) denote the gamma function.

**Lemma 3.3.** The marginal probability of the categorical random variable \( y \) is

\[
p(y) = \prod_{w=1}^{W} \Gamma \left( \sum_{f=1}^{F} \beta_{w,f} \right) \prod_{f=1}^{F} \Gamma \left( \sum_{w} \beta_{w,f} + m_{:,w}^{f} \right)
\]

\[\prod_{f=1}^{F} \Gamma \left( \beta_{w,f} \right) \prod_{f=1}^{F} \Gamma \left( \sum_{w} \beta_{w,f} + m_{:,w}^{f} \right),
\]

60
where the counts $m_{i,w}^f$ are defined in the appendix.

**Proof.** Let $\gamma = (\gamma_1, \ldots, \gamma_W)$ denote the collection of random variables for all weekhours. Below, we will augment the marginal probability with $\gamma$, and then factorize it based on the conditional independence assumptions made by our model:

$$
p(\gamma) = \int p(\gamma | \gamma) p(\gamma) \, d\gamma
= \int \left( \prod_{j \in M_{j,w}} p(y_j | \gamma) \right) \left( \prod_{w=1}^W p(\gamma_w) \right) \, d\gamma
= \int \left( \prod_{w=1}^W \prod_{j \in M_{j,w}} p(y_j | \gamma_w) \right) \left( \prod_{w=1}^W p(\gamma_w) \right) \, d\gamma
= \prod_{w=1}^W \int \left( p(\gamma_w) \prod_{j \in M_{j,w}} p(y_j | \gamma_w) \right) \, d\gamma_w
= \prod_{w=1}^W \int \left( \text{Dirichlet}_F(\gamma_w | \beta_w) \prod_{j \in M_{j,w}} \text{Categorical}(y_j | \gamma_w) \right) \, d\gamma_w. \quad (3.18)

Now, we substitute the probabilities in (3.18) with Dirichlet and categorical distri-
Lemma 3.4. The conditional probability of the categorical random variable \( z \) conditioned on \( y \) is

\[
p(z \mid y) = \prod_{u=1}^{U} \prod_{f=1}^{F} \frac{\Gamma (\alpha + m_{u}^{k,f})}{\Gamma (\alpha) \Gamma (\alpha + m_{u}^{k,f})}
\]

where the counts \( m_{u}^{k,f} \) and \( m_{u}^{k,f} \) are defined in the appendix.

Proof. Let \( \theta \) denote the collection of random variables for all users and components. Below, we will augment the conditional probability with \( \theta \), and then factorize it based on the conditional independence assump-

\[
p(y) = \prod_{w=1}^{W} \int \left( \text{Dirichlet}_F (\mathbf{y}_w \mid \mathbf{\beta}_w) \prod_{j \in M_w} \text{Categorical}(y_j \mid \mathbf{y}_w) \right) d\mathbf{y}_w
\]

\[
= \prod_{w=1}^{W} \int \left( \frac{1}{B(\mathbf{\beta}_w)} \prod_{f=1}^{F} \gamma_{w,f}^{\beta_{w,f} - 1} \right) \left( \prod_{f=1}^{F} \gamma_{w,f}^{\frac{m_{w}^{f}}{}} \right) d\mathbf{y}_w
\]

\[
= \prod_{w=1}^{W} \frac{1}{B(\mathbf{\beta}_w)} B \left( \beta_{w,1} + m_{w}^{1}, \ldots, \beta_{w,F} + m_{w}^{F} \right)
\]

\[
= \prod_{w=1}^{W} \frac{\Gamma \left( \sum_{f=1}^{F} \beta_{w,f} \right)}{\prod_{f=1}^{F} \Gamma (\beta_{w,f})} \frac{\prod_{f=1}^{F} \Gamma (\beta_{w,f} + m_{w}^{f})}{\prod_{f=1}^{F} \Gamma (\beta_{w,f})}
\]

\[
= \prod_{w=1}^{W} \frac{1}{\prod_{f=1}^{F} \Gamma (\beta_{w,f})} \frac{\prod_{f=1}^{F} \Gamma (\beta_{w,f} + m_{w}^{f})}{\prod_{f=1}^{F} \Gamma (\beta_{w,f})}
\]
tions made by our model:

\[
p(z \mid y) = \int p(z \mid y, \theta) p(\theta \mid y) \, d\theta
\]

\[
= \int \left( \prod_{j \in M'} p(z_j \mid y, \theta) \right) p(\theta) \, d\theta
\]

\[
= \int \left( \prod_{u=1}^U \prod_{f=1}^F \prod_{j \in M'_u,f} p(z_j \mid y_j, \theta_{u,f}) \right) \left( \prod_{u=1}^U \prod_{f=1}^F p(\theta_{u,f}) \right) \, d\theta
\]

\[
= \prod_{u=1}^U \prod_{f=1}^F \int \left( p(\theta_{u,f}) \prod_{j \in M'_u,f} p(z_j \mid y_j, \theta_{u,f}) \right) \, d\theta_{u,f}
\]

\[
= \prod_{u=1}^U \prod_{f=1}^F \int \left( \text{Dirichlet}_{K_u} \left( \theta_{u,f} \mid \alpha \right) \prod_{j \in M'_u,f} \text{Categorical} \left( z_j \mid \theta_{u,f} \right) \right) \, d\theta_{u,f} \tag{3.19}
\]

Now, we substitute the probabilities in (3.19) with Dirichlet and categorical distributions, which are defined in more detail in the appendix:

\[
p(z \mid y) = \prod_{u=1}^U \prod_{f=1}^F \int \left( \text{Dirichlet}_{K_u} \left( \theta_{u,f} \mid \alpha \right) \prod_{j \in M'_u,f} \text{Categorical} \left( z_j \mid \theta_{u,f} \right) \right) \, d\theta_{u,f}
\]

\[
= \prod_{u=1}^U \prod_{f=1}^F \int \left( \frac{1}{B(\alpha)} \prod_{k=1}^{K_u} \left( \theta_{u,k}^f \right)^{a-1} \left( \prod_{k=1}^{K_u} \left( \theta_{u,k}^f \right)^{m_{u,k}^f} \right) \right) \, d\theta_{u,f}
\]

\[
= \prod_{u=1}^U \prod_{f=1}^F \frac{1}{B(\alpha)} \Gamma \left( K_u \prod_{k=1}^{K_u} \Gamma \left( a + m_{u,k}^f \right) \right)
\]

\[
= \prod_{u=1}^U \prod_{f=1}^F \frac{1}{B(\alpha) \Gamma \left( (a + m_{u,k}^f) \right) \Gamma \left( a + m_{u,k}^f \right)}
\]

\[
= \prod_{u=1}^U \prod_{f=1}^F \frac{1}{B(\alpha) \Gamma \left( (a + m_{u,k}^f) \right) \Gamma \left( a + m_{u,k}^f \right)}
\]

\[
= \prod_{u=1}^U \prod_{f=1}^F \frac{1}{B(\alpha) \Gamma \left( (a + m_{u,k}^f) \right) \Gamma \left( a + m_{u,k}^f \right)}
\]

\[
= \prod_{u=1}^U \prod_{f=1}^F \frac{1}{B(\alpha) \Gamma \left( (a + m_{u,k}^f) \right) \Gamma \left( a + m_{u,k}^f \right)}
\]

\[
= \prod_{u=1}^U \prod_{f=1}^F \frac{1}{B(\alpha) \Gamma \left( (a + m_{u,k}^f) \right) \Gamma \left( a + m_{u,k}^f \right)}
\]

\[
= \prod_{u=1}^U \prod_{f=1}^F \frac{1}{B(\alpha) \Gamma \left( (a + m_{u,k}^f) \right) \Gamma \left( a + m_{u,k}^f \right)}
\]
For our final derivation, let $\Gamma_2$ denote the bivariate gamma function, and let $|\cdot|$ denote the determinant.

**Lemma 3.5.** The conditional probability of the observed locations $\ell$ conditioned on $z$ and $y$ is

$$ p(\ell | z, y) = \prod_{u=1}^U \prod_{k=1}^{K_u} \frac{\Gamma_2 \left( \frac{\nu_u}{2} \right) |\Lambda_k|^\frac{\nu}{2} p_k^u}{\pi^{n_{k,u}} \Gamma_2 \left( \frac{\nu}{2} \right) |\Lambda_u|^\frac{\nu}{2} \hat{p}_k^u}. $$

The parameters $\nu_u^k$, $\Lambda_u^k$, and $\hat{p}_k^u$ are defined in the proof, and the counts $m_{k,u}$ are defined in the appendix.

**Proof.** We will factorize the probability using the conditional independence assumptions made by the model, and then simplify the resulting probabilities by integrating out the means and covariances associated with the place clusters:

$$ p(\ell | z, y) = p(\ell | z) $$

$$ = \prod_{u=1}^U \prod_{k=1}^{K_u} p\left(\ell_{M_u^k} | z\right) $$

$$ = \prod_{u=1}^U \prod_{k=1}^{K_u} \int \int p\left(\ell_{M_u^k} | z, \phi_u^k, \Sigma_u^k\right) p\left(\phi_u^k, \Sigma_u^k\right) d\phi_u^k d\Sigma_u^k $$

$$ = \prod_{u=1}^U \prod_{k=1}^{K_u} \int \int p\left(\phi_u^k, \Sigma_u^k\right) \prod_{j \in M_u^k} p\left(\ell_j | z_j, \phi_u^k, \Sigma_u^k\right) d\phi_u^k d\Sigma_u^k $$

$$ = \prod_{u=1}^U \prod_{k=1}^{K_u} \int \int \mathcal{N}\left(\phi_u^k | \mu_u, \frac{\Sigma_u^k}{P_k} \right) IW\left(\Sigma_u^k | \Lambda_k, v\right) $$

$$ \times \prod_{j \in M_u^k} \mathcal{N}\left(\ell_j | \phi_u^k, \Sigma_u^k\right) d\phi_u^k d\Sigma_u^k. $$

We apply Equation 266 from Murphy [39], which describes the conjugacy properties of Gaussian distributions, to reformulate (3.20) into its final form:

$$ p(\ell | z, y) = \prod_{u=1}^U \prod_{k=1}^{K_u} \int \int \mathcal{N}\left(\phi_u^k | \mu_u, \frac{\Sigma_u^k}{P_k} \right) IW\left(\Sigma_u^k | \Lambda_k, v\right) \prod_{j \in M_u^k} \mathcal{N}\left(\ell_j | \phi_u^k, \Sigma_u^k\right) d\phi_u^k d\Sigma_u^k $$

$$ = \prod_{u=1}^U \prod_{k=1}^{K_u} \frac{\Gamma_2 \left( \frac{\nu_u}{2} \right) |\Lambda_k|^\frac{\nu}{2} p_k^u}{\pi^{n_{k,u}} \Gamma_2 \left( \frac{\nu}{2} \right) |\Lambda_u|^\frac{\nu}{2} \hat{p}_k^u}. $$
The definitions for $v_k^u$, $\Lambda_u^k$, and $\hat{p}_k^u$ are provided in (3.5).

Finally, we combine Lemmas 3.3, 3.4, and 3.5 to provide the log-likelihood of the samples $z$ and $y$ conditioned on the observations $\ell$.

**Theorem 3.6.** The log-likelihood of the samples $z$ and $y$ conditioned on the observations $\ell$ is

$$\log p(z, y | \ell) = \left( \sum_{w=1}^{W} \sum_{f=1}^{F} \log \Gamma \left( \beta_{w,f} + m_{w,f}^f \right) \right) + \left( \sum_{u=1}^{U} \sum_{f=1}^{F} -\log \Gamma \left( \alpha_k^u + m_{u,f}^f \right) \right) + \left( \sum_{k=1}^{K_u} \log \Gamma \left( \Lambda_k^u + m_{k,f}^u \right) \right)$$

$$+ \left( \sum_{u=1}^{U} \sum_{k=1}^{K_u} \left( \log \Gamma_2 \left( \frac{v_k^u}{2} \right) - m_{k,f}^u \log \pi - \frac{v_k^u}{2} \left( \log \hat{p}_k^u \right) \right) \right) + C,$$

where $C$ denotes the constant terms.

**Proof.** The result follows by multiplying the probabilities stated in Lemmas 3.3, 3.4, and 3.5, and applying the logarithm function.

### 3.3.3 Parameter estimation

In Subsection 3.3.1, we described a collapsed Gibbs sampler for sampling the posteriors of the categorical random variables. Below, Lemmas 3.7, 3.8, and 3.9 show how these samples, denoted as $y$ and $z$, can be used to approximate the posterior expectations of $\gamma$, $\theta$, $\phi$, and $\Sigma$.

**Lemma 3.7.** The expectation of $\gamma$ given the observed geographical coordinates and the posterior samples is

$$\hat{\gamma}_{w,f} = E[\gamma_{w,f} | y, z, \ell] = \frac{\beta_{w,f} + m_{w,f}^f}{\sum_f (\beta_{w,f} + m_{w,f}^f)},$$

where the counts $m_{w,f}^f$ are defined in the appendix.
Proof.

\[
p(y_w \mid y, z, \ell) = p(y_w \mid y_{M_{jw}}) = \frac{p(y_{M_{jw}} \mid y_w) p(y_w)}{p(y_{M_{jw}})} \frac{p(y_w) \prod_{j \in M_{jw}} p(y_j \mid y_w)}{p(y_{M_{jw}})} = \propto \text{Dirichlet}_F(y_w \mid \beta_w) \prod_{j \in M_{jw}} \text{Categorical}(y_j \mid y_w) = \text{Dirichlet}_F(y_w \mid \beta_{w,1} + m_{jw}^{-1}, \ldots, \beta_{w,F} + m_{jw}^{-F})
\]

\[
\Rightarrow \hat{y}_{w,f} = \mathbb{E}[y_{w,f} \mid y, z, \ell] = \frac{\beta_{w,f} + m_{jw}^{-f}}{\sum_f (\beta_{w,f} + m_{jw}^{-f})}
\]

Lemma 3.8. The expectation of \( \theta \) given the observed geographical coordinates and the posterior samples is

\[
\hat{\theta}_{u,k} = \mathbb{E}[\theta_{u,k} \mid y, z, \ell] = \frac{\alpha + m_{u,k}^{f}}{K_u \alpha + m_{u}^{f}}
\]

where the counts \( m_{u,k}^{k,f} \) and \( m_{u}^{f} \) are defined in the appendix.
Proof.

\[ p\left(\theta^f_u \mid y, z, \ell\right) = p\left(\theta^f_u \mid y, z\right) = p\left(\theta^f_u \mid z_{M_u^f}, y\right) = \frac{p\left(z_{M_u^f} \mid \theta^f_u, y\right) p\left(\theta^f_u \mid y\right)}{p\left(z_{M_u^f} \mid y\right)} = \frac{p\left(\theta^f_u\right) \prod_{j \in M_u^f} p\left(z_j \mid \theta^f_u, y\right)}{p\left(z_{M_u^f} \mid y\right)} \propto \text{Dirichlet}_{K_u}\left(\theta^f_u \mid \alpha\right) \prod_{j \in M_u^f} \text{Categorical}\left(z_j \mid \theta^f_u\right) = \text{Dirichlet}_{K_u}\left(\theta^f_u \mid \alpha + m^1_{M_u^f}, \ldots, \alpha + m^K_{M_u^f}\right) \Rightarrow \hat{\theta}^f_{u,k} = \mathbb{E}\left[\theta^f_{u,k} \mid y, z, \ell\right] = \frac{\alpha + m^k_{M_u^f}}{K_u \alpha + m^k_{M_u^f}}. \]

\[ \square \]

**Lemma 3.9.** The expectations of \( \phi \) and \( \Sigma \) given the observed geographical coordinates and the posterior samples is

\[ \hat{\phi}^k_u = \mathbb{E}\left[\phi^k_u \mid y, z, \ell\right] = \hat{\mu}^u_k \]

and

\[ \hat{\Sigma}^k_u = \mathbb{E}\left[\Sigma^k_u \mid y, z, \ell\right] = \frac{\hat{\Lambda}^k_u}{\hat{\phi}^u_k - 3}. \]

Parameters \( \hat{\mu}^u_k, \hat{\Lambda}^u_k, \) and \( \hat{\phi}^u_k \) are defined in the proof of Lemma 3.1.
Proof.

\[
p(\phi^k, \Sigma^k | y, z, \ell) = p(\phi^k, \Sigma^k | z, \ell) \\
= p(\phi^k, \Sigma^k | z, \ell M^k_u) \\
= \frac{p(\ell_{M^k_u} | \phi^k, \Sigma^k, z) p(\phi^k, \Sigma^k | z)}{\prod_{j \in M^k_u} p(\ell_j | \phi^k, \Sigma^k, z) p(\phi^k, \Sigma^k)} \\
= \frac{\mathcal{N}(\phi^k | \mu_u, \frac{\Sigma^k}{p^2_k}) \text{IW}(\Sigma^k | \Lambda^k_u, \nu) \prod_{j \in M^k_u} \mathcal{N}(\ell_j | \phi^k, \Sigma^k)}{p(\ell_{M^k_u} | z)} \\
\propto \mathcal{N}(\phi^k | \mu_u, \frac{\Sigma^k}{p^2_k}) \text{IW}(\Sigma^k | \Lambda^k_u, \nu) \prod_{j \in M^k_u} \mathcal{N}(\ell_j | \phi^k, \Sigma^k) \\
\implies p(\phi^k, \Sigma^k | y, z, \ell) = \mathcal{N}(\phi^k | \hat{\mu}^k, \frac{\hat{\Lambda}^k}{\hat{\nu}^k}) \text{IW}(\hat{\Sigma}^k | \hat{\Lambda}^k, \hat{\nu}^k) \\
\implies p(\phi^k | y, z, \ell) = \text{t}_{\hat{\nu}^k-1}\left(\phi^k | \hat{\mu}^k, \frac{\hat{\Lambda}^k}{\hat{\nu}^k} - 1\right) \\
\implies \hat{\phi}^k = \mathbb{E}[\phi^k | y, z, \ell] = \hat{\mu}^k \\
\implies p(\Sigma^k | y, z, \ell) = \text{IW}(\hat{\Sigma}^k | \hat{\Lambda}^k, \hat{\nu}^k) \\
\implies \hat{\Sigma}^k = \mathbb{E}[\Sigma^k | y, z, \ell] = \frac{\hat{\Lambda}^k}{\hat{\nu}^k - 3}.
\]

3.4 Experiments

We demonstrate the empirical capabilities of our models by applying them to two datasets, a dense cellular carrier dataset and a sparse mobile application dataset, whose density profiles are depicted in Figure 3.1. We assess our models quantita-
tively by measuring their location prediction performance and held-out likelihood. We also assess our models qualitatively by inspecting their inferred place clusters and the temporal distributions they associate with these clusters.

The sparse mobile application dataset consists of 5007 users and the dense cellular carrier dataset consists of 1000 users. For both dataset, each data point comprises a user ID, local time, and geographic coordinates (i.e. latitude and longitude). During preprocessing, we check if a user has logged multiple observations during the same hour, and if so, we replace these observations with their geometric median, computed using Weiszfeld’s algorithm. We then sort the data chronologically, and for each user, we partition the earlier 90% as training data and the subsequent 10% as test data.

When we applied CPM to our datasets, we observed that constraining all users to have the same place distribution yields worse predictive performance than baseline methods. Thus, we combined CPM and SPM to form a two-stage approach. In the first stage, we train a separate CPM for each user and each $K \in \{1, \ldots, 10\}$, where $K$ denotes the number of places, and for each user, we select the $K$ parameter where the corresponding model has the highest training likelihood. In the second stage, we train a single SPM, and for each user, we use the $K$ parameter that was determined in the previous stage.

We evaluate a method’s location prediction performance by measuring the distance between its predicted coordinate and the actual observed coordinate. Let $\hat{\gamma}$, $\hat{\theta}$, $\hat{\phi}$, and $\hat{\Sigma}$ be the posterior expectations computed in Section 3.3. Given a user $u$ and a weekhour $w$, our method makes a prediction by first computing the place cluster with the highest posterior weight, $\hat{k}_{u,w} = \arg\max_k \sum_f \hat{\gamma}_{w,f} \hat{\theta}_{u,k}^f$, and then outputting the posterior mean of that place cluster, $\hat{\phi}_{u,w}^{k_{u,w}}$. In contrast, the baseline method, which we call the user weekhour mode (UWM), partitions the coordinate space and assigns each observation in the training data to a grid cell. For a given user and weekhour, UWM makes a prediction by outputting the corresponding mode, which is the centroid of the grid cell with the highest number of historical observations. If a user has no previous observation at the query weekhour, UWM outputs the global mode for that user instead. The grid cell edge size is chosen optimally from $\{100, 250, 500\}$ to minimize the median distance error.

In Figure 3.3, we display the error distributions of SPM and UWM on held-out
Figure 3.3: Error distributions of UWM (top) and SPM (bottom) on held-out data. UWM yields a median distance error of 2546 meters, whereas SPM yields a median distance error of 2032 meters, a reduction in error of 20%. Furthermore, compared to UWM, SPM predicts many more examples with error less than 100 meters.

We evaluate our method qualitatively by analyzing the inferred place clusters and temporal distributions. We focus our analysis on a single user who gave us permission to dissect his location data publicly. CPM obtains its highest training likelihood when it partitions the user’s location data into 8 places. In Figure 3.4, we depict the user’s posterior place distributions, computed using $\sum_f \hat{\gamma}_{w,f} \hat{\theta}_{u}^f$. 
The top plot corresponds to CPM, and because location data is sparse, CPM learns place distributions that are jagged. In contrast, SPM copes with sparsity by sharing information across users, and consequently, learns place distributions that are smooth. In fact, SPM infers weekhour patterns that are reminiscent of home (Place 0) and work (Place 1), even though the user did not log any location data during most weekhours.

In Figure 3.5, we display the spatial cluster associated with place 0. The blue circles represent the user's historical observations and the blue pin represents the cluster center inferred by our model. Since our model is Bayesian and the predictive distribution associated with the cluster is a robust $t$-distribution, the outlier pin at the figure's far right side is not able to sway the cluster center too much. Furthermore, as depicted in Figure 3.4, the temporal distribution associated with place 0 is similar to a home distribution; it dips during typical work hours and peaks otherwise. We verified that the cluster center, represented as a blue pin, is within 50 meters of the user's apartment.

In Figure 3.6, we display the spatial cluster associated with place 1. The yellow circles represent the user's historical observations and the yellow pin represents the cluster center inferred by our model. In Figure 3.4, the temporal distribution associated with place 1 is similar to a work distribution; it peaks during typical work hours and dips otherwise. We verified that the cluster center, represented as a yellow pin, is within 50 meters of the user's work place.

In Figure 3.7, we display the spatial cluster associated with place 3. The orange circles represent the user's historical observations and the orange pin represents the cluster center inferred by our model. The posterior covariance associated with place 3 is much larger than the ones associated with previous places; the previous places covered only couple blocks whereas place 3 covers many more. A closer look at Figure 3.4 reveals that probability of place 3 peaks during the weekend. In fact, we verified that the user spends leisure time in this area.

When the goal is location prediction, our method always predicts a single geographic coordinate, the posterior mean of the most probable place cluster, associated with the query user and weekhour. However, our model actually constructs a much richer representation of how users relate to their geography and creates a probability distribution over the entire space of geographic coordinates. For ex-
Figure 3.4: Place distributions inferred by CPM (top) and SPM (bottom). Because of sparsity, CPM learns place distributions that are jagged. In contrast, SPM shares information across users and learns place distributions that are smooth. Furthermore, SPM infers weekhour patterns that are reminiscent of home (Place 0) and work (Place 1), even though the user was not observed during most weekhours.
Figure 3.5: User’s spatial cluster associated with place 0. The blue circles represent the user’s historical observations and the blue pin represents the cluster center inferred by our model. Since the predictive distribution associated with the cluster is a robust $t$-distribution, the outlier ping at the figure’s far right side is not able to sway the cluster center too much. Furthermore, we verified that the cluster center, represented by a blue pin, is within 50 meters of the user’s apartment.

Figure 3.6: User’s spatial cluster associated with place 1. The yellow circles represent the user’s historical observations and the yellow pin represents the cluster center inferred by our model. We verified that the cluster center, represented by a yellow pin, is within 50 meters of the user’s workplace.
Figure 3.7: User's spatial cluster associated with place 3. The orange circles represent the user's historical observations and the orange pin represents the cluster center inferred by our model. The posterior covariance associated with place 3 is much larger than the ones associated with previous places. We verified that the user spends leisure time in this area.

ample, for the user above, even though place 0 dominates all other place clusters during the weekend, the model nevertheless assigns a significant probability to place 2. Thus, even if we do not use the place clusters other than the top one for location prediction, learning a full probabilistic representation of users' whereabouts is important, as it enables many applications that make decisions based on rich geographic profiles rather than simple point estimates.

3.5 Discussion

In this chapter, we developed two unsupervised Bayesian graphical models, both of which construct a complete profile of users’ whereabouts based on sparse geographic coordinates. We demonstrated that our models outperform a simple but strong baseline with respect to predicting users’ future geographic coordinates. We also showed that, aside from location prediction, our models identify users’ significant places and infer temporal descriptions of how users spend their time in them. We provided a quantitative analysis of our method on a sparse mobile application
dataset, as well as qualitatively demonstrating the place clusters and hourly place distributions inferred by our method on a particular user.

3.6 Appendix

3.6.1 Miscellaneous notation

Throughout the chapter, we use various notations to represent sets of indices and their cardinalities. Vectors $y$ and $z$ denote the component and place assignments in SPM, respectively. Each vector entry is identified by a tuple index $(u, w, n)$, where $u \in \{1, \ldots, U\}$ is a user, $w \in \{1, \ldots, W\}$ is a weekhour, and $n \in \{1, \ldots, N_{u,w}\}$ is an iteration index.

For the subsequent notations, we assume that the random variables $y$ and $z$ are already sampled. We refer to a subset of indices using

$$M_{u_0,w_0}^{k_0,f_0} = \{(\hat{u}, \hat{w}, \hat{n}) \mid z_{\hat{u},\hat{w},\hat{n}} = k_0, y_{\hat{u},\hat{w},\hat{n}} = f_0, \hat{u} = u_0, \hat{w} = w_0\},$$

where $u_0$ denotes the user, $w_0$ denotes the weekhour, $k_0$ denotes the place, and $f_0$ denotes the component. If we want the subset of indices to be unrestricted with respect to a category, we use the placeholder “.”. For example,

$$M_{u_0,w_0}^{f_0} = \{(\hat{u}, \hat{w}, \hat{n}) \mid y_{\hat{u},\hat{w},\hat{n}} = f_0, \hat{u} = u_0, \hat{w} = w_0\}$$

has no constraints with respect to places.

Given a subset of indices denoted by $M$, the lowercase $m = |M|$ denotes its cardinality. For example, given a set of indices

$$M_{u_0,w_0}^{f_0} = \{(\hat{u}, \hat{w}, \hat{n}) \mid z_{\hat{u},\hat{w},\hat{n}} = \cdot, y_{\hat{u},\hat{w},\hat{n}} = f_0, \hat{u} = u_0, \hat{w} = w_0\},$$

its cardinality is

$$m_{u_0,w_0}^{f_0} = |M_{u_0,w_0}^{f_0}|.$$

For the collapsed Gibbs sampler, the sets of indices and cardinalities used in the derivations exclude the index that will be sampled. We use “~” to modify sets or cardinalities for this exclusion. Let $(u, w, n)$ denote the index that will be sampled,
then given an index set $M$, let $\tilde{M} = M - \{u, w, n\}$ represent the excluding set and let $\tilde{m} = |\tilde{M}|$ represent the corresponding cardinality. For example,

$$\tilde{M}^{f_0}_{u_0} = \tilde{M}^{f_0}_{u_0} - \{u, w, n\}$$

and

$$\tilde{m}^{f_0}_{u_0} = |\tilde{M}^{f_0}_{u_0}|.$$ 

In the proof of Lemma 3.1, parameters $\tilde{v}_u^k, \tilde{\theta}_u^k, \tilde{\delta}^u_k$, and $\tilde{p}_u^k$ are defined using cardinalities that exclude the current index $(u, w, n)$. Similarly, in the proof of Lemma 3.9, parameters $\hat{\theta}_u^k, \hat{\delta}^u_k$, and $\hat{v}_u^k$ are defined like their wiggly versions, but the counts used in their definitions do not exclude the current index.

### 3.6.2 Probability distributions

Let $\Gamma_2$ denote a bivariate gamma function, defined as

$$\Gamma_2(a) = \pi^\frac{1}{2} \prod_{j=1}^2 \Gamma \left( a + \frac{1-j}{2} \right).$$

Let $\nu > 1$ and let $\Lambda \in \mathbb{R}^{2 \times 2}$ be a positive definite scale matrix. The inverse-Wishart distribution, which is the conjugate prior to the multivariate normal distribution, is defined as

$$IW(\Sigma | \Lambda, \nu) = \frac{|\Lambda|^{\nu/2}}{2^\nu \Gamma_2(\nu/2)} |\Sigma|^{-\nu-3} \exp \left( -\frac{1}{2} \text{tr} \left( \Lambda \Sigma^{-1} \right) \right).$$

Let $\Sigma \in \mathbb{R}^{2 \times 2}$ be a positive definite covariance matrix and let $\mu \in \mathbb{R}^2$ denote a mean vector. The multivariate normal distribution is defined as

$$N(\ell | \mu, \Sigma) = (2\pi)^{-1} |\Sigma|^{-\frac{1}{2}} \exp \left( -\frac{1}{2} (\ell - \mu)^T \Sigma^{-1} (\ell - \mu) \right).$$

Let $\nu > 1$ and let $\Sigma \in \mathbb{R}^{2 \times 2}$, then the 2-dimensional $t$-distribution is defined as

$$t_\nu(x | \mu, \Sigma) = \frac{\Gamma(\nu/2 + 1)}{\Gamma(\nu/2)} \frac{|\Sigma|^{-\frac{1}{2}}}{\nu \pi} \left( 1 + \frac{1}{\nu} (x - \mu)^T \Sigma^{-1} (x - \mu) \right)^{-\frac{\nu+1}{2}}.$$
Let $K > 1$ be the number of categories and let $\alpha = (\alpha_1, \ldots, \alpha_K)$ be the concentration parameters, where $\alpha_k > 0$ for all $k \in \{1, \ldots, K\}$. Then, the $K$-dimensional Dirichlet distribution, which is the conjugate prior to the categorical distribution, is defined as

$$\text{Dirichlet}_K(x | \alpha) = \frac{1}{B(\alpha)} \prod_{k=1}^{K} x_k^{\alpha_k - 1},$$

where

$$B(\alpha) = \frac{\prod_{k=1}^{K} \Gamma(\alpha_k)}{\Gamma\left(\sum_{k=1}^{K} \alpha_k\right)}.$$

We abuse the Dirichlet notation slightly and use it to define the $K$-dimensional symmetric Dirichlet distribution as well. Let $\beta > 0$ be a scalar concentration parameter. Then, the symmetric Dirichlet distribution is defined as

$$\text{Dirichlet}_K(x | \beta) = \text{Dirichlet}_K(x | \alpha_1, \ldots, \alpha_K),$$

where $\beta = \alpha_k$ for all $k \in \{1, \ldots, K\}$. 

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Conclusion

In this thesis, we developed rigorous characterizations of three fundamental problems that involve location data, proposed novel machine learning methods for them, and justified these methods theoretically and empirically.

In Chapter 1, we presented a probabilistic graphical model that locates radio-tagged animals and learns their movement patterns. Our method incorporated both geographical and non-geographical spatial features and provided researchers with an interpretable model that enunciates the relationship between the animal and its environment. We showed that our model generalizes random walk and demonstrated empirically that a richer model improves animal location estimates. We also provided a fast parameter estimation algorithm and demonstrated its effectiveness both asymptotically and empirically. We applied our model to real datasets and demonstrated that it outperforms the most popular radio telemetry software package in ecology.

In Chapter 2, we formulated a new collaborative ranking framework, called Collaborative Local Ranking (CLR), which allowed us to formulate a wide variety of real-world ranking tasks that involve local and implicit feedback. We justified CLR with a bound on its generalization error. We also derived a simple alternating minimization algorithm and showed that each minimization step can be efficiently computed in time independent of the size of the training data. We applied CLR to a venue recommendation task and demonstrated that it outperforms state-of-the-art collaborative ranking methods, both in terms of generalization performance and running time.

In Chapter 3, we developed two unsupervised Bayesian graphical models, both of which construct a complete profile of users’ whereabouts based on sparse geographic coordinates. We demonstrated that our models outperform a simple but
strong baseline with respect to predicting users’ future geographic coordinates. We also showed that, aside from location prediction, our models identify users’ significant places and infer temporal descriptions of how users spend their time in them. We provided a quantitative analysis of our method on a sparse mobile application dataset, as well as qualitatively demonstrating the place clusters and hourly place distributions inferred by our method on specific users.

There are a multitude of ways that our research can be improved upon. In Chapter 2, we described CLR as a general collaborative ranking framework, and applying it to domains that do not involve location data would strengthen its case. In Chapter 3, we used a computationally expensive search procedure to determine the optimal number of places for each user, and designing non-parametric versions of our models with stochastic inference could improve both the efficiency and generalization capability of our methods. We leave the resolution of these open problems to future work.


