Stochastic Block Models with Multiple Continuous Attributes

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ABSTRACT

Stochastic block models (SBMs) are probabilistic models for community structure in networks, where nodes within a community are assumed to be connected to nodes within and between communities in a uniform, characteristic way. Typically, only the adjacency matrix is used to perform SBM parameter inference. In this paper, we consider circumstances in which nodes have an associated vector of continuous attributes. While this assumption is not realistic for every application, our model assumes that the attributes associated with the nodes in a network's community can be described by a particular multivariate Gaussian model. Moreover, in this augmented, attributed SBM, the objective is to learn the SBM connectivity probability parameters and the multivariate gaussian parameters describing each community. While there are recent examples in the literature that combine connectivity and attribute information to inform community detection, our model is the first version of the stochastic block model to handle multiple continuous attributes. The lack of labeled network data often makes community detection results difficult to validate, we highlight the usefulness of our model for two two network prediction tasks, link prediction and collaborative filtering. As a result of fitting this attributed stochastic block model, one can predict the attribute vector or connectivity patterns for a new node in the event of the complementary source of information (connectivity or attributes, respectively). We also highlight two biological examples where the attributed stochastic block model provides satisfactory power in the link prediction and collaborative filtering tasks.

Keywords

Stochastic Block Model, Networks, Community Detection

1. INTRODUCTION

Uncovering patterns in network data is a common pursuit across a range of fields, such as in biology [15], medicine [1, 11] and computational social science [10]. A powerful way to

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analyze mesoscale structural organization within a network is with community structure [20, 16, 21]. In this pursuit, the objective is to identify cohesive groups of nodes with a high density of within-group connections and few between-group connections. Numerous approaches exist to accomplish this task [20, 16], but typically only the adjacency matrix conveying the wiring of the graph is taken into account. In certain applications, each node in a network is equipped with additional information (or particular attributes) that was not implicitly taken into account in the construction of the network. For example, one could consider a collection of attributes measured for individuals in a social network (i.e., age, interest, level of education). The interplay between the connectivity-based (or structural) community organization of the network and attribute information has gained attention, as it is often unclear whether it is valid to assume that a structural community should necessarily correlate with an attribute-based functional community [13, 18, 26]. While these studies suggest that extreme caution should be taken in assuming a correlation between structural and functional communities, we limit our focus in this work to the assumption that a node's connectivity and attribute patterns can be jointly modeled based on its community assignment. In other words, we sought to develop an approach to assign node to communities based jointly on both sources of information, such that a community is defined as a group of nodes with similar connectivity and attribute patterns. Moreover, our objectives are two-fold. First, we seek to develop a probabilistic approach to jointly model connectivity and attributes. Second, we wish to ensure that our model can handle multiple, continuous attributes.

1.1 Related work in attributed networks

Recently, there have been numerous efforts to incorporate attribute information into the community detection problem [27, 17, 4, 13, 18]. Two classes of community detection methods are quality function-based and probabilistic. Quality function based methods define a quantity of interest that an ideal partition would satisfy, while probabilistic methods identify communities through likelihood optimization and focus on the underlying statistical distribution for the observed network. A recent quality function-based method to handle multiple attributes is I-louvain [4]. This method approaches the problem as an extension to the Louvain algorithm, which is the state-of-the-art scalable quality function modularity-based community detection method [2]. The traditional modularity-based approach to community detection defines a null model for community structure under the assumption that there is not substantial structural

organization in the network and seeks to identify a partition maximally different from this model through optimizing a quality function (modularity). The I-louvain method modifies the standard modularity quality function as what they call 'inertia-based modularity'. This quality function incorporates the euclidean distance between nodes based on their attributes. This work demonstrates that incorporating connectivity and attributes allows for a partition of nodes to communities that aligns with ground truth better than the results obtained based on connectivity or attributes in isolation.

Alternatively, there a variety of probabilistic approaches to handling attributed network data [17, 13, 18, 27]. Similar to our work in the sense that community membership is related to node attributes is CESNA [27]. The objective in this approach is to learn a set of propensities or affiliations for each node across all possible communities, such that two nodes with similar propensities towards communities should have more in common in terms of connectivity and attributes. In this model, each node has a vector multiple binary attributes. The affiliation model is useful and flexible because it does not enforce a hard partitioning of nodes into communities, which is useful in social network applications. In this inference problem, the connectivity and attribute information are used to infer a node's affiliations to communities and then models the probability of an edge between two nodes as a function of the similarity in their community affiliation propensities.

In contrast to the affiliation model, the stochastic block model [12] (at least the more standard variants of it), seeks to determine a hard partition for each nodes across communities and models edges between a pair of nodes according to their community assignments. The partition of nodes to communities through a stochastic block model framework is accomplished through maximum likelihood optimization. The adaptation of the stochastic block model was first explored by Clauset et al., [17], which adapts the classic stochastic block model to handle a single attribute with the assumption that metadata and communities are correlated. Next Hric et al., developed an attributed SBM [13] from a multilayer network perspective, with one layer modeling relational information between attributes and the other modeling connectivity. The authors then seek to assign nodes to communities maximizing the likelihood of the observed data in each layer. Finally, work of Peel et al. made important contributions in 1) establishing a statistical test to determine whether metadata actually correlates with community structure and 2) developing an SBM with flexibility in how strongly to couple metadata and community membership in the stochastic block model inference problem [18].

The model that we seek to develop in this work is distinguished by its ability to fit a stochastic block model to networks, where each node has multiple continuous attributes. This model is most appropriate for circumstances where there is domain-specific evidence that members of a community should exhibit similarities in attributes. We highlight two examples of this in section 5, where we apply our model to a protein interaction network and a microbiome subject similarity network. Our paper is organized as follows. First, we will define our attributed SBM and an inference technique for fitting the model. Next, we test this approach on a synthetic example as well as two biological network examples. Since community detection methods are often difficult

to validate due to the lack of ground truth information on the nodes, we use link prediction and collaborative filtering to quantify how well the attributed SBM represents the data.

1.2 Stochastic Block Models

Because our model is an extension to the widely-used stochastic block model [12], we provide a brief introduction here. This model assumes that edges within a community are connected within and between communities in a characteristic or probabilistic way. To fit this model to network data, the objective is to partition the nodes into communities, such that these assignments maximize the likelihood of the model according to the observed edges. In this inference problem for a network with N nodes and K communities, a $K \times K$ probability matrix, $\boldsymbol{\theta}$ and a N-length vector, \mathbf{z} of node-tocommunity assignments are learned. The matrix $\boldsymbol{\theta}$ described the probability of connections within and between communities, while z gives the node-to-community assignments. For a network with N nodes, K communities, adjacency matrix, $\bf A$ and a learned vector $\bf z$ of node-to-community assignments the SBM models an edge between nodes i and j as,

$$P(A_{ij} = 1) \sim \text{Bernoulli}(\theta_{z_i z_j})$$
 (1)

The node-to-community assignments (**z**) are inferred through likelihood optimization and will be outlined in section 2.3. Effective inference techniques for standard stochastic block model parameters are well explored [28, 19, 6] and are achieved with the EM algorithm, belief propagation, or with MCMC accept reject sampling.

2. MODEL

2.1 Objective

We seek to incorporate both connectivity (A) and attribute information (X) to infer node-to-community assignments, Z. Note that for a network with N nodes, K communities and p measured attributes, A, X, and Z have dimensions $N \times N$, $N \times p$ and $N \times K$, respectively. In particular, Z is a binary indicator matrix, where entry z_{ic} is 1 if and only if node i belongs to community c. We also define z to be the N-dimensional vector of node-to-community assignments. We assume that connectivity and attributes are conditionally independent, given the community membership label. The graphical model for the relationship between node-to-community labels, connectivity and attribute information is shown in Fig. 1.

To infer the \mathbf{Z} that best explains the data, we adopt a likelihood maximization approach. That is, we seek to find the partition of nodes to communities that best describes the observed connectivity and attribute information. Given the conditional independence assumption of \mathbf{X} and \mathbf{A} , we can express the log likelihood of the data, \mathcal{L} as the sum of connectivity and attribute log likelihoods, \mathcal{L}_A and \mathcal{L}_X , respectively, as

$$\mathcal{L} = \mathcal{L}_A + \mathcal{L}_X \quad . \tag{2}$$

This likelihood reflects the joint distribution of the adjacency matrix, \mathbf{A} , the attribute matrix, \mathbf{X} , and the matrix of node-to-community indicators, \mathbf{Z} ; formally, we have

$$\mathcal{L} = p(\mathbf{A}, \mathbf{X}, \mathbf{Z}) . \tag{3}$$

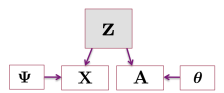


Figure 1: Modeling community membership in terms of attributes and connectivity. Node-to-community assignments specified by \mathbf{Z} are determined in terms of adjacency matrix information, \mathbf{A} and attribute matrix information, \mathbf{A} . \mathbf{A} and \mathbf{X} are assumed by be generated from a stochastic block model and a mixture of multivariate gaussian distributions, parameterized by $\boldsymbol{\theta}$ and $\boldsymbol{\Psi}$, respectively.

Given that \mathbf{Z} is a latent variable that we are trying to infer, we can approach the problem using the expectation maximization (EM) algorithm [8]. By doing this, we will alternate between estimating the posterior probability that a node i has community label c, or

$$p(z_{ic} = 1 \mid \mathbf{X}, \mathbf{A}) \tag{4}$$

and estimates for θ, Ψ , i.e., the model parameters specifying the adjacency and attribute matrices, respectively.

2.2 Attribute Likelihood

For a network with K communities, we assume that each particular community i has an associated p-dimensional mean μ_i and $p \times p$ covariance matrix, Σ_i . Note that these parameters uniquely identify a p-dimensional multivariate Gaussian distribution. To specify this model for all K communities, we define the parameter $\Psi = \{\mu_1, \mu_2, \dots \mu_k, \Sigma_1, \Sigma_2, \dots \Sigma_K\}$. Then, the probability of the i-th row of \mathbf{X} , denoted as \mathbf{x}_i , giving the values of the p attributes for node i, can be modeled as

$$p(\mathbf{x}_i \mid \mathbf{\Psi}) = \sum_{c=1}^{K} \pi_c p(\mathbf{x}_i \mid \boldsymbol{\mu}_c, \boldsymbol{\Sigma}_c) .$$
 (5)

Here, $p(\mathbf{x}_i \mid \boldsymbol{\mu}_c, \boldsymbol{\Sigma}_c)$ is the probability density function for the multivariate Gaussian and π_c is the probability that a node is assigned to community c.

2.3 Adjacency Matrix Likelihood

For the adjacency matrix, \mathbf{A} and the $K \times K$ matrix of stochastic block model parameters, $\boldsymbol{\theta}$, the probability of observing the connectivity pattern of node i, or the i-th row in the adjacency matrix, denoted as \mathbf{a}_i , is given by

$$\log p(\mathbf{a}_i \mid \mathbf{Z}, \boldsymbol{\theta}) = \sum_{c_1=1}^k z_{i,c_1} \left[\sum_{c_2=1}^k \sum_{j \in \mathcal{N}(i)} z_{j,c_2} \log(\boldsymbol{\theta}_{c_1,c_2}) - \sum_{j \notin \mathcal{N}(i)} z_{j,c_2} \log(1 - \boldsymbol{\theta}_{c_1,c_2}) \right]$$

$$(6)$$

2.4 Inference

To use EM to maximize the likelihood of the data, we break the process into the E-step and M-Step, and perform this step sequence iteratively until the estimates converge.

E-Step. During the E-step, we use the current value of learned model parameters, θ and Ψ to compute the posterior, given in Eq. (4). The posterior at each step, $\gamma(z_{ic})$, of node i belonging to community c, is given by

$$\gamma(z_{ic}) = p(z_{ic} = 1 \mid \mathbf{x}_i, \mathbf{a}_i)
= \frac{p(\mathbf{x}_i, \mathbf{a}_i, z_{ic})}{p(\mathbf{x}_i, \mathbf{a}_i)}
= \frac{p(\mathbf{x}_i \mid z_{ic})p(\mathbf{a}_i \mid z_{ic})\pi_c}{\sum_{c=1}^K p(\mathbf{x}_i \mid z_{ic})p(\mathbf{a}_i \mid z_{ic})\pi_c} .$$
(7)

M-Step. In the M-step, we can compute updates for $\boldsymbol{\theta}$ and $\boldsymbol{\Psi}$ using this expectation.

Since, the attributes follow a Gaussian mixture model, it can be shown that the update for the mean vector describing community c, μ_c , can be computed as

$$\mu_c = \frac{\sum_{i=1}^{N} \gamma(z_{ic}) \mathbf{x}_i}{\sum_{i=1}^{N} \gamma(z_{ic})} . \tag{8}$$

Similarly, the update for the covariance matrix describing a community, Σ_c , is computed as

$$\Sigma_c = \frac{\sum_{i=1}^{N} \gamma(z_{ic}) (\mathbf{x}_i - \boldsymbol{\mu}_c) (\mathbf{x}_i - \boldsymbol{\mu}_c)^T}{\sum_{i=1}^{N} \gamma(z_{ic})} . \tag{9}$$

To update the parameters of $\boldsymbol{\theta}$, we follow the method in [6] and update the probability of an edge existing between community q and l, given by $\boldsymbol{\theta}_{ql}$ as,

$$\boldsymbol{\theta}_{ql} = \frac{\sum_{i \neq j} \gamma(z_{iq}) \gamma(z_{jl}) x_{ij}}{\sum_{i \neq j} \gamma(z_{iq}) \gamma(z_{jl})}$$
(10)

We continue the process of iterating between the E-step and M-step until the change in the data log-likelihood, \mathcal{L} , is below a predefined tolerance threshold.

2.5 Initialization

Likelihood optimization approaches are often sensitive to initialization because it is easy to get stuck in a local optimum. As an initialization strategy for the nodes, we simply cluster the nodes in the network using the Louvain algorithm (cite louvain). We chose this approach because this algorithm is efficient and stable.

3. SYNTHETIC DATA RESULTS

We first test the performance of our model and inference procedure on a synthetic example. Here, we generated a network for a stochastic block model with N=200 nodes and K=4 communities . Edges in the adjacency matrix were generated according to a stochastic block model, parameterized as follows:

$$p(A_{ij} = 1) \sim \begin{cases} \text{Bernoulli}(.10), & \text{if } z_i \neq z_j \\ \text{Bernoulli}(.25), & \text{if } z_i = z_j \end{cases}$$
 (11)

Note that **z** is a 200-dimensional vector, where z_i identifies the community label for node i.

Fig. 2(A) shows the adjacency matrix for an example network generated according to this parametrization. The

black marks in the image indicate an edge. While this network has assertive structure with members of a community having more edges on average with each other than with other communities, there are still many noisy edges going between communities, making the correct community structure more difficult to discern.

To model attributes, for a community c, we randomly generated an 8-dimensional vector, μ_c , where each entry is from a Gaussian with 0-mean and unit variance. Associated with all c, $c \in \{1, 2, 3, 4\}$ is a 8×8 diagonal covariance, $\Sigma_c = \text{diag}(1.25)$. Moreover, using the μ_c and Σ_c , a sample attribute vector can be generated. That is, the attribute vector for node i, \mathbf{x}_i is generated as,

$$\mathbf{x}_i \sim \mathcal{N}(\boldsymbol{\mu}_{z_i}, \boldsymbol{\Sigma}_{z_i}) \tag{12}$$

where $\mathcal{N}(\cdot, \cdot)$ denotes a multivariate Gaussian.

Fig. 2(B) shows a PCA plot of the attribute vectors associated with each node in an example synthetic experiment and hence, each point represents a node. Since, the true dimension of these feature vectors is 8, this plot provides a projection to 2 dimensions that allows for visualization of the relatedness between the nodes, according to the attributes. One can observe that members of community 2 are overall nicely separated in attribute space but members of communities 3 and 4 are especially hard to discern.

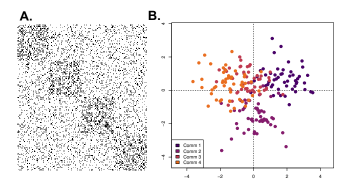


Figure 2: Synthetic Example. We generated a synthetic network with N=200 nodes, K=4 communities and an 8-dimensional multivariate gaussian for each community. A. A visualization of the adjacency matrix for this network where a black dot indicates an edge. We observe that there is an assortative block structure (blocks on the diagonal), but there are also many 'noisy' edges between communities making the true community structure with only a stochastic block model a bit harder to detect. B. We performed PCA on the $N \times p$ dimensional attribute array and plotted each of the N nodes in two dimensions. Points are colored by their true community assignments, z. Clustering the nodes according to only connectivity, only attributes, and with the attributed SBM, we quantified the partition accuracy with normalized mutual information. This gave results $NMI(\mathbf{z}, {\mathbf{z}^{connectivity}, \mathbf{z}^{attributes}, \mathbf{z}^{attribute sbm}})$ $\{0.65, 0.68, 0.83\}.$

To assess how well the attribute SBM approach performed in successfully assigning nodes to communities, we compared the results obtained from our model to clustering results obtained either only clustering the nodes based on connectivity, and to results of clustering nodes based only on their attribute information. We quantify the correctness of the partition with normalized mutual information (NMI) [5]. Letting z denote the true node-to-community assignments, then $\mathbf{z}^{\text{connectivity}}, \mathbf{z}^{\text{attributes}}, \text{ and } \mathbf{z}^{\text{attribute sbm}} \text{ denote the partition}$ of the nodes according to only the network connectivity only, attributes only, and with the attributes SBM. To cluster the network only according to connectivity, we fit a stochastic block model with 4 blocks. To cluster nodes with only attributes, we performed k-means clustering on only the attributes. Computing the NMI between ${\bf z}$ and each of these 3 cases, or $NMI(\mathbf{z}, \{\mathbf{z}^{connectivity}, \mathbf{z}^{attributes}, \mathbf{z}^{attribute sbm}\}) =$ $\{0.65, 0.68, 0.83\}$. These results show that by combining both sources of information, there is an improvement in the ability to correctly identify communities. To further probe this idea, we sought to empirically look closer at the so-called 'detectability limit'. Generally, detectability refers to the difficulty of correctly identifying clusters in data. Multiple works [7, 24] have explored these limits in stochastic block models observing the sharp phase transition that occurs in accuracy as soon as within-community probability (p_{in}) is sufficiently larger than the between-community probability $(p_{out}).$

Based on the results of the synthetic experiments in figure 2 where the attributes combined with connectivity lead to a more accurate partitioning of the nodes, we hypothesized that augmenting the network connectivity with attributes could somehow move this detectability limit. In figure 3, we explored how generating networks from a stochastic block model with varying ratios between p_{in} and p_{out} combined with the attributes used in figure 2 would affect the accuracy of the node-to-community partition. To do this, we considered values of p_{in} between 0.05 and 0.3 in increments of 0.05. For each of these p_{in} values, we found the corresponding value of p_{out} such that the mean degree was 20. For each of these p_{in} and p_{out} combinations, we generated 10 different networks using a stochastic block model. In figure 3 we plot the NMI between the true partition, z and the partitions using only the connectivity with the regular SBM $\mathbf{z}^{\text{connectivity}}$ and the attributed SBM $\mathbf{z}^{\text{attribute sbm}}$. These results are plotted in blue and pink, respectively. The shaded region around the points indicates standard deviation.

We see that while both inference approaches undergo a phase transition at a similar ratio of $p_{in}/p_{out}=3$, we notice that the curve for the attribute SBM model is slightly shifted to the left suggesting that the extra attribute information does positively impact the ability to correctly identify communities. Future work could focus on understanding detectability questions in relation to the parameters for the underlying multivariate gaussian distributions parametrizing each community. For example, how does the difference in means between a pair of communities shift the detectability curve?

4. USING THE FITTED ATTRIBUTED SBM FOR LINK PREDICTION AND COLLAB-ORATIVE FILTERING

One of the benefits of a generative network model is that it can be applied to prediction tasks. Most notably, in the absence of one source of information about a node (connec-

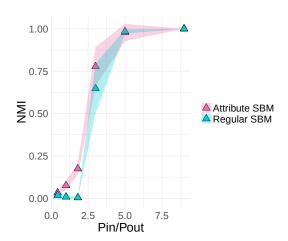


Figure 3: Detectability Analysis in Synthetic Example. To understand how attribute information can be combined with connectivity to assign nodes to communities accurately, we generated synthetic networks for withinprobabilities of p_{in} between 0.05 and 0.3 with corresponding p_{out} or between-community probabilities such that the mean degree of the network was 20. For each of these synthetic networks, we used the attributes from the analysis in figure 2 to fit the attributed SBM. Here, we plot the correctness of the node-to-community assignment with normalized mutual information using the partition obtained from regular SBM (blue) and the partition under the attributed SBM model fit (pink). For each combination of p_{in} and p_{out} , we generated 10 networks and hence the bands around the points denote standard deviation. Incorporating attributes with the attributes stochastic block model shifts the detectability limit slightly to the left.

tivity or attributes), the model can be used to predict the complementary information source (attributes or connectivity, respectively). By fitting an attributed SBM, we found that obtain successful performance in two fundamental network prediction tasks, link prediction and collaborative fil-

In the link prediction problem, when given two node stubs, the objective is to determine whether a link exists between them. Since we are modeling connectivity with a stochastic block model, we can predict links using the learned parameters. In particular, we highlight how this task can be performed using just the attribute information of the node stubs of interest. In the experiments to follow, we compare to 3 commonly-used link prediction methods. In all of these methods, a score is computed for all pairs of edge candidates and ultimately the top x set of edges with highest weights are kept (where x is some user-defined parameter). Let mand n be a pair of nodes and $\Gamma(m)$ denote the set of neighbors for a node m. Then, under the following 3 common link prediction methods [25], we can calculate the score of the potential link as, Score(m, n)

pe potential link as, Score(m, n) **Jaccard**: $Score(m, n) = \frac{\Gamma(m) \cap \Gamma(n)}{\Gamma(m) \cup \Gamma(n)}$ **Adamic Adar**: $Score(m, n) = \sum_{c \in (m) \cap \Gamma(n)} \frac{1}{\log |\Gamma(c)|}$ **Preferential Attachment**: Score $(m, n = |\Gamma(m)| \times |\Gamma(n)|$

Conversely, the collaborative filtering problem seeks to

predict a node's attributes based on its similarity to its neighbors. For some node of interest, we can use our fitted attributed SBM model to predict a node's attributes, given only the information about its connectivity. Formally, for node i, we seek to predict \mathbf{x}_i . In the following experiments, we compare our results to two common collaborative filtering approaches [22]. Let $\mathcal{N}^k(m)$ be the set of k-nearest neighbors in the network for node m. Let $\hat{\mathbf{x}}_i$ be the predicted attribute vector for node i and s_{ij} be a similarity measure between nodes i and j.

Neighborhood Avg:
$$\hat{\mathbf{x}}_i = \frac{1}{|\mathcal{N}^k(i)|} \sum_{j \in \mathcal{N}^k(i)} \mathbf{x}_j$$

Weighted Neighborhood Avg: $\hat{\mathbf{x}}_i = \frac{1}{\sum_{jin\mathcal{N}^k(i)} s_{ij}} \sum_{j \in \mathcal{N}^k(i)} s_{ij} \mathbf{x}_j$
We show results for these two tasks in two different bi-

ological network examples in section X. In particular, the experiments were designed in the following ways.

4.1 **Link Prediction Experiments**

For the link prediction tasks shown in figures 5 and 9, we performed a link prediction task by sampling pairs of nodes and utilizing the complementary source of attribute information. We sampled 10 different sets of 50 pairs of nodes. In each sample, 25 of the node pairs were those having an edge in the original network and 25 were pairs with no edge. For each of the 50 edges in each sample, we sought to predict whether an edge existed between the corresponding node pair in a leave one out manner. To do this, for each edge we fit the attributed SBM to the network with the pair of nodes (stubs) associated with the edge removed. We then use the nearest neighbor in attribute space of each stub as the the input to each of the 3 baseline community detection methods (Jaccard, Adamic Adar, and Preferential Attatchment). To use our attributed SBM in this link prediction task, we also consider the most commonly observed community among the 3 nearest neighbors for the stubs of the edge of interest. Again, using the nearest neighbors, which we denote by n and m of the stubs, then we define the link prediction score for the edge as θ_{z_n,z_m} , or the probability that an edge exists between nodes n and m according to the fitted model. After generating 10 samples of 50 edge pairs, this results in 500 total edge scores. Since we know the ground truth of whether or not these edges actually exist from the original network, we can construct an ROC curve for each method. From these curves we can plot area under the curve (AUC) to quantify the quality of the link prediction result. Using the attributed SBM is a way to incorporate community information into the link prediction problem which has previously shown to be effective [23].

Collaborative Filtering Experiments 4.2

In collaborative filtering experiments, the objective is to predict the vector of attributes for each node. In our experiments, we used leave-one-out validation to predict the attribute vector for each node. That is, for each node in the network, we created a single node test set. The training set, was then the rest of the network with the node to predict removed. For this single test set node, we identified neighbors it connects to in only connectivity space within the training set. For standard collaborative filtering approaches (Neighborhood average and weighted neighborhood average), the predicted attribute for the test set node is then the specified averaging of the neighbors. To use our model for this task, we first fit the attributed SBM model to the training set. Similar to the standard link prediction approaches, we identify the nearest neighbors for our test node in connectivity space within the training set. We then predict the community membership of our test node to be the most-frequently observed community among its neighbors. Using this community assignment, c, we then predict the attribute vector for our test node to be μ_c , or the mean vector that was learned to describe community c. The results of collaborative filtering experiments for the microbiome and protein network examples are shown in figures 6 and 10. For node a node i and its associated vector of attributes, \mathbf{x}_i we quantify the accuracy of the predicted attribute vector, $\hat{\mathbf{x}}_i$ with a a relative error measure, \mathcal{E} , such that

$$\mathcal{E} = \frac{||\hat{\mathbf{x}}_i - \mathbf{x}_i||_2}{||\mathbf{x}_i||_2}.$$
 (13)

Similar to the success of integrating community information for link prediction, an analogous effect has been observed using this information for collaborative filtering tasks [9].

5. APPLICATIONS IN BIOLOGICAL NET-WORKS

While the motivation for the development of this method was not motivated by problems in biological data, we evaluate the potential to combine similarity or relational information between a set of entities, whether that be proteins, or biological samples, and experimental data and metadata. Our application of this model to biological problems provides a framework to predict attribute or connectivity information about a new observation. Note that we do not intend to suggest any new biological insights, but rather that we can combine two sources of information for prediction tasks and alternative definitions of what constitutes a community in the data. Applying the attributed stochastic block model to integrate connectivity and attribute data provides a way to find a partition that takes into account two different sources of information, or a method to predict one source of information (connectivity, attributes) in the absence of the other (attributes, connectivity).

5.1 Microbiome Subject Similarity Results

Motivation

In the analysis of biological data, it is often useful to cluster subjects based on a set of their measure biological features and to then determine what makes each of the subgroups different. One type of biological data gaining much attention in recent years is metagenomic sequencing data, used to profile the composition of a microbiome. We refer to this as the 'metagenomic profile' and each feature is a count for each bacterial species, also known as operational taxonomic unit (OTU). Lahti et al. conducted a study among subjects across a variety of ethnicities, body mass (BMI) classifications, and age groups to understand differences in the intestinal microbiota [14]. Using metagenomic sequencing, the counts for 130 OTUs were provided for each subject. We created an experiment to test our model by seeing if we could overlay a similarity network between subjects with the individual OTU count vectors for each subject.

Pre-Processing The data were downloaded from http://datadryad.org/resource/doi:10.5061/dryad.pk75d.We extracted a subset of the subjects from Eastern Europe,Southern

Europe, Scandinavia, and the United States. Using only these subjects, a between-subject similarity network was constructed between the 121 individuals who had a BMI measurement. This resulted in a network of 121 nodes, where each edge is the pearson correlation between their microbial compositions. We then removed all edges in the network with weight (correlation) <0.7. Note that our attributed SBM does not allow for edge weights, so as input of this network into the model, we simply ignored the edge weights.

Constructing Node Attributes Since each node had a 130-dimensional vector of attributes (counts), we used this information to create a lower-dimensional attribute vector for each node by performing PCA and then representing each node with the first 5 principal components. Each dimension of this new attribute vector was then centered and scaled and has an approximately gaussian distribution.

We first visualized the differences in partitions obtained according to the classic and attributed stochastic block models in figure 4A-B, respectively. In both networks, nodes are colored by their community assignment. Using the classic stochastic block model and the model selection criteron described in [6], 7 blocks were identified. With the attributed stochastic block model, 6 blocks were identified. While we do not have ground truth labels on the nodes, it is visually apparent that adding the attributes to the inference problem helps to 'clean up' the partition. For example, in figure 4A there is mixing between the dark and lighter purple communities in the upper left of the network. In figure 4B, this mixing was reduced by assigning all of the nodes in the general region to the lighter purple community.

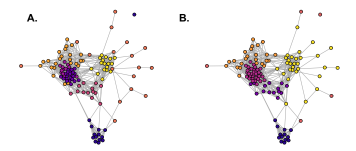


Figure 4: Microbiome subject similarity network: A visualization of the 121 node microbiome subject similarity network with nodes colored by the partition using the classic (A.) and attributed (B.) stochastic block model. A. Fitting the classic stochastic block model to the network, 7 communities were identified. B. Fitting the attributed stochastic block model to the network with the attributes being the first 5 principle components of each subject's OTU count vector (metagenomic profile), 6 communities were identified. Incorporating attributes in inferring this partition removed some of the noise in the partition on the network, specifically in the mixed purple community in the left of A.

Microbiome Link Prediction We performed link prediction on the microbiome subject similarity network as described in section 4.1. The associated ROC curves are plotted in figure 5. All four methods have satisfactory performance with the attributed stochastic block model giving

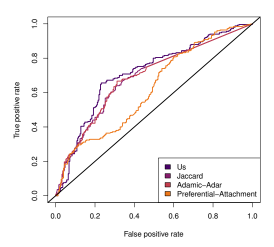


Figure 5: Link Prediction on the microbiome subject similarity network: The results for link prediction on the microbiome subject similarity network for the attributed SBM, jaccard, adamic-adar and preferntial attachment methods. The corresponding AUC values for these methods, respectively are, 0.71, 0.69, 0.69, and 0.62.

the best results. The AUC values for the attributed sbm, jaccard, agamic adar, and preferential attachment are 0.71, 0.69, 0.69, and 0.62, respectively.

Microbiome Collaborative Filtering We performed the collaborative filtering experiments on the microbiome subject similarity network in the manner described in section 4.2 to predict the 5-dimensional attribute vector for each node. The box plots indicate the distribution of relative errors over the 121 nodes for the attribute SBM (blue), neighbor average (pink) and weighted neighbor average (orange). While these distributions look similar, While the attributed SBM plotted has a similar distribution of relative errors with the standard collaborative filtering methods, the mean is slightly lower, at 0.21, compared to 0.26 and 0.27 in the neighbor average and weighted neighbor average, respectively.

5.2 Protein Interaction Network Results

We also apply our attributed SBM approach to the protein interaction network presented in [3]. This network represents interactions between proteins, predicted from the literature. Associated with each each node (protein), is a classification of one of 6 experimental modifications observed from the exposure of cancer cells to a chemotherapeutic drug. While communities in this network should reflect functional relatedness among proteins (i.e. similar biological functions, in general), we also expect that members of a community should share similarities in the observed modification type. Also associated with each of the 6 modification types is whether that particular type of modification became either more or less prominent after treatment with the drug. Since we have two types of labels associated with these nodes, we also sought to explore how these two labeling schemes (6 class vs. 2 class) aligned with the communities

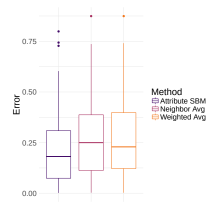


Figure 6: Collaborative Filtering Accuracy in Microbiome Subject Similarity Network: For each of the 121 nodes, we fit a model to the remaining 120 node network and given the node's closest neighbors (based on network connectivity) sought to predict its 5-dimensional attribute vector. The reported error is the relative error \mathcal{E} between the difference between the true attribute vector (\mathbf{x}_i) and its predicted attribute vector $(\hat{\mathbf{x}}_i)$. The mean error in \mathbf{x}_i is 0.21, as opposed to the neighbor average and weighted neighbor averages, having errors of 0.26 and 0.27, respectively.

returned by the algorithm.

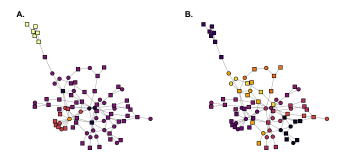


Figure 7: Protein interaction network. We visualize the 82 node protein interaction network under the classic stochastic block model **A.** and the attributed stochastic block model **B.** In both networks, nodes are colored by their community assignment and the node shape indicates whether the modification status increased (square) or decreased. **A.** Nodes colored according to the community partition under the stochastic block model. Nodes are assigned to one of five communities. **B.** Nodes are colored to the community partition under one of nine communities.

Data Pre-Processing: We downloaded the unweighted protein interaction network data and the modification information from the supplement of [3]. We removed 13 nodes that were not connected to the giant component of the network and considered only the 82 node giant connected component.

Constructing Node Attributes: As previously described, each node is classified with 1 of 6 possible modification type. For each node, we created an attribute vector that captured

the modification types of its neighbors. To do this, we considered the 4th order neighborhood of each node. That is, for each node, we collected its neighbors who were four hops or less away in the network. Then to define the value for attribute c of node i, or x_{ic} , we counted the number of fourth order neighbors of node i with label c. After defining these attributes across all nodes, for each of the 6 classes, we centered and scaled each attribute across all of the nodes.

Figure 7A-B show the results of fitting a classic SBM and attributed SBM, respectively. The 6 possible modifications exist for 3 biological processes that can either increase or decrease. The node shape reflects whether the experimental modification for a node increased (square) or decreased (circle) after treatment with the chemotherapeutic agent. Again by fitting an SBM with the model selection criterion in [6], 5 communities were identified. With our attributed SBM, 9 communities were identified.

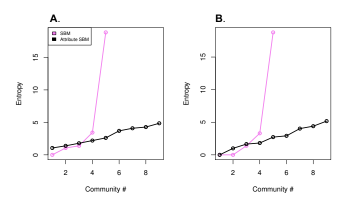


Figure 8: Community entropies in the protein interaction network. We studied the entropy of the 2 class and 6 class classifications of the nodes in $\bf A.$ and $\bf B.$, respectively under the classic SBM (black) and attributed SBM (purple) partitions. For $\bf A. - B.$ the horizontal axis denotes the community index for the particular partition. Nodes belonged to 1 of 5 communities under the classic SBM and belong to 1 of 9 communities with the attributed SBM. Incorporating attributes under both classifications succeeds in breaking up a high entropy community (5) from the classic SBM partition to lower entropy communities in the attributed SBM partition.

Using the partition of the nodes under the classic and attributed stochastic block models, we sought to use the two different classifications of the nodes (6 class modification type and 2 class increase/decrease) to compute entropy of labels within communities. The expectation is that by incorporating attribute information that is related to the functional protein information into the community detection problem, we should see a decrease in the entropy over the classification labels in communities. In figure 8A-B, we plot the entropy for the 2 class and 6 class node classifications, respectively. We define \mathbf{E}_c , the entropy for community c as,

$$\mathbf{E}_c = -\sum_k p_k \log(p_k). \tag{14}$$

Here, k is indexing the unique classifications found in community c and p_k is the probability that a node in community c belonged to classification k in community c. In these plots the black and purple curves correspond to the fits of the classic and attribute SBM fits, respectively. Using both types of node classifications to compute these entropy quantities, we see that the attribute SBM succeeds in breaking up one high entropy community (5) from the classic SBM partition into lower entropy communities.

Link Prediction in the Protein interaction network We performed link prediction on the protein interaction network using the procedure described in section 4.1. Given that this protein network is sparse, none of the link prediction methods performed particularly well. The AUC values for the attributed SBM, jaccard, adamic adar and preferential attachment were 0.61, 0.58, 0.58, and 0.54, respectively. The associated ROC curves are shown in figure 9.

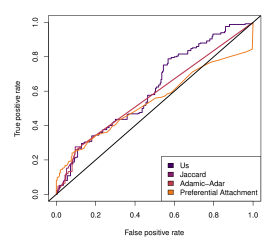


Figure 9: Link Prediction in the protein interaction network. Performing link prediction using the attributed SBM, jaccard, adamic adar, and preferential attachment. The corresponding AUC curves for these methods were 0.61, 0.58, 0.58, and 0.51, respectively.

Collaborative filtering in the protein interaction network Collaborative filtering was performed using the method described in section 4.2. Note that unlike the microbiome sample similarity network, the edges in this network are unweighted and hence the neighbor average and weighted neighbor average methods produce the same result. We note that performing collaborative filtering with the attributed stochastic block model results in a lower mean error of 0.21 compared to that of 0.48 when using the neighbor average. Similar to figure 5, the box plots represent the distribution of errors across each of the 82 nodes.

6. CONCLUSION AND FUTURE WORK

In this paper, we defined an attributed stochastic block model, where a node's community assignment determines its connectivity and its attribute vector. Our model builds on previous work with attributed stochastic block models because it can handle multiple continuous attributes. The con-

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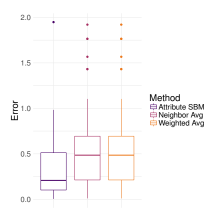


Figure 10: Collaborative filtering in the protein interaction network. For each of the 82 nodes, we fit a model to the remaining 81 node network and given the node's closest neighbors (based on network connectivity) sought to predict its 6-dimensional attribute vector. The reported error is the relative error \mathcal{E} between the difference between the true attribute vector (\mathbf{x}_i) and its predicted attribute vector $(\hat{\mathbf{x}}_i)$. The mean error in \mathbf{x}_i using the attributed SBM is 0.21, as opposed to the neighbor average error where it is 0.48.

tinuous attributes are modeled by a gaussian mixture model, where the assumption is that the attributes for members for each community are parameterized by a unique multivariate gaussian. Since community detection results are often difficult to validate due to the lack of unlabeled network data, or definition of what a correct partition means, we quantified the ability of the fitted attributed stochastic block model to represent a particular network by performing link prediction and collaborative filtering tasks. Applying link prediction and collaborative filtering to two biological networks, we observed that the attributed SBM is useful for these applications.

Future work could extend the model to handle a combination of multiple discrete and continuous attributes. Further, while the inference or understanding of fitting a stochastic block model to weighted networks is not well understood, figuring out how to integrate edge weights and attributes in determining community structure could be useful. Finally, we briefly discussed a detectability problem in figure 3. Interesting future work could aim to characterize the properties of the attributes and connectivity and how they relate to effective identification of community structure.

Networks used across fields are becoming increasingly complex, often with multiple sources of information to integrate in order to make a conclusion for the data. Our approach to an attributed SBM advances the understanding of how to jointly consider attribute and connectivity information in a probabilistic framework.

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