

Bayesian biclustering for microbial metagenomic sequencing data via multinomial matrix factorization

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SUMMARY

High-throughput sequencing technology provides unprecedented opportunities to quantitatively explore human gut microbiome and its relation to diseases. Microbiome data are compositional, sparse, noisy, and heterogeneous, which pose serious challenges for statistical modeling. We propose an identifiable Bayesian multinomial matrix factorization model to infer overlapping clusters on both microbes and hosts. The proposed method represents the observed over-dispersed zero-inflated count matrix as Dirichlet-multinomial mixtures on which latent cluster structures are built hierarchically. Under the Bayesian framework, the number of clusters is automatically determined and available information from a taxonomic rank tree of microbes is naturally incorporated, which greatly improves the interpretability of our findings. We demonstrate the utility of the proposed approach by comparing to alternative methods in simulations. An application to a human gut microbiome data set involving patients with inflammatory bowel disease reveals interesting clusters, which contain bacteria families *Bacteroidaceae*, *Bifdobacteriaceae*, *Enterobacteriaceae*, *Fusobacteriaceae*, *Lachnospiraceae*, *Ruminococcaceae*, *Pasteurellaceae*, and *Porphyromonadaceae* that are known to be related to the inflammatory bowel disease and its sub-types according to biological literature. Our findings can help generate potential hypotheses for future investigation of the heterogeneity of the human gut microbiome.

Keywords: Bayesian nonparametric prior; Compositional data analysis; Feature allocation; Mixture model; Phylogenetic Indian buffet process.

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1. INTRODUCTION

Microbes parasitic on various parts of the human body are inseparable from the well-being of their hosts. Recent studies have shown that microbiota have profound effects on the formation, development, and progression of numerous diseases like psoriasis (Benhadou *and others*, 2018), obesity (Castaner *and others*, 2018), inflammatory bowel disease (IBD, Franzosa *and others*, 2019), preterm birth (Fettweis *and others*, 2019), and diabetes (Tilg and Moschen, 2014). In this article, we focus on IBD, a chronic and complex disease that features heterogeneity at the microbiome level. As Lloyd-Price *and others* (2019) pointed out, the disease activity is accompanied by molecular disruptions in microbial transcription, variations with taxonomic shifts, and other genomic activities. The seemingly strong association between gut microbes and IBD urges scientists to investigate microbial composition profiles in patients, which can improve our understanding of disease etiology and potentially lead to personalized treatments.

The emergence of high-throughput sequencing technology such as deep metagenomic sequencing has generated a plethora of data that have enabled researchers to quantitatively study both taxonomic and functional effect of microbiota on hosts (Turnbaugh *and others*, 2007). However, due to the compositional, sparse, heterogeneous, and noisy nature of the microbiome abundance data, they pose serious challenges in statistical modeling.

Composition. Microbiome abundance data are inherently compositional (Gloor *and others*, 2017), in the sense that individual counts are restricted by a sum constrain due to tissue size or sequencing depth. The abundance of each microbial component is only coherently interpretable relative to others within that sample. As a consequence, models that treat microbial taxa as independent variables may lead to substantial biases (Buccianti, 2013).

Sparsity. Microbial counts are sparse. Taking our IBD data as an example, more than 45% of the observations are exact zeros, which greatly complicates the sampling distribution. Excessive zeros occur mainly for two reasons: (i) bacteria are not present in tested hosts and hence the zeros are true biological zeros and (ii) the sequencing depth is not enough to capture rare bacteria which is referred to as technical zeros. Often, approaches need to explicitly differentiate between these two types of zeros to reduce estimation biases, which are addressed by the two-part model, the tobit model, and their combination (Liu *and others*, 2019).

Heterogeneity. The composition of microbiota is heterogeneous and drastically different across hosts. Methods based on iid sampling are deemed unsuitable for microbiome data analysis. Individualized characterization is necessary to unravel genuine information and avoid spurious conclusions derived from a homogeneous modeling assumption.

Noisiness. Measurements from sequencing platforms contain high levels of noises due to the technical instability, which inevitably confounds with the biological variation that researchers strive to investigate. Methods that ignore the experimental noises are susceptible to false discoveries which will be propagated to downstream analysis and hinder scientific advancement.

Current statistical methodologies in the analysis of microbiome data are largely focused on the supervised learning framework. For example, in regression analysis where covariates are compositional, the linear log-contrast model with ℓ_1 regularization was adopted in Lin *and others* (2014) and Shi *and others* (2016) to select relevant covariates in the analysis of metagenomic data. However, they do not explicitly take into account the excessive zeros but replace them with arbitrary small numbers. When treating compositional data as response, a sparse Dirichlet-multinomial regression model was employed in Chen and Li (2013) to associate microbiome composition with environmental covariates. The method is able to account for over-dispersion of observed counts and select important covariates. Xia *and others* (2013) introduced an additive logistic normal multinomial regression model and selected significant covariates via a group ℓ_1 penalty. Chen and Li (2016) proposed a zero-inflated Beta regression model. The model includes a logistic regression component to model presence or absence of microbes in samples and a Beta regression component to model non-zero microbiome abundance. Wadsworth *and others* (2017) developed a Bayesian Dirichlet-multinomial regression model combined with spike-and-slab priors to select important covariates that are predictive of microbial abundances. Grantham *and others* (2020) proposed a Bayesian mixed-effects model for capturing the effects of treatment, covariates, and latent factors on microbial responses.

There are also a rising number of models focusing on revealing microbiome interactions. For example, Friedman and Alm (2012) proposed to estimate the Pearson correlations between log-transformed components of compositional data under the assumption of sparsity, which is later implemented more efficiently with parallel computing by Watts *and others* (2018). A composition-adjusted thresholding was proposed by Cao *and others* (2019a) to obtain a sparse correlation estimate. More recently, Cai *and others* (2019) developed a Markov random field model to detect differential microbial networks. A key step in their approach is to dichotomize microbial compositions into a binary matrix. However, the dichotomization in their approach is based on a fixed quantile, the choice of which is somewhat arbitrary and sensitive.

While the majority of microbiome data analyses are performed in a supervised manner, in this paper we focus on an unsupervised learning task, namely, the probabilistic matrix factorization of microbiome data which can also be interpreted as overlapping biclustering. Many matrix factorization techniques have been proposed to handle continuous matrices (Bhattacharya and Dunson, 2011; Ročková and George, 2016), non-negative matrices (Lee and Seung, 2000; Hoyer, 2004), count matrices (Zhou and others, 2012; Gopalan and others, 2014), and binary matrices (Meeds and others, 2007; Ni and others, 2019b; Wu and others, 2019). However, none of these methods is directly applicable to compositional microbiome data. To account for both sparsity and heterogeneity of microbiome data, a matrix factorization approach based on Dirichlet prior and low-dimensional representation was proposed in Shafiei and others (2015). Ren and others (2017) developed a Bayesian nonparametric ordination approach to capture the high-dimensional microbial dependencies via low-dimensional latent factors. Recently, a low rank approximation method was proposed by Cao and others (2019b) which minimizes the multinomial likelihood-based loss function combined with a nuclear norm regularization on the composition matrix. They focused on recovering the composition and matrix factorization rather than inferring latent clustering structure which is the main objective of this article. Xu and others (2020) developed a zero-inflated Poisson factor model with Poisson rates negatively related to inflated zero occurrences. Again, their main focus was on reducing the dimensionality of the microbiome data and a separate clustering algorithm is required to identify the clusters.

In this article, we propose a Bayesian multinomial matrix factorization (MMF) model that infers the latent clustering structure from compositional, sparse, heterogeneous, and noisy microbiome data. The proposed MMF introduces a mixture model representation of observations through a set of latent variables to indicate the relative abundance of taxa. In essence, this simple formulation of the sampling model adaptively dichotomizes the multinomial observations into a binary matrix, which is more robust to noise and does not require a separate treatment of excessive zeros. Given the binary indicator matrix, priors are imposed hierarchically to characterize the heterogeneity via latent features. Specifically, we construct the hierarchical model with a combination of latent logit model, phylogenetic Indian buffet process prior (pIBP, Miller and others, 2008; Chen and others, 2016), and beta-Bernoulli prior. Using pIBP, we are able to infer an unknown number of overlapping clusters/communities of the taxa. pIBP also takes into account the taxonomic relationships among the taxa, which gives rise to more interpretable and reliable results. Conditional on the clusters of taxa, the beta-Bernoulli prior are assigned to cluster hosts, again allowing overlaps. Moreover, the sparse nature of the pIBP and beta-Bernoulli priors leads to an identifiable matrix factorization under a mild condition. Using simulations, we demonstrate that the proposed MMF has favorable performance compared to competing methods and is relatively robust to the choice of hyperparameters and misspecified tree information. We then apply MMF to an IBD microbiome data set (Qin and others, 2010), which reveals interesting clusters containing bacteria families Bacteroidaceae, Bifidobacteriaceae, Enterobacteriaceae, Fusobacteriaceae, Lachnospiraceae, Ruminococcaceae, Pasteurellaceae,

and *Porphyromonadaceae* that are known to be related to the IBD and its subtypes according to biological literature. Despite the exploratory nature of this study, our findings can help generate hypotheses for further investigation of the heterogeneity of the human gut microbiome.

The rest of this article is organized as follows. We introduce the proposed MMF model in Section 2. Posterior inference based on Markov chain Monte Carlo (MCMC) sampling is described in Section 3. In Sections 4 and 5, we respectively illustrate our approach with simulation studies and the analysis of an IBD data set. This article is concluded with a brief discussion in Section 6.

2. Model

2.1. Classifying taxon abundance via adaptive dichotomization

Let x_{ij} denote the observed count of taxon j in host i, j = 1, ..., p and i = 1, ..., n. Let $\mathbf{x}_i = (x_{i1}, ..., x_{ip})^T$ and $N_i = \sum_{i=1}^p x_{ij}$. We assume \mathbf{x}_i follows a Dirichlet-multinomial distribution,

 $\boldsymbol{x}_i \sim \text{Multinomial}(N_i, \boldsymbol{\pi}_i)$

with host-specific relative abundances,

$$\boldsymbol{\pi}_i = (\pi_{i1}, \ldots, \pi_{ip})^{\mathsf{T}} \sim \mathsf{Dirichlet}(\boldsymbol{\eta}_i),$$

where $\boldsymbol{\eta}_i = (\eta_{i1}, \dots, \eta_{ip})^T$. Note that the Dirichlet-distributed relative abundances $\boldsymbol{\pi}_i$ can be equivalently represented as normalized gamma random variables $\boldsymbol{\pi}_i = \boldsymbol{\gamma}_i / \sum_{j=1}^p \gamma_{ij}$ with unnormalized relative abundances $\boldsymbol{\gamma}_i = (\gamma_{i1}, \dots, \gamma_{ip})^T$ and $\gamma_{ij} \stackrel{\text{ind}}{\sim} \text{Gamma}(\eta_{ij}, 1)$, where the gamma distribution is parameterized as $\text{Gamma}(x; a, b) = \frac{b^a}{\Gamma(a)} x^{a-1} e^{-bx}$.

We introduce a latent indicator variable z_{ij} to classify whether a taxon *j* is significantly *present* or *absent* in host *i*. However, there is no consensus on the classification of taxa based on absolute or relative abundances. In supervised tasks, the classification rule may be chosen to minimize certain objective functions. For example, when the goal is to predict a response variable with microbiome covariates, one can potentially find an optimal dichotomization that minimizes the prediction error. Lack of such gold standard in matrix factorization, we propose a mixture model to probabilistically classify raw taxa counts into states of high presence versus low presence and absence,

$$\gamma_{ij} \sim I(z_{ij} = 1) \text{Gamma}(s_j, 1) + I(z_{ij} = 0) \text{Gamma}(t_j, 1) \text{ with } s_j > t_j.$$
 (2.1)

In words, due to the constraint $s_j > t_j$, $z_{ij} = 1$ indicates high relative abundance and $z_{ij} = 0$ indicates low relative abundance. The choice of the two-component mixture model is motivated by the fact that the distribution of microbial abundances tend to be overdispersed and bi-modal (Koren *and others*, 2013; Lahti *and others*, 2014). If we further constrain $t_j < 1$, the prior (2.1) becomes a spike-and-slab prior with Gamma(t_j , 1) as the spike distribution, assigning an infinite mass at zero. Through the multinomial sampling and adaptive discretization, the zero counts would naturally fall into the category of low abundance with high probability. Therefore, we do not need an extra zero-inflated component to explicitly deal with the zero counts in our model. In addition, the adaptive dichotomization also accounts for sequencing errors as in Parmigiani *and others* (2002). The induced distribution of x_i is a discrete mixture of Dirichletmultinomial distributions with 2^p components, with each component corresponding to one configuration of $(z_{i1}, \ldots, z_{ip})^T$. The latent variable z_{ij} can be viewed as a denoised version of the raw observations x_{ij} . A similar idea of denoising was recently used by Cai *and others* (2019) where they assumed that taxa with relative abundances lower than 0.001% are due to noise or sequencing errors and adopted the 0.25 quantile as a hard cutoff for more abundant taxa. Our approach differs from theirs in that we do not need to fix a cutoff and the proposed method adaptively dichotomizes the data. We assign hyperpriors on the unknown parameters (s_i, t_i) ,

$$p(s_i, t_i) = \text{Gamma}(s_i | \alpha_s, \beta_s) \times \text{Gamma}(t_i | \alpha_t, \beta_t) \times I(s_i > t_i),$$
(2.2)

with $\alpha_s = \alpha_t = \alpha = 1$ and $\beta_s = \beta_t = \beta = 0.1$. Sensitivity analyses will be performed on the choice of all the hyperparameters in Section 4 and Section B of the Supplementary material available at *Biostatistics* online.

The mixture model in (2.1) can reliably classify taxa with well separated relative abundances into two states. However, the classification can have greater uncertainties for taxa with less variable relative abundances across observations. In Section 2.2, we will introduce latent structures on $\mathbf{Z} = (z_{ij})$ that stabilize uncertain classifications, reduce the dimensionality, and induce overlapping cluster structure for both hosts and microbial taxa.

2.2. Biclustering taxa and hosts via binary matrix factorization

We introduce lower-dimensional matrices to characterize the heterogeneity of both rows and columns of Z. In particular, we let $A = (a_{ik}) \in \{0, 1\}^{n \times K}$ and $B = (b_{jk}) \in \{0, 1\}^{p \times K}$ denote the host-cluster and taxon-cluster matrices with K clusters. The clustering interpretations of A and B will be elaborated in Section 2.3. The number K of columns of A and B is usually much smaller than the dimensions of the original data (n and p). We link A and B to z_{ij} by a latent logit model

$$logit\{Pr(z_{ij} = 1)\} = c_j + \sum_{k=1}^{K} a_{ik} w_{jk} b_{jk},$$
(2.3)

where $logit(p) = log\{p/(1-p)\}$. If a group of hosts have a common activated biological pathway (related to normal body functions or diseases) that involves a common set of taxa, then these taxa are likely to have significant presence in those host samples. Therefore, we choose to constrain w_{jk} to be positive, although in principle they can take any values; similar considerations in a different context were made in Wood *and others* (2006). Parameter c_j represents the log odds ratio of baseline probability of the presence of taxon *j*. We assume weakly informative priors on w_{jk} and c_j , $w_{jk} \sim Gamma(\alpha_w, \beta_w)$, and $c_j \sim N(\mu_c, \sigma_c^2)$, with $\mu_c = 0, \sigma_c^2 = 100, \alpha_w = 1, \text{ and } \beta_w = 0.1$.

2.3. Indian buffet process and taxonomic rank tree

The host-cluster matrix A and taxon-cluster matrix B can be interpreted as clustering of rows and columns of Z, respectively. Host i (taxon j) belongs to cluster k if the corresponding $a_{ik} = 1$ ($b_{jk} = 1$). Since we do not constrain A and B to having unit row sums, clusters can have overlaps. This is useful in microbiome applications because a taxon can be active in multiple communities and likewise a host can also belong to more than one group. To make inference on these two matrices, we will impose a Bayesian nonparametric prior on B that can automatically determine the number K of clusters.

The Indian buffet process (IBP, Griffiths and Ghahramani, 2005) has been widely used as a Bayesian nonparametric prior on binary matrices with potentially unbounded number of columns. IBP assumes the rows of the binary matrix are exchangeable. This assumption becomes a limitation when the rows (taxa) are seemingly dependent as in our case. For instance, the relationships between taxa are commonly organized as a taxonomic rank tree. Taxa with smaller distances on the tree tend to have similar biological functions and therefore are expected to have higher probability of being in the same cluster. To incorporate this prior knowledge, we adopt the phylogenetic IBP (pIBP, Miller *and others*, 2008) to encourage taxonomically similar taxa to form clusters.



Fig. 1. A graphical representation of the model. Dashed edges and squares are deterministic, and solid edges and circles are stochastic.

To describe the generating process of pIBP, we first assume a fixed and finite number \tilde{K} of clusters and will later relax it. Conditional on \tilde{K} , we associate a parameter p_k to each column of \boldsymbol{B} , which is assigned a Beta $(m/\tilde{K}, 1)$ prior. We put a Gamma(1, 1) prior on m to infer its value from data. While the columns of \boldsymbol{B} are still independent as in IBP, entries within each column are generated jointly, with the pattern of dependence characterized by a stochastic process on a taxonomic rank tree. The tree has p taxa of interest as leaves and higher taxonomic ranks as internal/root nodes. Assume the path from every leaf up to the root contains (L-1) internal nodes, and each edge has length 1/L so that the total length of the path from every leaf to the root is 1. This implies that the marginal prior probability of $b_{jk} = 1$ is the same across taxa $j = 1, \ldots, p$.

To generate the entries of the *k*th column, we proceed as follows: (i) assign value zero to the root node of the tree; (ii) along any path from the root to a leaf, let the value change to one with an exponential rate $-\log(1-p_k)/L$; (iii) once the value has changed to one along a path from the root, all leaves below that change point are assigned value one; and (iv) set the entries in the *k*th column of **B** to the values of the corresponding leaves. By construction, leaves that are closer on the tree tend to receive identical values (of zeros or ones) in each column and therefore the corresponding taxa are more likely to fall in the same cluster. Note that the marginal prior probability of $b_{jk} = 1$ is p_k , as given in the original paper of Miller *and others* (2008). To remove the dependency of the generating process from a fixed \tilde{K} , we let \tilde{K} go to infinity and obtain the pIBP. Hereafter, we omit empty columns and denote the number of non-empty columns by K.

Conditional on taxon-cluster matrix **B** (only via K), each element a_{ik} in **A** follows an independent beta-Bernoulli distribution $a_{ik} \stackrel{\text{iid}}{\sim} \text{Bernoulli}(\rho)$, and $\rho \sim \text{Beta}(\alpha_{\rho}, \beta_{\rho})$ with $\alpha_{\rho} = \beta_{\rho} = 1$. The complete hierarchical model is represented as directed acyclic graph in Figure 1.

Our choice of the nonparametric pIBP prior allows for flexible modeling of the latent structures. First, the number of clusters is potentially unbounded (i.e., it can increase as the sample size grows) and can be inferred from data. Second, given the number of clusters, the prior model assigns positive mass on any taxon-cluster matrix \boldsymbol{B} (via pIBP) and any host-cluster matrix \boldsymbol{A} (via independent Bernoulli's conditional on \boldsymbol{B}). Third, through the logit link, the prior model on \boldsymbol{A} and \boldsymbol{B} also induces a flexible prior on the latent abundance matrix \boldsymbol{Z} .

2.4. Identifiability of the proposed model

Matrix factorization is often non-identifiable without additional assumptions. For example, model (2.3) can be written in a matrix form,

$$\boldsymbol{Q} = \boldsymbol{C} + \boldsymbol{A}\boldsymbol{B}^{\mathsf{T}},$$

where $Q = (q_{ij})$ with $q_{ij} = \text{logit}\{\Pr(z_{ij} = 1)\}$, $C = \mathbf{1}_p c^{\mathsf{T}}$ with $\mathbf{1}_p = (1, \dots, 1)^{\mathsf{T}}$ and $c = (c_1, \dots, c_p)^{\mathsf{T}}$, and, slightly abusing the notation, $B = (w_{jk}b_{jk})$ absorbs the weights w_{jk} . Let $\widetilde{A} = AP$ and $\widetilde{B} = BP$ for any $K \times K$ orthogonal matrix P. It is obvious that $\widetilde{AB}^{\mathsf{T}} = APP^{\mathsf{T}}B^{\mathsf{T}} = AB^{\mathsf{T}}$. Consequently, (A, B) and $(\widetilde{A}, \widetilde{B})$ would lead to the same Q and the same sampling distribution and are therefore non-identifiable in general. However, the fact that A is binary makes the proposed matrix factorization identifiable up to column permutations under a mild condition.

PROPOSITION 1 If A is a binary matrix and there exists an integer matrix $R \in \mathbb{Z}^{K \times n}$ such that RA = I, then A and B is uniquely identifiable up to column permutation.

Proof. Let $\widetilde{A} = AP$ with an orthogonal matrix P. We will show that P must be a permutation matrix if \widetilde{A} is a binary matrix. We have

$$\widetilde{RA} = RAP = P.$$

Since both R and \widetilde{A} are integer matrices, P must be an integer matrix. This implies that each row of P is a unit vector and P is therefore a permutation matrix. \Box

The condition is, in our opinion, mild. For example, it is satisfied if for any k = 1, ..., K, there exists i = 1, ..., n such that $a_i = e_k$ where a_i is the *i*th row of A and e_k is a unit vector with 1 at its *k*th entry (in this case R would simply be a binary matrix that acts to select those K rows of A). In words, the proposed model is identifiable if for any cluster k, there exists at least one member of this cluster that does not belong to any other clusters. Below, for completeness, we give a non-identifiable example when the condition of Proposition 1 is not met.

EXAMPLE 1 Suppose

$$\boldsymbol{AP} = \begin{pmatrix} 0 & 0 & 1 & 1 \\ 1 & 1 & 0 & 0 \\ 0 & 1 & 0 & 1 \\ 1 & 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} 1/2 & 1/2 & 1/2 & -1/2 \\ 1/2 & 1/2 & -1/2 & 1/2 \\ -1/2 & 1/2 & 1/2 & 1/2 \end{pmatrix} = \begin{pmatrix} 0 & 0 & 1 & 1 \\ 1 & 1 & 0 & 0 \\ 0 & 1 & 0 & 1 \\ 0 & 1 & 1 & 0 \end{pmatrix} = \widetilde{\boldsymbol{A}}.$$

Then, the matrix **P** satisfies $PP^{\mathsf{T}} = I$ but is not a permutation matrix.

3. POSTERIOR INFERENCE

The proposed MMF is parameterized by $\{A, B, Z, \{\gamma_i\}_{i=1}^n \{w_i, c_j, s_j, t_j\}_{j=1}^p, \{p_k\}_{k=1}^K, m, \rho\}$. We carry out the posterior inference by MCMC simulation. To improve mixing, we marginalize out unnormalized relative abundance parameters γ_i 's. While other parameters are trivial to update with Gibbs or Metropolis–Hastings (M–H), care must be taken in updating B and $\{p_k\}_{k=1}^K$, details of which are provided below. The updating procedures of other parameters are presented in Section A of the Supplementary material available at *Biostatistics* online. We let b_k and b_k^{-j} respectively denote the *k*th column of B and the *k*th column of B without the *j*th entry. Sequentially for $j = 1, \ldots, p$, we cycle through the following three steps.

Step i. Update existing (non-empty) columns k = 1, ..., K of **B**. For j = 1, ..., p, we sample the binary b_{jk} from the full conditional distribution,

$$p(b_{jk}|\cdot) \propto p(b_{jk}|\boldsymbol{b}_{k}^{-j},p_{k}) \prod_{i=1}^{n} p(z_{ij}|\{a_{ik},b_{jk},w_{jk}\}_{k=1}^{K},c_{j}\}$$

While an analytic form of $p(b_{jk}|\boldsymbol{b}_k^{-j}, p_k)$ is hard to obtain, it can be computed by the sum-product algorithm exactly and efficiently. Details of the sum-product algorithm can be found in Bishop (2006). Importantly, if a column becomes all zeros after update, we delete that column and reduce *K* by 1.

Step ii. Update p_k for existing columns of **B**. Suppose b_{jk} is any non-zero entry in the *k*th column. The full conditional of p_k is given by

$$p(p_k|\boldsymbol{b}_k,m) \propto p(p_k|b_{jk},m) p(\boldsymbol{b}_k^{-j}|p_k,b_{jk}),$$

where the first factor is a standard uniform distribution (Miller *and others*, 2008) and the second factor can be efficiently computed by decomposing it into a series of univariate conditional distributions using the chain rule. For example, without loss of generality, assuming j = 1, then

$$p(\boldsymbol{b}_{k}^{-1}|p_{k},b_{1k}) = p(b_{2k}|p_{k},b_{1k}) p(b_{3k}|p_{k},b_{1k},b_{2k}) \cdots p(b_{pk}|p_{k},b_{1k},\ldots,b_{p-1,k}),$$

where each factor can be computed again using the sum-product algorithm. Since we only know the full conditional up to a normalization constant, we draw p_k by a M-H step, where a new value is proposed from $p_k^* \sim q(p_k^*|p_k) = N(p_k, \sigma_k^2)$ and is accepted with probability

$$\min\left\{1, \frac{q(p_k|p_k^*) p(\boldsymbol{b}_k^{-j}|p_k^*, b_{jk})}{q(p_k^*|p_k) p(\boldsymbol{b}_k^{-j}|p_k, b_{jk})} I(p_k^* \in [0, 1])\right\}$$

Following the default choice in Miller *and others* (2008), we choose $\sigma_k^2 = cp_k(1 - p_k) + \delta$, with c = 0.06 and $\delta = 0.08$.

Step iii. Propose new columns. After all the existing columns are updated, we propose to add new columns. We first draw

$$K^* \sim \text{Poisson} \left(m \left\{ \psi \left((P-1)/L + 1 \right) - \psi \left((P-2)/L + 1 \right) \right\} \right),$$

where $\psi(\cdot)$ is the digamma function and *P* is the total number of nodes in the tree. If $K^* = 0$, we will go to the next step. Otherwise, we propose a set of new parameters $a_k^* = (a_{1k}^*, \ldots, a_{nk}^*)^T$ and w_{jk}^* from their prior distributions, $k = K + 1, \ldots, K + K^*$. We accept new columns and the associated new parameters with probability

$$\min\left\{1, \frac{\prod_{i=1}^{n} p(z_{ij}|\{a_{ik}, b_{jk}, w_{jk}\}_{k=1}^{K}, \{a_{ik}^{*}, b_{jk}^{*}, w_{jk}^{*}\}_{k=K+1}^{K+K^{*}}, c_{j})}{\prod_{i=1}^{n} p(z_{ij}|\{a_{ik}, b_{jk}, w_{jk}\}_{k=1}^{K}, c_{j})}\right\}$$

where $b_{j,K+1} = \ldots = b_{j,K+K^*} = 1$. Lastly, if new columns are accepted, we increase *K* by K^* and sample p_k for the new columns by a M–H step,

$$p(p_k|\boldsymbol{b}_k) \propto \{1 - (1 - p_k)^{1/L}\}(1 - p_k)^{(P-2)/L}/p_k$$

To summarize the posterior distribution based on the Monte Carlo samples, we proceed by first calculating the maximum a posteriori estimate \hat{K} of K from the marginal posterior distribution. Conditional on \widehat{K} , we find an estimate of **B** by the following procedure. For any matrices $B, \widetilde{B} \in \{0, 1\}^{p \times \widehat{K}}$, we define a distance

$$d(\boldsymbol{B}, \widetilde{\boldsymbol{B}}) = \min_{\boldsymbol{\pi}} H(\boldsymbol{B}, \boldsymbol{\pi}(\widetilde{\boldsymbol{B}})), \qquad (3.4)$$

where $\pi(\tilde{B})$ denotes a permutation of the columns of \tilde{B} and $H(\cdot, \cdot)$ is the Hamming distance between two binary matrices, i.e., counting the number of different entries between the two matrices. A point estimator \tilde{B} of B is then obtained as

$$\widehat{\boldsymbol{B}} = \operatorname*{arg\,min}_{\widetilde{\boldsymbol{B}}} \int d(\boldsymbol{B}, \widetilde{\boldsymbol{B}}) \, \mathrm{d}p(\boldsymbol{B}|\cdot),$$

where $p(\boldsymbol{B}|\cdot)$ denotes the marginal posterior distribution of \boldsymbol{B} given \widehat{K} . Empirically, both the integration and the optimization can be approximated using the available Monte Carlo samples. Specifically, we define the posterior mode $\widehat{\boldsymbol{B}}$ as

$$\widehat{\boldsymbol{B}} = \operatorname*{arg\,min}_{\widetilde{\boldsymbol{B}}\in\mathcal{B}} \frac{1}{S} \sum_{s=1}^{S} d(\boldsymbol{B}^{(s)}, \widetilde{\boldsymbol{B}}),$$

where $\mathcal{B} = \{B^{(s)}, s = 1, ..., S\}$ is the set of posterior samples of **B** and the distance function is given in (3.4). Conditional on \hat{B} , we continue to run the Markov chain for a while. Then, the point estimates of other parameters are obtained as the posterior means computed from the new Monte Carlo samples.

4. SIMULATION

In our simulation study, we considered a dataset with n = 300 hosts, p = 46 taxa, and K = 6 true clusters; similar in size to the later application. For k = 1, ..., K, we first set $a_{ik} = 1, i = 50(k - 1) + 1, ..., 50k$, and 0 all the others. Then, we randomly changed 10% of zero entries in the host-cluster matrix A to one. We used the same taxonomic rank tree as in later application to generate the taxon-cluster matrix B, which had L + 1 = 5 levels. Furthermore, cluster-specific probability parameters p_k were all set to 0.3. The resulting true A and B, along with the phylogenetic tree are shown in Figure S.1 of the Supplementary material available at *Biostatistics* online. By construction, each taxon or host was allowed to belong to multiple clusters. Latent indicators z_{ij} were generated from the logit model (2.3) with $w_j =$ $w = (2.0, 2.5, 3.0, 3.5, 4.0, 4.5)^T$ and $c_j = \log 0.5$. For the unnormalized relative abundance γ_{ij} , we simulated them from the gamma mixture model (2.1) with varying degrees of separation of mixture components, $(s_j, t_j) = (s, t) = (2, 0.7)$, (3, 0.6), and (5, 0.5). Among these three simulation scenarios, (s, t) = (2, 0.7) was the most difficult as it induced the least separation between the two mixture component. The observations were finally generated from the multinomial sampling model for which the total counts were drawn from the discrete uniform distribution U(50, 500).

We ran the MCMC algorithm of MMF for 5000 iterations with 10 random initial clusters. The first 2500 iterations were discarded as burn-in and posterior samples were retained every 5th iteration after burn-in. On average, it took 3.8 h on a 2.3 GHz Quad-Core Intel Core i7 laptop. To evaluate the recovery accuracy, we calculated the estimation errors for both A and B. Specifically, we computed the Hamming distance between the estimated and true A and B, normalized by the respective total number of elements. When the estimated number of clusters was different from the truth, we padded the smaller matrix with columns of zeros, making the resulting matrices comparable in dimension.

Table 1. Simulation results of the proposed MMF and competing methods. Average errors in estimating *A* and *B* are quantified as the Hamming distance between the estimated and true matrices, normalized by the respective total number of elements. The numbers in the parentheses are standard deviations. The smallest errors are in boldface. The competing methods are low rank approximation (LRA), non-negative matrix factorization (NNMF), zero-inflated Poisson factor model (ZIPFM), and two-step multinomial matrix factorization (TSMF)

(s,t)	(2, 0.7)		(3, 0.6)		(5, 0.5)	
	Error A	Error B	Error A	Error B	Error A	Error B
MMF	0.373 (0.062)	0.167 (0.029)	0.171 (0.022)	0.055 (0.021)	0.117 (0.023)	0.057 (0.028)
LRA	0.298 (0.042)	0.269 (0.023)	0.205 (0.052)	0.185 (0.017)	0.203 (0.058)	0.165 (0.021)
NNMF	0.351 (0.049)	0.279 (0.030)	0.288 (0.062)	0.247 (0.021)	0.256 (0.059)	0.208 (0.014)
ZIPFM	0.425 (0.014)	0.258 (0.031)	0.291 (0.008)	0.249 (0.023)	0.246 (0.002)	0.232 (0.022)
TSMF	0.382 (0.092)	0.253 (0.033)	0.325 (0.057)	0.132 (0.043)	0.237 (0.092)	0.089 (0.018)

Method evaluation. The results under three sets of true values of (s, t) are summarized in Table 1 based on 50 repeated simulations. As expected, the performance improved as the two mixture components in (2.2) became more separated, from (s, t) = (2, 0.7) to (5, 0.5). The proposed MMF was able to identify the correct number K of clusters at least 95% of the time. Figure S.1 of the Supplementary material available at *Biostatistics* online depicts the estimated host-cluster and taxon-cluster matrices \hat{A} and \hat{B} of the proposed MMF from one simulation result with the worst error rate in the scenario (s, t) = (5, 0.5)after adjusting for label switching and dropping redundant columns. They are visually quite close to the truth, indicating that the proposed method was able to consistently and accurately identify the clusters of hosts and taxa.

Comparisons with competing methods. Matrix factorization has been studied extensively in the literature. We compared the proposed MMF with three existing alternative matrix factorization methods, the low rank approximation (LRA, Cao *and others* 2019b), the non-negative matrix factorization (NNMF, Cai *and others* 2017), and the zero-inflated Poisson factor model (ZIPFM, Xu *and others* 2020).

In order to compare the performance of biclustering, the overlapping clustering method, fuzzy c-means (Bezdek *and others*, 1984), was applied to the latent factors or low rank matrices obtained from the competing methods. The dimension of latent factors was chosen by their default optimization procedure. The number of clusters was set to the truth K = 6 for competing methods whereas it was estimated for the proposed MMF. The estimation errors were defined by first converting clustering results to binary host–cluster or taxon–cluster matrix, and then calculating the distance between the estimated and true matrices as was done for the proposed method. In addition, we also considered a two-step approach that was similar to the proposed MMF, denoted by TSMF. Specifically, instead of joint modeling, the two-step approach first dichotomized observations using a default cutoff as suggested in Cai *and others* (2019) and then applied the same Bayesian nonparametric binary matrix factorization method as in MMF to the binary data. The results of all the methods are reported in Table 1. The proposed MMF consistently outperformed the competing methods in most settings (keeping in mind that the number of clusters was

Table 2. Simulation results of the misspecified model. Average errors in estimating A and B are quantified as the Hamming distance between the estimated and true matrices, normalized by the respective total number of elements. The numbers in the parentheses are standard deviations. The competing methods are low rank approximation (LRA), non-negative matrix factorization (NNMF), zero-inflated Poisson factor model (ZIPFM), and two-step multinomial matrix factorization (TSMF).

	MMF	LRA	NNMF	ZIPFM	TSMF
Error rate A	0.323	0.301	0.351	0.368	0.329
	(0.072)	(0.044)	(0.062)	(0.015)	(0.083)
Error rate B	0.157	0.293	0.319	0.382	0.191
	(0.036)	(0.032)	(0.023)	(0.019)	(0.047)

set to truth for LRA, NNMF, and ZIPFM), especially for the taxon-cluster matrix **B**. Although when (s, t) was specified to be (2, 0.7), the estimation error of **A** in LRA was smaller, the significantly more accurate result of estimating **B** in the proposed MMF shows the benefit of using the phylogenetic tree information.

Additional simulations with different values of w. We performed additional simulation for $w = (0.5, 0.6, 0.7, 0.9, 1.1, 1.2)^T$ and $(1.0, 1.2, 1.5, 1.7, 2.0, 2.3)^T$ in Section B of the Supplementary material available at *Biostatistics* online, which led to similar conclusion as above.

Misspecified model. For fairer comparison, we mimicked the generating process of microbial metagenomic sequencing data, which was different from the proposed model. In this experiment, *A* and *B* were first generated as before. Then, the true counts $Y = (y_{ij})$ were simulated from a negative binomial model,

$$\mathrm{NB}(y_{ij};\mu_{ij},\kappa_{ij}) = \frac{\Gamma(\kappa_{ij}+y_{ij})}{\Gamma(\kappa_{ij})y_{ij}!} \left(\frac{\kappa_{ij}}{\kappa_{ij}+\mu_{ij}}\right)^{\kappa_{ij}} \left(\frac{\mu_{ij}}{\kappa_{ij}+\mu_{ij}}\right)^{y_{ij}},$$

where $\mu_{ij} = \kappa_{ij} = \exp(\sum_k a_{ik} w_{jk} b_{jk} + c_j)$ such that $\mathbb{E}(y_{ij}) = \mu_{ij}$ and $\operatorname{Var}(y_{ij}) = 2\mu_{ij}$. The counts were then proportionally down-sampled from a multinomial distribution with sequencing depth uniformly chosen from (50, 500). We subsequently applied the proposed MMF and the competing methods to the down-sampled dataset. The results based on 50 repetitions are summarized in Table 2, which show the overall competitive performance of the proposed MMF over the alternatives, especially in estimating **B**.

Sensitivity analyses. We performed two sets of sensitive analyses regarding the choice of all the hyperparameters as well as the impact of misspecified tree information. The inference under MMF was relatively robust, in our opinion. Details are provided in Section B of the Supplementary material available at *Biostatistics* online. In practice, if no prior knowledge is available, we recommend to use the default non-informative prior specification in Sections 2.1–2.3.

5. Real data

Gut microbes actively interact with their hosts and have profound relevance to inflammatory bowel disease (IBD) which is a very heterogeneous disease at the microbiome level. The goal of this case study was to investigate the heterogeneous microbial profiles in relation to IBD in an unsupervised data-driven manner. We applied the proposed MMF to an IBD microbiome dataset (Qin *and others*, 2010). The data were obtained by sequencing fecal specimens collected from IBD patients as well as healthy adult controls using the Illumina's Genome Analyzer (metagenomic sequencing); details of the data generating procedure can be found in Qin *and others* (2010). The dataset contained n = 372 observations with 240 healthy hosts



Fig. 2. Real data. Heatmaps of estimated clusters using the proposed MMF. Colored cells are ones and black cells are zeros. Rows of A are hosts arranged in a block-diagonal-liked form. Rows of B are taxa which are arranged according to the taxonomic rank tree. Each column represents a cluster with overlaps. The red/green cells in the heatmap of A represents patients/controls with ones.

and 132 IBD patients, and provided information on microbial compositions at various taxonomic levels (kingdom, phylum, class, order, family, genus, and species) of these samples. We chose to work with the family level counts because lower levels (e.g., the specie level) had extremely large number of zeros (more than 80% elements in the count matrix were zeros). In addition, we filtered out families that appear in less than 10% of samples (i.e., taxa with more than 90% of zeros) and preserved only families belonging to kingdom bacteria. The resulting data had p = 46 taxa for subsequent analysis. The relationships of taxa can be naturally represented by a taxonomic rank tree. Taxa that are closer on the tree tend to have similar activities and functions. We depict the tree of 46 taxa along with their higher taxonomic ranks, kingdom, phylum, class, and order in Figure 2. This taxonomic rank tree was used as prior information to encourage the clustering of taxa that are taxonomically similar.

We ran two separate Markov chains of MMF for 10 000 iterations. The first 5000 iterations were discarded as burn-in and posterior samples were retained every 5th iteration after burn-in. It took 8.2 h on a 2.3 GHz Quad-Core Intel Core i7 laptop. To monitor the MCMC convergence, we computed the Gelman and Rubin's potential scale reduction factor (PSRF, Gelman and Rubin, 1992) for key parameters. The MCMC diagnostic did not show a sign of lack of convergence: the PSRF was 1.01 for number K of clusters and the median PSRF was < 1.1 (with stdev 0.1) for $c_j + \sum_{k=1}^{K} a_{ik}w_{jk}b_{jk}$, the quantity on the right-hand side of (2.3). The Monte Carlo samples from the two Markov chains were combined for subsequent analysis.

To check the model fit adequacy (measure of "lack-of-fit"), we performed within-sample prediction that compared the observed composition (i.e., x_i/N_i) with the posterior predictive mean. The scatter plot of predicted versus observed relative abundance of taxa is given in Figure S.2 (a) of the Supplementary material available at *Biostatistics* online showing that the within-sample prediction was accurate. The correlation between two matrices was 0.94, which indicated an adequate model fit.

Figure S.2 (b) of the Supplementary material available at *Biostatistics* online shows the posterior distribution of the number K of clusters. The posterior mode occurred at K = 6. Conditional on K, the posterior estimates of A and B are shown in Figure 2. In Figure 2a, black cells are 0, green cells are 1 for

controls, and red cells are 1 for patients. Samples that did not belong to any clusters are omitted in the figures.

Cluster 1 contained predominantly IBD patients (\sim 70%). The biclustering nature of the proposed MMF allowed us to investigate the corresponding subset of taxa that were related to these IBD patients. For example, the cluster 1 contains family *Enterobacteriaceae*, part of class *Gammaproteobacteria*, which have been reported to increase in relative abundance in patient with IBD (Lupp and others, 2007). The fact that it exclusively belonged to patient-dominated cluster 1 is consistent with its biological relevance to IBD. Moreover, genus *Fusobacterium*, a member of the family *Fusobacteriaceae*, have been found to be at a higher abundance in patients with ulcerative colitis (UC, a subtype of IBD) relative to control subjects (Ohkusa *and others*, 2002). *Fusobacteriaceae* family was also contained in cluster 1 only, which again signified the importance of this family with relevance to IBD. Generally, phyla *Proteobacteria* and *Actinobacteria* are expected to increase in IBD patients (Matsuoka and Kanai, 2015), which is consistent with our result in cluster 1. Apart from the findings that were confirmed by the existing literature, cluster 1 includes some families in phylum *Firmicutes*, which are known to play major anti-inflammatory roles and therefore their abundances are expected to decrease in IBD patients. Further biological investigation is required to validate this new finding.

Likewise, most hosts in cluster 6 were IBD patients as well. This cluster shared quite a few taxa with cluster 1, which was not surprising as they both contained predominantly IBD patients. However, they also had distinct taxa that are biologically meaningful. On the one hand, cluster 6 uniquely contained the family *Pasteurellaceae*, of which the abundances tend to increase in patients with Crohn's disease (CD, Gevers *and others*, 2014), another subtype of IBD. This suggested the possibility of these patients belonging to the CD subtype. On the other hand, cluster 1, as discussed earlier, uniquely contained the family *Fusobacteriaceae*, which suggested the possibility of these patients belonging to the UC subtype (Ohkusa *and others*, 2002).

Cluster 2 was dominated by control samples (healthy hosts). It was associated with families *Bifidobac-teriaceae* and *Ruminococcaceae*. Their members, genera *Bifidobacterium* and *Faecalibacterium*, have been shown to be protective of the host from inflammation via several mechanisms (Sokol *and others*, 2008), including the stimulation of the anti-inflammatory cytokine and down-regulation of inflammatory cytokines. A reduced abundance of genus *Odoribacter*, which belongs to family *Porphyromonadaceae*, has been discovered in the most severe form of UC called pancolitis (Morgan *and others*, 2012). It also contained families in class *Betaproteobacteria*, whose relationship with IBD is yet to be established.

Clusters 3, 4, and 5 had a mix of patients and controls. They contained families *Bacteroidaceae* and *Lachnospiraceae*. Their members, genera *Bacteroides* and *Roseburia*, have been shown to decrease in IBD patients (Machiels *and others*, 2014; Zhou and Zhi, 2016).

We have reported results that were confirmed by the biological literature. Our biclustering results also provided novel insights into the relationships between microbial abundances and IBD, which need to be further verified by biological experiments. Our discoveries were potentially useful as a guidance to design and conduct more targeted and focused experiments.

For comparison, we applied MMF without the tree information (i.e., using the ordinary IBP prior) to this dataset. The result is shown in Figure S.3 of the Supplementary material available at *Biostatistics* online. It identified four clusters, and most clusters were dominated by control samples. Without tree information, we failed to identify the cluster associated with IBD patients and two IBD-related bacteria families *Enterobacteriaceae* and *Fusobacteriaceae*, which were successfully discovered when prior knowledge regarding the taxonomic ranks were incorporated in the analysis. Additionally, taxa from the same cluster were much less similar taxonomically: the log probability of generating this matrix from the taxonomic tree was -119.95, whereas the log probability of generating the matrix inferred from the pIBP was -91.48, which indicated that the results from the pIBP prior were substantially more consistent with the taxonomic rank tree. Without using the tree information, the lack of taxonomic similarity within the identified

clusters made it hard to interpret the results biologically. Although pIBP imposed structures on taxa only, the interpretation of the clusters of hosts was significantly enhanced as we have demonstrated earlier.

As suggested by an anonymous referee, in Figure 3, we report the posterior mean of Z from (a) the proposed MMF, (b) modified MMF with host-specific s_{ij} and t_{ij} rather than s_j and t_j , and (c) modified MMF with independent Bernoulli prior on Z instead of pIBP. Additionally, as a reference, we also plot the thresholded data in Figure 3(d) by following the rule in Cai *and others* (2019). While there is no gold standard in unsupervised learning, we found that the posterior mean of Z from the proposed MMF captures the latent abundance pattern better than (b) and (c) by comparing with the deterministic reference (d).

6. DISCUSSION

In this article, we have developed a novel identifiable sparse MMF method to simultaneously cluster microbes and hosts. The proposed approach accounts for the compositional, sparse, heterogeneous, and noisy nature of microbiome data, and describes the data generating process by a hierarchical Bayesian model, which allows for probabilistic characterization of latent structures (i.e., overlapping clusters) through full posterior inference. The incorporation of taxonomic knowledge can facilitate the interpretability and reproducibility of the inferred clusters when the prior information resembles the truth. Our simulation results demonstrate the advantage of utilizing prior information to assist inference on latent clusters. In analyzing a human gut microbiome dataset, we find latent microbial communities that are closely related to IBD and its subtypes.

There are four directions that can be taken to extend this work. First, zero-inflation exists in other data types such as single-cell RNA-seq data. It is far less common to treat single-cell data as multinomial counts and therefore the proposed MMF cannot be directly applied. However, with a minor modification of the sampling distribution (e.g., zero-inflated Poisson distribution), the method can be generalized for biclustering single-cell data. Second, the joint modeling approach can be used for many other tasks beyond matrix factorization. For example, microbial networks can be inferred by replacing the matrix factorization model with a graphical model such as Markov random fields and Bayesian networks on the latent binary indicators Z. Third, MCMC allows for full posterior inference but is not scalable to large and high-dimensional data. The current inference algorithm can be substantially accelerated by using consensus Monte Carlo algorithms for big-data clustering (Ni and others, 2019a, 2020) without sacrificing much accuracy. Fourth, the overlapping clusters can be restricted to non-overlapping clusters if desired by considering random partition models including various extensions of the Dirichlet process (Lijoi and others, 2007; Favaro and Teh, 2013; De Blasi and others, 2015).

7. Software

R code, together with a complete documentation, is available on request from the first author (fangtingzhou@tamu.edu). The data that support the findings of this study are openly available in the R package curatedMetagenomicData which can be downloaded at https://github.com/waldronlab/curatedMetagenomicData/.

SUPPLEMENTARY MATERIAL

Supplementary material is available online at http://biostatistics.oxfordjournals.org.

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Conflict of Interest: None declared.



Fig. 3. The posterior mean of the latent binary matrix Z obtained from (a) MMF, (b) modified MMF with host-specific s_{ij} and t_{ij} , and (c) modified MMF with independent Bernoulli prior on Z. The hard thresholded data are given in (d). Colors change from black to green indicating 0 to 1.

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