

Covariate-dependent negative binomial factor analysis of RNA sequencing data

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Abstract

Motivation: High-throughput sequencing technologies, in particular RNA sequencing (RNA-seq), have become the basic practice for genomic studies in biomedical research. In addition to studying genes individually, for example, through differential expression analysis, investigating coordinated expression variations of genes may help reveal the underlying cellular mechanisms to derive better understanding and more effective prognosis and intervention strategies. Although there exists a variety of co-expression network based methods to analyze microarray data for this purpose, instead of blindly extending these methods for microarray data that may introduce unnecessary bias, it is crucial to develop methods well adapted to RNA-seq data to identify the functional modules of genes with similar expression patterns.

Results: We have developed a fully Bayesian covariate-dependent negative binomial factor analysis (dNBFA) method—dNBFA—for RNA-seq count data, to capture coordinated gene expression changes, while considering effects from covariates reflecting different influencing factors. Unlike existing co-expression network based methods, our proposed model does not require multiple *ad-hoc* choices on data processing, transformation, as well as co-expression measures and can be directly applied to RNA-seq data. Furthermore, being capable of incorporating covariate information, the proposed method can tackle setups with complex confounding factors in different experiment designs. Finally, the natural model parameterization removes the need for a normalization preprocessing step, as commonly adopted to compensate for the effect of sequencing-depth variations. Efficient Bayesian inference of model parameters is derived by exploiting conditional conjugacy via novel data augmentation techniques. Experimental results on several real-world RNA-seq datasets on complex diseases suggest dNBFA as a powerful tool for discovering the gene modules with significant differential expression and meaningful biological insight.

Availability and implementation: dNBFA is implemented in R language and is available at <https://github.com/siamakz/dNBFA>.

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1 Introduction

High-throughput sequencing technology has emerged as a powerful tool for life science research (Wang *et al.*, 2009). In particular, RNA sequencing (RNA-seq), which measures the expression of each gene (transcript) or genomic feature of interest by counting the number of sequence reads mapped to them, has been widely adopted for genotype-phenotype association studies. A large body of statistical tools and frameworks have been developed for identifying the genes that are differentially expressed between different groups of samples, either based on their genotypes, phenotypic traits, or treatment

conditions (Dadaneh *et al.*, 2017; Law *et al.*, 2014; Love *et al.*, 2014; Robinson *et al.*, 2010).

Living systems are complex and their behavior is coordinated by multiple components. Especially, when studying complex disease, phenotypic changes have been shown to be associated with coordinated regulation in functional modules of interacting genes (pathways or sub-networks) rather than statistically significant changes in individual genes (Nam and Kim, 2008). Therefore, a class of approaches has been developed to detect genes with similar expression patterns as potential functional modules. Weighted Gene

Co-Expression Network Analysis (WGCNA; Langfelder and Horvath, 2008), a popular tool for gene co-expression network analysis, first constructs an adjacency matrix based on the pairwise co-expression measures, for example, based on the correlation between gene expressions across samples; then it assigns genes to different modules using the hierarchical clustering algorithm. DiffCoEx (Tesson et al., 2010) builds on WGCNA, and by computing the matrix of adjacency differences between different experiment conditions, aims at identifying differentially co-expressed genes. Several *targeted* methods also have been proposed for studying co-expression changes across conditions, relying on pre-defined gene modules (Cho et al., 2009; Choi et al., 2005; Choi and Kendzierski, 2009). For instance, Choi and Kendzierski (2009) focuses on the analysis of modules based on known gene annotations, such as gene ontology categories.

All of the aforementioned methods were proposed for data generated from microarray based experiments; and thus there remains a lack of tools for gene module detection specifically designed for RNA-seq count data. Furthermore, the existing methods often require prior knowledge from either manual annotations or other module identification methods. They need to be supplied with prepared lists of genes as candidate functional modules. For example, Lei et al. (2017) have proposed a network module-based generalized linear model for identifying differentially expressed pre-defined gene sets.

A suitable method for gene module identification based on RNA-seq data should explicitly model highly over-dispersed count data that are often skewed (Datta and Nettleton, 2014) to avoid potential bias introduced by inappropriate modeling. One of the most popular solutions to account for over-dispersion due to biological variability is using the negative binomial (NB) distribution, which possesses a quadratic variance-mean relationship. More importantly, the number of *ad-hoc* choices in modeling and data analytics should be minimized. Many existing methods, which often take two stages to first construct co-expression networks based on expression profile data and then identify co-expressed modules based on different clustering methods, may lead to uncertain results sensitive to different choices. Last but not least, when dealing with RNA-seq data, the variability of the sequencing depths across samples needs to be taken into account.

In this paper, we propose a novel covariate-dependent NB factorization model for identifying gene modules in RNA-seq experiments. The proposed method, directly applied to gene counts from RNA-seq, obviates the need for multiple *ad-hoc* steps as required in co-expression network analyses of WGCNA (Langfelder and Horvath, 2008) and DiffCoEx (Tesson et al., 2010). In addition, by employing a flexible regression model for the scale parameter of the gamma distribution in our fully Bayesian NB factor analysis model, dNBFA is capable of tackling RNA-seq experiments with complex confounding factors and quantifies the impact of these factors on the identified modules. Finally, similar to the mechanisms employed in Dadaneh et al. (2017), this new approach does not require an *ad-hoc* normalization step, as the model accounts for the sequencing-depth heterogeneity of different samples automatically.

For posterior inference of the model parameters of dNBFA, closed-form Gibbs sampling update equations are derived by exploiting novel data augmentation techniques. More specifically, we apply the augmentation technique of Zhou and Carin (2015) for the NB distribution, and the Polya-Gamma (PG) distributed auxiliary variable technique of Polson et al. (2013) for the closed-form inference of regression coefficients, removing the need for non-trivial Metropolis-Hastings correction steps (Chib and Greenberg, 1995).

Experimental results on various RNA-seq datasets demonstrate the superior performance of dNBFA in finding more biologically significant modules, in comparison to the *two-stage* methods, such as WGCNA and DiffCoEx.

The remainder of this paper is organized as follows. In Section 2, we introduce the model and its Gibbs sampling inference for covariate-dependent NB factor analysis of RNA-seq data. In Section 3, we present our experimental results on multiple The Cancer Genome Atlas (TCGA) cancer datasets as well as a Autism study dataset, which demonstrate that not only the modules identified by dNBFA are more differentially expressed, but also the incorporation of confounding factors in dNBFA may increase the chance for relevant biological discoveries tailored to the designed experiments. We conclude the paper in Section 4.

2 Materials and methods

2.1 NB factor analysis

In this section, we first present the Negative Binomial Factor Analysis (NBFA) method for count data (Zhou, 2017), and demonstrate how it can be applied in the context of RNA-seq data analysis for the identification of gene modules. Let n_{vj} denote the number of sequencing reads mapped to gene $v \in \{1, \dots, V\}$ in sequencing sample $j \in \{1, \dots, J\}$, and let the $V \times 1$ vector \mathbf{n}_j contain all the gene counts for sample j . The NB distribution is a popular choice to model RNA-seq count data, allowing one to account for over-dispersion due to technical and biological variations (Anders and Huber, 2010; Dadaneh et al., 2017; Robinson et al., 2010). Under the NBFA model (Zhou, 2017), the sample counts are factorized as

$$\mathbf{n}_j \sim \text{NB}(\Phi\theta_j, p_j), \quad (1)$$

where $n \sim \text{NB}(r, p)$ denotes the NB distribution with the probability mass function (PMF) $f_N(n) = \frac{\Gamma(n+r)}{n! \Gamma(r)} p^r (1-p)^n$, where $\Gamma(\cdot)$ is the gamma function and $n \in \{0, 1, 2, \dots\}$. $\Phi = (\phi_1, \dots, \phi_K) \in \mathbb{R}_+^{V \times K}$ represents the factor loading matrix, $\Theta = (\theta_1, \dots, \theta_J) \in \mathbb{R}_+^{K \times J}$ represents the factor score matrix and $\mathbb{R}_+ = \{x : x \geq 0\}$. For each latent factor, $\phi_k = (\phi_{1k}, \dots, \phi_{V_k})^T$ encodes the weights of the V genes associated with factor k and θ_j encodes the popularity of K factors in sample j . The NBFA can be augmented as

$$n_{vj} = \sum_{k=1}^K n_{vjk}, \quad n_{vjk} \sim \text{NB}(\phi_{vk}\theta_{kj}, p_j). \quad (2)$$

From biological perspectives, K factors can correspond to the underlying biological processes (BPs) or functional modules related to genotypic, phenotypic, or treatment condition changes. The corresponding sub-counts n_{vjk} can be viewed as the result of the contribution of underlying BP k to the expression of gene v in sample j . The probability parameter p_j , which only depends on the sample index, can be considered as a parameter reflecting the potential heterogeneity of counts, due to the variation of the sequencing depths across different samples.

More precisely, using Equation (2) and the formula for the mean of the NB distribution, the expected expression of gene v in sample j can be expressed as $\mathbb{E}[n_{vj}] = \left(\sum_{k=1}^K \phi_{vk}\theta_{kj} \right) \frac{p_j}{1-p_j}$. The term $\frac{p_j}{1-p_j}$ can be interpreted as the effect of the sequencing-depth heterogeneity of sample j on the corresponding gene expression in this sample. This approach removes the need for an *ad-hoc* normalization step, as the model accounts for the sequencing-depth heterogeneity of different samples automatically, similar to the mechanisms employed in Dadaneh et al. (2017). The remaining term in this expectation, $\sum_{k=1}^K \phi_{vk}\theta_{kj}$, can represent the true abundance of gene v in sample j .

Specifically, it comprises of contributions from all latent factors, where each contribution is encoded as the product of the gene association with latent factors as modules and the contribution of those modules to sample j .

NBFA proceeds by placing the Dirichlet and gamma prior distributions on ϕ_{vk} and θ_{kj} , respectively, and appropriate prior distributions on the other model parameters. A Gibbs sampling algorithm that exploits novel data augmentation techniques has been derived for inferring the model parameters (Zhou, 2017).

2.2 Covariate-dependent NBFA

In real-world RNA-seq experiments, it is often desirable to identify the functional modules corresponding to critical BPs specific to the behavior of interest by the design of experiments. Often, the presence of potential confounding factors also requires that the developed factor analysis method based on RNA-seq data can take them into account (when the corresponding conditions are given) to derive correct functional module results. The aforementioned NBFA model neglects such information about sequencing samples from designed experiments. In order to empower the NBFA model in tackling the setups with complex experiment design, we extend its framework to make it capable of incorporating the external covariate information (e.g. phenotypes, treatments and other confounding factors) into the factor analysis model to derive the new covariate-dependent NBFA (dNBFA) model.

The graphical representation of dNBFA is illustrated as a hierarchical model in Figure 1. In the first layer of dNBFA, similar to NBFA, the gene counts are modeled using the same NB distribution as in Equation (2). Then, in the next layer we place a gamma prior distribution on θ_{kj} as

$$\theta_{kj} \sim \text{Gamma}\left(r_k, e^{\beta_k^T \mathbf{x}_j}\right), \quad (3)$$

where \mathbf{x}_j is the $P \times 1$ vector of covariates for sample j , reflecting the corresponding experiment design. In this model, both numerical and categorical covariates can be used.

Employing the law of total expectation, and removing the sequencing depth effect by the related terms containing p_j , we have $\mathbb{E}[n_{vjk}] \propto \phi_{vk} r_k e^{\beta_k^T \mathbf{x}_j}$. This new layer of model, splits the effect of the latent factor k on sample j into two parts; r_k , which can be considered as representing the baseline expression of the factor k across all samples, and the exponential term $e^{\beta_k^T \mathbf{x}_j}$, which adjusts the effect of the latent factor on the sample according to its traits. We note that including an intercept in $\beta_k^T \mathbf{x}_j$ may weaken the identifiability of r_k , as in the expectation of the count n_{vjk} a product term $r_k e^{\beta_{k0}}$ depending only on latent factor k appears. Thus in all subsequent experiments a separate intercept term is not used when considering covariate effects. The parameters of the dNBFA model with their interpretations in the context of RNA-seq experiments are presented in Table 1.

We place independent zero-mean normal distributions on the components of the regression coefficient parameters as

$$\beta_k \sim \prod_{p=1}^P \text{N}\left(0, \alpha_p^{-1}\right), \quad (4)$$

where α_p is the precision parameter of the normal distribution. By assuming identical precisions for components of the regression coefficients across all latent factors, dNBFA burrows statistical strengths to infer these precision parameters.

Similar to NBFA, a Dirichlet prior distribution with the smoothing parameter η is imposed on the gene-module association parameters ϕ_{vk} :

$$(\phi_{1k}, \dots, \phi_{V_k}) \sim \text{Dir}(\eta, \dots, \eta). \quad (5)$$

The Dirichlet smoothing parameter η controls the sparsity of the inferred latent factors. Generally speaking, the smaller η is, the more sparse and specific the inferred factors are encouraged to be.

A challenge in NB factorization is how to determine the number of latent factors K . To address this issue, one can employ a reasonably large K and then according to the inference step for r_k [refer to Equation (12) below], the baseline expression inferred for non-important latent factors vanishes as the number of assigned gene sub-counts to it decreases.

We complete the model by placing conjugate priors on hyperparameters. Specifically, we exploit the gamma-Poisson conjugacy, beta-NB conjugacy with respect to the probability parameter and gamma-gamma conjugacy with respect to the scale parameter of the gamma distribution. The complete dNBFA model is presented below:

$$\begin{aligned} n_{vj} &= \sum_{k=1}^K n_{vjk}, \quad n_{vjk} \sim \text{NB}(\phi_{vk} \theta_{kj}, p_j), \\ \theta_{kj} &\sim \text{Gamma}\left(r_k, e^{\beta_k^T \mathbf{x}_j}\right), \quad r_k \sim \text{Gamma}(\gamma_0/K, 1/c_0), \\ (\phi_{1k}, \dots, \phi_{V_k}) &\sim \text{Dir}(\eta, \dots, \eta), \quad \beta_k \sim \prod_{p=1}^P \text{N}\left(0, \alpha_p^{-1}\right), \\ \gamma_0, c_0, \alpha, \eta &\sim \text{Gamma}(e_0, 1/f_0), \quad p_j \sim \text{Beta}(a_0, b_0). \end{aligned} \quad (6)$$

Throughout the experiments in this paper, we set the hyperparameters as $e_0 = f_0 = 0.01$ and $a_0 = b_0 = 1$. In the following section, we provide an efficient inference algorithm that adopts novel data augmentation techniques tailored to our dNBFA model.

2.3 Inference via Gibbs sampling

By utilizing a few data augmentation techniques (Polson *et al.*, 2013; Zhou *et al.*, 2012; Zhou and Carin, 2015), we derive an efficient Gibbs sampling algorithm for inferring the model parameters in Equation (6), as described below. Algorithm 1 summarizes all the steps in the Gibbs sampling algorithm.

Sample ϕ_{vk} and θ_{kj} . We start with the data augmentation technique developed for inferring the NB dispersion parameter (Zhou and Carin, 2015). More precisely, the NB random variable $n \sim \text{NB}(r, p)$ can be generated from a compound Poisson distribution as

$$n = \sum_{t=1}^{\ell} u_t, \quad u_t \sim \text{Log}(p), \quad \ell \sim \text{Pois}(-r \ln(1-p)),$$

where $u \sim \text{Log}(p)$ corresponds to the logarithmic random variable (Johnson *et al.*, 2005), with the PMF $f_U(u) = -\frac{p^u}{u \ln(1-p)}$, $u \in \{1, 2, \dots\}$. As shown in Zhou and Carin (2015), given n and r , the distribution of ℓ is a Chinese Restaurant Table (CRT) distribution, $(\ell|n, r) \sim \text{CRT}(n, r)$, which can be generated as $\ell = \sum_{t=1}^n b_t$, $b_t \sim \text{Bernoulli}\left(\frac{r}{r+t-1}\right)$.

Utilizing the above data augmentation technique, for each observed count n_{vj} , a latent count is sampled as

$$(\ell_{vj}|-) \sim \text{CRT}\left(n_{vj}, \sum_{k=1}^K \phi_{vk} \theta_{kj}\right). \quad (7)$$

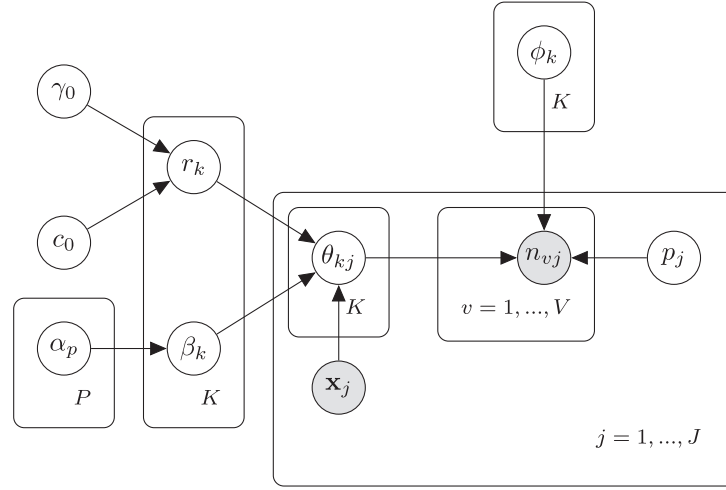


Fig. 1. Graphical representation of the dNBFA model

Table 1. Parameters of covariate-dNBFA and their interpretations in the context of RNA-seq data

Parameter	Constraint	Interpretation
r_k	$r_k > 0$	Module baseline expression
p_j	$0 < p_j < 1$	Sequencing depth
ϕ_{vk}	$\sum_{v=1}^V \phi_{vk} = 1, \phi_{vk} > 0$	Gene-module association
θ_{kj}	$\theta_{kj} > 0$	Popularity of factor k in sample j
β_{kp}	$\beta_{kp} \in \mathbb{R}$	Impact of covariate p on expression of factor k

Note: The inputs of dNBFA are gene counts n_{vj} and vector of covariates \mathbf{x}_j .

These counts are then further split into latent sub-counts [Proposition 3 of Zhou (2017)] using a multinomial distribution:

$$(\ell_{vj1}, \dots, \ell_{vjK} | -) \sim \text{Mult} \left(\ell_{vj}; \left(\frac{\phi_{v1}\theta_{1j}}{\sum_{k=1}^K \phi_{vk}\theta_{kj}}, \dots, \frac{\phi_{vK}\theta_{Kj}}{\sum_{k=1}^K \phi_{vk}\theta_{kj}} \right) \right). \quad (8)$$

These latent counts can be considered as being generated as $\ell_{vj} \sim \text{Pois}(q_j \phi_{vk} \theta_{kj})$, where $q_j := -\ln(1 - p_j)$. Hence, using gamma-Poisson conjugacy, ϕ_{vk} and θ_{kj} are updated as

$$\begin{aligned} (\phi_{1k}, \dots, \phi_{V,k} | -) &\sim \text{Dir}(\eta + \ell_{1,k}, \dots, \eta + \ell_{V,k}) \\ (\theta_{kj} | -) &\sim \text{Gamma} \left(r_k + \ell_{\cdot,kj}, \frac{1}{q_j + e^{-\beta_k^T \mathbf{x}_j}} \right), \end{aligned} \quad (9)$$

where $\ell_{v,\cdot} = \sum_{j=1}^J \ell_{vj}$ and $\ell_{\cdot,kj} = \sum_{v=1}^V \ell_{vj}$.

Sample r_k and γ_0 . Let us denote $\psi_{kj} := \beta_k^T \mathbf{x}_j + \ln q_j$. Starting with $\ell_{\cdot,jk} \sim \text{Pois}(q_j \theta_{kj})$, marginalizing out θ_{kj} leads to

$$\ell_{\cdot,jk} \sim \text{NB} \left(r_k, \frac{1}{1 + e^{-\psi_{kj}}} \right). \quad (10)$$

Employing the CRT augmentation technique as

$$(\tilde{\ell}_{jk} | -) \sim \text{CRT}(\ell_{\cdot,jk}, r_k), \quad (11)$$

the Gibbs sampling update for r_k can be written as

$$(r_k | -) \sim \text{Gamma} \left(\gamma_0 / K + \tilde{\ell}_{\cdot,k}, \frac{1}{c_0 + \sum_j \ln(1 + e^{\psi_{kj}})} \right). \quad (12)$$

Following a similar procedure for γ_0 , first we draw

$$(\tilde{\ell}_k | -) \sim \text{CRT}(\tilde{\ell}_{\cdot,k}, \gamma_0 / K), \quad (13)$$

and then we update the conditional posterior of γ_0 as

$$(\gamma_0 | -) \sim \text{Gamma} \left(e_0 + \sum_k \tilde{\ell}_k, \frac{1}{f_0 - \sum_k \ln(1 - \tilde{p}_k) / K} \right), \quad (14)$$

where $\tilde{p}_k := \frac{\sum_j \ln(1 + e^{\psi_{kj}})}{c_0 + \sum_j \ln(1 + e^{\psi_{kj}})}$.

Sample β_k . For the regression coefficients modeling potential covariate effects, the lack of conditional conjugacy precludes immediate closed-form inference. Therefore, we adopt another data augmentation technique, specifically designed for dNBFA, to infer the regression coefficients β_k , relying on the PG data augmentation of (Polson et al., 2013; Zhou et al., 2012). Denote ω_{kj} as a random variable drawn from the PG distribution as $\omega_{kj} \sim \text{PG}(\ell_{\cdot,jk} + r_k, 0)$.

Since $\mathbb{E}_{\omega_{kj}}[\exp(-\omega_{kj} \psi_{kj}^2 / 2)] = \cosh(\ell_{\cdot,jk} + r_k) (\psi_{kj}^2 / 2)$, the likelihood of ψ_{kj} in Equation (10) can be expressed as

$$\begin{aligned} \mathcal{L}(\psi_{kj}) &\propto \frac{(e^{\psi_{kj}})^{\ell_{\cdot,jk}}}{(1 + e^{\psi_{kj}})^{\ell_{\cdot,jk} + r_k}} \\ &\propto \exp \left(\frac{\ell_{\cdot,jk} - r_k}{2} \psi_{kj} \right) \mathbb{E}_{\omega_{kj}} \left[\exp \left(-\omega_{kj} \psi_{kj}^2 / 2 \right) \right]. \end{aligned} \quad (15)$$

Exploiting the exponential tilting of the PG distribution in Polson et al. (2013), we draw ω_{kj} as

$$(\omega_{kj} | -) \sim \text{PG}(\ell_{\cdot,jk} + r_k, \psi_{kj}). \quad (16)$$

Given the values of the auxiliary variables ω_{kj} for $j = 1, \dots, J$ and the prior in Equation (6), the conditional posterior of β_k can be updated as

$$(\beta_k | -) \sim \text{N}(\mu_k, \Sigma_k), \quad (17)$$

where $\Sigma_k = \left(\text{diag}(\alpha_1, \dots, \alpha_p) + \sum_j \omega_{kj} \mathbf{x}_j \mathbf{x}_j^T \right)^{-1}$ and $\mu_k = \Sigma_k \left[\sum_j \left(\frac{\ell_{\cdot,jk} - r_k}{2} - \omega_{kj} \ln(q_j) \right) \mathbf{x}_j \right]$.

Sample η . To derive the update steps for Dirichlet hyperparameters, we note that the likelihood for $\{\phi_k\}$ is

$$\mathcal{L}(\{\phi_k\}) \propto \prod_{k=1} \text{Mult}(\ell_{1,k}, \dots, \ell_{V,k}; \ell_{\cdot,k}, \phi_k). \quad (18)$$

Marginalizing out $\{\phi_k\}$ from Equation (18), the likelihood for η can be expressed as

$$\mathcal{L}(\eta) \propto \prod_{k=1} \text{DirMult}(\ell_{1,k}, \dots, \ell_{V,k}; \ell_{\cdot,k}, \eta, \dots, \eta), \quad (19)$$

where DirMult denotes the Dirichlet-Multinomial distribution (Zhou, 2017). Since the product of $\ell(\eta)$ and $\prod_k \text{Beta}(q_k; \ell_{\cdot,k}, \eta V)$ can be written as

$$\mathcal{L}(\eta) \prod_k \text{Beta}(q_k; \ell_{\cdot,k}, \eta V) \propto \prod_k \prod_v \text{NB}(\ell_{v,k}; \eta, q_k), \quad (20)$$

we can further apply the data augmentation technique for the NB distribution of Zhou and Carin (2015) to derive closed-form update equations for η as

$$\begin{aligned} (q_k | -) &\sim \text{Beta}(\ell_{\cdot,k}, \eta V), \quad (u_{vk} | -) \sim \text{CRT}(\ell_{v,k}, \eta) \\ (\eta | -) &\sim \text{Gamma}\left(e_0 + \sum_{v,k} u_{vk}, \frac{1}{f_0 - V \sum_k \ln(1 - q_k)}\right). \end{aligned} \quad (21)$$

Sample α_p, p_j and c_0 . Using appropriate conditional conjugacies, we can sample the remaining parameters as

$$\begin{aligned} (\alpha_p | -) &\sim \text{Gamma}\left(e_0 + K/2, \frac{1}{f_0 + \sum_k \beta_{kp}^2/2}\right) \\ (p_j | -) &\sim \text{Beta}\left(a_0 + \sum_{v=1}^V n_{vj}, b_0 + \sum_{k=1}^K \theta_{kj}\right) \\ (c_0 | -) &\sim \text{Gamma}\left(e_0 + \gamma_0, \frac{1}{f_0 + \sum_k r_k}\right). \end{aligned} \quad (22)$$

The Gibbs sampling steps in Equations (7) to (22) are summarized in Algorithm 1.

Algorithm 1 dNBFA model inference

Inputs: RNA-seq counts, design matrix of covariate effects, N

Outputs: gene module membership matrix

Initialize model parameters

Do Gibbs sampling:

for $iter = 1$ to N do

 Sample ℓ_{vjk} using the CRT distribution [Equation (7)]

 Update ϕ_{vk} and θ_{kj} using the gamma-Poisson conjugacy [Equation (9)]

 Sample $\tilde{\ell}_{jk}$ using the CRT distribution [Equation (11)]

 Update r_k and γ_0 using the gamma-Poisson conjugacy [Equation (12),(14)]

 Sample auxiliary variables ω_{kj} , using the PG distribution [Equation (16)]

 Update regression coefficients [Equation (17)]

 Update η using auxiliary beta distributed random variables [Equation (21)]

 Update α_p, p_j and c_0 [Equation (22)]

end for

3 Results

We evaluate our dNBFA for covariate-dependent factor analysis based on two sets of real-world RNA-seq data studying complex diseases, and compare its performance with those of WGCNA (Langfelder and Horvath, 2008) and DiffCoEx (Tesson *et al.*, 2010), two commonly adopted two-stage co-expression network based methods.

The first set of RNA-seq data was extracted from TCGA [Cancer Genome Atlas (TCGA) Research Network and others, 2008], including three datasets on breast invasive carcinoma (BRCA), lung squamous cell carcinoma (LUSC) and kidney renal clear cell carcinoma (KIRC). These data were retrieved using the TCGA2STAT R package (Wan *et al.*, 2015). Using TCGA data we expect to illustrate the higher differential expression significance of gene modules identified by dNBFA with respect to the disease factor compared to the results from WGCNA and DiffCoEx.

The second experiment was performed on a RNA-seq dataset of the Autism study in Gupta *et al.* (2014), where samples were obtained from three brain regions: the cerebral cortex Brodmann area (BA) 19, anterior prefrontal cortex (BA10) and a part of the frontal cortex (BA44). For this dataset, we demonstrate how incorporating covariate information may enhance the chance of achieving meaningful biological discoveries.

For both TCGA and Autism experiments, dNBFA was run using 3000 MCMC iterations, where after the first 1000 burn-in iterations, the posterior samples with the highest likelihood were collected as the point estimates of model parameters. The total number of latent factors for both TCGA and Autism were initially set as $K = 250$, and after the parameter inference, only the top 100 factors with non-negligible baseline expressions were kept for further analyses. In addition, to determine module membership, for each latent factor k , only the top 20 genes with highest ϕ_{vk} were considered as members of module k . It should be noted that when an evaluation metric that can take advantage of the whole association matrix Φ exists, this *ad-hoc* step of using a cut-off for the gene-association parameter can be avoided.

For WGCNA, the adjacency matrix was built by first computing the pairwise Pearson correlation coefficients between gene expression profiles and then applying the soft threshold $\beta = 6, 9$ for TCGA and Autism data, respectively. The gene modules were identified by applying a hierarchical clustering algorithm to the derived topological overlap dissimilarity matrix (Ravasz *et al.*, 2002). A similar procedure was followed for DiffCoEx, except that the topological overlap matrix was built upon the matrix of adjacency difference (Tesson *et al.*, 2010). Our experiments show that the discovered modules by WGCNA and DiffCoEx comprise of large lists of genes, where no further modeling capability is provided to narrow down the gene sets for more consequent exploratory analysis.

3.1 TCGA data

For all TCGA datasets, we have filtered out the genes whose total read counts across all samples are less than 50, resulting in roughly 20000 genes in each dataset. The total numbers of samples for BRCA, LUSC and KIRC datasets are, respectively, 40, 34 and 40, where in each case the number of primary tumor and normal samples are equal.

Based on the resulting RNA-seq count data, dNBFA, WGCNA and DiffCoEx have been applied to derive functional gene modules

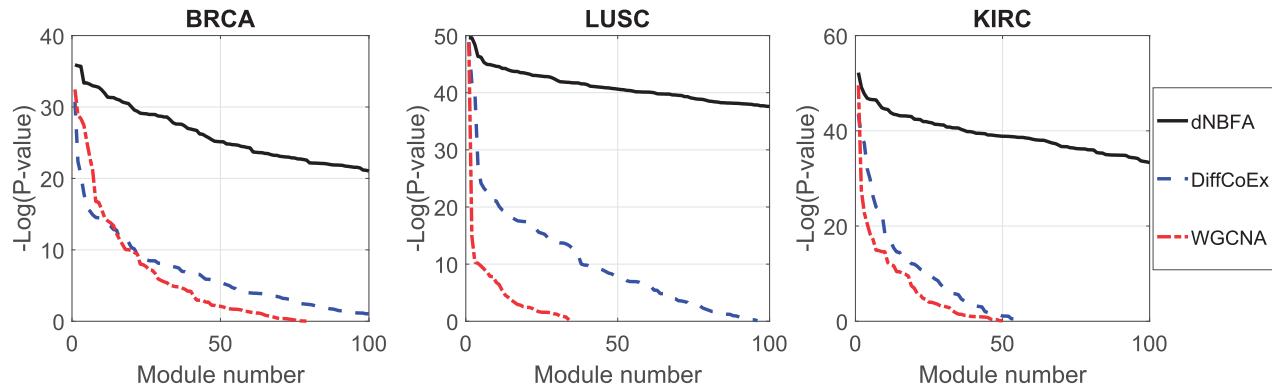


Fig. 2. Significance of differential expression for eigengenes associated with gene modules identified by dNBFA, WGCNA and DiffCoEx applied to three TCGA datasets. The panels show the sorted negative logarithm of P -values of the derived modules. P -values are calculated using the student's t -test on association between module eigengene expression and the samples' condition factor (cancerous versus normal)

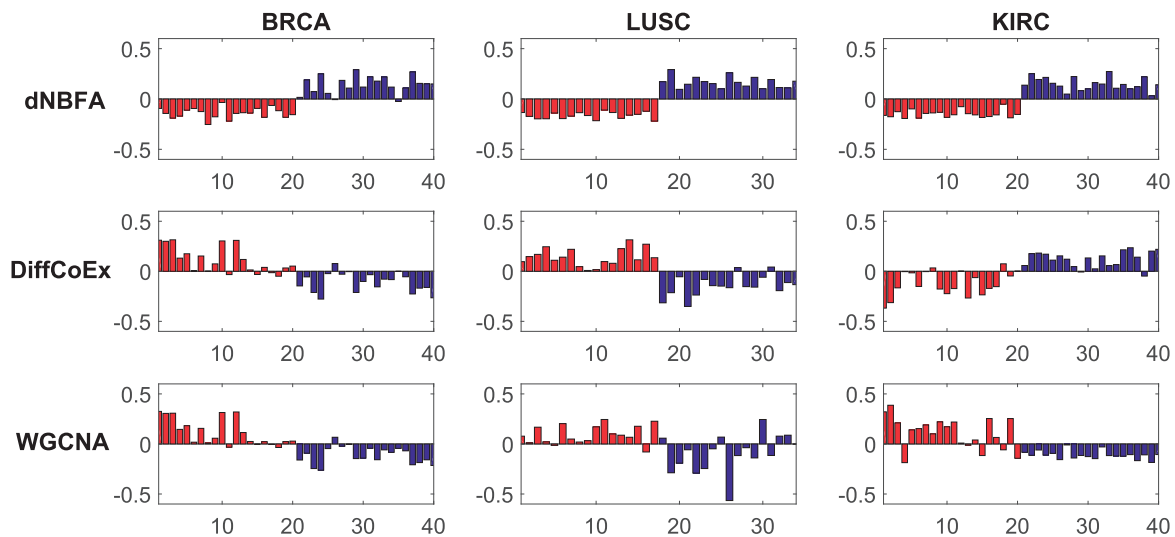


Fig. 3. Per-sample eigengene expression of modules with the 10th lowest P -values discovered by dNBFA, WGCNA and DiffCoEx, across cancerous and normal samples for the three TCGA datasets. In each figure the y-axis is the eigengene expression, and the x-axis is the sample number. Red and blue bars correspond to the normal and cancer groups, respectively. Figures in top, middle and bottom rows are the results of dNBFA, DiffCoEx and WGCNA, respectively. Figures in left, middle and right columns correspond to BRCA, LUSC and KIRC datasets, respectively

using the aforementioned settings. To assess the significance of differential expression of identified modules with respect to the disease status of samples, we follow the framework of Langfelder and Horvath (2008). More precisely, for each detected module, first the *eigengene* (Langfelder and Horvath, 2008) is computed via the first principal component of the expression matrix of the corresponding derived module. The module eigengene is used to summarize and represent the expression profiles of the module genes (Horvath and Dong, 2008). Then, the association of the eigengene expression with the disease status is evaluated and finally the significance of the association is assessed based on the student's t -test.

We calculate the P -values for gene modules identified by dNBFA, WGCNA and DiffCoEx, applied to the three TCGA datasets. The sorted P -values [based on $-\log(P\text{-value})$] are illustrated in Figure 2. The eigengenes of the modules detected by dNBFA are remarkably more differentially expressed than those detected by WGCNA and DiffCoEx in all three TCGA datasets. To further investigate the results, we present the per-sample eigengenes of the module ranked 10th for differential expression, which was identified by dNBFA, WGCNA and DiffCoEx for the three TCGA datasets in Figure 3. The per-sample eigengenes of

dNBFA modules are more consistently differentially expressed with respect to the disease status covariate for all three TCGA datasets, while per-sample eigengenes of WGCNA and DiffCoEx demonstrate higher variations within each group of samples with the same disease status. To ensure that the gene modules detected by dNBFA are not redundant, we also have examined the modules for significant overlap. Except a minor overlap between two modules, the rest of the modules identified by dNBFA are completely disjoint. These results suggest that dNBFA can be a powerful untargeted module identification tool, without predefined gene lists, for genomic experiments that study coordinated gene expression pattern changes across multiple groups.

To further verify the advantages of dNBFA that it avoids overfitting when the initial number of modules K is set high, we present in Figure 4 the learned r_k 's, representing the baseline expression associated with the derived modules, for three TCGA datasets. Only the top 40 r_k 's are included in this figure. For all datasets, only a fraction of modules have significantly large baseline expression; and thus in practice, a threshold can be used to extract the modules that contribute significantly to coordinated gene expression changes specific to the experiment design factors of interest.

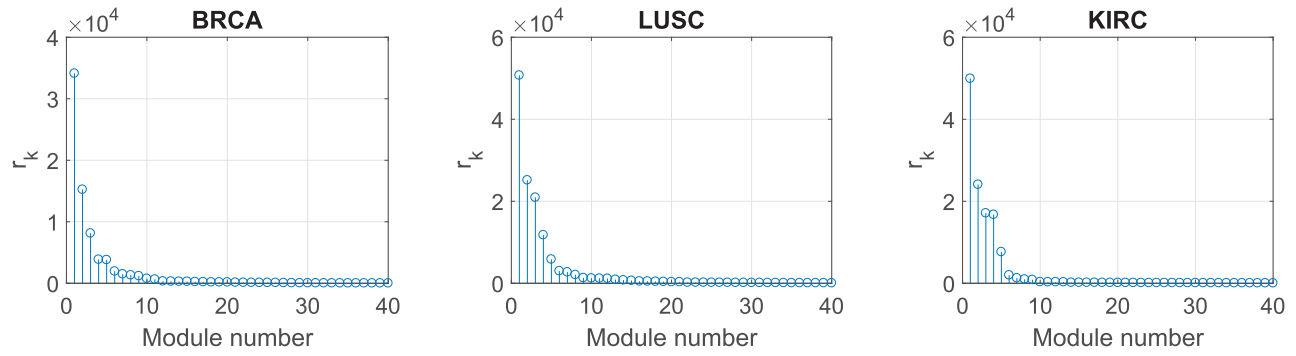


Fig. 4. Inferred baseline expression r_k for modules detected by dNBFA in the three TCGA datasets. Only the top 40 r_k s are included in this figure

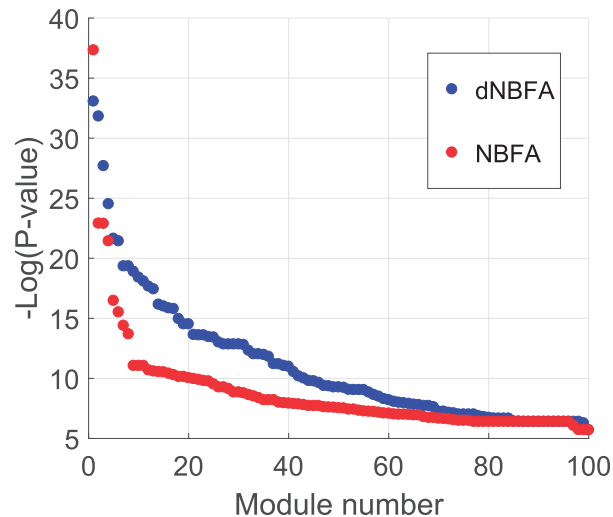


Fig. 5. Negative logarithm of P -values for GO term enrichment analysis of modules detected by dNBFA and NBFA, applied to Autism RNA-seq data. For dNBFA, site of sample collection, age, sex and brain region are used as covariate information, while no such information is incorporated for NBFA

For the analysis of the real-world TCGA dataset on a single cluster node with Intel Xeon 2.5 GHz E5-2670 v2 processor, on average it took around 8 h for both the dNBFA and NBFA methods with 3000 MCMC iterations and about 1 h for both WGCNA and DiffCoEx.

3.2 Autism data

Autism is a neuro-developmental disorder, in which the affected individuals are characterized by impairments in social and communicative developments (Gupta *et al.*, 2014). To apply dNBFA to the RNA-seq dataset of the Autism study in Gupta *et al.* (2014), we first discard the samples with low sequencing depths, resulting in a dataset with 36 samples from the control group and 23 Autism samples. The following analyses are performed using a subset of 12 010 genes that have a count of at least three per sample across 90% of the samples. In this experiment, site of sample collection, age, sex and brain region are available as the covariate information in factor analysis. To examine how this information can alter the NB factor analysis results, in the first set of experiments we use the covariates to apply dNBFA; and in the second set of experiments, we neglect all covariate information and run the naive NBFA on the dataset.

We perform gene set enrichment analysis on the discovered modules by applying dNBFA and NBFA, respectively, to the Autism

Table 2. Top enriched GO terms identified by dNBFA algorithm applied to Autism RNA-seq data

GO-ID	Aspect	Term	P -value
GO: 0060337	BP	Type I interferon signaling pathway	4.377782e-15
GO: 0043209	CC	Myelin sheath	1.522628e-14
GO: 0002460	BP	^a See blow	9.487407e-13
GO: 0061024	BP	Membrane organization	2.250911e-11
GO: 0044456	CC	Synapse part	4.010908e-10
GO: 0005575	CC	Cellular component	4.950009e-10
GO: 0033693	BP	Neurofilament bundle assembly	3.982179e-09
GO: 0031720	MF	Haptoglobin binding	3.982179e-09
GO: 0000982	MF	^b See blow	6.267732e-09
GO: 0001504	BP	Neurotransmitter uptake	1.015843e-08

^aAdaptive immune response based on somatic recombination of immune receptors built from immunoglobulin superfamily domains.

^bTranscription factor activity, RNA polymerase II core promoter proximal region sequence-specific binding.

data, covering molecular function (MF), cellular component (CC) and BP ontology domains. We calculate the significance of GO terms using Fisher's exact test and depict the sorted negative logarithm of P -values for both dNBFA and NBFA in Figure 5. The modules detected by dNBFA have, in general, lower P -values than those identified by NBFA without covariates, suggesting that incorporating covariate information may increase the chance of discovering biologically meaningful modules.

To investigate the gene ontology results more thoroughly, the top 10 GO terms with the lowest P -values are presented in Tables 2 and 3 for dNBFA and NBFA methods, respectively. In these tables, each row is the most significant GO term corresponding to one module identified by dNBFA or NBFA. The top modules discovered by dNBFA provide more explicit connections to neural system. Especially, the top module identified by dNBFA, which was not detected by NBFA, is associated with GO term 'type I interferon signaling pathway', where type I Interferon responses in the brain are classically attributed to viral infections (Delhaye *et al.*, 2006), which in turn are connected to Autism (Patterson, 2011). Another important module detected only by dNBFA, the third module in Table 2, is related to adaptive immune response which is closely correlated to the development of Autism spectrum disorders (Ashwood *et al.*, 2006; Heuer *et al.*, 2008). More precisely, this module includes the human leukocyte antigen (HLA) genes that play an instrumental role in many innate and adaptive immune responses

Table 3. Top enriched GO terms identified by NBFA algorithm applied to Autism RNA-seq data

GO-ID	Aspect	Term	P-value
GO: 0022625	CC	Cytosolic large ribosomal subunit	6.175728e-17
GO: 0097458	CC	Neuron part	1.134089e-10
GO: 0006735	BP	NADH regeneration	1.153145e-10
GO: 0005575	CC	Cellular component	4.950009e-10
GO: 0051050	BP	Positive regulation of transport	7.042487e-08
GO: 0007399	BP	Nervous system development	1.841155e-07
GO: 0065010	CC	Extracellular membrane-bounded organelle	5.614232e-07
GO: 0017111	MF	Nucleoside-triphosphatase activity	1.144582e-06
GO: 0048630	BP	Skeletal muscle tissue growth	1.586369e-05
GO: 0071208	MF	Histone pre-mRNA DCP binding	1.586369e-05

(Torres *et al.*, 2012). Many reports have provided the evidence on associations between Autism and HLA genes/haplotypes, suggesting an underlying dysregulation of the immune system mediated by HLA genes (Torres *et al.*, 2002; Torres *et al.*, 2012; Warren *et al.*, 1996). A third important module identified only by dNBFA is associated with GO term ‘neuron differentiation’ (GO: 0030182, P -value = 1.4×10^{-08}). Specifically, this module includes calmodulin 1 (CALM1) gene. Significant defects in CALM1 interaction modules, which regulate voltage-independent calcium-activated action potentials at the neuronal synapse, are reported in autistic patients (Hadley *et al.*, 2014).

Other GO terms directly related to the nervous system associated with the top modules discovered by dNBFA include ‘Myelin sheath’, ‘synapse part’, ‘neurofilament bundle assembly’ and ‘neurotransmitter uptake’. Specifically, the decreased thickness of myelin in the orbitofrontal cortex region is closely related to Autism disorders (Zikopoulos and Barbas, 2010). In addition, the module detected by dNBFA corresponding to GO term ‘synapse part’ has the highest association with the gene SNAP-25, whose reduced expression level is responsible for the cognitive deficits in children affected by Autism spectrum disorders (Braidă *et al.*, 2015).

Examining the detected modules by both dNBFA and NBFA, we observe that multiple GO terms relevant to Autism, such as ‘myelin sheath’, ‘NADH regeneration’ and ‘nervous system development’, are revealed by both algorithms. NADH is mainly involved in catabolic reactions (energy metabolism and mitochondrial function), whose decreased level has been reported in some children with Autism (Adams *et al.*, 2011). On the other hand, defects in Autism appear closely tied to late developmental steps of nervous system that depend on synaptic activity and activity-dependent transcriptional changes (Walsh *et al.*, 2008). Hence the relevance of the discovered GO terms by both dNBFA and NBFA to Autism is confirmed.

Finally, by examining the trace plots of model parameters, such as c_0 and r_B , we find that the Markov chains for the dNBFA method converge fast and mix well, supporting the practice of performing downstream analysis with 3000 MCMC iterations.

In summary, both NBFA and dNBFA methods emerge as useful module identification tools in RNA-seq data analysis, as in comparison to other available methods for gene module detection, they require minimum user adjustments. Specifically, the experimental results on the Autism dataset show that the incorporation of

covariate information by dNBFA may lead to the discovery of more significant Autism-relevant modules, which otherwise would be missed by NBFA.

4 Conclusions

We propose a novel Bayesian covariate-dependent negative binomial factor analysis (dNBFA) method for analyzing RNA-seq count data. Our experimental results on real-world RNA-seq data demonstrate that dNBFA is capable of handling complex experiments involving multiple factors. What’s more, dNBFA does not require any *ad-hoc* data normalization, data preprocessing, or co-expression network construction steps. By taking advantage of novel data augmentation techniques, dNBFA possesses efficient closed-form Gibbs sampling update equations. Experimental results on multiple RNA-seq data studying complex diseases, both cancer and Autism, demonstrate that our dNBFA can be directly applied to RNA-seq data to derive meaningful functional modules and it has potential advantages over existing two-stage co-expression network based methods.

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