NOVEL COMPUTATIONAL TOOLS AND DATABASES

ZERO-INFLATED NEGATIVE BINOMIAL REGRESSION FOR DIFFERENTIAL ABUNDANCE TESTING IN MICROBIOME STUDIES

None declared.

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Abstract

Motivation: The human microbiome plays an important role in human health and disease. The composition of the human microbiome is influenced by multiple factors and understanding these factors is critical to elucidate the role of the microbiome in health and disease and for development of new diagnostics or therapeutic targets based on the microbiome. 16S ribosomal RNA (rRNA) gene targeted amplicon sequencing is a commonly used approach to determine the taxonomic composition of the bacterial community. Operational taxonomic units (OTUs) are clustered based on generated sequence reads and used to determine whether and how the abundance of microbiome is correlated with some characteristics of the samples, such as health/disease status, smoking status, or dietary habit. However, OTU count data is not only overdispersed but also contains an excess number of zero counts due to undersampling. Efficient analytical tools are therefore needed for downstream statistical analysis which can simultaneously account for overdispersion and sparsity in microbiome data.

Results: In this paper, we propose a Zero-inflated Negative Binomial (ZINB) regression for identifying differentially abundant taxa between two or more populations. The proposed method utilizes an Expectation Maximization (EM) algorithm, by incorporating a two-part mixture model consisting of (i) a negative binomial model to account for overdispersion and (ii) a logistic regression model to account for excessive zero counts. Extensive simulation studies are conducted which indicate that ZINB demonstrates better performance as compared to several state-of-the-art approaches, as measured by the area under the curve (AUC). Application to two real datasets indicate that the proposed method is capable of detecting biologically meaningful taxa, consistent with previous studies.

Availability: The software implementation of ZINB is available at: http://www.ssg.uab.edu/bhglm/.

Supplementary information: Supplementary data are available at Journal of Bioinformatics and Genomics online.

Keywords: Differential Abundance Testing, EM Algorithm, Human Microbiome, Metagenomics, OTU, Zero-inflated Negative Binomial.

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1. Introduction

The advent of next-generation sequencing (NGS) technology enables the generation of large volume of metagenomic sequencing data (Gilbert, Meyer, & Bailey, 2011). This opens a new era of genomics study to explore microbial communities sampled directly from environments without isolation and cultivation (Cho & Blaser, 2012; Hugenholtz, 2002; Wooley & Ye, 2009). One of the environment is human or mammalian body which harbors a dense microbial population across different body sites, containing taxa across the tree of life including bacteria, viruses, micro-eukaryotes, and archaea (Dethlefsen, McFall-Ngai, & Relman, 2007; Whitman, Coleman, & Wiebe, 1998). The combination of microbiota, their genomes (metagenome), and the host environment forms human or mammalian microbiome (Cho & Blaser, 2012). An important research interest in human microbiome study is to assess whether and how two or more microbiome communities differ from each other. Many factors can influence the human microbiome composition (Turnbaugh et al., 2007). These factors include the host genotype (Spor,
Koren, & Ley, 2011), host physiological status such as aging (Biagi et al., 2010), host pathophysiological status (Turnbaugh et al., 2009), host lifestyle such as dietary habit (De Filippo et al., 2010; Wu et al., 2011), and host environment (Domínguez-Bello et al., 2010). Different studies have investigated the association between the human microbiome and human diseases such as obesity (Turnbaugh et al., 2006), diabetes (Samuel & Gordon, 2006), inflammatory bowel disease (IBD) (Frank et al., 2007), and cancers (Holes, Li, Athanasiou, Ashrafian, & Nicholson, 2011). The findings from these studies demonstrated that the human microbiome has extraordinary potential implications in new therapeutic targets or biomarkers for disease prevention and early diagnosis (Collison et al., 2012; Knights, Parfrey, Zaneveld, Lozupone, & Knight, 2011; Segata et al., 2011; Virgin & Todd, 2011).

16S ribosomal RNA (rRNA) gene targeted amplicon sequencing is a commonly used approach to determine the taxonomic composition and species diversity of the bacterial community (Matsen, Kodner, & Armbrust, 2010). Hypervariable regions within the gene are amplified and sequenced, and sequence reads are clustered into operational taxonomic units (OTUs) based on sequence similarity (Ghods, Liu, & Pop, 2011). Representative sequences from each cluster are then classified taxonomically by alignment against a database of previously characterized 16S ribosomal DNA (rDNA) reference sequences. The resulting OTU counts are then used to determine whether and how the abundance of microbiome is correlated with some characteristics of the samples, such as health/disease status, smoking status, or dietary habit.

However, the OTU/taxa data are high-dimensional with added complexity leading to several statistical challenges. The first challenge of microbiome data is how to properly account for variability induced by differences in sequencing depth across samples. This is due to our inability to accurately specify the exact number of sequences to be measured on a sample using currently available technology. Although the total number of sequences for a given sample is not associated with any biological feature of the sample, it affects the OTU counts, and hence, should be accounted for in downstream bioinformatic and statistical analysis. A common approach to account for this variation in the total number of sequences is the total-sum scaling (TSS) normalization, i.e. conversion of the sequence counts to relative abundance (i.e., taxon counts/total counts) within a particular sample (Wagner, Robertson, & Harris, 2011). However, using total counts as the normalization/scaling factor may be problematic and may lead to biases in differential abundance estimates (Knights et al., 2011; Kostic et al., 2012; Paulson, Stine, Bravo, & Pop, 2013). To adjust for differential sequencing depths, different approaches such as cumulative sum scaling (CSS) (Paulson et al., 2013), trimmed mean of M-values (TMM) (Robinson & Oshlack, 2010), and relative log expression (RLE) (Anders & Huber, 2010), have been proposed in the literature. Furthermore, due to the association between detected number of features (OTUs) and the depth of coverage in different samples, only a few OTUs are shared in various samples, whereas the rest are only present in a small proportion of samples, resulting in excess of zero counts in the OTUs count matrix (Paulson et al., 2013; Peng, Li, & Liu, 2015).

Second, OTU counts are over-dispersed, meaning the variance of the counts varies with the value of the mean. For such overdispersed count data, standard methods such as Poisson regression can result in high false positives (Peng et al., 2015; White, Nagarajan, & Pop, 2009). The overdispersed data have been widely studied in differential expression analysis in microarray and high-throughput sequencing (serial analysis of gene expression (SAGE) (Velculescu, Zhang, Vogelstein, & Kinzler, 1995) and RNA-seq) (McMurdie & Holmes, 2014). Differential abundance analysis in microbiome is a direct analogy to differential expression analysis. Many analytical tools (edgeR and DESeq) developed for differential expression analysis of RNA-Seq data can be adapted to differential abundance analysis of OTUs (Anders & Huber, 2010; Robinson, McCarthy, & Smyth, 2010). For example, White et al. (2009) extended the methods in differential expression analysis to microbiome studies by converting the raw abundances to proportions, which represent the relative contribution of each feature for each individual. This method was implemented in the software Metastats. However, microbiome data have the distinct characteristic of zero-inflation, which is notably less severe in RNA-seq data. To this end, various zero-inflated models have been proposed to correct for sparse counts in microbiome measurements. Paulson et al. (2013) proposed a zero-inflated Gaussian mixture model for modeling the CSS-normalized log-transformed count data using an Expectation Maximization (EM) algorithm, implemented in the Bioconductor package metagenomeSeq. Sohn et al. (2015) implemented a ratio approach for identifying differential abundance (RAIDA) by utilizing the ratio between features in a modified zero-inflated lognormal model. Peng et al. (2015) proposed a zero-inflated beta regression (ZIBSeq) approach for modeling the TSS-normalized relative abundance data. All these zero-inflated methods are designed to analyze microbiome data after a suitable normalization and/or transformation. However, differential abundance estimates based on these methods may not be interpretable on the original scale, leading to challenges in future prediction tasks and replication studies.

To address the above limitations, we propose a Zero-inflated Negative Binomial (ZINB) mixture model, which directly models the raw OTU counts. We implemented our model in R package BHGLM by utilizing an EM algorithm incorporating a two-part mixture model consisting of (i) negative binomial model to account for over-dispersion and (ii) a logistic regression model to account for zero-inflation. In our simulations, we show that ZINB outperforms DESeq, edgeR, and metagenomeSeq in various sparse scenarios in terms of Area under the Curve (AUC) estimates. Application to two real microbiome data sets reveal biologically significant taxa, which are consistent with previous studies. The software implementation of ZINB is freely available at: http://www.ssg.uab.edu/bhglm/.

2. Methods

Assume that there are \( n \) samples and \( m \) features. In 16S rRNA microbiome data, the features refer to OTUs, species, or genus, etc. Let \( C_{ij} \) be the observed count for \( i \)-th sample and \( j \)-th feature, and \( T_i \) be the total read for \( i \)-th sample (also referred to as depths of coverage or library size) or a linear scaling factor that accounts for its library size. Let \( X_i \) be a factor indicating host health/disease status, physiological status such as aging, or lifestyle such as dietary habit, etc. In differential abundance analysis, the goal is to determine differentially abundant microbial features between groups defined by the
host factor. We directly model the raw count data with the zero-inflated negative binomial model. To adjust for differential sequencing depth, the natural logarithm of total counts (per sample) are included as offset in the negative binomial model. This allows us to handle the raw data matrix without normalization, leading to interpretable differential abundance estimates. In the following subsection, we describe our model and algorithm.

2.1. Zero-inflated Negative Binomial (ZINB) Model

ZINB models assume that observed zero counts may come from either a degenerate distribution having the point mass at zero or a negative binomial distribution and observed non-zero counts come exclusively from the negative binomial distribution. Therefore, the count response for feature $j$, i.e., $y_i$, follows the mixture distribution:

$$y_i \sim \begin{cases} 0 & \text{with probability } p_i, \\ NB(y_i | \mu_i, \theta) & \text{with probability } 1 - p_i, \end{cases}$$

(1)

where $\mu_i$ and $\theta$ are the mean and the shape parameter of the negative binomial distribution $NB(y_i | \mu_i, \theta)$, respectively, and $p_i$ are the mixture probability parameters. The mean parameters $\mu_i$ are related to the variables $X_i$ (including the intercept) via the link function logit:

$$\logit(p_i) = \log \left( \frac{p_i}{1 - p_i} \right) = Z_i \gamma$$

(3)

where $Z_i$ includes variables that are potentially associated with the zero state. We can include only intercept in $Z_i$ (as in the simulation study and real data analysis). However, our method can be applied to general cases. The log-likelihood function for the parameters $(\beta, \theta)$ and $\gamma$ is given by

$$L(\beta, \theta, \gamma; y) = \sum_{y_i = 0}^{n} \log \left[ p_i + (1 - p_i)NB(y_i = 0 | \mu_i, \theta) \right]$$

$$+ \sum_{y_i > 0}^{n} \log \left[ (1 - p_i)NB(y_i = k | \mu_i, \theta) \right]$$

(4)

The first term in the log-likelihood includes both $(\beta, \theta)$ and $\gamma$ and thus constitutes the maximization of the log-likelihood. In principle, this likelihood can be optimized directly by the Newton-Raphson algorithm (as implemented by the `zeroinfl` function in the R package `pscl`). However, the Newton-Raphson algorithm for fitting this model is known to be unstable and have severe non-convergence issues in small samples (Mallick & Tiwari, 2016). We thus propose an EM algorithm, which is stable and efficient, leading to accurate estimation and inference.

2.2. EM algorithm for fitting the ZINB model

The EM algorithm introduces a vector of latent indicator variables $\xi = (\xi_1, \cdots, \xi_N)$ to distinguish the zero state from the negative binomial state, where $\xi_i = 1$ when $y_i$ is from the zero state, and $\xi_i = 0$ when $y_i$ is from the negative binomial state.

With the indicator variables, the ZINB model can be expressed as

$$y_i \sim \begin{cases} 0 & \text{if } \xi_i = 1, \\ NB(y_i | \mu, \theta) & \text{if } \xi_i = 0, \end{cases}$$

and

$$p(\xi_i = 1) = \frac{\exp(Z_i \gamma)}{1 + \exp(Z_i \gamma)}$$

(5)

The log-likelihood with the complete data $(y, \xi)$ is given by

$$L(\beta, \theta, \gamma; y, \xi) = \sum_{i=1}^{n} \log \left\{ p_i^\xi (1 - p_i)^{1 - \xi_i} \right\}$$

$$+ \sum_{i=1}^{n} (1 - \xi_i) \log \left( NB(y_i | \mu, \theta) \right)$$

(6)

The EM algorithm replaces the indicator variables $(\xi_i)$ by their conditional expectations $(\hat{\xi}_i)$ (E-step), and then updates the coefficients $\beta$ and $\gamma$ and the shape parameter $\theta$ by maximizing $L(\beta, \theta, \gamma; y, \hat{\xi})$ (M-step). When the EM algorithm converges, we obtain the estimate $(\hat{\beta}, \hat{\theta}, \hat{\gamma})$ that maximizes the log-likelihood $L(\beta, \theta, \gamma; y)$. In the E-step, we calculate the conditional expectations of the indicator variables $(\hat{\xi}_i)$. The conditional expectation of $\xi_i$ can be easily calculated as:

$$\hat{\xi}_i = p(\xi_i = 1 | \beta, \theta, \gamma, y) = \frac{p(y_i \mid \beta, \theta, \hat{\xi}_i = 1) p(\hat{\xi}_i = 1 | \gamma)}{p(y_i \mid \beta, \theta, \hat{\xi}_i = 0) p(\hat{\xi}_i = 0 | \gamma) + p(y_i \mid \beta, \theta, \hat{\xi}_i = 1) p(\hat{\xi}_i = 1 | \gamma)}$$

(7)

If $y_i > 0$, we have $p(y_i \mid \beta, \theta, \hat{\xi}_i = 1) = 0$, and thus $p(\hat{\xi}_i = 0 | \beta, \theta, \gamma, y) = 1$. We then update $p(\hat{\xi}_i = 1 | \beta, \theta, \gamma, y)$ by

$$p(\hat{\xi}_i = 1 | \beta, \theta, \gamma, y) = \left[ \exp(-s_i \gamma)NB(y_i = 0 | \mu, \theta) + 1 \right]^{-1}.$$
U_i = \hat{\beta}_i / \sqrt{\text{Var}(\hat{\beta}_i)}, which approximately follow the standard normal distribution.

2.3. IWLS algorithm for fitting the negative binomial model

The above EM algorithm needs fitting a weighted negative binomial model. We extend the IWLS (Iterative Weighted Least Square) algorithm for generalized linear models to fit the weighted negative binomial model. Our algorithm is based on the well-known fact that given the shape parameter $\theta$ the negative binomial density has the exponential form:

$$ NB(y_i | \mu_i, \theta) = \frac{\Gamma(y_i + \theta)}{\Gamma(\theta)\Gamma(y_i)!} \left( \frac{\mu_i}{\mu_i + \theta} \right)^{\theta} \left( \frac{\theta}{\mu_i + \theta} \right)^{y_i} $$

(8)

$$ = \exp \left\{ \frac{y_i \theta - b(\theta)}{\phi} + c(y_i, \phi) \right\} $$

where $\phi = \log \frac{\mu_i}{\mu_i + \theta}$, $\phi = 1$, $c(y_i, \phi) = \log \left( \frac{\Gamma(y_i + \theta)\theta^\theta}{\Gamma(\theta)\Gamma(y_i)!} \right)$, and $b(\theta) = -\theta \log \left( 1 - e^{-\theta} \right) = -\theta \log \left( 1 - e^{\phi} \right)$. Therefore, the negative binomial model is a special case of generalized linear models (GLMs) for any fixed $\theta$.

The IWLS algorithm for fitting the negative binomial model proceeds as follows. Given the shape parameter $\theta$, the estimate of $\beta$ can be obtained by fitting the weighted normal linear model:

$$ z_i = \log(T_i) + X_i \beta + e_i $$

(9)

where $e_i \sim N(0, (1 - \hat{\gamma})^{-1} \lambda^{-1})$. The pseudo-response $z_i$ and pseudo-weights $w_i$ are calculated by

$$ z_i = \widehat{\eta}_i - L'(y_i | \widehat{\eta}_i, \widehat{\theta}) \quad \text{and} \quad w_i = -L''(y_i | \widehat{\eta}_i, \widehat{\theta}) $$

(10)

where $\widehat{\eta}_i = \log(T_i) + X_i \hat{\theta}$, $L'(y_i | \hat{\eta}_i, \hat{\theta}) = \log NB(y_i | \hat{\mu}_i, \hat{\theta})$, $L''(y_i | \hat{\eta}_i, \hat{\theta}) = dL'(y_i | \hat{\eta}_i, \hat{\theta}) / d\hat{\eta}_i$, and $\hat{\beta}$ and $\hat{\theta}$ are the current estimates of $\beta$ and $\theta$, respectively. Conditional on $\beta$, the shape parameter $\theta$ can be updated by maximizing the NB likelihood using the standard Newton-Raphson algorithm.

3. Results

3.1. Simulation Studies

We performed extensive simulations to benchmark our method as compared to several state-of-the-art approaches including DESeq, edgeR, and metagenomeSeq. To simulate count data for each feature, we generated OTU counts, $c_{ij}$, from the zero-inflated negative binomial distribution defined in (1). The ranges of the parameters in the simulation studies are described below.

We simulated OTU-level count data with $1000$ features for $n$ samples in each of two groups. The sample size $n$ in each group was set to be 50, 100, and 150. The effects of the first 50 features were set to be non-zero, and the others had zero effects. In this setting, the parameter $T_i$ denotes the scaling factor for sample $i$. We randomly set the offset $\log(T_i)$ from the range $[0.1, 3.5]$, the coefficient $\beta$ from three values $0.5, 1.0, 1.5$, and the shape parameter $\theta$ from the range $[0.1, 8]$. To simulate excess amount of zeros for each feature, we randomly generated from a Bernoulli trial with a preset proportion of zeros. We tested five proportions of zeros by varying $p_z = (0.1, 0.2, 0.3, 0.4, 0.5)$. We set these ranges of the parameters similar to those in Sohn et al. (2015). The ranges of all the parameters used in the simulation are summarized in Table 1.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>Logarithm of Scaling Factor $\log(T)$</td>
<td>Uniform $[0.1, 3.5]$</td>
</tr>
<tr>
<td>Shape Parameter $\theta$</td>
<td>Uniform $[0.1, 8]$</td>
</tr>
<tr>
<td>Coefficient Estimate $\beta$</td>
<td>$[0.5, 1.0, 1.5]$</td>
</tr>
<tr>
<td>Proportion $p_z$</td>
<td>$[0.1, 0.2, 0.3, 0.4, 0.5]$</td>
</tr>
</tbody>
</table>

We carried out differential abundance analysis for each simulated feature by using the following methods:

1. *edgeR* – exactTest - an exact binomial test with negative binomial model preceded by TMM normalization (Robinson et al., 2010).
2. *DESeq* – nbinomTest – a negative binomial conditioned test similar to *edgeR* but preceding RLE normalization (Anders & Huber, 2010).
3. *metagenomeSeq* – *fitZig* and *fitFeatureModel*. An Expectation-Maximization estimate of the posterior probabilities of differential abundance based on a zero inflated Gaussian model and zero-inflated log-normal mixture model respectively preceding a CSS normalization and log-transformation (Paulson et al., 2013).

For each combination of parameters, the procedure was repeated 100 times. All tests were corrected for multiple testing using Benjamini-Hochberg method to control the False Discovery Rate at the alpha level of 0.05. AUC, power, and false positive rate were calculated to compare the performance of each method. The simulations and real data analyses rely on the following R packages: *BhGLM*, *DESeq*, *edgeR*, *ggplot2*, *metagenomSeq*, *phyloseq*, *reshape2*, and *ROCR*. The reproducible simulation code in R is available upon request from the first authors.

Curves comparing AUC, power, and type I error rates are shown in Figures 1-3 to illustrate the performance of *BhGLM*, *ZINB*, *DESeq*, *edgeR*, and *metagenomeSeq* (*fitZig* and *fitFeatureModel*) on synthetic data with different proportions of zeros in detecting significant differentially abundant features for three different sample sizes $n = 100, 200$, and 300 respectively. Corresponding AUC, power, and type I error quantities are also listed in Tables A.1-A.3 (see Appendix).
Among them, we have listed the top ten significant taxa selected by the method metagenomeSeq-fitZig. The first three taxa are Neisseria polysaccharea, Neisseria meningitides, and Neisseria elongate. Neisseria meningitides has been reported as an uncommon cause of pneumonia (Winstead, McKinsey, Tasker, De Groote, & Baddour, 2000). On the other hand, an infective endocarditis caused by Neisseria elongata has been reported by (Haddow et al., 2003). Porphyromonas sp. has been isolated from both Cystic Fibrosis lung infections and non-small cell lung cancer patients (Rogers et al., 2004; Sato et al., 2015). Prevotella intermedia has also been investigated for its synergic effects to induce Severe Bacteremic Pneumococcal Pneumonia in mice (Nagaoka et al., 2014).

As shown in Figures 1-2 and Tables A.1-A.2, zero-inflated methods (BhGLM-ZINB, and metagenomeSeq-fitZig) performed similarly on simulated zero inflated data with respect to AUC. When the sample size is low (n=100) and the zero-inflation is 50%, the AUCs of BhGLM-ZINB and metagenomeSeq-fitZig are 0.862 and 0.858, respectively. With the same sample size but 10% zero-inflation, the AUCs of BhGLM-ZINB, and metagenomeSeq-fitZig were 0.921 and 0.910, respectively. With respect to empirical power, metagenomeSeq-fitZig performs slightly well than BhGLM-ZINB for ≥30% zero-inflation, while BhGLM-ZINB performs slightly better than metagenomeSeq-fitZig when the zero-inflation is 10%. On the other hand, other three methods, DESeq, edgeR, and metagenomeSeq-fitFeatureModel, have much lower power regardless of the sparsity level. As the proportion of zeroes increases, the empirical power of these methods worsens considerably. In terms of Type-I error rates, BhGLM-ZINB performs consistently better than metagenomeSeq-fitFeatureModel across all zero proportions. When the amount of zero-inflation is 50% and n = 100, BhGLM-ZINB controls the type I error rate at 0.026, while metagenomeSeq-fitFeatureModel and edgeR both have a much higher type I error rate (0.041 and 0.040 respectively). In all scenarios, the type I error rates are higher for DESeq, edgeR, and metagenomeSeq-fitFeatureModel, whereas for the proposed method, it is well-controlled, always lower than 0.03. In summary, our simulation indicates that the proposed ZINB method is consistently high in power along with well-controlled type I error.

3.2. Real Data Analysis

3.2.1. Lung Microbiome Data

We applied our method to a lung microbiome data from Charlson et al. (2011). The data was sampled in the respiratory flora from six healthy individuals. Among them, three individuals were smokers and the other three were non-smokers. The two-bronchoscope procedure was used to sample the respiratory tract, followed by a serial bronchoalveolar lavage and lower-airway protected brushes. A more detailed description of the lung microbiome samples, collection, and protocols is available in Charlson et al. (2011). The processed OTU level data was acquired directly from the R package metagenomeSeq. After the rare features were trimmed, we only include 259 features in our analysis. Differential abundance analysis to determine the associations between the remaining features with smoking status of 66 samples was carried out.

Among them, we have listed the top ten significant taxa in Figure 4. Some of the features are consistent with the significant taxa selected by the method metagenomeSeq-fitZig. The first three taxa are Neisseria polysaccharea, Neisseria meningitides, and Neisseria elongate. Neisseria meningitides has been reported as an uncommon cause of pneumonia in small cell lung cancer patients (Rogers et al., 2004; Sato et al., 2015). Prevotella intermedia has also been investigated for its synergic effects to induce Severe Bacteremic Pneumococcal Pneumonia in mice (Nagaoka et al., 2014).
3.2.2. Colorectal Cancer Microbiome Data

We further applied our method to a colorectal cancer gut microbiome data originally published by Kostic et al. (2012). They studied a secondary cohort of 95 individuals with both tumor and normal tissue acquired. A more detailed description about the gut microbiome samples, collection, and protocols is available in Kostic et al. (2012). We included 274 features with zero proportion not greater than 0.8 for 185 samples in our analysis. The differential abundance analysis was to explore the association of the features between normal and tumor samples.

The original study investigated the association between *Fusobacterium* and colorectal carcinoma. We have listed the coefficient estimates for the significant OTUs by phylum for each genus in Figure 5. The significant differentially abundant phylum for tumor samples includes *Firmicutes*, *Proteobacteria*, *Tenericutes*, *Actinobacteria*, *Bacteroidetes*, and *Fusobacteria*. Consistent with the original study and Wang et al. (2012), *Fusobacteria* in cancer samples are significantly enriched. Besides, the changes in *Bacteroidetes* and *Firmicutes* phyla are similar to the original study. It is presumed in the original study that *Fusobacterium* species contribute to the evolvement of tumor microenvironment, leading to an altered microbiota in accordance with the ‘alpha-bug’ hypothesis introduced by Sears and Pardoll (2011). 12 of the *Fusobacterium* species are differentially abundant between tumor and normal samples and four of them are enriched in tumor samples, both consistent with Kostic et al. (2012).

We also include the coefficients and p-values of top significantly abundant species in Figure 6. *Bacteroides uniformis* was reported as an enriched OTU-related species in healthy volunteers by Wang et al. (2012). Furthermore, based on the genera of the significant species, genera *Lactococcus*, *Bacteroides*, *Fusobacterium*, *Prevotella*, and *Streptococcus* exhibited more enriched in cancerous tissues than normal tissues, which is consistent with Gao et al. (2015). It is worth noting that two important species among the significant species in Figure 6 have been reported as antitumor bacteria. *Streptococcus thermophiles* has been known as an effective probiotic in deactivating risk factors of colon cancer (Wollowski, I., Rechkemmer, G., & Pool-Zobel, B. L., 2001).

*Ruminococcus bromii* was investigated for its ability to degradation of resistant starch in the human colon to potentially prevent colon cancer (Ze, X., Duncan, S. H., Louis, P., & Flint, H. J., 2012). This confirms that ZINB is able to replicate findings from previous reports, and can be used as an efficient tool for differential abundance analysis in future microbiome studies.

**Fig. 4 - Coefficients and P-values for the Top Significant Differentially Abundant Taxa between Smokers and Non-smokers in Lung Microbiome Data.** The left column is names of taxa; the right column is p-values. The ranges of coefficients are presented in the middle column.

**Fig. 5 - Coefficients Plot for Different Genus by Phylum in Colorectal Cancer Microbiome Data.**

**Fig. 6 - Coefficients and P-values for the Top Significant Differentially Abundant Species between Normal and Tumor Samples in Gut Microbiome Data.** The left column is names of taxa; the right column is p-values. The ranges of coefficients are presented in the middle column.

4. Discussion

Existing differential abundance analysis methods in the literature can be divided into three main categories: (i) methods that address the non-negativity and over-dispersion of the microbiome counts (typically based on negative binomial distribution) typically preceded by normalization, i.e. DESeq, DESeq2, and edgeR (Anders & Huber, 2010; Li, Witten, Johnstone, & Tibshirani, 2012; Peng et al., 2015; Robinson et al., 2010; White et al., 2009), (ii) methods that rely on some transformation of the normalized counts (e.g. metagenomeSeq), and (iii) methods that handle zero-inflation by formulating some mixture model (e.g. metagenomeSeq, ZIBSeq). While most of these methods rely on some normalization scheme such as CSS, TMM, or TSS prior to modeling, the proposed method bypasses the need of normalization and accounts for sparsity.
and overdispersion simultaneously by directly modeling the raw counts. This makes ZINB a flexible modeling strategy, leading to interpretable differential abundance estimates in the original measurement scale. We have shown the consistency of ZINB on various types of samples in the simulation study as well as in two real human microbiome datasets.

In the lung microbiome data and some other previous studies, we have found that a common problem with the existing methods is that they fail to consider clustered structure (if any) in the data. For example, lung microbiome data has 78 samples taken from six individuals which were treated as independent samples in the previous analyses. Therefore, a future direction of research is to incorporate random effects in the proposed ZINB model that can account for clustered structure in the observations. We have optimized our simulations to the analysis of differential abundance between two conditions of samples such as healthy versus diseased in this article. However, our method can be easily extended to more than two conditions. Finally, even though we have developed our method for microbiome data, ZINB should be applicable to other similar types of count data such as RNA-Seq. This strength significantly broadens the impact of ZINB among researchers in the biological community.

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ОТРИЦАТЕЛЬНАЯ БИНОМИАЛЬНАЯ РЕГРЕССИЯ С НУЛЕВЫМ РАЗДУВОМ ДЛЯ ОЦЕНКИ ОТНОСИТЕЛЬНОГО СОДЕРЖАНИЯ В МИКРОБИОМНЫХ ИССЛЕДОВАНИЯХ

Конфликт интересов

Не указан.

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Аннотация

Мотивация: Человеческий микробиом играет важную роль в нормальном и патологическом состоянии человека. Состав микробиома человека формируется под влиянием многих факторов, и понимание этих факторов имеет решающее значение для установления роли микробиома в нормальном и патологическом состоянии, а также для разработки новых диагностических или терапевтических целей на основании микробиома. Целевое секвенирование ампликона гена 16S рибосомальной РНК (рРНК) является широко используемым подходом для определения таксономического состава бактериального сообщества. Операционные таксономические единицы (OTU) группируются в кластеры на основе сгенерированной последовательности ридов и используются для определения того, каким образом относительное содержание микробиома коррелирует с некоторыми характеристиками испытуемых, такими как здоровое/патологическое состояние, курение или пищевые привычки. Тем не менее, данные подсчета OTU не только имеют чрезвычайно сильный разброс, но также содержат избыточное количество нулевых результатов подсчета из-за недостаточности выборки. Поэтому для последующего статистического анализа необходимы эффективные аналитические инструменты, которые могут одновременно учитывать избыточную дисперсию и разреженность данных микробиома.

Результаты: В данной статье предложена отрицательная биномиальная регрессию с нулевым раздувом (ZINB) для выявления различий в относительном содержании таксонов между двумя или более популяциями. Предлагаемый способ использует алгоритм максимизации ожидания (EM-алгоритм) путем включения смешанной модели из двух частей, состоящей из (i) отрицательной биномиальной модели для учета избыточной дисперсии и (ii) регрессионной логистической модели для учета избыточного количества нулевых результатов подсчета. Проводятся обширные симуляционные исследования, которые показывают, что ZINB демонстрирует более высокую эффективность по сравнению с несколькими современными подходами на основании измерения площади под кривой (AUC). Применение предлагаемого метода к двум реальным наборам данных показывает, что он способен обнаруживать биологически значимые таксоны и сопоставлять их с предыдущими исследованиями.


Дополнительная информация: Дополнительные данные доступны в Интернете в Журнале биоинформатики и геномики.

Ключевые слова: Дифференциальная оценка относительного содержания, EM-алгоритм, микробиом человека, метагеномика, OTU, отрицательная биномиальная [регрессия] с нулевым раздувом.