Transformation and differential abundance analysis of microbiome data incorporating phylogeny

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Genome analysis

Transformation and differential abundance analysis of microbiome data incorporating phylogeny

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Outline: Overview of paper

- Extend Dirichlet-tree multinomial to zero-inflated Dirichlet tree multinomial to model multivariate microbial counts.
- Use an Empirical Bayes based posterior mean transformation to convert raw counts into non-zero relative abundances that sum to 1.
- Introduce phylogenetically informed DA procedure on transformed data - adaptive analysis of composition of microbiomes for DA testing (adaANCOM)- log-ratios adaptively on the tree for each taxon.

Introduction

- Differential abundance tests need to account for the high dimensional, sparse, compositional, negatively and positively correlated, phylogenetically structured microbiome data.
- Scaling methods to correct for compositional bias (TMM, DESeq, CSS) - assume that most taxa are not differentially abundant, when the count matrix is sparse it can over or underestimate diversity, distort correlations

Review of Dirichlet-Multinomial distribution

- The Dirichlet-multinomial (DM) distribution is commonly used for OTU counts and is a compound distribution comprised of Dirichlet and Multinomial distributions
- Let $\mathbf{y} = (y_1, \dots, y_K)^T$ be the count vector for a sample with K OTUs.

•
$$y^+ = \sum_{k=1}^K y_k$$
, $p = (p_1, \dots, p_K)^T$, $\sum_{k=1}^K p_k = 1$, $\alpha^+ = \sum_{k=1}^K \alpha_k$

Multinomial pdf:

Dirichlet pdf:

$$f_M(\mathbf{y}; \mathbf{p}) = \frac{\Gamma(\mathbf{y}^+ + 1)}{\prod\limits_{k=1}^K \Gamma(\mathbf{y}_k + 1)} \prod\limits_{k=1}^K p_k^{\mathbf{y}_k}, \qquad f_D(\mathbf{p}; \mathbf{\alpha}) = \frac{\Gamma(\alpha^+)}{\prod\limits_{k=1}^K \Gamma(\alpha_k)} \prod\limits_{k=1}^K p_k^{\alpha_k},$$

• Combining both we get the Dirichlet-Multinomial distribution:

$$= \frac{f_{DM}(\mathbf{y}; \mathbf{\alpha}) = \int f_{M}(\mathbf{y}; \mathbf{p}) f_{D}(\mathbf{p}; \mathbf{\alpha}) d\mathbf{p}}{\Gamma(\mathbf{y}^{+} + 1)\Gamma(\alpha^{+})} \prod_{k=1}^{K} \frac{\Gamma(\mathbf{y}_{k} + \alpha_{k})}{\Gamma(\mathbf{y}_{k} + 1)\Gamma(\alpha_{k})}$$

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Relationship between Gamma, Beta, & Dirichlet distributions

- Let Z_1, \ldots, Z_K independent $Z_k \sim \mathsf{Gamma}(\alpha_k, \lambda)$,
- $X_k = \frac{Z_k}{\sum_{j=1}^K Z_j}$
- $W_k = \frac{Z_k}{\sum_{j=k}^K Z_j}$
- Joint distribution of $\boldsymbol{X} = (X_1, \dots, X_K)^T$ is Dirichlet distributed with parameter α
- $W_k \sim \text{Beta}(\alpha_k, \alpha_k^+)$ independent
- $X_k = W_k \prod_{j=1}^{k-1} (1 W_j)$
- We can rewrite the Dirichlet-Multinomial distribution as

$$f_{DM}(\mathbf{y}; \boldsymbol{\alpha}) = \int f_M(\mathbf{y}; h(\boldsymbol{w})) \prod_{k=1}^{K-1} f_B(\boldsymbol{w}_k; \alpha_k, \alpha_k^+) d\boldsymbol{w}$$

• $h(\cdot)$ is the transformation from W to X.

Zero Inflated Dirichlet Multinomial

 We can introduce zero inflation by defining the Zero Inflated Dirichlet Multinomial (ZIDM) as:

$$f_{ZIDM}(\boldsymbol{y}; \boldsymbol{\pi}, \boldsymbol{\alpha}) = \int f_{M}(\boldsymbol{y}; h(\boldsymbol{w})) \prod_{k=1}^{K-1} f_{ZIB}(\boldsymbol{w}_{k}; \pi_{k}, \alpha_{k}, \alpha_{k}^{+}) d\boldsymbol{w}$$

where

$$f_{ZIB}(w_k; \pi_k, \alpha_k, \alpha_k^+) = \pi_k \delta(0) + (1 - \pi_k) f_B(w_k; \alpha_k, \alpha_k^+).$$

is a zero-inflated binomial distribution

• π_k is the probability of zero-inflation in the kth component $\delta(\cdot)$ is the Dirac delta function

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Introducing the Dirichlet Tree Multinomial

- Suppose the relationships between OTUs are encoded in a tree $\mathcal T$ which is composed of internal nodes $\mathcal V$ and leaf nodes $\mathcal L$
- Leaf nodes are OTUs, internal nodes are the node to branch on the phylogenetic tree
- For each $v \in \mathcal{V}$, denote C_v as the set of child nodes of v, \boldsymbol{y}_v the vector of counts corresponding to C_v , and $y_v^+ = \sum_{u \in C_v} y_u$.

Dirichlet Tree Multinomial & Zero-Inflated Dirichlet Tree Multinomial

• Assume that \mathbf{y}_{v} conditional on y_{v}^{+} are independent across internal nodes, the Dirichlet Tree Multinomial (DTM) distribution is the product of DM distributions that factorize over the tree

$$f_{\mathrm{DTM}}(\mathbf{y}; \mathbf{\alpha}_{\nu}, \nu \in \mathcal{V}) = \prod_{\nu \in \mathcal{V}} f_{\mathrm{DM}}(\mathbf{y}_{\nu}; \mathbf{y}_{\nu}^{+}, \mathbf{\alpha}_{\nu})$$

 To extend the DTM to the ZIDTM, we replace the Dirichlet multinomial with Zero-Inflated Dirichlet Multinomial

$$f_{ZIDTM}(\mathbf{y}; \mathbf{\pi}_{\nu}, \mathbf{\alpha}_{\nu}, \mathbf{v} \in \mathcal{V}) = \prod_{v \in \mathcal{V}} f_{ZIDM}(\mathbf{y}_{v}; \mathbf{y}_{v}^{+}, \mathbf{\pi}_{\nu}, \mathbf{\alpha}_{v})$$

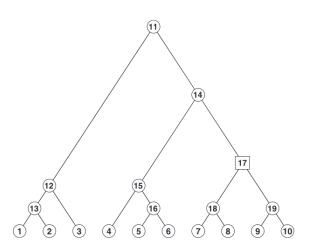


Fig. 1. A binary tree with K = 10 leaves. Here $\mathcal{L} = \{1, 2, ..., 10\}$, $\mathcal{V} = \{11, 12, ..., 19\}$. For illustration, $\mathcal{C}_{17} = \{18, 19\}$, $y_{17} = (y_{18}, y_{19})^T$ and $y_{17}^+ = y_{18} + y_{19}$. Given y_{17}^+ , y_{17} has a DM or ZIDM distribution. The factorization over internal nodes means that these conditional distributions are independent

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Maximum likelihood estimation for ZIDTM

- Use EM-algorithm to estimate unknown parameters: $\theta = \{\alpha_{\nu}, \pi_{\nu}, \nu \in \mathcal{V}\}$
- Assume $C_v = \{1, \dots, K_v\}$, where $K_v = |C_v|$
- Consider n observations y^1, \ldots, y^n , then the log-likelihood function (without constant terms) is:

$$l(\boldsymbol{\theta}) = \sum_{i=1}^{n} \sum_{v \in \mathcal{V}} \sum_{k=1}^{K_{v}-1} \{ l_{1}(\delta_{vk}^{i}, \pi_{vk}) + l_{2}(\delta_{vk}^{i}, w_{vk}^{i}, \alpha_{vk}, \alpha_{vk}^{+}) \}$$

with

$$\begin{split} l_1(\delta^i_{vk},\pi_{vk}) &= \delta^i_{vk} \log \pi_{vk} + (1-\delta^i_{vk}) \log (1-\pi_{vk}) \\ l_2(\delta^i_{vk},w^i_{vk},\alpha_{vk},\alpha^+_{vk}) &= (1-\delta^i_{vk}) \{ -\log \mathcal{B}(\alpha_{vk},\alpha^+_{vk}) \\ &+ (\alpha_{vk}-1) \log w^i_{vk} + (\alpha^+_{vk}-1) \log (1-w^i_{vk}) \} \end{split}$$

ullet $\delta^i_{\it vk}$ is the indicator of zero-inflation ${\cal B}()$ is the beta function

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E-step

• Compute expectation of $l(\theta)$ with respect to posterior distribution of $(\delta^i_{vk}, w^i_{vk})|\mathbf{y}^i_{v}$, indexed by current value of θ , which gives the Q-function:

$$Q(\boldsymbol{\theta}) = \sum_{i=1}^{n} \sum_{v \in \mathcal{V}} \sum_{k=1}^{K_{v}-1} E\{l_{1}(\delta_{vk}^{i}, \pi_{vk}) + l_{2}(\delta_{vk}^{i}, w_{vk}^{i}, \alpha_{vk}, \alpha_{vk}^{+})\}$$

E-step: Definitions

- $\bullet \ \delta_{vk}^{i*} = E(\delta_{vk}^i | \boldsymbol{y}_v^i)$
- $R_{vk}^{i*} = E(\log w_{vk}^i | \mathbf{y}_v^i, \delta_{vk}^i = 0)$
- $S_{vk}^{i*} = E\{\log(1 w_{vk}^i)|\mathbf{y}_v^i, \delta_{vk}^i = 0\}$
- Then,

$$\delta_{vk}^{i*} = egin{cases} 0, & y_{vk}^{i} > 0, \ \dfrac{\pi_{vk}}{\pi_{vk} + (1 - \pi_{vk}) \dfrac{\mathcal{B}(lpha_{vk}^{i*}, lpha_{vk}^{i*+})}{\mathcal{B}(lpha_{vk}, lpha_{vk}^{+})}, & y_{vk}^{i} = 0, \end{cases}$$

$$S_{\nu k}^{i*} = \psi(\alpha_{\nu k}^{i*+}) - \psi(\alpha_{\nu k}^{i*} + \alpha_{\nu k}^{i*+}) \\ R_{\nu k}^{i*} = \psi(\alpha_{\nu k}^{i*}) - \psi(\alpha_{\nu k}^{i*+})$$

• $\alpha_{vk}^{i*} = \alpha_{vk} + y_{vk}^i$, $\alpha_{vk}^{i*+} = \sum_{j=k+1}^{K_v} \alpha_{vk}^{i*}$, $\psi(\cdot)$ is the digamma function.

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E-step: Q-function

• We can rewrite $Q(\theta)$ as:

$$Q(\boldsymbol{\theta}) = \sum_{i=1}^{n} \sum_{v \in \mathcal{V}} \sum_{k=1}^{K_{v}-1} \{Q_{1}(\pi_{vk}, \delta_{vk}^{i}) + Q_{2}(\alpha_{vk}, \alpha_{vk}^{+}, R_{vk}^{i*}, S_{vk}^{i*})\}$$

where

$$Q_1(\pi_{\nu k}, \delta_{\nu k}^i) = \delta_{\nu k}^{i*} \log(\pi_{\nu k}) + (1 - \delta_{\nu k}^{i*}) \log(1 - \pi_{\nu k})$$

and

$$\begin{split} Q_2(\alpha_{vk},\alpha_{vk}^+,R_{vk}^{i*},S_{vk}^{i*}) &= (1-\delta_{vk}^{i*})\{-\log\mathcal{B}(\alpha_{vk},\alpha_{vk}^+)\\ &+ (\alpha_{vk}-1)R_{vk}^{i*} + (\alpha_{vk}^+-1)S_{vk}^{i*}\} \end{split}$$

M-step

- Maximize $Q(\theta)$ with respect to θ
- ullet The parameters in Q_1 and Q_2 can be optimized separately
- Model depends on ordering of OTUs at each internal node (matching problem)
- Fit separate ZIDM model for each possible ordering of $|C_v|$ taxa, and select the best fitted model.
- Computational cost is $O(\sum_{v \in \mathcal{V}} |C_v|!)$ compared to $O(|\mathcal{L}|!)$ for GDM or ZIDM models

Binary trees

- From now on, consider binary trees
- i.e. $|C_v| = 2$ for all $v \in \mathcal{V}$
- Dirichlet, multinomial, dirichlet multinomial, and zero inflated dirichlet multinomial are reduced to beta, binomial, beta-binomial and zero-inflated beta-binomial.
- The ZIDTM is then a product of ZIBBs

Posterior mean transformation

- Estimate underlying proportions from Bayesian perspective (since Dirichlet is conjugate to multinomial)
- Consider the posterior mean for the Dirichlet Multinomial:

$$E_{DM}(p_k|\mathbf{y}) = \frac{\alpha_k + y_k}{\sum\limits_{j=1}^K (y_j + \alpha_j)}$$

- We estimate unknown parameters by maximizing data likelihood
- Use the estimated parameters as "pseudo data" and combine with real observed data.
- This produces non-zero proportions for zero counts
- Alternative to using a pseudo-count for sample proportions
- Becomes difficult in presence of zero-inflation

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Solving posterior mean transformation

- When we have a binary tree, there is an explicit closed-form solution to the zero-inflation
- At each internal node $v \in \mathcal{V}$,

$$E_{BB}(p_{\nu 1}|\mathbf{y}_{\nu}) = \frac{\alpha_{\nu 1} + y_{\nu 1}}{\alpha_{\nu 1} + \alpha_{\nu 2} + y_{\nu 1} + y_{\nu 2}},$$

and

$$E_{ZIBB}(p_{\nu 1}|\mathbf{y}_{\nu}) = \frac{(1 - B_{\nu 0})B_{\nu 1}}{B_{\nu 0}\mathcal{B}(\alpha_{\nu 1}, \alpha_{\nu 2}) + (1 - B_{\nu 0})B_{\nu 2}},$$

- $B_{v0} = \pi I(y_{v1} = 0)$
- $B_1 = \mathcal{B}(1 + \alpha_{v1} + y_{v1}, \alpha_{v2} + y_{v2})$
- $B_2 = \mathcal{B}(\alpha_{v1} + y_{v1}, \alpha_{v2} + y_{v2})$



Likelihood-ratio test & count transformation

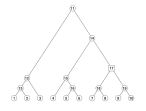
- Having a correct model specification at internal nodes effects quality of posterior estimates.
- Perform a two-stage likelihood-ratio test
- First, assume count data at $v \in \mathcal{V}$ are not zero-inflated, fit beta-binomial model and test for over-dispersion.
- Counts at nodes without over-dispersion are transformed into sample proportions after adding constant of .5, equivalent to using E_{BB} with $\alpha_{v1}=\alpha_{v2}=0.5$
- Second, Nodes with over-dispersion are refit using a ZIBB model, and tested again for zero-inflation
- Counts are then transformed using $E_{ZIBB}(p_{v1}|\mathbf{y}_{v})$.
- ullet If there is no zero-inflation, counts are transformed using $E_{BB}(p_{v1},oldsymbol{y}_v)$
- $\hat{\boldsymbol{p}}_{v} = (\hat{p}_{v1}, 1 \hat{p}_{v1})^{T}$ is the posterior estimate

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Path-level definitions - Phylogeny-aware normalization

- To ensure that the normalization is 'phylogeny-aware', we consider the path level instead of focussing on individual internal nodes.
- Define A_u as the ancestor node set of u for each node $u \in \mathcal{L} \cup \mathcal{V}$ that contains all internal nodes in the path from the root node to u
- \bullet \mathcal{L}_u is the set of leaves in the same path

$$\begin{array}{l} \bullet \ \, \mathcal{A}_1 = \mathcal{A}_2 = \{11,12,13\}, \\ \mathcal{A}_3 = \{11,12\}, \ \mathcal{L}_{11} = \mathcal{L}, \mathcal{L}_{12} = \\ \{1,2,3\}, \mathcal{L}_{13} = \{1,2\} \end{array}$$



- Define $q_u = \prod_{v \in A_u} \hat{p}_v$
- \mathcal{U} is the set of nodes such that $\bigcup_{u \in \mathcal{U}} \mathcal{L}_u = \mathcal{L}$, $\mathcal{L}_u \cap \mathcal{L}_{u'} = \emptyset$ when $u \neq u'$.
- ullet Then, $\sum_{u\in\mathcal{U}}q_u=1$ and $\sum_{l\in\mathcal{L}}q_l=1$

adaANCOM

- Detect differentially abundant OTUs at ecosystem level
- Similar to ANCOM
- Difference is how we use the phylogenetic tree to incorporate this information
- Tests differences between two groups
- Speed up ANCOM by using phylogenetic tree to construct adaptive log-ratios

Hypothesis tests - ANCOM

• Test the hypothesis

$$H_{0I}: \log \mu_I^1 = \log \mu_I^2$$

for $I \in \mathcal{L} = \{1, \dots, K\}$, and where μ_I^g is the mean absolute abundance of the /th OTU from gth group, and g = 1, 2

• In ANCOM, we test K-1 hypothesis using each OTU as the denominator in the additive log ratio

$$H_{0lm}: \log(\mu_l^1/\mu_m^1) = \log(\mu_l^2/\mu_m^2)$$

for all $m \neq I$

- ullet ANCOM counts the number of rejections across the K-1 hypothesis tests, and compares to the empirical distribution of counts to determine if the /th OTU is differentially abundant.
- If K is high, ANCOM suffers from a high FDR. Additionally the computational burden is high

adaANCOM log-ratios & hypothesis tests

- Assume that abundance difference on the log scale at an internal node is passed down to child nodes
- adaANCOM is comprised of a two-step process
 - Test the internal node-level hypothesis

$$H_{0\nu}: \log(\mu_{\nu 1}^1/\mu_{\nu 2}^1) = \log(\mu_{\nu 1}^2/\mu_{\nu 2}^2)$$

for $v \in \mathcal{V}$, with μ_{v1}^g and μ_{v2}^g as mean absolute abundances of children of v from the gth group.

For a given $\alpha, \mathcal{D}_{\mathcal{V}}$ is the set of internal nodes which the hypotheses are rejected

② For each leaf node $I \in \mathcal{L}$, calculate the log-ratio Define ref_I as the sibling node of I if $\mathcal{A}_I \cap \mathcal{D}_{\mathcal{V}} = \emptyset$ and the child node of v not in \mathcal{A}_I otherwise, when v is the node in $\mathcal{A}_I \cap \mathcal{D}_{\mathcal{V}}$ closest to the root node.

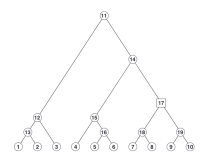
The null hypothesis is:

$$H_{0l}^{ada}: \log(\mu_l^1/\mu_{ref_l}^1) = \log(\mu_l^2/\mu_{ref_l}^2)$$

• adaANCOM rejects the null hypothesis H_{0I} if H_{0I}^{ada} is rejected

adaANCOM example

- Assume $\mathcal{D}_{\mathcal{V}} = 17$.
- $ref_1 = 2$, $ref_2 = 1$ $ref_3 = 13$, $ref_4 = 16$, $ref_5 = 6$, $ref_6 = 5$, $ref_7 = ref_8 = 19$, $ref_9 = ref_{10} = 18$



Accounting for log-ratios of zero y_u counts

- H_{0v} and H_{0l}^{ada} are based on the log-ratios of q_u s for $u \in \mathcal{L} \cup \mathcal{V}$, but when the original y_u counts have many zero counts, log-ratios give abnormal results, and test statistics are sensitive to these abnormal values
- ullet For internal node v, define ϕ_v as the maximum of

$$|\log(\hat{p}_{v1}/\hat{p}_{v2})|$$

that have $y_{v1} > 0$ and $y_{v2} > 0$.

• Data with $y_{v1}=0$ or $y_{v2}=2$ are removed if $|\log(\hat{p}_{v1}/\hat{p}_{v2})|>\phi_v$

Comparisons to ANCOM

- Similar to ANCOM, adaANCOM uses relative abundance data, constructs log-ratios, and performs t-tests or Wilcoxon-rank-sum tests.
- The advantage of adaANCOM is the computation time, as the number of tests is reduced by a factor of $|\mathcal{L}|$.
- adaANCOM also has an advantage of interpretability since we are guided by the tree, so we result in both DA leaves and nodes.
- Additionally, adaANCOM controls FDR better

adaANCOM algorithm

Algorithm 1: adaANCOM

```
Input: A binary tree T = (\ell, V), posterior-mean-transformed data
          \{\hat{p}_{v}, v \in \mathcal{V}\}, group information, and a testing procedure;
Output: DA internal nodes \mathcal{D}_{\mathcal{V}}, and DA leaf nodes \mathcal{D}_{\ell};
Step 1:
Set \mathcal{D}_{\mathcal{V}} = \emptyset, for \nu \in \mathcal{V} do
   Construct the log-ratios \log(\hat{p}_{v1}/\hat{p}_{v2}), and remove the outliers;
   Test H_{0\nu}, and if rejected, update \mathcal{D}_{\mathcal{V}} = \mathcal{D}_{\mathcal{V}} \cup \{\nu\};
end
Step 2:
Set \mathcal{D}_{\ell} = \emptyset, for l \in \ell do
   Search ref_l, compute q_l and q_{ref_l};
   Construct the log-ratios \log(q_l/q_{ref_l}), and remove the
   outliers;
   Test H_{0l}^{ada}, and if rejected, update \mathcal{D}_{\ell} = \mathcal{D}_{\ell} \cup \{l\};
end
```