

BRIGHAM AND WOMEN'S HOSPITAL

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Introduction

Microbes are everywhere including in and on our bodies, and have been shown to play key roles in a variety of prevalent human diseases. Consequently, there has been intense interest in the design of bacteriotherapies or "bugs as drugs", which are communities of bacteria administered to patients for specific therapeutic applications. Central to the design of such cocktails is the knowledge (or inference) of a causal microbial interaction network and prediction of the population dynamics of the organisms. In this work we present a Bayesian nonparametric model and associated efficient inference algorithm that addresses the key conceptual and practical challenges of learning microbial dynamics from time series microbe abundance data.

Interpretability and the Microbiome

Challenges associated with inference in the microbiome:

- High-dimensional (300+ strains of bacteria in the gut, **potentially 100,000** microbe-microbe interactions).
- Temporally sparse and non-uniformly sampled data.
- High measurement noise
- Nonlinear and physically non-negative dynamics.

Many potential interactions: how do we decide what interactions are "real" and how do we simplify the interaction landscape so as to be interpretable?

Contribution:

- 1. Introduction of a temporally varying auxiliary variable technique to enable efficient inference by relaxing hard non-negativity constraint
- 2. Clustering of microbes into redundant interaction modules and structural learning of a compact interaction network among modules.



Robust and Scalable Models of Microbiome Dynamics

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Lotka-Volterra Dynamics Abundance of Microbe *i* at time t: $X_{t,i}$ growth parameter $\frac{\mathrm{d}\mathbf{x}_{t,i}}{\mathrm{d}t} = \mathbf{\alpha}_i \mathbf{x}_{t,i} + \mathbf{\beta}_{ii} \mathbf{x}_{t,i}^2 + \sum_{j \neq i} \mathbf{\beta}_{ij} \mathbf{x}_{t,i} \mathbf{x}_{t,j} + \frac{\mathrm{d}\mathbf{w}_{t,i}}{\mathrm{d}t}$ self limiting

Discrete approximation to the Lotka-Volterra dynamics

 $\mathbf{x}_{k+1,i} = \mathbf{x}_{k,i} + \left(\boldsymbol{\alpha}_i \mathbf{x}_{k,i} + \boldsymbol{\beta}_{ii} \mathbf{x}_{k,i}^2 + \sum_{i \neq i} \boldsymbol{\beta}_{ij} \mathbf{x}_{k,i} \right)$: discrete time index

- Cluster the microbes into interaction modules (**Dirichlet Process** prior) • No interactions within cluster, only between clusters.
- Edge Selection: add indicator variables for cluster-cluster interactions (Bayesian variable selection, structure learning for graphical models) • Insert auxiliary variable q between the measurements y, Q and the state x

Comment: not allowing within cluster interactions dramatically reduces the number of inferred interactions and is consistent with the biologically important scenario of redundant functionality among sets of microbes

Additional Model Components

Dirichlet Process (clustering)

cluster $m{\pi_{c}} \mid m{lpha} \sim \mathtt{Stick}(m{lpha})$ assignment $\mathbf{C}_i \mid \boldsymbol{\pi}_{\mathbf{c}} \sim \texttt{Multinomial}(\boldsymbol{\pi}_{\mathbf{c}})$ $\mathbf{b}_{\mathbf{c}_i,\mathbf{c}_i} \mid \boldsymbol{\sigma}_{\mathbf{b}} \sim \texttt{Normal}(0, \boldsymbol{\sigma}_{\mathbf{b}}^2)$

cluster interactions coefficient

Dynamics

 $\mathbf{x}_{k+1,i} \mid \mathbf{x}_k, \mathbf{a}_i, \mathbf{b}, \mathbf{c}, \mathbf{z}, oldsymbol{\sigma_w} \sim \mathbf{a}_i$

$$\operatorname{Normal}\left(\mathbf{x}_{k,i} + \mathbf{x}_{k,i}\left(\mathbf{a}_{i,1} + \mathbf{a}_{i,2}\mathbf{x}_{k,i} + \sum_{\mathbf{c}_j \neq \mathbf{c}_i} \mathbf{b}_{\mathbf{c}_i,\mathbf{c}_j} \mathbf{z}_{\mathbf{c}_i,\mathbf{c}_j} \mathbf{x}_{k,j}\right), \Delta_k \boldsymbol{\sigma}_{\mathbf{w}}^2\right)$$

Measurement Model

auxiliary variable	$\mathbf{q}_{k,i} \mid \mathbf{x}_{k,i} \sim \texttt{Normal}(k)$
reads	$\mathbf{y}_{k,i} \mid \mathbf{q}_{k,i} \sim \mathtt{NegBin}($
qPCR	$Q_k \mid q_{k,i} \sim \texttt{Normal}$ (

Introduction of **q** allows for

- Efficient Gibbs/collapsed Gibbs sampling
- Posterior distributions for coefficients a, **b** are Gaussian, (direct sampling from posterior)
- •Can Marginalize out in closed form the interaction coefficients **b** for both module learning and structure learning

Without q, dynamics would have been distribution. modeled truncated with resulting in the posteriors of a, b being truncated as well and not allowing for marginalization elsewhere in the model.

[1] Bucci, Vanni, et al. "MDSINE: Microbial Dynamical Systems INference Engine for microbiome time-series analyses." *Genome Biology* (2016)

Dynamical Model

interaction coefficient

disturbance term

$$(k, i \mathbf{x}_{k,j}) \Delta_{k} + (\mathbf{w}_{k+1,i} - \mathbf{w}_{k,i}) \sqrt{\Delta_{k}}$$

discrete time step size

Edge Selection

- $\mathbf{Z}_{\mathbf{C}_i,\mathbf{C}_i} \mid \boldsymbol{\pi}_{\mathsf{z}} \sim \texttt{Bernouli}(\boldsymbol{\pi}_{\mathsf{z}})$ indicator variable
- for cluster interaction

Self Interaction

$$\mathbf{a}_{i,1}, \mathbf{a}_{i,2} \mid \boldsymbol{\sigma}_{\mathbf{a}} \sim \texttt{Normal}(0, \boldsymbol{\sigma}_{\mathbf{a}}^2)$$

$$egin{aligned} \mathbf{x}_{k,i}, oldsymbol{\sigma}_{\mathbf{q}}^2) \ \phi(\mathbf{q}_k), \epsilon(\mathbf{q}_k) \end{pmatrix} \ (\sum_i \mathbf{q}_{k,i}, \sigma^2_{\mathbf{Q}_k}) \end{aligned}$$

 $\mathbf{q}_{k,i} \sim \texttt{Uniform}[\mathsf{0},\mathtt{L})$





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HARVARD MEDICAL SCHOOL



 Colonize mice with human fecal sample containing 300+ bacteria