Combinatorial Mixture Models for Single Cell Assays

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March 21, 2012

Abstract

Immunological endpoints in vaccine trials are measured through a number of different assays that often provide single-cell measurements of multiple, specific intracellular or cell surface proteins, or mRNA expression levels of genes in single cells from specific cell sub-populations. While these measurements are usually continuous, they are often discretized and analyzed as counts of single cells. For example, in flow cytometry, individual cells are often classified as either positive or negative for a marker based on a predetermined threshold. One such assay is the intracellular cytokine staining assay that is used to assess an individual's immune response to a vaccine by measuring the abundance antigen-specific T-cell subpopulations producing different cytokines. Cells are classified as cytokine positive or negative and the resulting cell counts are used to identify experimental conditions leading to an increase in cytokine producing cells. ICS assays are generally used to identify "vaccine responders". These are individuals whose immune system produces significantly more cytokine positive cells in response to antigen stimulation than at baseline. The rarity of these cell populations makes maximizing the sensitivity and specificity of the assay a primary concern. The standard approach to analyze such data, Fisher's exact test, is problematic for two reasons. It can be overly conservative in detecting true differences for small counts, and the data generally do not meet the assumptions of the test. Specifically, the total cell counts across conditions are generally not fixed since these are generated by independent experiments. In this paper we present a coherent framework based on mixtures of Beta-binomial or Dirichlet-Multinomial distributions for analyzing count data derived from single-cell assays. Using the ICS assay as a motivating example, our method models cytokine-specific T-cell response across all individuals simultaneously, sharing information across observations using empirical Bayes methods. The model represents the responder and non-responder observations as separate components in the model, stimulated and unstimulated cell counts are modelled independently. Using simulations and real-world vaccine trial data, we show that our model increases the sensitivity and specificity for positivity calls in ICS assays compared to Fisher's exact test.

1 Introduction

Single—cell assays are an important tool in immunology, providing a functional and phenotypic snapshot of the immune system at a given time. These assays typically measure multiple variables simultaneously on individual cells in a homogeneous mixture such as whole blood. These variables are used to classify individual cells in the mixture into more homogeneous sub—populations based on phenotypic or functional differences. Such single—cell assays are used for immune monitoring of disease, vaccine research, and diagnosis of haematological malignancies [1–3].

A motivating example from vaccine trial research is the flow cytometric intracellular cytokine staining (ICS) assay, which is used to identify individuals whose immune system responds to a vaccine. Upon vaccination, antigen in the vaccine is taken up and presented to CD4 or CD8 T-cels via antigen presenting cells. Not all T-cells can recognize all antigens. Those that recognize antigens in the vaccine become activated and produce a variety of cytokines that further promote the immune response. After activation, this antigen-specific subpopulation proliferates and can persist in the immune system for some time providing *memory* that can more rapidly recognize the same antigen again in the future. None the less, these antigen-specific T-cell subpopulations a very small fraction of the total number of CD4 and CD8 T-cells. The ICS assay measures the number of antigen-specific T-cells in whole blood by measuring cytokine production in response to activation following stimulation by an antigen that was present in the original vaccine. Individual cells are labelled using fluorescently conjugated antibodies against phenotypic markers (CD3, CD4, and CD8) and functional markers (cytokines) of the cell subpopulations of interest [2, 4, 5]. A sufficiently large number of cells must be collected to ensure that the rare cell populations can be detected. Subsequently, each individual cell is classified as either positive or negative for each maker based on predetermined thresholds, then the number of cells matching each subpopulation phenotype is counted. These counts are compared between antigen stimulated and unstimulated samples from an individual to identify significant differences. Assessing a broad T cell response to a vaccine is particularly important in HIV vaccine trials, where the search for immune correlates of protection against HIV progression and infection is ongoing [4, 6, 7].

The ICS assay in flow cytometry is just one example of the applications of single—cell technologies to immunology. Single—cell gene expression technologies such as Fluidigm are enabling researchers to measure the expression of 96 genes in 96 single cells from a homogeneous population sorted by flow cytometry[8]. This technology has been used to identify signatures of immune cell sub—populations that correlate with different types of vaccine [9]. To date, published analysis of Fluidigm data ignore cells where transcripts are undetected, focusing solely on continuous gene expression measurements. However, the proportion of single cells expressing individual genes also carries information and should be evaluated.

Here we present a combinatorial mixture model framework for the analysis of single-cell assay data in which multivariate measurement are made individual cells. The model is used to contrast two conditions (i.e. treatment and control) and identify observations where a significant differences exists between the treatment and control samples with respect to the proportions of cells expressing different combinations of proteins, genes, or other measured properties of the cells.

We develop our framework in the context of ICS data. While there is no standard approach to analyzing ICS assay data, Fisher's exact test, as well as more ad-hoc methods, are commonly applied on a per-cytokine basis to identify observations where the number of cytokine producing cells is greater in the stimulated than the control data [4, 10, 11]. The drawbacks of Fisher's exact test applied to 2 x 2 contingency table data when the assumptions of the test are not met, have been reviewed in the literature [12]. Briefly, for single-cell assay data the assumptions of fixed margins are not met, since the observations for treatment and control conditions arise from independent experiments, thus the p-values from Fisher's exact test are incorrect. More importantly, Fisher's exact test does not share information across observations, resulting in low power when counts are small. The framework developed in this paper addresses these issues explicitly.

Materials and Methods

Vaccine Trial ICS Dataset Description

HVTN054 is a phase 1 (safety and efficacy) trial of an adenoviral vector vaccine in individuals without prior immunity [13]. The vaccine vector expressed Gag, Pol and Env proteins from multiple HIV clades [13]. Vaccine was given at two increasing doses, as well as a placebo. T–cell responses to antigens in the vaccine were measured via the ICS assay [4, 13]. The cytokines measured were IFNg (Interferon– γ), IL2 (Interleukin–2), TNFa (Tumor necrosis factor– α) and IL4 (Interleukin 4) [4]. The sample size consisted of 20 vaccine and four placebo recipients. Statistical analysis of the original positivity calls is described in the original publication [13].

Data Import, Preprocessing, and Gating

The gated ICS assay data was imported into R from the original flowJo workspaces (version 6, TreeStar Inc, Ashland, OR) using the BioConductor tool, *flowWorkspace* (v 1.1.6) and ncdfFlow (v 1.1.4). Data were preprocessed using the flowJo–defined compensation matrices and data transformations extracted from the workspace file, and gated using methods from the flowCore package (v 1.19.2) to extract counts of cytokine positive and negative T–cells for each sample and stimulation [14].

Statistical Analysis of Responder and Non-responder calls

Below, we summarize the methods for statistical analysis of responder and non-responder calls in the published trial, as well as the methods compared in this paper.

Statistical Analysis in the Published Trial

The methodology for statistical analysis and calling responders and non-responders in the original vaccine trial is described in the original publication [13]. In general, a participant is called a "responder" to an antigen stimulation if, for a given cytokine, the number of cytokine-positive T-cells

in the antigen–stimulated sample is significantly greater (for some statistical measure of significance) than the number of cytokine–positive T-cells for the negative control (unstimulated) sample from the same individual. In the original trial, significance was measured via one–sided Fisher's exact test for each participant and cytokine, comparing stimulated against unstimulated samples from that individual. A discrete Bonferroni adjustment for multiple comparisons was applied, and stimulations with an adjusted p–value below $\alpha=0.00001$ were called positive.

Statistical Analysis of Responder and Non-responder Calls for Direct Comparison Against the Bayesian Mixture Model Approach

Positivity calls for vaccine responders and non–responders depend upon the selection of an appropriate threshold. Therefore, to compare different methods of analysis, comparable thresholds for positivity must be selected for the methods. Our mixture modelling approach is fit within each stimulation, and we make positivity calls based on a false discovery rate calculated across individuals, within each stimulation, whereas the originally published analysis makes multiple testing adjustments within individuals, across cytokines. In order to have comparable response rates, we reanalyzed the ICS data using Fisher's one–sided exact test (as described in the original publication) but made positivity calls based on the false discovery rate computed across individuals within each stimulation.

Two Competing Beta-Binomial Models

Our approach to modelling an individual's response to vaccine using ICS data takes a Bayesian approach. We model all observations (individuals) simultaneously for each combination of cytokine and stimulation (including the unstimulated samples). For a given cytokine, we let n_s be the number of cytokine–positive cells in the stimulated sample, N_s the total number of cells in the stimulated sample, and n_u , N_u , the number of cytokine–positive and total number of cells in the unstimulated sample, respectively. Note that usually, $N_u \neq N_s$. The observed count data \mathbf{y} is a matrix of size $4 \times P$, where P is the number of participants. For the i'th individual $\mathbf{y}_i = \langle N_{si}, n_{si}, N_{ui}, n_{ui} \rangle$, and can be represented as the following contingency table:

Table 1: 2 x 2 contingency table of counts for cytokine positive and cytokine negative events between stimulated and unstimulated conditions

	Cytokine		
	Negative	Positive	
Stimulated	$N_{si} - n_{si}$	n_{si}	
Unstimulated	$N_{ui} - n_{ui}$	n_{ui}	

The positive cell counts for stimulated and unstimulated samples from the same individual are

modelled as:

if
$$p_{si} = p_{ui} \equiv p_{0i}$$
; $n_{si} \sim \text{Bin}(N_{si}, p_u)$; $n_{ui} \sim \text{Bin}(N_{ui}, p_{ui})$ (1)

if
$$p_{si} > p_{ui}$$
; $n_{si} \sim \text{Bin}(N_{si}, p_{si})$; $n_{ui} \sim \text{Bin}(N_{ui}, p_{ui})$ (2)

Where p_{si} and p_{ui} are the unobserved proportions. Equation (1) represents the *null* hypothesis where there is no difference between the stimulation and the control. Equation (2) represents the alternate hypothesis where the cytokine response is stronger in the stimulation than in the control. We place a common Beta prior on the p_{si} and p_{ui} across individuals, as shown:

$$p_{si} \sim \text{Beta}(\alpha_s, \beta_s)$$
 (3)

$$p_{ui} \sim \text{Beta}(\alpha_u, \beta_u)$$
 (4)

If $p_{si} = p_{ui}$ we assume that $\alpha_s = \alpha_u$ and $\beta_s = \beta_u$, thus sharing the hyper-parameters between the null and alternative model for the unstimulated samples, such that β_u , α_u hyper-parameters are equal for both the stimulated and unstimulated models. Given this formulation, the posterior probability of the data given that it is generated by model (1), is:

$$\Pr(y_i|\alpha_u,\beta_u) = \binom{N_{si}}{n_{si}} \binom{N_{ui}}{n_{ui}} \frac{B(n_{si} + n_{ui} + \alpha_u, N_{si} - n_{si} + N_{ui} - n_{ui} + \beta_u)}{B(\alpha_u,\beta_u)}$$
(5)

with marginal log-likelihood:

$$\mathcal{L}(\alpha_u, \beta_u | \mathbf{y}) = \sum_{i=1}^{P} \left[\log \binom{N_{si}}{n_{si}} + \log \binom{N_{ui}}{n_{ui}} + \log \left(\frac{N_{ui}}{n_{ui}} \right) + \log \left(\frac{N_{ui}}{n_{ui}} \right) + \log \left(\frac{N_{ui}}{n_{ui}} + \alpha_u, N_{si} - n_{si} + N_{ui} - n_{ui} + \beta_u \right) \right] - P \log \left(\frac{N_{ui}}{n_{ui}} \right)$$
(6)

Thus, in the case of no response to stimulation, the counts for the stimulated and unstimulated samples are modelled as draws from the same "unstimulated" Beta-binomial distribution. Note that the unobserved parameters, p_{si} , p_{ui} have been integrated out to give the marginal log-likelihood.

If the data is generated by model (2), the posterior probability of the data is given by:

$$\begin{split} \Pr(y_{i} | \alpha_{u}, \beta_{u}, \alpha_{s}, \beta_{s}) &= \binom{N_{ui}}{n_{ui}} \binom{N_{si}}{n_{si}} \frac{\mathbf{B}(n_{ui} + \alpha_{u}, N_{ui} - n_{ui} + \beta_{u})}{\mathbf{B}(\alpha_{u}, \beta_{u})} \frac{\mathbf{B}(n_{si} + \alpha_{s}, N_{si} - n_{si} + \beta_{s})}{\mathbf{B}(\alpha_{s}, \beta_{s})} \cdot \\ & \int_{p_{ui}=0}^{1} \left(\frac{1}{\mathbf{B}(n_{ui} + \alpha_{u}, N_{ui} - n_{ui} + \beta_{u})} p_{ui}^{n_{ui} + \alpha_{u} - 1} (1 - p_{ui})^{N_{ui} - n_{ui} + \beta_{u} - 1} \right) \left(\mathbf{I}_{1 - \mathbf{p}_{ui}} (N_{s}^{i} - n_{s}^{i} + \beta_{s}, n_{s}^{i} + \alpha_{s}) \right) dp_{ui} \\ & \int_{p_{ui}=0}^{1} \left(\frac{1}{\mathbf{B}(\alpha_{u}, \beta_{u})} p_{ui}^{\alpha_{u} - 1} (1 - p_{ui})^{\beta_{u} - 1} \right) \left(\mathbf{I}_{1 - \mathbf{p}_{ui}} (\beta_{s}, \alpha_{s}) \right) dp_{ui} \end{split}$$

(7)

with marginal log-likelihood:

$$\mathcal{L}(\alpha_{s}, \alpha_{u}, \beta_{s}, \beta_{u} | \mathbf{y}) = -P \log \left(\mathbf{B}(\alpha_{u}, \beta_{u}) \right) - P \log \left(\mathbf{B}(\alpha_{s}, \beta_{s}) \right) +$$

$$\sum_{i=0}^{P} \left\{ \log \binom{N_{ui}}{n_{ui}} + \log \binom{N_{si}}{n_{si}} + \log \left(\mathbf{B}(n_{ui} + \alpha_{u}, N_{ui} - n_{ui} + \beta_{u}) \right) + \log \left(\mathbf{B}(n_{si} + \alpha_{s}, N_{si} - n_{si} + \beta_{s}) \right) +$$

$$\log \left[\int_{p_{ui}=0}^{1} \left(\frac{1}{\mathbf{B}(n_{ui} + \alpha_{u}, N_{ui} - n_{ui} + \beta_{u})} p_{ui}^{n_{ui} + \alpha_{u} - 1} (1 - p_{ui})^{N_{ui} - n_{ui} + \beta_{u} - 1} \right) \right.$$

$$\left. \left(\mathbf{I}_{1-\mathbf{p}_{ui}} (N_{si} - n_{si} + \beta_{s}, n_{si} + \alpha_{s}) \right) dp_{ui} \right]$$

$$\left. - \log \left[\int_{p_{ui}=0}^{1} \left(\frac{1}{\mathbf{B}(\alpha_{u}, \beta_{u})} p_{ui}^{\alpha_{u} - 1} (1 - p_{ui})^{\beta_{u} - 1} \right) \right.$$

$$\left. \left(\mathbf{I}_{1-\mathbf{p}_{ui}} (\beta_{s}, \alpha_{s}) \right) dp_{ui} \right] \right\}$$

The ratio of integrals in (7) accounts for the different normalizing constants due to the constraints $p_{si} > p_{ui}$ on the prior and the posterior distributions. We call this the *constrained* model.

The term $I_{1-p_{ui}}(\beta_s, \alpha_s) = 1 - I_{p_{ui}}(\alpha_s, \beta_s) = Pr(p_{si} > p_{ui}; \alpha_s, \beta_s)$, is just the CDF of Beta distribution with parameters α_s , β_s , leaving a 1-dimensional integration for the ratio of normalizing constants.

Without constraints, $(p_s \neq p_u)$, the ratio of integrals for the normalizing constant in (7) is dropped. We call this the *unconstrained* model.

1.1 The Mixture of Beta-Binomials

Although we have specified the two models for the data, we do not know which observation was generated by which model. Clearly, not all individuals are expected to exhibit an immune response to a stimulation. Any individual observation, y_i , could either be generated by model (1) or by model (2). We capture this uncertainty with a mixture framework of the two competing beta-binomial models. The likelihood for the mixture is given by:

$$L(\alpha_{s}, \beta_{s}, \alpha_{u}, \beta_{u}, \pi_{k} | \mathbf{y}) = \prod_{i=1}^{P} \left[\pi_{1} f_{1}(y_{i} | \theta_{1}) + \pi_{2} f_{2}(y_{i} | \theta_{2}) \right],$$

$$\sum_{k=1}^{2} \pi_{k} = 1$$
(9)

Where $\theta_1 = \{\alpha_u, \beta_u\}, \theta_2 = \{\alpha_u, \beta_u, \alpha_s, \beta_s\}, \pi_1$ is the fraction of observations exhibiting no response to stimulation, π_2 the fraction of observations exhibiting a response to stimulation, and $f_1 = Pr(y_i | \alpha_u, \beta_u), f_2 = Pr(y_i | \alpha_u, \beta_u, \alpha_s, \beta_s)$ from (5) and (7), above.

The unobserved component memberships are treated as missing data and modelled as random variables $\mathbf{z}_i = \{z_{i1}, (1-z_{i1})\}$

$$z_{ik} = \left\{ \begin{array}{ll} 1 & \text{if observation } i \text{ is from the } k\text{'th model (component)} \\ 0 & \text{otherwise} \end{array} \right.$$

Each \mathbf{z}_i follows an independent multinomial distribution with one trial and parameters $\pi = \{\pi_1, 1 - \pi_1\}$. Given the z_i 's, the complete data log–likelihood is:

$$\mathcal{L}_c(\alpha_s, \beta_s, \alpha_u, \beta_u, \pi_k | \mathbf{y}, \mathbf{z}) = \sum_{i=1}^P \sum_{k=1}^2 z_{ik} \left[\log \pi_k + \log f_k(y_i | \theta_k) \right]$$
(10)

In this form, we use the expectation–maximization (EM) algorithm [15] to fit the model.

E-step

Given the model parameters $\Psi = \{\alpha_u, \beta_u, \alpha_s, \beta_s, \pi_k\}$, and the data \mathbf{y} , we estimate the unobserved component memberships, $\mathbf{Z_i}$ by computing the conditional expectation of the $\mathbf{Z_i}$'s, $\mathbb{E}_{\Psi}(\mathbf{Z_i}|\mathbf{y}_i)$:

$$\tilde{z}_{ik} = \frac{\pi_k f_k(\mathbf{y}_i | \theta_k)}{\sum\limits_{k=1}^{2} \pi_k f_k(\mathbf{y}_i | \theta_k)}$$
(11)

M-step

Finally, given the \tilde{z}_{ik} , we update the estimates of the model parameters to maximize the conditional expectation of the complete–data log–likelihood. The mixing proportions are given by:

$$\hat{\pi}_k = \frac{\sum_i^P \tilde{z}_{ik}}{n} \tag{12}$$

There is no closed form for the model hyper–parameters, α_u , β_u , α_s , β_s , and they are estimated via numerical optimization using R's *optim* function. For this purpose they are re–parameterized as $\mu_u = \frac{\alpha_u}{\alpha_u + \beta_u}$ and $S = \alpha_u + \beta_u$ (likewise for the α_s , β_s), corresponding to the mean and sample size of the prior distributions.

Initialization

We initialize the z_{ik} 's using Fisher's exact test to assign each observation to either the $p_{si} = p_{ui}$ or $p_{si} > p_{ui}$ components. We then use the \hat{z}_i 's to initialize the hyper–parameters to their method–of–moments estimates:

$$\hat{\alpha} = \hat{\mu} \left(\frac{\hat{\mu}(1-\hat{\mu})}{\hat{\sigma}^2} - 1 \right) \tag{13}$$

$$\hat{\beta} = \left(1 - \hat{\mu} \left(\frac{\hat{\mu}(1-\hat{\mu})}{\hat{\sigma}^2} - 1\right)$$
 (14)

Where $\hat{\mu}$ and $\hat{\sigma}^2$ are the sample mean and sample variance estimates, given the z_{ik} 's.

Generalization to Multiple Cytokines and Polyfunctionality with the Multinomial Dirichlet

The model can be generalized to handle multiple cytokines in a single stimulation, in order to assess polyfunctional cytokine responses of T-cells. We use the Multinomial-Dirichlet family of distributions to model counts of events in two *different* 2x2 contingency tables. Here we consider only the *unconstrained* case ($p_s \neq p_u$). The observed data can be represented in the following way:

Where the vector of observed counts for individual i in the stimulated or unstimulated sample is denoted: $\bar{n}_{\{s,u\}i} = \{n_{\{s,u\}ij}\}; j \in \{1\dots 4\}$, and j indexes the cells of the appropriate contingency table shown in Table 2. The counts are modelled as draws from different multinomial distributions:

if
$$\bar{p}_{si} = \bar{p}_{ui}$$
; $\bar{n}_{ui} \sim \mathcal{M}(\bar{p}_{ui}, N_{ui}); \bar{n}_{si} \sim \mathcal{M}(\bar{p}_{ui}, N_{si})$ (15)

if
$$\bar{p}_{si} \neq \bar{p}_{ui}$$
; $\bar{n}_{ui} \sim \mathcal{M}(\bar{p}_{ui}, N_{ui}); \bar{n}_{si} \sim \mathcal{M}(\bar{p}_{si}, N_{si})$ (16)

with Dirichlet priors on the proportions:

$$\bar{p}_{si} \sim \text{Dir}(\bar{\alpha}_s); \bar{p}_{ui} \sim \text{Dir}(\bar{\alpha}_u)$$
 (17)

Table 2: Contingency tables for counts of cells expressing two cytokines between stimulated and unstimulated conditions. $n_{\{s,u\}j}$ denotes observed counts for stimulated or unstimulated table cell j, and individual i

Stimulated		U	Unstimulated		
	Cytokine A			Cytokine A	
	Negative	Positive		Negative	Positive
Cytokine B			Cytokine B		
Negative	n_{si1}	n_{si2}	Negative	n_{ui1}	n_{ui2}
Positive	n_{si3}	n_{si4}	Positive	n_{ui3}	n_{ui4}

For the null component, where $\bar{p}_s = \bar{p}_u$ the marginal likelihood is given by:

$$L(\bar{n}_s, \bar{n}_u, N_s, N_u | \bar{\alpha}_u) = \prod_{i=0}^{P} \frac{B_j(\bar{\alpha}_u + \bar{n}_{ui} + \bar{n}_{si})}{B_j(\bar{\alpha}_u)} \cdot \frac{N_{si}!}{\prod_{j=1}^{J} n_{sij}!} \cdot \frac{N_{ui}!}{\prod_{j=1}^{J} n_{uij}!}$$
(18)

Where B_j is the j-dimensional Beta function: $\frac{\prod_{j=1}^J \Gamma(\alpha_j)}{\Gamma(\sum \alpha_j)}$. The marginal likelihood for a component where $p_{sj} \neq p_{uj}$ for all j, is given by:

$$L(\bar{n}_{s}, \bar{n}_{u}, N_{s}, N_{u} | \bar{\alpha}_{u}, \bar{\alpha}_{s}) = \prod_{i=0}^{P} \frac{B_{j}(\bar{\alpha}_{u} + \bar{n}_{ui})B_{j}(\bar{\alpha}_{s} + \bar{n}_{si})}{B_{j}(\bar{\alpha}_{s})B_{j}(\bar{\alpha}_{u})} \cdot \frac{N_{si}!}{\prod_{j=1}^{J} n_{sij}!} \cdot \frac{N_{ui}!}{\prod_{j=1}^{J} n_{uij}!}$$
(19)

Without loss of generality, if only some p_i are different between stimulated and unstimulated samples, the appropriate components of α_i can be substituted in the calculation of the likelihood

The marginal likelihood of the non-responding component, given in eq (18) can be rewritten

$$l(\bar{\alpha}_{u}|D) = \sum_{i} \left[\log \Gamma(\sum_{j} \alpha_{uj}) - \log \Gamma(\sum_{j} \alpha_{uj} + n_{uij} + n_{sij}) + \sum_{j} (\log \Gamma(\alpha_{uj} + n_{uij} + n_{sij}) - \log \Gamma(\alpha_{uj})) + \log \Gamma(N_{si} + 1) + \log \Gamma(N_{ui} + 1) - \left[\sum_{j} (\log \Gamma(n_{sij} + 1) + \log \Gamma(n_{uij} + 1)) \right] \right]$$

$$(20)$$

The gradient is given by:

$$\frac{\partial l}{\partial \alpha_{uj}} = \sum_{i} \Psi(\sum_{j} \alpha_{uj}) - \Psi(\sum_{j} \alpha_{uj} + n_{uij} + n_{sij}) + \Psi(\alpha_{uj} + n_{uij} + n_{sij}) - \Psi(\alpha_{uj}) \quad (21)$$

and the Hessian is:

$$j \neq k; \quad \frac{\partial^{2}l}{\partial\alpha_{uj}\alpha_{uk}} = \sum_{i} \Psi'(\sum_{j} \alpha_{uj}) - \Psi'(\sum_{j} \alpha_{uj} + n_{uij} + n_{sij})$$

$$j = k; \quad \frac{\partial^{2}l}{\partial\alpha_{uj}^{2}} = \sum_{i} \Psi'(\sum_{j} \alpha_{uj}) - \Psi'(\sum_{j} \alpha_{uj} + n_{uij} + n_{sij}) + \Psi'(\alpha_{uj} + n_{uij} + n_{sij}) - \Psi'(\alpha_{uj})$$
(23)

The marginal likelihood of the responding component in eq 19 can be rewritten as:

$$\sum_{i} \left[\sum_{j} \left[\log \Gamma(\alpha_{uj} + n_{uij}) + \log \Gamma(\alpha_{sj} + n_{sij}) - \log \Gamma(\alpha_{uj}) - \log \Gamma(\alpha_{sj}) \right. \right.$$

$$\left. - \log \Gamma(n_{sij} + 1) - \log \Gamma(n_{uij} + 1) \right] + \log \Gamma(\sum_{j} \alpha_{sj}) + \log \Gamma(\sum_{j} \alpha_{uj})$$

$$\left. - \log \Gamma(\sum_{j} \alpha_{uj} + n_{uij}) - \log \Gamma(\sum_{j} \alpha_{sj} + n_{sij}) + \log \Gamma(N_{si} + 1) + \log \Gamma(N_{ui} + 1) \right]$$

$$\left. - \log \Gamma(\sum_{j} \alpha_{uj} + n_{uij}) - \log \Gamma(\sum_{j} \alpha_{sj} + n_{sij}) + \log \Gamma(N_{si} + 1) + \log \Gamma(N_{ui} + 1) \right]$$

$$\left. - \log \Gamma(\sum_{j} \alpha_{uj} + n_{uij}) - \log \Gamma(\sum_{j} \alpha_{sj} + n_{sij}) + \log \Gamma(N_{si} + 1) + \log \Gamma(N_{ui} + 1) \right]$$

with gradient:

$$\frac{\partial l}{\partial \alpha_{\phi j}} = \sum_{i} \Psi(\alpha_{\phi j} + n_{\phi ij}) - \Psi(\alpha_{\phi j}) + \Psi(\sum_{j} \alpha_{\phi j}) - \Psi(\sum_{j} \alpha_{\phi j} + n_{\phi ij}) : (\phi \in s, u) \quad (25)$$

and Hessian:

$$\frac{\partial^2 l}{\partial \alpha_{\phi j}^2} \stackrel{k=j}{=} \sum_i \Psi'(\alpha_{\phi j} + n_{\phi ij}) - \Psi'(\alpha_{\phi j}) + \Psi'(\sum_j \alpha_{\phi j}) - \Psi'(\sum_j \alpha_{\phi j} + n_{\phi ij})$$
 (26)

$$\frac{\partial^2 l}{\partial \alpha_{\phi j} \alpha_{\phi k}} \stackrel{k \neq j}{=} \sum_i \Psi'(\sum_j \alpha_{\phi j}) - \Psi'(\sum_j \alpha_{\phi j} + n_{\phi ij})$$
(27)

Mixture Model Complexity

We may wish to detect any of $2^3 = 8$ different possible scenarios where the proportion of events in corresponding cells of the contingency tables are either equal or unequal between stimulated and unstimulated conditions. Such a model would have 8 components and 55 parameters. However, if we recognize that the models can be nested, i.e. that parameters can be shared across components with similar outcomes, then the number of parameters can be reduced to 19, and further to 15 if we only consider components where any one cell of the tables differs between stimulated and unstimulated conditions. This is outlined in Table 3.

Table 3: Nesting of models and parameter counts. Each row is a model component. The three columns correspond to cells two, three, and four of the contingency tables shown in Table 2. An open circle at a position indicates that the component models $p_{sj} = p_{uj}$, and a filled circle indicates that the component models $p_{sj} \neq p_{uj}$. The number of additional parameters that need to be estimated by including each additional component in the mixture model is in the fourth column (number of parameters for proportions + number of parameters for component weights).

Cell of Table			
cell 2	cell 3	cell 4	# of parameters
0	0	0	6+1
0	0	•	2+1
0	•	0	2+1
•	0	0	2+1
0	•	•	0+1
•	•	0	0+1
•	0	•	0+1
•	•	•	0

Simulation Studies

We examined the performance of the constrained $(p_s > p_u)$ and unconstrained $(p_s \neq p_u)$ beta-binomial mixture models via simulations. Using hyper parameters estimated from the model fit of the constrained model to data from Gag1 stimulated CD4-positive, IL2 expressing T-cells on day 28 from the HVTN054 data set, we simulated data from the constrained model with 500 observations, a response rate of 40%, an N of 10K, 20K, 30K, 50K, 75K, 100K, and 150K events, and ten independent realizations for each N. The constrained model was fit to this data and the sensitivity and specificity of the model's ability to correctly identify observations from the "responder" and "non-responder" components was evaluated through ROC curve analysis and compared against Fisher's exact test. (Figure 1). The nominal vs observed false discovery rate was also examined for the models and for Fisher's exact test, as well as the accuracy of the estimated prior distributions. In order to assess whether the unconstrained model could be used to effectively fit data from a constrained model and benefit from faster computations using exact closed form expressions rather than Monte-Carlo integration, we fit the same data simulated from the constrained model using the unconstrained model (Figure 1.

To assess the sensitivity of the model to deviations from model assumptions, we repeated the simulations with the cell proportions drawn from truncated normal distributions on (0,1), rather than beta distributions. The means and variances of the truncated normal distributions were set to the MLE estimates of the beta distributions defined by the α , β hyper parameters estimated from the HVTN054 data set (Figure 2).

Results

Simulations Show MIMOSA Outperforms Fisher's Exact Test Event When Model Assumptions are Violated

We ran simulation studies to assess the performance of the constrained and unconstrained models, as described in the Methods. We found that the both the constrained and unconstrained MIMOSA models out–performed Fisher's exact test with respect to sensitivity and specificity at all values of N, and that the false discovery rate observed for the mixture model more closely reflected the nominal false discovery rate than Fisher's exact test (Figure 1). Furthermore, both models gave reasonable estimates of the true hyper–parameters (Figure 1).

Since the constrained model relies on Monte–Carlo integration, which can be computationally costly, to estimate the normalizing constant in the likelihood calculation, we examined whether the unconstrained model could be used to accurately fit data generated from the constrained model. We found that the unconstrained model performed as well as the constrained model when fitting data generated from the constrained model (Figure 1).

In order to assess the sensitivity of the model to deviations from model assumptions, we fit the unconstrained model to simulated data where the proportions were drawn from truncated normal distributions (Figure 2). Even in these circumstances the unconstrained MIMOSA model outperformed Fisher's exact test and in fact performed about as well as the constrained model fit to constrained data.

MIMOSA Outperforms Fisher's Exact Test in Real-World Vaccine Trial Data from HVTN054

We tested our method on ICS data from HVTN054, a safety and immunogenicity vaccine trial for a replication defective Adenovirus vector HIV vaccine. The data set consisted of 48 individuals who received two doses of vaccine or placebo, and ICS time points for these individuals were available at day 0 and day 28 after vaccination. Responders for each cytokine and stimulation combination were classified using Fisher's exact test and the constrained MIMOSA model at the 1% FDR level, then the response rate within the control and treatment groups were calculated for both methods. Figure 3 shows the response rates for a subset of the ICS data (Gag2 and Pol2 stimulations, for the IL2 and IFNg cytokines, at days 0 and 28, in both CD4 and CD8 T–cell subpopulations). On day 0, we found that the response rates from the MIMOSA model were zero, and equal to response rates computed from Fisher's exact test across all treatment and control groups. The same was true for the response rate in the control groups at day 28. In the treatment groups, at day 28, the response rates for the MIMOSA model were generally higher or equal to those for Fisher's exact test, demonstrating the increased sensitivity and specificity of our method.

The MIMOSA framework generates predictions for individual samples. To better understand the differences between the predictions from the MIMOSA model and Fisher's exact test, we examined the model fit from Gag2 stimulated CD4+ T-cells at day 28 producing IFNg (Figure 4). We

see that two individuals are predicted as responders by the MIMOSA model which are not detected by Fisher's exact test. Both individuals have a larger proportion of IFNg producing antigen specific T-cells in antigen stimulated than unstimulated samples, as measure by the MAP or ML estimates of the proportions (Figure 4).

Testing Multifunctional Response via the Multinomial-Dirichlet model

We applied the Multinomial–Dirichlet extension of MIMOSA to the HVTN054 trial data to identify observations with cells expressing polyfunctional cytokine profiles. As a proof of principle we limited the model to the two–component case (Figure 3), with one component for the null model where we expect no difference between stimulations, and one component for the alternative model where we expect a difference between stimulations. We fit the model to the IL4 and TNFa cytokines for Env3 and Env1 stimulated CD4 T–cells from the day zero and day 28 time points (Figures 5 and 6).

The day 28 stimulations show that the posterior probability of response is generally higher for the multivariate MIMOSA model than for the univariate models, and the day zero stimulation (Figure 5) shows that the multivariate model has fewer false positives than the univariate MIMOSA model fit to distinct cytokine combinations. We compared multivariate MIMOSA against Fisher's exact test. Fisher's exact test was used to test whether the number of cells expressing a specific combination of cytokines (i.e TNFa+/IL4-, TNFa+/IL4+, and so forth) was significantly higher in the stimulated samples than in the unstimulated samples. In this comparison, Fisher's exact test detected only one of the 8 responders detected by the MIMOSA model in the Env3 stimulated, day 28 data, and detected two false positives in the day zero data. Fisher's exact test also failed to identify four of the 12 responders identified by the MIMOSA model in the day 28, Env1 stimulated data.

Discussion

Conclusions

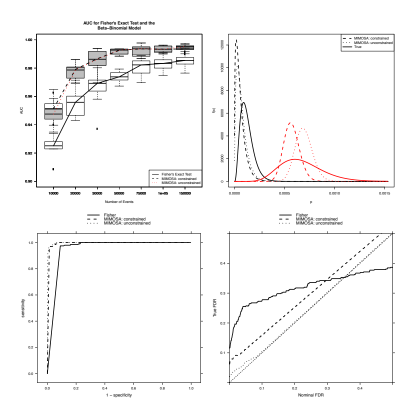


Figure 1: Performance of the constrained and unconstrained Beta-binomial mixture model vs Fisher's exact test on simulated data. Data were simulated from a constrained model with hyper-parameters estimated from a real data set of Gag1 stimulated, CD4+, IL2 expressing T-cells on day 28 from the HVTN054 trial. For total cell counts from 10,000 to 150,000, we simulated ten data sets each of 500 observations with a response rate of 40%. The performance, measured by the AUC (area under the curve), of the constrained and unconstrained Beta-binomial mixture model compared to Fisher's exact test is shown in the first panel, as a function of increasing number of cells. The beta distributions for the estimated and true hyper parameters are shown in the second panel for one simulated data set, with N=150,000 events. The ROC curve for Fisher's exact test and the constrained and unconstrained Beta-binomial model for the same simulation are shown in the third panel. The observed vs expected false discovery rate for Fisher's exact test and the constrained and unconstrained Beta-binomial model are shown in the fourth panel.

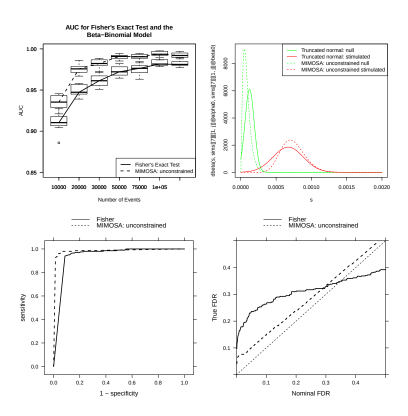


Figure 2: Sensitivity to departures from model assumptions. We generated data from variant of the constrained model $(p_s > p_u)$ where the proportions were simulated from truncated normal distributions on (0,1), rather than from beta distributions. The mean and variance of the normal distributions was given by $\mu = \alpha/(\alpha + beta)$, $\sigma^2 = (\alpha\beta)/((\alpha + beta)^2(\alpha + \beta + 1))$, where α, β are the Beta-prior hyper parameters for the MIMOSA model estimated from real data (s,u) subscripts omitted for brevity). The AUC for the unconstrained MIMOSA model and Fisher's exact test as a function of event count are shown in the first panel. The beta distributions for the estimated and true hyper parameters are shown in the second panel. The ROC curves for one data set and the observed and true false discovery rates for the model and Fisher's exact test are shown in the third and fourth panels.

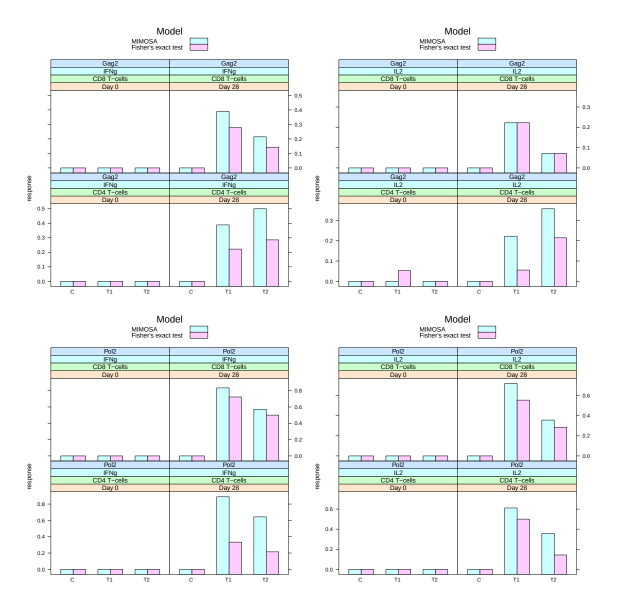


Figure 3: Comparison of MIMOSA and Fisher's exact test for calling responders in ICS data. Response rates for IL2 and IFNg cytokine positivity in Gag2 and Pol2 stimulated samples at days 0 and 28 in the CD4 and CD8 T—cell subpopulations of the HVTN054 ICS data. In the treatment groups, response rates for the MIMOSA model are greater than or equal to those for Fisher's exact test at day 28 but are equal (and zero) at day 0 as expected. In the control groups, response rates are equal (and zero) between Fisher's exact test and the MIMOSA model at day 0 and at day 28, as expected.

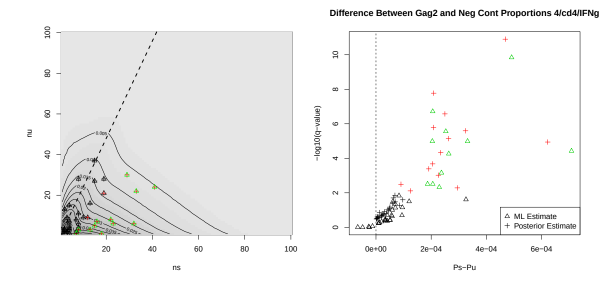


Figure 4: Likelihood surface and volcano plot for IFNg producing, CD4+ T-cells in Gag2 stimulated vs control samples on day 28. A significant difference between control and stimulated samples is called at the 1% FDR threshold (red) for the Beta-binomial model, and at 1% for FDR adjusted p-values from Fisher's exact test (green). The likelihood surface shows the observed counts from stimulated and unstimulated samples. The volcano plot shows the difference between the proportion of cytokine positive cells in the stimulated and unstimulated samples, for the maximum likelihood estimates of the proportions (triangles) and for the MAP estimates (crosses). The effect of shrinking the MAP estimates towards zero can be seen in the volcano plot.

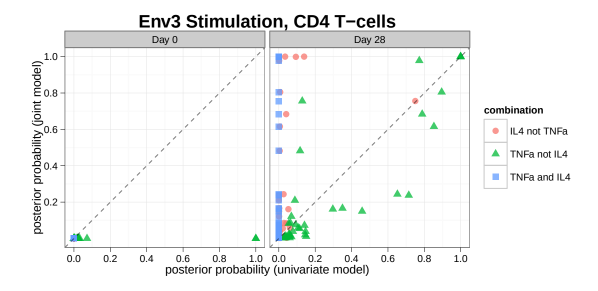


Figure 5: Posterior probabilities from the four–dimensional, Multivariate–Dirichlet, MIMOSA model, compared against posterior probabilities from univariate MIMOSA for each combination of the cytokines IL4 and TNFa, on days zero and 28 for Env–2 peptide stimulated CD4+ T–cells.

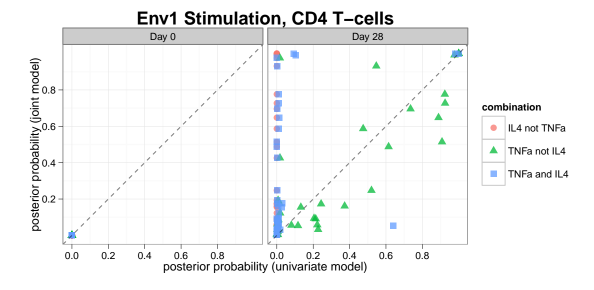


Figure 6: Posterior probabilities from the four–dimensional, Multivariate–Dirichlet, MIMOSA model, compared against posterior probabilities from univariate MIMOSA for each combination of the cytokines IL4 and TNFa, on days zero and 28 for Env–1 peptide stimulated CD4+ T–cells.

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