Accounting for correlations in competitive gene set test for improved interpretation of genome-scale data

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SUMMARY: Competitive gene set test is a widely used tool for interpreting high-throughput biological data, such as gene expression and proteomics data. It aims at testing categories of genes for enriched association signals in a list of genes inferred from genome-wide data. Most conventional enrichment testing methods ignore or do not properly account for the widespread correlations among genes, which, as we show, can result in inflated type I error rates and power loss. We propose a new framework, MEQLEA, for gene set test based on a mixed effects quasi-likelihood model, where the data are not required to be Gaussian. Our method effectively adjusts for completely unknown, unstructured correlations among the genes. It uses a score test approach and allows for analytical assessment of p-values. Compared to existing methods such as GSEA and CAMERA, our method enjoys robust and substantially improved control over type 1 error and maintains good power in a variety of correlation structure and association settings. We also present two real data analysis to illustrate our approach.

KEY WORDS:

1. Introduction

What is enrichment analysis? Why would people care about that?

Gene set test is a statistical framework of studying the association between a test set—a prior set of biologically related genes—and a set of genes that are significantly correlated with treatment or experimental design variables. A key task of gene expression analysis involves the detection of differentially expressed genes. Differential expression (DE) analysis evaluates each individual gene separately, and therefore it fails to provide insight into the relation between treatment variables and the prior gene set under study. Gene set test helps researchers better understand the underlying biological processes in terms of ensembles of genes.

What are the differences between self-contained and competitive test? And how does they work?

Depending on the definition of the null hypothesis, there are two types of gene set test: the *self-contained* test and the *competitive* test (Goeman and Bühlmann, 2007). A self-contained test examines a set of genes by a fixed standard without reference to other genes in the genome (see, for example, Goeman et al. (2004, 2005); Tsai and Chen (2009); Wu et al. (2010); Huang and Lin (2013)). A competitive test compares DE genes in the test set to those not in the test

set (Tian et al., 2005; Wu and Smyth, 2012; Yaari et al., 2013). Many methods, regardless of the type of test, perform a three-stage analysis (Khatri et al., 2012): on the first stage, a gene-level statistic is calculated for each gene in the whole genome to measure the association between the expression profiles and the experimental design variables; such genelevel statistic includes, among others, signal-to-noise ratio (Subramanian et al., 2005), ordinary t-statistic (Tian et al., 2005) or moderated t-statistic (Smyth, 2004), log fold change (Kim and Volsky, 2005) and Z-score (Efron, 2007). On the second stage, a set-level statistic is obtained by utilizing the gene-level statistics from the first stage and their membership with respect to the test set (i.e., whether the gene belongs to the test set). Examples of the set-level statistic are enrichment score (Subramanian et al., 2005), maxmean statistic (Efron and Tibshirani, 2007), and statistic derived from joint distribution of gene-level statistics (Yaari et al., 2013), to name a few. On the third stage, a p-value is assigned to the test set by comparing the set-level statistic to its reference distribution. The competitive gene set test is much more popular among genomic literatures (Goeman and Bühlmann, 2007; Gatti et al., 2010).

Independent gene set test

Many competitive gene set tests rely on independence of gene-

level statistics which further depends on independence among genes. Those tests are parametric or rank-based procedures that assume the gene-level statistics to be independent and identically distributed, or gene permutation procedures that generate the same approximate null for the set-level statistics. For example, PAGE (Kim and Volsky, 2005) conducts onesample z-test by comparing the mean of gene-level statistics (i.e., the mean of log fold changes) in the test set to a normal distribution under the null. The 2×2 contingency-table-based tests examine the significance of the test set by dichotomizing the outcomes of DE analysis and cross-classifying the genes according to whether they are indicated as DE and whether they are in the test set (see Huang et al. (2009) for a review and references therein). sigPathway (Tian et al., 2005) and "geneSetTest" in the limma package (Smyth, 2004) evaluate the set-level p-values by permuting gene labels. However, tests assuming independence of genes may result in inflated false discovery rate (Efron and Tibshirani, 2007; Goeman and Bühlmann, 2007; Gatti et al., 2010; Wu and Smyth, 2012; Yaari et al., 2013), as genes within a gene set are often coexpressed and function together.

Tests that account for inter-gene correlation

A handful of methods have been proposed to account for intergene correlation in competitive gene set test. One attempt is to evaluate the set-level statistic by permuting the biological sample labels (see, for example, Subramanian et al. (2005); Efron and Tibshirani (2007)). Permuting sample labels does not require an explicit understanding of the underlying correlation structure among genes and thus protects the test against such correlation. Since permuting sample labels is computationally inefficient, Zhou et al. (2013) proposed an analytic approximation to permutations for set-level score statistics, which preserves the essence of permutation gene set analysis with greatly reduced computational burden. However, an unavoidable problem arising from sample permutation approach is that it implicitly alters the null hypothesis being tested and it is therefore difficult to characterize the null and the alternative hypotheses (Goeman and Bühlmann. 2007; Khatri et al., 2012; Wu and Smyth, 2012). Another attempt is to use set-level statistic that directly includes intergene correlation estimated from the data. For example, CAM-ERA (Wu and Smyth, 2012) calculates a variance inflation factor (VIF) from sample correlation (after the treatment effect removed), and then incorporates it into their set-level statistics to account for inter-gene correlations. A recent extension to CAMERA, QuSAGE (Yaari et al., 2013), also used the same VIF to adjust for inter-gene correlations but includes the VIF in a different set-level statistic. The VIF is a crucial factor and valid estimation of it relies on the assumption that correlation between any two local statistics are almost the same as correlation between their corresponding expression profiles. This assumption has been demonstrated (??? a better word???) by simulation (Barry et al., 2008) for several gene-level statistics (e.g., t-statistic, Wald-type statistic for regressing expression on censored time-to-event data through a Cox proportional hazards model). However, this assumption holds only for the case where all the hypotheses are under the null (i.e., no gene is DE), and the correlation among gene-level statistics (e.g., t-statistics) can be badly estimated by sample

correlation when a fraction of genes are DE (Zhuo and Di, unpublished work).

What do we propose?

We propose a new framework for enrichment analysis that we will call Mixed Effects Quasi-Likelihood Enrichment Analysis (MEQLEA). Our strategy is to avoid the discrepancy between correlations among expression profiles and those among genelevel statistics caused by the presence of DE genes. Instead, we use differences in mean as gene-level statistics for a two group comparison experiment. We model the covariance of genelevel statistics by two variance components, one attributable to correlations among samples after treatment effect removed, and the other attributable to the DE effect associate with the treatment. The benefit of quasi-likelihood is that the data are not required to be Gaussian. Our method effectively adjusts for completely unknown, unstructured correlations among the genes. MEQLEA uses a score test approach and allows for analytical assessment of p-values. Compared to existing methods including GSEA and CAMERA, MEQLEA enjoys robust and improved control over type I error and maintains good power in a variety of correlation structure and association settings.

What is the plan of this paper?

The rest of the paper is organized as follows: in Section 2 we describe the methodology and then the simulation setup of MEQLEA, and summarize related existing methods; in Section 3 we present results from comparison of MEQLEA to other existing methods by simulation study, and illustrate the application of our method by two real data sets; in Section 4 we conclude and also specifies the future work.

2. Methods

We consider a gene expression (e.g. RNA-Seq or microarray) experiment, in which we compare the gene expression profiles of samples from two groups: a treatment group with n_1 samples referred to as "cases" and a control group with n_2 samples referred to as "controls" $(n_1, n_2 \ge 3)$. Suppose the expression profiles of a set of m genes are observed for each sample. An unknown subset of these genes are DE between cases and controls, with varying sign and magnitude of DE effects. The genes are also allowed to have (negatively or positively) correlated expression profiles. In enrichment analysis, we are interested in a pre-defined set of genes, for example, from a known pathway or given by a functional annotation term from a database such as KEGG (Kanehisa and Goto, 2000) or GO (Ashburner et al., 2000). Our goal is to test whether this known gene set is enriched with differential expression signals. Let G be an m-dimensional vector defining the gene set of interest, where $G_i = 1$ if and only if the i^{th} gene is in the set and $G_i = 0$ otherwise. Our analysis will condition on ${m G}$ and test if ${m G}$ is associated with enhanced DE effects. In the following sections, we will first construct a hierarchical model for the gene expression data incorporating possible correlations among the m genes, from which we will derive a quasi-likelihood model for the gene-level DE statistics jointly for all the genes. Based on this model, we will then present our enrichment test, and discuss its connections with CAMERA. Finally, we will describe our simulation studies used to evaluate our method. For the rest of **Methods**, our presentation of the method is conditional on G unless otherwise indicated.

2.1 MEQLEA

2.1.1 A hierarchical model for the gene expression data. We will start by presenting the hierarchical model for the observed gene expression data, which will incorporate the following features. Firstly, for a given sample, the expression profiles of different genes are allowed to be correlated. We further assume that the correlation structure is the same across samples. Secondly, different genes may have different baseline expression profiles, where "baseline" refers to the average among controls. Thirdly, for any given gene, its mean expression profile in the treatment group can be either higher, lower or the same compared to the control group, depending on whether the gene is up-regulated, down-regulated, or not DE. For the genes that are differentially expressed, their DE effects are modeled additively and are allowed to have heterogeneous signs and magnitudes. Finally, given a gene, and its DE effect, the expression profile is allowed to vary independently across samples, which captures measurement error and sample-level variability.

To present our model formally, we first introduce some notation. Let $n = n_1 + n_2$ be the total sample size. Let X be an n-dimensional known vector of 1's and 0's denoting the casecontrol membership of the samples, with $X_i = 1$ for a case and $X_i = 0$ for a control. Let **Y** be an n by m matrix representing the expression data, in which each row is the expression profile for a sample and Y_{ij} $(1 \le i \le n, 1 \le j \le m)$ is the expression profile of sample i at gene j. Let μ_j $(1 \leqslant j \leqslant m)$ be the baseline expression profile for gene j. μ_j 's are treated as nuisance parameters and as we will see later do not contribute to our analysis. Let $\Delta = (\Delta_1, \dots, \Delta_m)^T$ be a vector for the additive DE effects for the genes. Gene j is not DE if $\Delta_j = 0$, up-regulated if $\Delta_j > 0$ and down-regulated if $\Delta_j < 0$. We model Δ as a random effect, for which we will detail our assumptions later. Given μ_i and Δ_i , the mean expression profile for the control group and the treatment group are μ_j and $\mu_j + \Delta_j$, respectively. Given these means, the noise in the observed expression data for the i^{th} sample is denoted by the mean zero error vector $\epsilon_i = (\epsilon_{i1}, \dots, \epsilon_{im})^T$, $1 \leqslant i \leqslant n$. We assume $\epsilon := (\epsilon_1, \cdots, \epsilon_m)$ to be independent of Δ and to have mean zero. Without loss of generality, we also assume $Var(\epsilon_{ij}) = 1$ for all genes and samples. For a real gene expression data set typically not satisfying this assumption, we can standardize the data by each gene to ensure its empirical variance equals one before implementing our method. For the covariance structure of ϵ , we assume

$$\epsilon_{i_1} \text{ and } \epsilon_{i_2} \text{ are independent}, \quad i_1 \neq i_2,$$
(1)

$$Cov(\epsilon_i|\mathbf{G}) = \mathbf{C}, \ 1 \leqslant i \leqslant n,$$
 (2)

where C is an m by m inter-gene correlation matrix shared by all samples. C is generally unknown.

Putting these elements together, we obtain the following model for the expression data Y given X and G

$$Y_{ij} = \mu_j + X_i \cdot \Delta_j + \epsilon_{ij}, \tag{3}$$

for $1 \leqslant i \leqslant n, 1 \leqslant j \leqslant m$. G enters this model via Δ_j and possibly μ_j .

2.1.2 Assumptions on the DE effects Δ_j . Conditional on G, we assume that the Δ_j 's are mutually independent and

come from either of the two distributions, \mathcal{D}_1 and \mathcal{D}_2 , depending on whether $G_j = 0$ or 1. We denote the expected values of \mathcal{D}_1 and \mathcal{D}_2 by β_0 and $\beta_0 + \beta_1$, respectively, and their variances by σ_1^2 and σ_2^2 , respectively. It follows that

$$E(\Delta|G) = \beta_0 + \beta_1 G$$
, $var(\Delta|G) = \sigma_1^2 I_1 + \sigma_2^2 I_2$, (4)

where I_1 and I_2 are diagonal matrices of dimension m with 0's and 1's on their diagonals. The 1's in the diagonal of I_1 correspond to the genes with $G_j = 1$ and those for I_2 to the genes with $G_j = 0$.

Aside from the conditions in Equation (4) on the first two moments, we do not impose any specific distributional assumptions such as normality on Δ . For example, the distribution of a given Δ_j can put positive mass on zero, which allows for the highly likely event that some of the genes are not DE. To further motivate our general framework for Δ , we present a simple model included by Equation (4) as a special case. Suppose the m genes are independently sampled to be either DE or not. The probability for gene j to be DE is p_t if $G_j = 1$ or p_b if $G_j = 0$. For DE genes, their DE effects are sampled independently from a common distribution with mean μ_{δ} and variance σ_{δ}^2 . Under these assumptions,

$$E(\Delta_j | \mathbf{G}) = p_j \mu_\delta, \text{ var}(\Delta_j | \mathbf{G}) = p_j \sigma_\delta^2 + p_j (1 - p_j) \mu_\delta^2,$$
 (5)

where $p_j = p_t$ if $G_j = 1$ and $p_j = p_b$ if $G_j = 0$. It can be shown that this model is a special case of Equation (4).

2.1.3 Model for gene-level statistics. For each gene j, we consider the DE statistic (gene-level statistic???) given by

$$U_j = \frac{\sum_{i:X_i=1} Y_{ij}}{n_1} - \frac{\sum_{i:X_i=0} Y_{ij}}{n_2},$$
 (6)

which is sample mean difference in the expression profile between cases and controls. Given our assumption that ϵ_j has variance 1, U_j provides a DE metric for gene j. We will construct a quasi-likelihood model for $\boldsymbol{U} = (U_1, \dots, U_m)^T$ by deriving the mean and covariance structures of \boldsymbol{U} from the model for \boldsymbol{Y} described in Sections 2.1.1 and 2.1.2. We first observe that combining Equations (6) and (3) yields

$$U_j = \Delta_j + \eta_j$$
, where $\eta_j = \frac{1}{n_1} \sum_{i:X_i=1} \epsilon_{ij} - \frac{1}{n_1} \sum_{i:X_i=0} \epsilon_{ij}$. (7)

It can be shown based on Equations (1), (2) and (4) that

$$E(\boldsymbol{U}|\boldsymbol{G}) = \beta_0 + \beta_1 \boldsymbol{G},\tag{8}$$

$$\Sigma := \operatorname{var}(\boldsymbol{U}|\boldsymbol{G}) = \sigma_0^2 \boldsymbol{C} + \sigma_1^2 \boldsymbol{I}_1 + \sigma_2^2 \boldsymbol{I}_2, \tag{9}$$

where $\sigma_0^2 = 1/n_1 + 1/n_2$ is a known parameter. We note that in Equation (9), the covariance structure of U has three components, a component with C which accounts for the contribution from sample-level noise ϵ , and two additional components from the DE effect Δ . It is noteworthy that both the C component and the Δ components contribute to the variance of U_j 's, whereas only the C component contributes to the correlation among U_j 's.

2.1.4 The set-level test statistic. For a competitive gene set test, it is often unclear what the hypothesized null is and what is being tested (Barry et al., 2008; Wu and Smyth, 2012). In our approach, to detect patterns of the DE signals in the

gene set of interest that stand out compared with genes not in the set, we test $H_0: \mathscr{D}_0 = \mathscr{D}_1$ against $H_1: \mathscr{D}_0 \neq \mathscr{D}_1$. For example, for the special scenario given by Equation (5), this amounts to testing $p_b = p_t$ against $p_b \neq p_t$. To construct the test statistic, we focus on the part of the alternative space where $E(\mathscr{D}_0) \neq E(\mathscr{D}_1)$, or equivalently $\beta_1 \neq 0$. We first consider the less interesting case with uncorrelated genes, in which C equals I, an m-dimensional identity matrix. Under the quasi-likelihood model for U given in Section 2.1.3, the quasi-score statistic for β_1 has the form $S \propto \mathbf{G}^T(\mathbf{U} - \hat{\beta}_0 \mathbf{1}_m)$, where $\hat{\beta}_0 = \overline{U}$ is an estimate for β_0 and $\mathbf{1}_m$ is a m-dimensional vector of 1's. To perform a quasi-score test, one would divide S^2 by its estimated variance under H_0 and the assumption that C = I. The resulting test statistic is

$$T_{\mathrm{u}} = \frac{S^2}{\widehat{\mathrm{Var}}_{0,\boldsymbol{C}=\boldsymbol{I}}(S|\boldsymbol{G})} = \frac{[\boldsymbol{G}^T(\boldsymbol{U} - \hat{\beta}_0)\mathbf{1}_m]^2}{\boldsymbol{G}^T(\boldsymbol{I} - \boldsymbol{H})\boldsymbol{G}}, \quad (10)$$

where $\boldsymbol{H}=\frac{1}{n}\mathbf{1}_{m}\mathbf{1}_{m}^{T}$. The subscript "u" stands for "uncorrelated genes." For the case of interest when inter-gene correlation is present, \boldsymbol{C} is a non-trivial correlation (???Covariance) matrix. We will again form our test statistic based on S. But for the denominator of the statistic, the null variance of S will be evaluated under the quasi-likelihood model with non-trivial \boldsymbol{C} . By Equation (9), the variance of S is given by $\mathrm{Var}(S|\boldsymbol{G}) = \boldsymbol{G}^T(\boldsymbol{I} - \boldsymbol{H})\Sigma(\boldsymbol{I} - \boldsymbol{H})\boldsymbol{G}$. Note that $H_0: \mathscr{D}_0 = \mathscr{D}_1$ implies $\sigma_1^2 = \sigma_2^2$. Thus, under H_0 , $\Sigma := \mathrm{Var}_0(\boldsymbol{U}|\boldsymbol{G}) = \sigma_0^2\boldsymbol{C} + \sigma_1^2\boldsymbol{I}$, where $\sigma_0 = 1/n_1 + 1/n_2$ is known and σ_1^2 is an unknown parameter. To estimate σ_1^2 under H_0 , we observe that $\mathrm{var}_0(U_j) = \sigma_0^2 + \sigma_1^2$ and use $\hat{\sigma}_1^2 = \sum_{j=1}^m (U_j - \overline{U})^2/(m-1) - \sigma_0^2$. Therefore, assuming \boldsymbol{C} is known, we can obtain the MEQLEA test statistic given by

$$T = \frac{S^2}{\widehat{\text{Var}}_0(S|\mathbf{G})} = \frac{[\mathbf{G}^T(\mathbf{U} - \hat{\beta}_0 \mathbf{1}_m)]^2}{\mathbf{G}^T(\mathbf{I} - \mathbf{H})\hat{\mathbf{\Sigma}}(\mathbf{I} - \mathbf{H})\mathbf{G}}, \quad (11)$$

where $\hat{\Sigma} = (1/n_1 + 1/n_2)C + \hat{\sigma}_1^2 I$ is a null estimate of Σ . Under suitable regularity conditions, significance of the test could then be assessed by comparing T to a χ_1^2 distribution.

In practice, the inter-gene covariance matrix C is usually unknown. So we substitute C with \hat{C} , the empirical covariance matrix of the expression data after controlling for possible DE effects by centering the expression profiles of cases and controls separately around zero. Formally, \hat{C} is given by $\hat{C}_{jk} = \frac{1}{n} \sum_{i=1}^{n} (Y_{ij} - \alpha_{ij})(Y_{ik} - \alpha_{ik})$ where $\alpha_{ij} = \sum_{i':X_{i'}=X_i} Y_{i'j} / \sum_{i'=1}^{n} 1\{X_{i'}=X_i\}$ is the average expression profile at gene j for all samples from the same group (either treatment or control) as sample i. In real data sets, the number of genes, m, is usually much greater than the sample size n, in which case C is a high-dimensional parameter that cannot be efficiently estimated by \hat{C} . Interestingly, however, we find (note???) that the test statistic T relies not on the accurate estimation of the entire C, but only on three parameters involving C, which can be much more realistically estimated by a moderate sample size. To demonstrate this, we re-arrange the order of the rows and columns of C to allow the partition $C = \begin{bmatrix} C_{11} & C_{12} \\ C_{12}^T & C_{22} \end{bmatrix}$, where C_{11} is the correlation matrix for genes in the test set, C_{22} is that for gene in the

background set (i.e., the complement of the test set), and

 C_{12} is the cross-correlation matrix between the two classes of genes. (To be continued....)

$2.2\ Simulation\ Methods$

2.2.1 Simulation Setup. In this section, we present results from type I error and power simulations. Since a standardization procedure is required by MEQLEA for preprocessing data, we will simulate the standardized expression profiles for method illustration purpose.

Let Y_i be the expression profile of gene i and $\operatorname{Cov}(Y_{i_1},Y_{i_2})=\rho_{i_1,i_2}$ for any two genes i_1 and i_2 . We assume that $\operatorname{Cov}(Y_{i_1},Y_{i_2})=\rho_1$ if genes i_1 and i_2 are both from the test set (i.e., $G_{i_1}=G_{i_2}=1$), $\operatorname{Cov}(Y_{i_1},Y_{i_2})=\rho_2$ if they are both from the background set (i.e., $G_{i_1}=G_{i_2}=0$), and $\operatorname{Cov}(Y_{i_1},Y_{i_2})=\rho_3$ if i_1 is from the test set and i_2 is from the background set (i.e., $G_{i_1}=1,G_{i_2}=0$). We examine five different correlation structures, listed as follows:

- (a): $\rho_1 = \rho_2 = \rho_3 = 0$; that is, the genes are independent of each other.
- (b): $\rho_1 = 0.1$, $\rho_2 = \rho_3 = 0$; that is, only the genes in the test set are correlated.
- (c): $\rho_1 = \rho_2 = \rho_3 = 0.1$; that is, all genes are correlated, with an exchangeable correlation structure.
- (d): $\rho_1 = 0.1$, $\rho_2 = 0.05$, $\rho_3 = 0$; that is, genes are correlated within the test set and within the background set, but any two genes, one from the test set and the other from the background set, are independent.
- (e): $\rho_1 = 0.1$, $\rho_2 = 0.05$, $\rho_3 = -0.05$; that is, all genes are correlated, but the correlation between two genes depend on whether they belong to the test set or not.

The simulations run as follows: first, we generate an entire gene set containing m=500 genes, from which we randomly sample $m_1=100$ genes to represent those in the test set, and the remaining $m_2=400$ genes those in the background set; second, for gene $i=1,\ldots,m$, we set the DE size δ_i to be 1 and simulate the DE indicator Z_i from $\text{Binom}(1,p_i)$, where $p_i=p_t$ if gene i belongs to the test set and $p_i=p_b$ otherwise, and then the DE effect Δ_i is the product of Z_i and δ_i ; third, we set the "true" mean expression values $\mu_1=\mathbf{0}_m$ and $\mu_2=\Delta$, respectively, for the control and treatment groups; fourth, we simulate n_1 samples from $\text{MVN}(\mu_1, \Sigma)$ for the control group and n_2 samples from $\text{MVN}(\mu_2, \Sigma)$ for the treatment group, where the covariance $\Sigma = [\text{Cov}(Y_{i_1}, Y_{i_2})]_{m \times m}$ may be one of the correlation structures in (a)-(e).

We have mentioned in the Introduction part that the test statistics correlations among genes are not equal to their sample correlations when at least one gene is truly DE (under two sample t-test???). Therefore, if there are true DE genes in the entire gene set, approaches assuming almost equality of correlations among gene-level statistics and those among expression values may not perform well. To illustrate this point, we performed two groups of simulations for each of (a)-(e) correlation structures. In both type I error and power simulations, we set the DE probability to be 0% in group A_1 and 10% in group A_2 for genes in the background set. In the type I error simulation, we have $p_t = p_b$ under the null. In the power simulation, we considered four different alternative scenarios S_1 - S_4 : for genes in the test set, we set DE probability to be 5%, 10%, 15% and 20% in group A_1 , and 15%, 20%, 25%

and 30% in group A_2 . Table 1 summarizes the simulation setup for the two groups.

Table 1: Parameter setup for type I error and power simulations. S_0 repersents scenario for type I error simulation. S_1 - S_4 represent the four alternatives regarding power simulation.

Group	p_b		p_t					
	S_0	S_1	S_2	S_3	S_4			
A_1	0%	5%	10%	15%	20%			
A_2	10%	15%	20%	25%	30%			

 p_b : DE probablility for genes in the background set.

 p_t : DE probability for genes in the test set.

2.2.2 Other methods considered. We will compare MEQLEA to six existing gene set tests: GSEA (Subramanian et al., 2005), two versions of the CAMERA procedure —CAMERA-modt and CAMERA-rank (Wu and Smyth, 2012), SigPathway (Tian et al., 2005), MRSGE (Michaud et al., 2008), and QuSAGE (Yaari et al., 2013). Except SigPathway and MRSGE, all methods account for correlation among genes. GSEA calculates an enrichment score for the test set by examining the ranking (according to some metric, for example, signal-to-noise ratio) of its member genes, and determines the significance of the enrichment score by randomly permuting sample labels. CAMERA-modt uses moderated t-statistics (Smyth, 2004) as gene-level statistics and estimate a VIF to account for inter-gene correlations in the set-level statistic, and CAMERA-rank is the rank version of the CAMERA-modt. The SigPathway is slightly different from its original version of Tian et al. (2005) in that it uses moderated t-statistics rather than the ordinary t-statistics as gene-level statistics. MRSGE is also known as the rank version of SigPathway (Wu and Smyth, 2012). QuSAGE generates from t-test a probability density function (PDF) for each gene, combines the individual PDFs using convolution, and quantifies enrichment of the test set with a complete PDF.

For software implementation, GSEA is modified from the original R-GSEA script (http://software.broadinstitute.org/gsea/index.jsp) to accommodate single gene set test. CAMERA and MRSGE are implemented in the limma package (Smyth, 2005) in the Bioconductor project (Gentleman et al., 2004), QuSAGE is available in the Bioconductor package of the same name, and SigPathway is implemented by ourselves. Because GSEA and MEQLEA do not support linear models, the implementations are restricted to two-group comparisons.

In terms of type I error control and power, we expect some of the six tests to have different performances between group A_1 and A_2 simulations under one or more correlation structures.

3. Results

According to the simulation setup in Section 2.2.1, the test set is not enriched if DE probabilities are the same for genes in the test set and for those in the background set (i.e., $p_t = 0\%$

for group A_1 and $p_t=10\%$ for group A_2), in which case we evaluate the type I error. As to power, we set DE probability according to each of alternative scenarios S_1 - S_4 (see Table 1) and calculate the proportion of data sets for which a test would reject at a given level α . The results are based on 1,000 simulated data sets.

3.1 Type I error simulations

We report the type I error simulation results for group A_1 and A_2 simulations. Figure 1 shows the uniform quantile-quantile (QQ) plots of p-values for the seven approaches (MEQLEA, SigPathway, MRSGE, CAMERA-modt, CAMERA-rank, GSEA and QuSAGE) under each of the five correlation structures. The left column of Figure 1 shows the p-values of group A_1 simulations. Each plot, from top to bottom, corresponds accordingly to correlation structures (a)-(e). GSEA and MEQLEA hold the size of type I error rate correctly for all five correlation structures, with simulated p-values uniformly distributed on [0,1]. The two versions of CAMERA control type I errors correctly for correlation structures (a) and (b), yet they are too conservative for the case of (c) and anti-conservative for correlation structures (d) and (e). SigPathway and MRSGE procedures have well-calibrated type I error for correlation structures (a) and (c), but are anti-conservative the case of (b), (d) and (e). QuSAGE has good type I error control for only (b), and is too conservative for (a), (d) and (e), and anti-conservative for (c).

The right column of Figure 1 shows type I error rate of group A_2 simulations. MEQLEA continues to hold the size of type I error rate, whereas GSEA is skewed towards small p-values, under all five correlation structures. The two versions of CAMERA control type I error rate correctly for (a) where genes are simulated to be independent, but may liberal in other situations (conservative in (b) (c), and (d), and slightly anti-conservative in (e)). SigPathway, MRSGE and QuSAGE have similar performances as they do, respectively, in group A_1 simulations.

Explain why this happens

MEQLEA shows consistent accuracy for type I error control across all simulations, but the accuracy of the other six methods may be affected by two factors: the inter-gene correlation structures, and DE probability of each gene. MEQLEA controls the size of type I error well because it uses difference in mean as gene-level statistic, and the correlations among genelevel statistics are fully reflected in sample correlation. GSEA evaluates the enrichment score of a test set by generating its null distribution from sample permutation. When there's no DE genes such as in the case of group A_1 simulations, GSEA performs extremely well since permuting sample labels won't change the underlying correlation structure. When DE genes exist, however, sample permutation will destroy the inter-gene correlation structure, which explains the complete failure of GSEA in controlling type I error for the case of group A_2 simulations. For CAMERA and QuSAGE, the VIF of the gene-level statistics (moderated t-test in Wu and Smyth (2012)) may be over-estimated when a fraction of genes are DE (Zhuo and Di, unpublished work), and therefore the setlevel test statistic is under-estimated. The performances of those methods—QuSAGE and two versions of CAMERAare subject to the underlying correlation structures. Moreover,

the performance of CAMERA is complicated by the fact that the set-level statistic takes into account only the inter-gene correlation in the test set without addressing that in the background set.

Different from the five methods mentioned above, Sig-Pathway and MRSGE rely on independence between genes. It's not surprising that gene permutation method, such as SigPathway and MRSGE, controls type I error correctly when genes are "equally-correlated": in (a) genes are simulated to be independent, and in (c) genes are simulated to have an exchangeable correlation structure. However, both SigPathway and MRSGE fail to hold type I error size for the remaining three correlation structures. These simulations show that even small inter-gene correlations will inflate the false discovery rate when the test does not account for intergene correlations.

3.2 Power simulation

We compare the power of MEQLEA to those of the other six methods under different correlation structures. Since some of these tests are not well calibrated at the sample size considered (see results in Section 3.1), we report calibrated power. For calibrated power, the critical value $c(\alpha)$ is chosen so that when the null hypothesis is true, exactly $100 \cdot \alpha\%$ of the resulting p-values are less than $c(\alpha)$; that is, $c(\alpha)$ is the α quantile of null distribution of p-values, where the null distribution is generated from simulation. Calibrated power allows a more fair comparison among tests, as tests that are too conservative under the null hypothesis will have greater power due to the tendency to produce small p-values, yet this apparent power does not truly distinguish between the null and the alternative.

Table 2 summarizes the calibrated power for the two groups of simulations (i.e., A_1 and A_2 in Table 1). We only report the results for correlation structure (a) where genes are simulated to be independent. For A_1 simulations, GSEA has the highest, and rank based methods MRSGE and CAMERArank have the lowest, calibrated power across all four alternative scenarios (the data for S_4 not shown). CAMERAmodt and MEQLEA have no systematic difference in the calibrated power. In group A_2 simulations, GSEA shows virtually no power. MEQLEA, CAMERA-modt, SigPathway and QuSAGE have indistinguishable calibrated power and are among the best.

Figure 2 shows for MEQLEA, the variations in power according to different correlation structures across four alternative scenarios S_1 - S_4 . For each correlation structure and each alternative, we report the power (without recalibration) at a significance level of 0.05. The top is the power for group A_1 , and the bottom for group A_2 . The powers for case (a) and (c) are very similar, and are among the highest under each of the four alternatives. It's not surprising because they correspond to the simplest correlation structures: gene expression values in (a) are simulated to be independent and in (c) are simulated to have the same correlation 0.1. As the correlation structure becomes more complex, from (b) to (d) then to (e), the power decreases under every alternative scenario. The power under correlation structure (e) is the lowest for both A_1 and A_2 simulations.

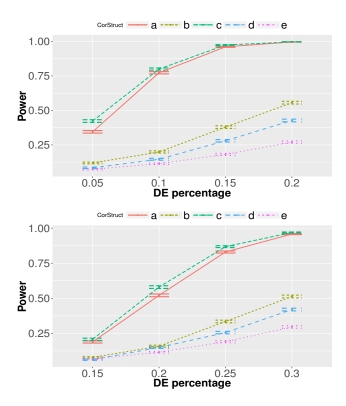


Figure 2: Power for MEQLEA under correlation structures (a)-(e) of Section 2.2.1. The top corresponds to group A_1 simulations, and the bottom to group A_2 simulations (see Table 1). The error bars are the 95% CIs based on 1,000 simulations.

3.3 Real Data

We applied MEQLEA to two example data sets, and compared the enriched gene sets to those obtained by GSEA and by CAMERA-modt. In both examples, MEQLEA were able to identify more gene sets as enriched. Our results lend credence to previous studies in finding potential gene sets correlated with Huntington's disease and those correlated with chromosome Y and Y bands in lymphoblastoid cells.

3.3.1 Huntington's Disease Data. We examined the Huntington's Disease (HD) RNA-Sequencing (RNA-Seq) data (Labadorf et al., 2015) to identify which gene sets are enriched among DE genes in HD. The mRNA expression profiles in human prefrontal cortex were obtained from 20 Huntington's Disease samples and 49 neurologically normal controls. Expression values were normalized and filtered as described in the methods section of Labadorf et al. (2015). The data, containing 28,087 genes, is available as a series GSE64810 in the GEO database (http://www.ncbi.nlm.nih.gov/geo/). We performed enrichment analysis using the MsigDB (Subramanian et al., 2005) C2 Canonical Pathways gene sets (February 5, 2016, data last accessed). The C2 Canonical Pathway gene sets have a collection of 1330 gene sets, with an average set size of 50 (the set sizes range from 3 to 1028, and the median is 29). Since the genes in C2 are named by HGNC symbols and by ensembl IDs in the HD expression data set, we converted the ensembl IDs in the expression data into HGNC sym-

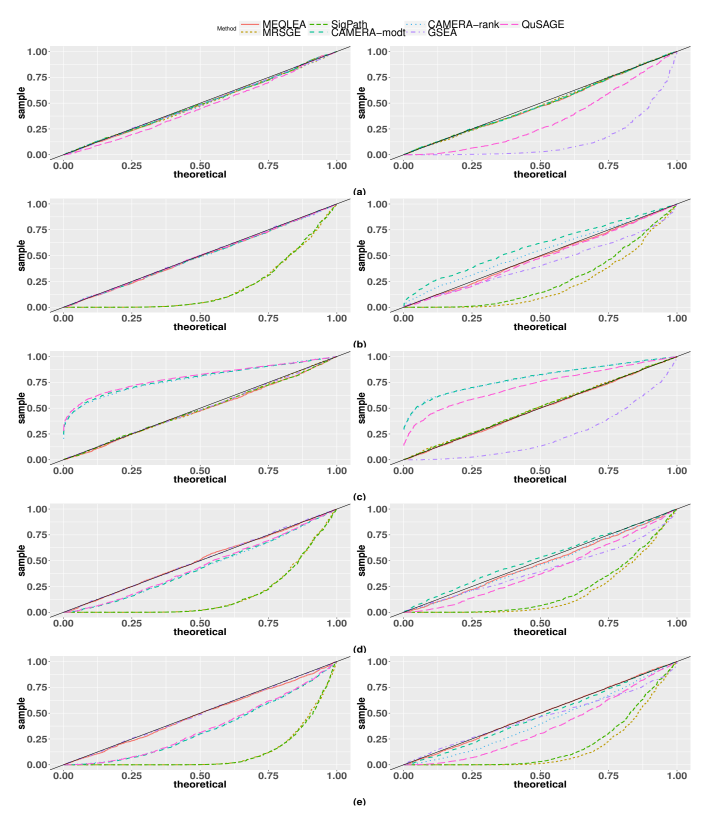


Figure 1: Uniform quantile-quantile plots for p-values by different methods. Each plot from top to bottom corresponds to correlation structures (a)-(e), respectively. The left column is for group A_1 simulation, and the right column for group A_2 simulation (see Table 1 for detail). Results are based on 1,000 simulations.

Table 2: Recalibrated power (standard error) for different methods. The powers are summarized under three alternatives S_1 - S_3 in each of the group A_1 and A_2 simulations (see Table 1 for detail). Results are based on 1,000 simulations.

Group	Method	S_1	S_2	S_3	S_4
A_1	MEQLEA	0.335(0.015)	0.765(0.013)	0.959(0.006)	0.996(0.002)
	MRSGE	0.129(0.011)	0.344(0.015)	0.609(0.015)	0.839(0.012)
	SigPathway	0.350(0.015)	0.778(0.013)	0.961(0.006)	0.995(0.002)
	CAMERA-modt	0.347(0.015)	0.767(0.013)	0.960(0.006)	0.995(0.002)
	CAMERA-rank	0.119(0.010)	0.331(0.015)	0.602(0.015)	0.834(0.012)
	GSEA	0.486(0.016)	0.891(0.010)	0.996(0.002)	0.998(0.001)
	QuSAGE	0.387(0.015)	0.804(0.013)	0.977(0.005)	0.996(0.002)
	MEQLEA	0.199(0.013)	0.538(0.016)	0.836(0.012)	0.959(0.006)
A_2	MRSGE	0.121(0.010)	0.334(0.015)	0.626(0.015)	0.866(0.011)
	SigPathway	0.201(0.013)	0.534(0.016)	0.846(0.011)	0.961(0.006)
	CAMERA-modt	0.197(0.013)	0.531(0.016)	0.837(0.012)	0.960(0.006)
	CAMERA-rank	0.118(0.010)	0.329(0.015)	0.619(0.015)	0.859(0.011)
	GSEA	0.000(0.000)	0.000(0.000)	0.000(0.000)	0.000(0.000)
	QuSAGE	0.197(0.013)	0.536(0.016)	0.846(0.011)	0.960(0.006)

bols using *BioMart* (http://uswest.ensembl.org/biomart/martview/). We retained 26,941 genes that have corresponding HGNC symbols. For MEQLEA, we standardized the data in the way as described in Section....

We used three test procedures (MEQLEA, GSEA and CAMERA-modt) to run enrichment analysis for the entire C2 Canonical Pathway gene sets, and compared the three tests in terms of resulting enriched gene sets. MEQLEA found 176 out of 1330 gene sets to be enriched using the Benjamini-Hochberg (BH) procedure at a false discovery rate (FDR) of 0.05 (for multiple hypothesis testing, unless specified otherwise, all pvalues in Section 3.3 were adjusted by BH procedure). GSEA found 9 enriched gene sets—8 of them were also among the 176 gene sets we identified (the one that was not significant according to MEQLEA had a p-value of 0.008 and FDR 0.057). CAMERA-modt found no enriched gene sets. In Figure 3 we present pairwise p-value plots for MEQLEA, GSEA and CAMERA-modt. When plotted against p-values of GSEA, for MEQLEA, smaller p-values (e.g., less than 0.1) are more likely to cluster—as compared to larger p-values; that is, MEQLEA produces more small p-values than GSEA does while MEQLEA and GSEA do not differ much in producing larger p-values. The p-values of CAMERA-modt are overwhelmingly larger than their counterparts of MEQLEA, even if p-values of the two methods are highly correlated (Pearson's correlation is 0.96). This is consistent with our earlier simulation (see results in Section 3.1) that CAMERA-modt could be too conservative. There is no systematic difference in p-values of GSEA and those of CAMERA-modt.

We report the top 30 enriched gene sets in Table 3. Five enriched gene sets identified by GSEA are also present (noted by "*" in the table). Originally, Labadorf et al. (2015) used the same HD data set to conduct enrichment analysis using topGo (Alexa and Rahnenfuhrer, 2010). They found that the enriched gene sets they identified show a clear immune response and inflammation-related pattern, including "REACTOME INNATE IMMUNE SYSTEM, PID IL4 2PATHWAY", and "PID NFKAPPAB CANONICAL PATHWAY". These three gene sets rank 6,10 and 2 respectively in Table 3.

Many of our enriched gene sets have been shown to be

closely related to HD pathogenesis. For example, the top enriched gene set by MEQLEA, "PID SMAD2 3NUCLEAR PATHWAY" (see Table 3), is responsible for regulation of nuclear SMAD2/3 signaling. Katsuno et al. (2010) showed that nuclear SMAD2/3 are related to polyglutamine disease, which includes HD. The second enriched gene set, "PID NFKAPPAB CANONICAL PATHWAY", is a canonical NFkappaB pathway, and its dysregulation causes HD immune dysfunction (Träger et al., 2014). Also, Marcora and Kennedy (2010) found that reduced transport of NF-kappaB out of dendritic spines and its activity in neuronal nuclei may contribute to the etiology of HD. Another gene set, "REACTOME IN-NATE IMMUNE SYSTEM", contributes to HD pathogenesis (Träger et al., 2014; Labadorf et al., 2015). For AKT signaling pathway, "BIOCARTA AKT PATHWAY", Humbert et al. (2002) demonstrated that huntingtin is a substrate of AKT and that phosphorylation of huntingtin by AKT is crucial to mediate the neuroprotective effects of IGF-1. They also showed that AKT is altered in Huntingtons disease patients. Chiang et al. (2010) demonstrated that the systematic downregulation of PPAR γ , related to "BIOCARTA" PPARA PATHWAY", seems to play a critical role in the dysregulation of energy homeostasis observed in HD, and that PPAR γ is a potential therapeutic target for this disease. For "REACTOME SIGNALING BY TGF BETA RECEPTOR COMPLEX", Kandasamy et al. (2011) demonstrated that TGF-beta1 signaling appears to be a crucial modulator of neurogenesis in HD pathology and it can be a promising target for endogenous cell-based regenerative therapy. For "PID P53 DOWNSTREAM PATHWAY", Ghose et al. (2011) showed the likely involvement of NFkB (RelA), p53 and miRNAs in the regulation of cell death in HD pathogenesis.

3.3.2 Male vs Female Lymphoblastoid Cells Data. We analyzed the mRNA expression profiles from lymphoblastoid cell lines derived from 17 females and 15 males. Subramanian et al. (2005) examined this data set with their GSEA method, testing the enrichment of the cytogenetic gene sets (C1). C1 includes 24 sets, one for each of the 24 human chromosomes, and 295 sets corresponding to cytogenetic bands. For the

Table 3: Top 30 enriched gene sets using MEQLEA for HD data. Gene sets are ranked by their associated p-values. FDR is the adjusted p-value using Benjamini-Hochberg (BH) procedure.

Gene Set	Size	ρ_1	ρ_2	ρ_3	p-value	FDR	
PID SMAD2 3NUCLEAR PATHWAY	79	0.071	0.011	0.017	7.5E-07	9.9E-04	*, **
PID NFKAPPAB CANONICAL PATHWAY	22	0.124	0.011	0.020	2.4E-06	1.6E-03	**
REACTOME YAP1 AND WWTR1 TAZ STIMU-	23	0.130	0.011	0.018	4.4E-06	1.7E-03	**
LATED GENE EXPRESSION							
REACTOME SIGNALING BY TGF BETA RECEP-	60	0.045	0.011	0.015	7.3E-06	1.7E-03	**
TOR COMPLEX							
BIOCARTA NTHI PATHWAY	23	0.124	0.011	0.024	7.5E-06	1.7E-03	
REACTOME INNATE IMMUNE SYSTEM	209	0.048	0.011	0.010	7.8E-06	1.7E-03	**
KEGG PATHWAYS IN CANCER	311	0.029	0.011	0.010	8.9E-06	1.7E-03	
REACTOME DOWNSTREAM TCR SIGNALING	31	0.095	0.011	0.013	1.2E-05	1.9E-03	**
KEGG NOD LIKE RECEPTOR SIGNALING PATH-	55	0.054	0.011	0.010	1.3E-05	1.9E-03	**
WAY							
PID IL4 2PATHWAY	59	0.086	0.011	0.012	1.4E-05	1.9E-03	**
KEGG TGF BETA SIGNALING PATHWAY	82	0.062	0.011	0.013	2.7E-05	3.3E-03	**
BIOCARTA 41BB PATHWAY	14	0.095	0.011	0.023	3.2E-05	3.4E-03	
PID P53 DOWNSTREAM PATHWAY	131	0.052	0.011	0.013	3.4E-05	3.4E-03	**
REACTOME TCR SIGNALING	48	0.098	0.011	0.016	3.6E-05	3.5E-03	**
REACTOME ACTIVATED TLR4 SIGNALLING	87	0.027	0.011	0.010	4.9E-05	4.2E-03	**
REACTOME TOLL RECEPTOR CASCADES	110	0.038	0.011	0.010	5.2E-05	4.2E-03	**
REACTOME TRANSCRIPTIONAL REGULATION	69	0.015	0.011	0.010	5.4E-05	4.2E-03	
OF WHITE ADIPOCYTE DIFFERENTIATION							
BIOCARTA TID PATHWAY	18	0.125	0.011	0.017	5.7E-05	4.2E-03	**
BIOCARTA ALK PATHWAY	34	0.064	0.011	0.011	7.4E-05	5.1E-03	*
REACTOME SMAD2 SMAD3 SMAD4 HET-	25	0.102	0.011	0.021	7.6E-05	5.1E-03	*
EROTRIMER REGULATES TRANSCRIPTION							
REACTOME TRANSCRIPTIONAL ACTIVITY OF	36	0.079	0.011	0.021	8.3E-05	5.1E-03	
SMAD2 SMAD3 SMAD4 HETEROTRIMER							
BIOCARTA AKT PATHWAY	20	0.023	0.011	0.010	8.8E-05	5.1E-03	*
ST TUMOR NECROSIS FACTOR PATHWAY	28	0.039	0.011	0.016	9.0E-05	5.1E-03	*
PID ANGIOPOIETIN RECEPTOR PATHWAY	50	0.082	0.011	0.013	9.3E-05	5.1E-03	**
KEGG P53 SIGNALING PATHWAY	65	0.037	0.011	0.007	9.7E-05	5.1E-03	
KEGG APOPTOSIS	82	0.041	0.011	0.009	1.0E-04	5.1E-03	
BIOCARTA PPARA PATHWAY	53	0.026	0.011	0.008	1.1E-04	5.2E-03	
REACTOME MYD88 MAL CASCADE INITIATED	78	0.026	0.011	0.010	1.1E-04	5.2E-03	**
ON PLASMA MEMBRANE							
PID BCR 5PATHWAY	64	0.064	0.011	0.016	1.2E-04	5.3E-03	
PID HIF1 TFPATHWAY	64	0.067	0.011	0.011	1.2E-04	5.3E-03	

 $[\]rho_1$: average sample correlation between genes in the test set.

comparison "male VS female", they expected to find gene sets on chromosome Y, not on chromosome X. We run enrichment analysis with three tests (MEQLEA, GSEA and CAMERAmodt). In Table 4, we summarized all the gene sets with nominal p-value ≤ 0.01 in at least one test. Three gene sets, one from chromosome Y and two Y bands, were found to be enriched by all three tests at FDR level 0.05. Interestingly, MEQLEA identified another Y band, chrYp22, as enriched. In fact, the four gene sets called significant by MEQLEA are the only four containing at least 3 genes in C1 and corresponding to chromosome Y or Y bands. MEQLEA did not produce

small p-value (≤ 0.01) for the remaining three gene sets in Table 4, which was just as expected in that study.

4. Conclusion and Discussion

(Conclusion) MEQLEA is a mixed effects quasi-likelihood model for competitive gene set test. It effectively adjusts for completely unknown, unstructured correlations among the genes. It uses a score test approach and allows for analytical assessment of p-values. Compared to existing approaches, MEQLEA controls type I error correctly and maintains good power under different correlation structures.

 $[\]rho_2$: average sample correlation between genes in the background set.

 $[\]rho_3$: average sample correlation between two genes, one from the test set and the other from the background set.

^{*:} enriched gene sets identified by GSEA.

^{**:} enriched gene sets identified by MRSGE.

Table 4: Summary of gene sets for lymphoblastoid cells data. Reported are gene sets with p-value ≤ 0.01 for at least one of the MEQLEA, GSEA, and CAMERA-modt methods. The FDR is the adjusted p-value using Benjamini-Hochberg (BH) procedure.

		MEQLEA		GS	GSEA		CAMERA-modt	
Gene set	Size	<i>p</i> -value	FDR	<i>p</i> -value	FDR	<i>p</i> -value	FDR	
chrY	40	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001	0.002	
chrYq11	16	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001	
chrYp11	18	< 0.001	< 0.001	< 0.001	< 0.001	< 0.001	0.028	
chrYp22	8	< 0.001	0.036	0.012	0.503	0.010	0.762	
chr7p11	8	0.049	0.835	0.006	0.352	0.101	0.998	
chr11p12	5	0.065	0.835	0.008	0.388	0.115	0.998	
chrXp22	76	0.072	0.835	0.004	0.295	0.581	0.998	

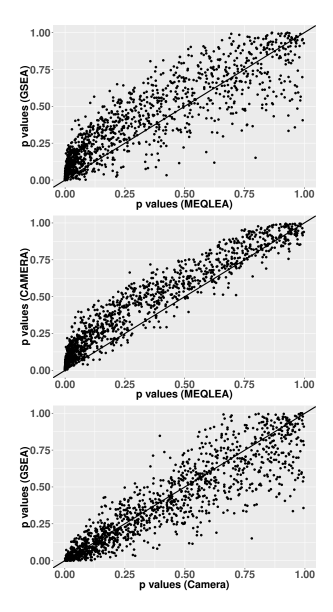


Figure 3: Pairwise comparisons of p-values for MEQLEA, GSEA, and CAMERA-modt. The p-values are reported from enrichment test of each gene set in the C2 Canonical Pathway gene sets.

(What we proposed) Under competitive gene set test framework, a number of methods have been proposed to account for correlation among genes. One approach is to evaluate the set-level statistic by permuting sample labels to generate the null distribution, as adopted by the widely used GSEA (Subramanian et al., 2005). However, sample permutation method has been criticized for altering the null hypotheses being tested (Goeman and Bühlmann, 2007; Khatri et al., 2012). Instead, CAMERA (Wu and Smyth, 2012) proposed to correct for the correlation among genes by estimating a VIF directly from the data. This approach has also been used by Yaari et al. (2013) in their QuSAGE procedure. The major shortcoming with this approach is that it tries to estimate correlations among gene-level test statistics directly from sample correlation (is it clear??). Zhuo and Di have argued (unpublished work) that the correlations among gene-level statistics are not necessarily equal to those among samples due to the presence of DE genes. The estimated VIF could be biased without taking into account such a discrepancy and thus undermines the performance of CAMERA and QuSAGE. MEQLEA avoids the discrepancy by using the differences in mean as gene-level statistics for a two group comparison experiment. It models the covariance of gene-level statistics by two variance components, one attributable to correlations among samples after treatment effect removed, and the other attributable to the DE effect associate with the treatment. We note that for MEQLEA, the estimation of covariance among gene-level statistics need not be exact: MEQLEA uses a score test that involves linear combinations of the entries in the covariance matrix. The denominator in the score test statistic (REF EQ) can usually be accurately approximated given the high dimensionality of the covariance matrix. MEQLEA is based on quasi-likelihood, therefore it does not require normal assumption of expression data, and could be applied to both microarray and RNA-Seq experiments.

(Summarize the results) We compared MEQLEA to other existing approaches in both simulation study and real data analysis. In the simulation study, we examined the performance of MEQLEA and other six method (SigPathway, MRSGE, CAMERA-modt, CAMERA-rank, GSEA and QuSAGE) in terms of type I error control and power. We demonstrated that under a variety of correlation structures, MEQLEA holds correct type I error size and also maintains good power. In the real data analysis, MEQLEA was able to identify more gene sets as enriched, some of which, in the

corresponding studies, are insightful yet not found by methods such as GSEA or CAMERA.

(Future work) Currently, MEQLEA only supports enrichment test for two-group comparisons. In many gene expression experiments, however, researchers might use more complex design to study different comparisons of interest, in which case a linear model would be more appropriate. Our future work will focus on generalizing MEQLEA to allow for more complicated design structures.

The R codes for reproducing results in this paper are available at https://github.com/zhuob/EnrichmentAnalysis.

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Appendix

Standardization Standardization for each gene: first, we obtain the residuals by subtracting off the means within each treatment group;

$$r_{ijk} = y_{ijk} - \sum_{j=1}^{n_k} y_{ijk} / n_k;$$
 (12)

then we calculate the pooled standard deviation from the residuals,

$$s_i = std(r_{ijk}); (13)$$

next we get the standardized expression by dividing the original expression profiles by the standard deviation,

$$y_{ijk}^* = y_{ijk}/s_i \tag{14}$$

We perform the standardization procedure to every gene in the data set.

First $E(\Delta_i) = E(Z_i\delta_i) = E(Z_i)E(\delta_i) = p_i\mu_{\delta}$. Next note that

$$\operatorname{Var}(\Delta_i) = E[(Z_i \delta_i)^2] - [E(Z_i \delta_i)]^2$$

$$= \operatorname{Var}(Z_i)[E(\delta_i)]^2 + [(EZ_i)^2 + \operatorname{Var}(Z_i)] \operatorname{Var}(\delta_i)$$

$$= p_i \sigma_\delta^2 + p_i (1 - p_i) \mu_\delta^2$$

Let $T_i = \bar{Y}_{i,2} - \bar{Y}_{i,1}$ be the difference in mean expression profiles between the treatment group and the control group. We have

$$E(T_i) = E(\bar{Y}_{i,2}) - E(\bar{Y}_{i,1}) = E(\Delta_i) = E(Z_i\delta_i) = p_i\mu_{\delta}$$

The covariance between two genes i_1 and i_2 is given by (I HAVE CONCERNS HERE, IS IT VALID TO ASSUME THAT DE EFFECTS ARE INDEPENDENT BETWEEN GENES? WE SEE CO-EXPRESSION!! OR WE'VE ALREADY TAKEN THAT INTO ACCOUNT BY CORRELATION BETWEEN GENES"),

$$Cov(T_{i_1}, T_{i_2}) = E\left[Cov(T_{i_1}, T_{i_2} | \Delta_{i_1}, \Delta_{i_2})\right] + Cov\left[E(T_{i_1} | \Delta_{i_1}), E(T_{i_2} | \Delta_{i_2})\right]$$

$$= E\left(\frac{1}{n_1}\rho_{i_1, i_2} + \frac{1}{n_2}\rho_{i_1, i_2}\right) + Cov(\Delta_{i_1}, \Delta_{i_2})$$

$$= \left(\frac{1}{n_1} + \frac{1}{n_2}\right)\rho_{i_1, i_2}$$
(15)

For gene i, the variance $Var(T_i) = Var(\bar{Y}_{i,1}) + Var(\bar{Y}_{i,2})$, with

$$\operatorname{Var}(\bar{Y}_{i,1}) = \frac{1}{n_1}$$

$$\operatorname{Var}(\bar{Y}_{i,2}) = \frac{1}{n_2^2} \left[\sum_{j=1}^{n_2} \operatorname{Var}(Y_{ij2}) + 2 \sum_{1 \leq j_1 < j_2 \leq n_2} \operatorname{Cov}(Y_{ij_12}, Y_{ij_22}) \right]$$

$$= \frac{1}{n_2} \operatorname{Var}(Y_{ij2}) + \frac{n_2 - 1}{n_2} \operatorname{Cov}(Y_{ij_12}, Y_{ij_22})$$

$$= \frac{1}{n_2} \left[E\left(\operatorname{Var}(Y_{ij2} | \Delta_i) \right) + \operatorname{Var}\left(E(Y_{ij2} | \Delta_i) \right) \right]$$

$$+ \frac{n_2 - 1}{n_2} E\left(\operatorname{Cov}(Y_{ij_12}, Y_{ij_22} | \Delta_i) \right)$$

$$+ \frac{n_2 - 1}{n_2} \operatorname{Cov}\left(E(Y_{ij_12} | \Delta_i), E(Y_{ij_22} | \Delta_i) \right)$$

$$= \frac{1}{n_2} + \operatorname{Var}(\Delta_i)$$
(16)

Therefore $\operatorname{Var}(T_i) = \frac{1}{n_1} + \frac{1}{n_2} + \operatorname{Var}(\Delta_i)$, and it follows that

$$Cov(T) = D + \sigma_2^2 C$$
 (17)

where D is a diagonal matrix with $Var(\Delta_i) = p_i \sigma_\delta^2 + p_i (1 - p_i) \mu_\delta^2$ as its *i*th diagonal element, and $\sigma_2^2 = \left(\frac{1}{n_1} + \frac{1}{n_2}\right)$.