Higher-level Analysis of RNA-Seq Experiment: Multiple Data Sets and Multiple Genes

Ву

Bin Zhuo

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Differential expression (DE) analysis is a key task in gene expression study, because it uncovers the association between expression levels of a gene and the covariates of interest. This dissertation pertains to two particular aspects of DE analysis—identifying stably expressed genes for count normalization and accounting for correlation between DE test statistics in gene-set test. RNA-Sequencing (RNA-Seq) has become the tool of choice for measuring gene expression over the past few years, and data generated from RNA-Seq experiments are the focus of this thesis.

Identifying stably expressed genes is useful for count normalization and DE analysis. We examined RNA-Seq data on 211 biological samples from 24 different experiments conducted by different labs, and identified genes that are stably expressed across samples, treatment conditions, and experiments. We fit a Poisson log-linear mixed-effect model to the count data, and decomposed the total variance into between-sample, between-treatment and between-experiment variance components. The variance component analysis that we explore here is a first step towards understanding the sources and nature of the RNA-Seq count variation. The stability ranking of genes, when quantified by a numerical stability measure, is dependent on several factors: the background sample set and the reference gene set used for count normalization, the

technology used to measure gene expression, and the specific stability measure. Since DE is measured by relative frequencies, we argue that DE is a relative concept. We advocate using an explicit reference gene set for count normalization to improve interpretability of DE results, and recommend using a common reference gene set when analyzing multiple RNA-Seq experiments to avoid potential inconsistent conclusions.

We investigate the relationship between correlation among test statistics and the correlation of underlying observed data. For false discovery control (FDR) procedures and gene-set tests, pooling DE test statistics together is a frequently used idea and the correlation among test statistics needs to be taken into account. The sample correlation of observed data are often used to approximate the test statistics correlation. We show, however, that such an approximation is only valid under limited settings. In particular, we derive a formula for correlation between test statistics when they take a specific form, and as a special case, we present the exact expression of test statistics correlation for equal-variance two sample t-test statistic under bivariate normal assumption. We conclude that test statistics correlation is no more greater in absolute value than the correlation of underlying observed data (normally distributed) in the context of equal-variance two-sample t-test.

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, MEACA, 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 con-

trol over type I 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: RNA-Seq, stably expressed genes, correlation, gene-set test

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<u>Doctor of Philosophy</u> dissertation of <u>Bin Zhuo</u> presented on <u>June 22, 2016</u>
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Major Professor, representing Statistics
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1. Introduction

In Section 1.1, we give an overview of the three research questions that will be addressed in this thesis. In Section 1.2, we provide necessary background on RNA-Sequencing (RNA-Seq) experiments and RNA-Seq data processing. RNA-Seq experiments are the focus of this thesis. In Section 1.3, we review generalized linear mixed models that will be used for variance component analysis of RNA-Seq data.

1.1. Overview of research questions

In this thesis, we investigate three questions pertaining to gene differential expression (DE) analysis: 1. Identifying stably expressed genes from multiple RNA-Seq data sets by a numerical stability measure (Chapter 2); 2. Investigating the relationship between the correlations among DE test statistics and the correlations among underlying observed data (Chapter 3); 3. Accounting for correlations among DE test statistics in competitive gene-set test (Chapter 4).

1.1.1. Identifying stably expressed genes

DE analysis is a key task in gene expression study, because it uncovers the association between expression levels of a gene and the covariates of interest. The covariates could either be categorical (e.g., treatment/control status, cell types), or continuous (e.g., reagent concentration, time). For example, to understand the effect of a drug, one might ask which genes are *up-regulated* (increased expression levels) or *down-regulated* (decreased expression levels) between treatment and control groups. Finding these genes will help researchers understand which genes or pathways are involved in the drug's functions.

Despite the optimistic claim that DE analysis of RNA-Seq data does not need sophisticated normalization [127], many works have shown that normalization of count data is highly desirable to account for various sources of bias between samples [3,

28, 48, 104, 105, 106]. Many of the current normalization methods, for example, TMM [106] and DESeq [3] normalizations, assume that the majority of genes are not DE within the experiment under investigation. However, this assumption could be violated in some experiments where over 50% of the genes' expression levels are altered by the treatments [78, 130]. The consequence with such assumption can be alleviated if one could identify a set of reference genes whose expression levels are stable across different experimental conditions. This motivates us to examine a large number of existing RNA-Seq data sets and to identify reference (stably expressed) genes.

In microarray studies, there have been many attempts to find reference genes for normalization. Traditionally, the house-keeping genes are used as reference genes. However, a number of works have shown that house-keeping genes are not necessarily stably expressed according to numerical stability measures (see, for example, [24, 56]). Another choice, the *spike-in genes*, is not reliable for normalization due to the same issue [104]. A popular approach has been to search from large sets of experiments for reference genes [24, 25, 38, 46, 115] whose expression stability are evaluated by some numerical stability measure. Validation experiments (e.g. reverse transcription-PCR) show that reference genes identified by numerical methods generally outperform house-keeping genes or spike-in genes in terms of expression stability [24, 52]. We will follow the strategy of quantifying gene expression stability by numerical measures and identify stably expressed genes.

Identifying stably expressed genes not only helps count normalization, but also improves interpretability and comparability of RNA-Seq experiments in integrative analysis. Since genes are measured by relative frequencies, we argue that DE is a relative concept: when a normalization procedure is applied to a single data set, it effectively uses an implicit reference set of genes. Furthermore, making the reference set explicit will be beneficial during DE analysis, because often times biologists compare results from one experiment to those from others experiments whose data are

publicly available.

1.1.2. Correlation among test statistics

DE analysis involves simultaneous hypothesis testing for an ensemble of genes. One common feature of testing an ensemble of genes is that a test statistic quantifying the magnitude of DE is calculated for each gene, and then all the test statistics are pooled together and treated as a sample from which to estimate sampling distribution of test statistics. Since the observed data (expression levels) are correlated, the test statistics derived from them are also correlated, and this inter-dependency of test statistics cannot be ignored when testing a set of genes [10, 31, 132].

However, without replicating the experiment, we cannot directly compute the sample correlation of the test statistic values between genes since only a single test statistic value is available for each gene. The sample correlation between observed data is often used to approximate the sample correlation between two test statistic values computed from these data. Efron [31] used this idea to approximate the distribution of test statistics correlation, and uses the resulting distribution to derive the false discovery proportion in a multiple hypothesis testing procedure. Barry et al. [10], Wu and Smyth [132] and Yaari et al. [134] also used correlation of observed data to summarize test statistics correlation in their gene-set test procedures to account for correlation among test statistics. However, it has only been shown by simulation under limited settings that the correlations among test statistics are almost the same as the correlations among underlying observed data [10, 31]. In Chapter 3, we will investigate when and to what degree that the correlation of test statistics can be approximated by the correlation of observed data.

1.1.3. Accounting for correlations of test statistics in competitive gene set test

DE analysis evaluates each individual gene separately. Single-gene analysis may fail to provide insights into biological mechanisms when multiple genes function together. Gene-set analysis is a commonly-used technique that provides a higher-level interpretation of the DE test results. For a given set of genes, a gene-set test will evaluate whether there is an overabundance of DE genes in the set as compared to DE genes in a background set (such a gene-set test is sometimes called a "competitive test" [42]). Commonly considered gene sets are gene pathways, genes assigned to a common ontology term, and so on.

Many existing gene-set tests do not take possible correlation between test statistics into consideration, such as SigPathway [121], PAGE [61], MRGSE [87] or the 2 × 2 contingency-table-based tests [1, 54, 135]. However, it has been argued that such test procedures will result in inflated type I error [33, 39, 42, 132, 134], as genes within a gene set are often co-expressed and function together. Those that do consider test-statistic correlation often use observed-data correlation to approximate test-statistic correlation [10, 132, 134]. As we will show in Chapter 3, such an approximation is problematic when there is DE among genes. In Chapter 4, we introduce a gene-set test that correctly handles the test-statistic correlation among genes.

1.2. Background on RNA-Sequencing experiments

Gene is a piece of DNA that encodes a functional RNA or protein product, and is the basic physical and functional unit of heredity. The process by which genes are used to synthesize functional gene products is called *gene expression*. A gene is considered to be expressed in a cell or group of cells when a gene product is detected. These products can be transcribed messenger RNA (mRNA) and proteins for protein coding genes, or functional RNA species such as transfer RNA (tRNA) or small nuclear RNA

(snRNA) for non-protein coding genes. Since the information encoded in a gene is first transcribed into RNA molecules, which is then used to make functional gene products, the RNAs transcribed in a certain condition reflect the current state of the cell.

In a typical gene expression experiment, researchers are usually interested in comparing expression levels of one or more genes under different experiment/environmental conditions. Factors for comparison could be before vs after effect in a drug treatment, tumor vs normal tissues in clinical study, or wild type vs mutant strains in plant research. Another important factor is the time-course, where cells/tissues at different stages are sampled with the purpose of discovering temporal pattern of gene expression. There are many other types of experiment, each with specific factors of interest to be studied.

The expression levels of a gene can be measured using techniques such as complementary DNA (cDNA) libraries, microarray analysis, RNA fingerprinting by arbitrary primed PCR (RAP-PCR), expressed sequence tag (EST) sequencing, serial analysis of gene expression (SAGE), and RNA sequencing (RNA-Seq) (see [21] for a review). RNA-Seq, also known as whole transcriptome shotgun sequencing [90], is a next-generation sequencing (NGS) technology used to uncover the presence and quality of RNA in a biological sample. It is rapidly becoming technology of choice for transcriptome profiling over the past few years. The standard procedure of an RNA-Seq experiment runs as follows [36]: first, the RNAs in the biological sample are fragmented and reverse-transcribed into cDNAs; second, the cDNA fragments are amplified and sequenced in a high-throughput sequencing platform (e.g., Illumine 3000, http://www.illumina.com) to generate (up to) hundreds of millions of reads; third, those reads are mapped to a reference genome or a reference transcriptome. It is the number of reads aligned to each gene (referred to as "read count") on the reference genome/transcriptome that quantifies the genes' expression levels.

RNA-Seq technology offers several key advantages over other methods for mea-

suring gene expression [127], the most important of which are that it does not require prior knowledge of an organism for detecting transcripts, and that it is sensitive to genes expressed at either low or higher levels and thus provides higher dynamic range. The sequencing of RNA allows researchers to study the entire transcriptome of a species using only small amount of RNA. It has been demonstrated that a coordinated effort between RNA-Seq and real time PCR (RT-PCR) is one of the most effective ways to identify new exons [51]. However, one major challenge of this technique is data processing: RNA-Seq experiment produces a huge amount of reads (up to hundreds of millions per sample) and obtaining the expression profiles requires fast read mapping tools as well as a lot of computing resource [65, 70].

Preprocessing RNA-Seq data consists of two main steps: 1) mapping reads to the reference genome/transcriptome, and 2) summarizing read counts at given genomic feature (e.g., exon, gene or transcript) level. Read mapping is the first computational, and usually, the most time-consuming step in RNA-Seq data analysis. Currently, there are many alignment tools available, for example, Bowtie [64, 65], BWA [68, 69, Subread [111] and STAR [29]. In all situations, an index of either the reference genome/transcriptome or the reads is built at the beginning using hash tables or Burrows-Wheeler transform (BWT) [18]. The index allows fast retrieval of the set of positions in the reference sequence where the reads are more likely to align. Once those positions are decided, alignment is performed in the candidate regions. The precision and speed of the alignment is mainly determined by the algorithm used in the alignment tool (see [49] or [71] for a review). After the reads have been aligned, the numbers of reads mapped to each unit of a specified genomic feature are counted, giving the estimate of the corresponding expression levels. This can be done using HTSeq [2] or featureCounts [72], among other options. Finally, a read count matrix is obtained in which each row represents a genomic feature unit and each column corresponds to a biological sample.

In this dissertation, we assemble an in-house pipeline to process RNA-Seq data

sets based on the R [100] platform. This pipeline, modified from a standard procedure given by Anders et al. [4], is designed to work for sequencing data available at the National Center for Biotechnology Information (NCBI, http://www.ncbi.nlm.nih.gov/). It uses the SRA (Sequence Read Archive) Toolkit [67] to convert SRA files to FASTQ files, the Subread aligner [73] to map reads to the reference genome, and then the featureCounts [72] to summarize counts (see Figure 1.1 for the work flow). We will use this pipeline to process multiple RNA-Seq data sets that are needed in Chapter 2.

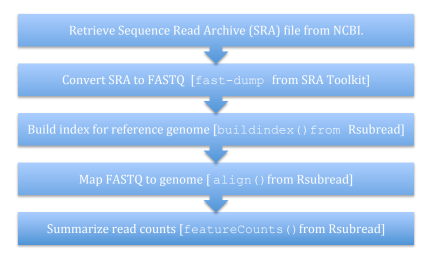


Figure 1.1: Work flow of RNA-Seq data preprocessing: from raw reads sequencing data to read counts. Raw data in this workflow are retrieved from the NCBI respository. Data processing is carried out based on two softwares—the SRA Toolkit [67] and the Rsubread aligner [73].

1.3. Background on generalized linear mixed models

In recent years, many statistical tools have been developed for DE detection (methods review can be found in [101, 110, 114]) in RNA-Seq experiments. Generalized linear models (GLMs) are a natural choice for analyzing RNA-Seq expression data because RNA-Seq data are present in the form of counts. Most of those approaches are based on Poisson [82, 126] or Negative Binomial (NB) distribution [3, 26, 93, 107, 133]. The NB distribution based models are more popular for their flexibility to deal with

over-dispersion (a.k.a. extra-Poisson variation) that are often observed in RNA-Seq expression data. In addition, to account for random effects in biological experiments, GLMs are sometimes extended to generalized linear mixed models (GLMMs). In this section, we will first describe the formulation GLMMs, and then review commonly used methods for parameter estimation under this framework.

1.3.1. Framework of generalized linear mixed models

GLMMs are a natural generalization of classical linear models. To illustrate this point, we will begin with classical linear models, and discuss how to generalize them to linear mixed models and then to GLMMs by relaxing different layers of assumptions.

1.3.1.1 Classical linear models

In a classical linear model, a vector \boldsymbol{y} of n observations is assumed to be a realization of random variable \boldsymbol{Y} whose components are identically distributed with mean $\boldsymbol{\mu}$. The systematic part of this model is a specification of the mean $\boldsymbol{\mu}$ over a few unknown parameters [84]. In the context of classical linear models, the mean is a function of p covariates $\boldsymbol{X}_1, \ldots, \boldsymbol{X}_p$,

$$\boldsymbol{\mu} = \beta_0 + \sum_{i=1}^p \beta_i \boldsymbol{X}_i \tag{1}$$

where $\boldsymbol{\beta} = (\beta_0, \beta_1, \dots, \beta_p)^T$ are unknown parameters and need to be estimated from data. For jth observation Y_j , we specify a random term ϵ_j to allow for measurement error. Assuming a linear relationship between response Y_j and predictors (x_{1j}, \dots, x_{pj}) , we present the linear model

$$Y_j = \beta_0 + \beta_1 x_{1j} + \ldots + \beta_p x_{pj} + \epsilon_j \tag{2}$$

It is often required that ϵ_i 's meet Gauss-Markov assumption,

$$E(\epsilon_i) = 0, \text{ Var}[\epsilon_i] = \sigma^2 < \infty, \text{ Cov}[\epsilon_i, \epsilon_j] = 0, \forall i \neq j.$$
 (3)

In practice, the error term is frequently, if not always, assumed to be normally distributed,

$$\epsilon \sim N(0, \sigma^2 \mathbf{I}).$$
 (4)

1.3.1.2 Linear mixed models

The Gauss-Markov assumption in equation (3) is vulnerable in practice, for example, in the case of nonconstant variance or correlated data where $Cov[\epsilon_i, \epsilon_j] \neq 0$. Equation (2) in either case, without loss of generality, can be expressed in matrix form as

$$Y = X\beta + \epsilon, \ E[\epsilon] = 0, \ Cov[\epsilon] = V$$
 (5)

where V is a known positive definite matrix. Let $Y^* = V^{-1/2}Y = V^{-1/2}X\beta + V^{-1/2}\epsilon$. It follows that $Cov(Y^*) = I$ and the techniques in classical linear models are readily applicable to estimate β . However, this method relies on the assumption that V is known which is rarely, if ever, given. On the other hand, the structure of V, which depends on experiment setup, can often be specified by a few unknown parameters.

Nonindependence can occur in the form of serial correlation or cluster correlation [103, chapter 17]. Serial correlation usually exists in experiments with repeated measurements—multiple measurements taken from a response variable on the same experimental unit. Several covariance structures are available for implementation (for more details, see Littell et al. [74, chapter 5]). Cluster correlation is present when measurements of a response variable are grouped in some way. In many situations, the covariance of cluster correlated data can be specified using an extension of standard linear model by

$$Y = X\beta + Z_1u_1 + \dots + Z_qu_q + \epsilon \tag{6}$$

Equation (6) differs from equation (5) only in the $\mathbf{Z}_{i}\mathbf{u}_{i}$ terms, which is the key part of linear mixed models. The \mathbf{Z}_{i} 's are known $n \times p_{i}$ full rank matrices, usually

used to specify membership of predictors in various subgroups. The most important innovation in this model is that instead of estimating \mathbf{u}_i 's as fixed parameters, we assume them to be unknown random quantities, and $E[\mathbf{u}_i] = 0$, $\text{Cov}[\mathbf{u}_i] = \sigma_i^2 \mathbf{I}_{p_i}$ for i = 1, ..., q. It is, in many cases, reasonable to require that \mathbf{u}_i be mutually independent, and that \mathbf{u}_i be independent of $\boldsymbol{\epsilon}$ for i = 1, ..., q. If we further impose normal distribution on the random terms and errors, then equation (6) can be casted in a Bayesian framework,

$$y|u_1, \dots, u_q \sim N_n(X\beta + \sum_{i=1}^q Z_i u_i, \sigma^2 I_n),$$

$$u_i \sim N_{p_i}(0, \sigma_i^2 I_{p_i}).$$
(7)

The modeling issues are: (a) estimation of variance components σ_i^2 and σ^2 ; (b) estimation of random effects u_i if needed. For the variance component estimation, there are primarily three approaches: (i) procedures based on expected mean squares from analysis of variance (ANOVA); (ii) maximum likelihood (ML); and (iii) restricted/residual maximum likelihood (REML). For more details, see Littell et al. [74, Chapter 1].

1.3.1.3 Generalized linear models

We can take a different perspective of classical linear models by arranging equations (1)–(3) into three parts, following the notations of McCullagh and Nelder [84, Chapter 2],

- (i) the random component Y_j has constant variance σ^2 and $E[Y_j] = \mu_j$.
- (ii) the *systematic component*—the linear predictor η_j is modeled by covariates $\boldsymbol{x}_j =: x_{1j}, \dots, x_{pj},$

$$\eta_j = \sum_{i=1}^p \beta_i x_{ij} = \boldsymbol{x_j} \boldsymbol{\beta}. \tag{8}$$

(iii) the link function relates the random components and the systematic components

$$\eta_j = g(\mu_j). \tag{9}$$

The classical linear models fit within this framework if we assume that the random components Y_j 's are independent and normally distributed, and that the link function is identity (i.e., $g(\mu_j) = \mu_j$).

We can extend part (i)—by allowing Y_j to come from an exponential family (e.g., Poisson, Gamma or Binomial distribution), and part (iii)— by requiring the link function to be monotonic differentiable (e.g., $g(\mu_j) = \log \mu_j$). These two extensions lead to the *generalized linear models* (GLMs), a framework that is especially suitable when a normal distribution is no longer appropriate to be assumed on the response.

1.3.1.4 Generalized linear mixed models

Generalized linear mixed models (GLMMs) is a further extension of GLMs that incorporates random components into part (ii), represented in a matrix notation

$$\eta = X\beta + \sum_{i=1}^{q} Z_i u_i \tag{10}$$

where \mathbf{Z}_i and \mathbf{u}_i are specified in equation (6).

To formally present GLMMs, we start with the conditional distribution of \boldsymbol{y} given \boldsymbol{u} . It is typical to assume that vector \boldsymbol{y} consists of conditionally independent elements, each coming from the exponential family (or similar to the exponential family),

$$y_{j}|\boldsymbol{u} \sim \text{ indep. } f_{Y_{j}|\boldsymbol{u}}(y_{j}|\boldsymbol{u}),$$

$$f_{Y_{j}|\boldsymbol{u}}(y_{j};\boldsymbol{\theta},\phi|\boldsymbol{u}) = \exp\left[\frac{y_{j}\theta_{j} - b(\theta_{j})}{a_{j}(\phi)} + c(y_{i},\phi)\right]. \tag{11}$$

It can be verified that the conditional mean of y_j is related to θ_j in equation (11) by the identity $\mu_j = \partial b(\theta_j)/\partial \theta_j$. The transformation of the mean allows us to model the fixed and the random factors by a linear model

$$E[y_j|\mathbf{u}] = \mu_j,$$

$$g(\mu_j) = \eta_j = \mathbf{X}_j \mathbf{\beta} + \mathbf{Z}_j \mathbf{u}.$$
(12)

Finally, we assign a distribution to the random effects

$$U \sim \phi_U(u),$$
 (13)

which completes the specification of GLMMs. It is often, if not always, assumed that \boldsymbol{u} come from a normal distribution.

1.3.1.5 An example—Poisson log-linear mixed-effect model

We will illustrate one specific type of GLMMs—the Poisson log-linear mixed-effect model in the context of RNA-seq experiments. Suppose we have RNA-Seq expression profiles (in the form of counts) randomly selected from three experiments conducted in three different labs. For each experiment, there are two treatments and two biological replicates within each treatment. We are not interested in the specific levels of treatment, but focus more on the overall variation of treatments. In this sense, the treatment effects are also considered as random. For a single gene, let $Y_{jkl} \sim \text{Poisson}(\mu_{jkl})$ be the read count for jth biological sample from kth treatment of lth experiment. The link function $\eta_{jkl} = \log(\mu_{jkl})$ relates the mean μ_{jkl} to the linear predictors by equation (12)

$$\log(\mu_{jkl}) = \log(N_{jkl}R_{jkl}) + \xi + a_j + b_{k(j)} + \epsilon_{jkl}, \tag{14}$$

where $N_{jkl}R_{jkl}$ is the normalized library size (total number of read counts mapped to the genome), j = 1, 2, 3, k = 1, 2 and l = 1, 2; the random terms $a_j \sim N(0, \sigma_1^2), b_{k(j)} \sim N(0, \sigma_2^2)$ and $\epsilon_{jkl} \sim N(0, \sigma_0^2)$ represent the experimental, treatment and sample ef-

fects respectively, and are mutually independent. If the observations are sorted by experiment and then by treatment nested in experiment, we can present the model in the form of equation (10), with $\boldsymbol{\beta} = (\log[N_{111}R_{111}] + \xi, \dots, \log[N_{223}R_{223}] + \xi)^T$, $\boldsymbol{u} = (\boldsymbol{a}, \boldsymbol{b}, \boldsymbol{\epsilon})$ and

Then it follows that

$$oldsymbol{\Sigma} = \sigma_1^2 oldsymbol{Z_1} oldsymbol{Z_1} oldsymbol{I_1} oldsymbol{Y_2} + \sigma_2^2 oldsymbol{Z_2} oldsymbol{Z_2} oldsymbol{Y_2} + \sigma_0^2 oldsymbol{I_{12}} = \left[egin{array}{ccc} oldsymbol{\Sigma}_d & oldsymbol{O} & oldsymbol{\Sigma}_d & oldsymbol{O} \\ oldsymbol{O} & oldsymbol{O} & oldsymbol{\Sigma}_d \end{array}
ight],$$

where \mathbf{O} is a 4×4 matrix of 0 and

$$oldsymbol{\Sigma}_d = \left[egin{array}{ccccc} \sigma_1^2 + \sigma_2^2 + \sigma_0^2 & \sigma_1^2 + \sigma_2^2 & \sigma_1^2 & \sigma_1^2 \ \sigma_1^2 + \sigma_2^2 & \sigma_1^2 + \sigma_2^2 + \sigma_0^2 & \sigma_1^2 & \sigma_1^2 \ \sigma_1^2 & \sigma_1^2 & \sigma_1^2 + \sigma_2^2 + \sigma_3^2 & \sigma_1^2 + \sigma_2^2 \ \sigma_1^2 & \sigma_2^2 & \sigma_1^2 + \sigma_2^2 & \sigma_1^2 + \sigma_2^2 + \sigma_0^2 \ \end{array}
ight]$$

The challenge due to the complexity of GLMM is the estimation of parameters. In the next section, we will summarize current available methods for estimating parameters and variance components.

1.3.2. Parameter estimation of generalized linear mixed models

There are three general approaches for estimating parameters under GLMM settings [91, Chapter 7]: (i) using numerical method to approximate the integrals for the likelihood functions and obtaining the estimating equations; (ii) linearization of the conditional mean and then iteratively applying linear mixed model techniques to the approximated model; (iii) Bayesian approach.

In the following discussion, we assume conditional distribution of Y given u is $f_Y(y|\beta, u)$, the link function is $\eta = g(\mu)$, and η relates the covariates by equation (12). We also assume that the random term u have some distribution $U \sim \phi_U(u|\Sigma)$.

1.3.2.1 Likelihood function approach

It is straightforward to write down the likelihood function of Y by first obtaining the joint likelihood of (Y, u) and then integrating out the random term u,

$$L(\mathbf{Y}|\boldsymbol{\beta}, \boldsymbol{\Sigma}) = \int f(\mathbf{y}|\boldsymbol{\beta}, \boldsymbol{u}) \phi(\boldsymbol{u}|\boldsymbol{\Sigma}) d\boldsymbol{u}.$$
 (15)

A major challenge in estimating GLMMs is the integration in equation (15) over the *n*-dimensional distribution of \boldsymbol{u} . Numerical approximation are usually used in evaluating the integral. In this part we will discuss the *Gauss-Hermite* (GH) quadrature which is recognized as a higher order Laplace approximation [77]. Gauss-Hermite quadrature is used for integrals of the form $\int_{-\infty}^{\infty} f(x)e^{-x^2}dx$ that can be approximated by a weighted sum of f(x):

$$\int_{-\infty}^{\infty} f(x)e^{-x^2}dx \approx \sum_{i=1}^{m} w_i f(x_i)$$
(16)

In equation (16), x_i 's are the zeros of mth order Hermite polynomial

$$H_m(x) = (-1)^m \exp(\frac{x^2}{2}) \frac{d^m}{dx^m} \exp(-\frac{x^2}{2})$$

and w_i are the corresponding weights. For a Hermite polynomial of degree m, x_i and w_i can be calculated as

$$x_i = i \text{th zero of } H_m(x), \quad w_i = \frac{2^{m-1} m! \sqrt{\pi}}{m^2 [H_{m-1}(x_i)]^2}.$$
 (17)

Equation (16) gives the exact numerical value for all polynomials up to degree of 2m-1. An improved version of the regular Gauss-Hermite quadrature is to center and scale the quadrature points by the empirical Bayes estimate of the random effects and the Hessian matrix from the Bayes estimate suboptimization [77]. This procedure is called *Adaptive Gauss-Hermite* (AGH) quadrature [96].

The AGH quadrature starts with maximizing the integrand $h(\boldsymbol{u}|\boldsymbol{y},\boldsymbol{\beta},\boldsymbol{\Sigma}):=f(\boldsymbol{y}|\boldsymbol{\beta},\boldsymbol{u})\phi(\boldsymbol{u}|\boldsymbol{\Sigma})$ in equation (15) with respect to the random term \boldsymbol{u} . The resulting estimate $\hat{\boldsymbol{u}}^{(n)}$ at iteration n is the joint posterior modes for the random effects. Because $\boldsymbol{\beta}$ and $\boldsymbol{\Sigma}$ are unknown, they are replaced by the current estimates $\hat{\boldsymbol{\beta}}^{(n)}$ and $\hat{\boldsymbol{\Sigma}}^{(n)}$. The Hessian matrix $\hat{\boldsymbol{H}}^{(n)}$ can be obtained by evaluating the second order partial derivatives of $\log(h(\boldsymbol{u}|\boldsymbol{y},\hat{\boldsymbol{\beta}}^{(n)},\hat{\boldsymbol{\Sigma}}^{(n)}))$ at $\hat{\boldsymbol{u}}^{(n)}$. Consequently, $\hat{\boldsymbol{\Omega}}^{(n)}=-\hat{\boldsymbol{H}}^{(n)}$ is the estimated covariance matrix for the random effects posterior modes. It follows from equation (15) that for the ith cluster

$$L(\mathbf{Y}_{i}|\boldsymbol{\beta},\boldsymbol{\Sigma}) = \int f(\boldsymbol{y}_{i}|\boldsymbol{\beta},\boldsymbol{u})\phi(\boldsymbol{u}|\boldsymbol{\Sigma})d\boldsymbol{u} = \int \frac{f(\boldsymbol{y}_{i}|\boldsymbol{\beta},\boldsymbol{u})\phi(\boldsymbol{u}|\boldsymbol{\Sigma})}{\phi(\boldsymbol{u}|\hat{\boldsymbol{u}}^{(n)},\hat{\boldsymbol{\Omega}}^{(n)})}\phi(\boldsymbol{u}|\hat{\boldsymbol{u}}^{(n)},\hat{\boldsymbol{\Omega}}^{(n)})d\boldsymbol{u}.$$
(18)

Let m be the number of quadrature points (i.e., the order of the Hermite polynomial) in each dimension for each random effect term, and Q the number of random effects. If $\mathbf{x} = (x_1, \dots, x_m)$ are the nodes for standard Gauss-Hermite quadrature, and $\mathbf{x}_j^* = (x_{j_1}, \dots, x_{j_Q})$ is a point on the Q dimensional quadrature grid, then the centered and

scaled nodes are

$$a_j^* = \hat{u}^{(n)} + \sqrt{2}[\hat{\Omega}^{(n)}]^{1/2}x_j^*$$
 (19)

The centered and scaled nodes, along with the Gauss-Hermite quadrature weights $\mathbf{w} = (w_1, \dots, w_m)$ are used to construct the Q dimensional integral of equation (18), approximated by

$$L(\boldsymbol{y}_{i}|\boldsymbol{\beta},\boldsymbol{\Sigma}) \approx \sum_{j_{1}=1}^{m} \cdots \sum_{j_{Q}=1}^{m} \frac{f(\boldsymbol{y}_{i}|\boldsymbol{\beta},\boldsymbol{a}_{j}^{*})\phi(\boldsymbol{a}_{j}^{*}|\boldsymbol{\Sigma})}{\phi(\boldsymbol{a}_{j}^{*}|\hat{\boldsymbol{u}}^{(n)},\hat{\boldsymbol{\Omega}}^{(n)})} w_{j_{1}} \cdots w_{j_{Q}}$$

$$= (2)^{Q/2} |\hat{\boldsymbol{\Omega}}^{(n)}|^{1/2} \sum_{j_{1}=1}^{m} \cdots \sum_{j_{Q}=1}^{m} \left[f(\boldsymbol{y}_{i}|\boldsymbol{\beta},\boldsymbol{a}_{j}^{*})\phi(\boldsymbol{a}_{j}^{*}|\boldsymbol{\Sigma}) \prod_{k=1}^{Q} w_{jk} \exp(x_{jk}^{2}) \right].$$

$$(20)$$

Thus the multidimensional unbounded integral is approximated by a finite summations. Now that the likelihood has the form of equation (20), a number of numerical methods (e.g. Newton-Raphson or Fisher's scoring) can be used to estimate (β, Σ) .

It should be noted, however, that as the number of dimension Q increases, the computational burden for approximating equation (20) grows exponentially since the total number of nodes is m^Q . Therefore it is difficult to implement AGH procedure with more than three random effects [14].

1.3.2.2 Estimation based on linearization

Under GLMM framework, we have some conditional distribution of Y given u. Without loss of generality, we assume

$$E[\mathbf{Y}|\mathbf{u}] = \boldsymbol{\mu} = g^{-1}(\boldsymbol{\eta}) = g^{-1}(\mathbf{X}\boldsymbol{\beta} + \mathbf{Z}\mathbf{u}),$$

$$Var[\mathbf{Y}|\mathbf{u}] = \mathbf{S},$$
(21)

where $\boldsymbol{u} \sim N(\boldsymbol{0}, \boldsymbol{\Sigma})$. The linearization is done by Taylor expansion of equation (21) about estimates $\boldsymbol{\eta}$. Two approaches proposed by Breslow and Clayton [16]—the penalized quasi-likelihood (PQL) and the marginal quasi-likelihood (MQL)—may be

used for this purpose.

Penalized Quasi-likelihood The PQL procedure uses a first order Taylor expansion of $\boldsymbol{\beta}$ and \boldsymbol{u} , at $\tilde{\boldsymbol{\beta}}$ and $\tilde{\boldsymbol{u}}$, respectively

$$g^{-1}(\boldsymbol{\eta}) \approx g^{-1}(\hat{\boldsymbol{\eta}}) + \tilde{\Omega}_{PQL}(\boldsymbol{\eta} - \tilde{\boldsymbol{\eta}}),$$
 (22)

where $\tilde{\Omega}_{PQL}$ is an $n \times n$ diagonal matrix whose (i, i) entry is $\partial g^{-1}(\eta_i)/\partial \eta_i$ evaluated at $\tilde{\eta} = X\tilde{\beta} + Z\tilde{u}$. Multiplying both sides by $\tilde{\Omega}_{PQL}^{-1}$, we can rearrange equation (22) as

$$\boldsymbol{X}\boldsymbol{\beta} + \boldsymbol{Z}\boldsymbol{u} \approx \tilde{\boldsymbol{\Omega}}_{PQL}^{-1}[g^{-1}(\boldsymbol{\eta}) - g^{-1}(\tilde{\boldsymbol{\eta}})] + \boldsymbol{X}\tilde{\boldsymbol{\beta}} + \boldsymbol{Z}\tilde{\boldsymbol{u}}.$$
 (23)

Note that the right hand side of equation (23) is just the expected value, conditioning on $\tilde{\beta}$ and \tilde{u} , of the pseudo-response

$$\tilde{\mathbf{Y}} = \tilde{\mathbf{\Omega}}_{POL}^{-1}[\mathbf{Y} - g^{-1}(\tilde{\boldsymbol{\eta}})] + \mathbf{X}\tilde{\boldsymbol{\beta}} + \mathbf{Z}\tilde{\boldsymbol{u}}, \tag{24}$$

whose variance-covariance matrix given u is

$$\operatorname{Var}[\tilde{\boldsymbol{Y}}|\boldsymbol{u}] = \tilde{\Omega}_{POL}^{-1} \operatorname{Var}[\boldsymbol{Y}|\boldsymbol{u}] \tilde{\Omega}_{POL}^{-1} = \tilde{\Omega}_{POL}^{-1} \boldsymbol{S} \tilde{\Omega}_{POL}^{-1}.$$
(25)

Then we can consider the model

$$\tilde{Y} = X\beta + Zu + \epsilon \tag{26}$$

which is a linear mixed model with pseudo response $\tilde{\boldsymbol{Y}}$ with covariance matrix

$$\boldsymbol{W} = \operatorname{Var}[\tilde{\boldsymbol{Y}}|\boldsymbol{u}] = \boldsymbol{Z}\boldsymbol{\Sigma}\boldsymbol{Z'} + \tilde{\boldsymbol{\Omega}}_{PQL}^{-1}\boldsymbol{S}\tilde{\boldsymbol{\Omega}}_{PQL}^{-1}. \tag{27}$$

Model (26) has exactly the same form as the linear mixed models (see Section 1.3.1.2), except that an estimate of (β, u) is needed for calculating the pseudo-response \tilde{Y} in

equation (24). An iterative procedure can be used to estimate the parameters in model (26) by substituting raw data \boldsymbol{y} for $\tilde{\boldsymbol{y}}$ and identity matrix \boldsymbol{I} for \boldsymbol{S} as starting values. Techniques for fitting LMM such as REML can be readily applied to estimate variance components $\boldsymbol{\Sigma}$, upon which $\hat{\boldsymbol{W}}$ is calculated. The estimate for $\boldsymbol{\beta}$ is given by

$$\hat{\boldsymbol{\beta}} = (\boldsymbol{X}^T \hat{\boldsymbol{W}}^{-1} \boldsymbol{X})^{-1} \boldsymbol{X}^T \hat{\boldsymbol{W}}^{-1} \boldsymbol{X} \tilde{\boldsymbol{y}}, \tag{28}$$

and the estimate for the random effects is

$$\hat{\boldsymbol{u}} = \hat{\boldsymbol{\Sigma}} \boldsymbol{Z} \hat{\boldsymbol{W}}^{-1} (\tilde{\boldsymbol{y}} - \boldsymbol{X} \hat{\boldsymbol{\beta}}). \tag{29}$$

Then the pseudo-response is updated and the procedure is repeated until convergence is reached for fixed effects and variance components. Note that equation (29) estimates a vector of random effects. For this reason, PQL is also referred to as subject-specific estimate procedure.

Marginal Quasi-likelihood One of the motivation for MQL is that usually one is more interested in estimating the marginal mean of the response than estimating the conditional mean as is done by equation (29) in PQL. Since $E[\boldsymbol{\eta}|\boldsymbol{u}] = \boldsymbol{X}\boldsymbol{\beta} + \boldsymbol{Z}\boldsymbol{u}$, the unconditional mean is $E[\boldsymbol{\eta}] = E[E(\boldsymbol{\eta}|\boldsymbol{u})] = \boldsymbol{X}\boldsymbol{\beta}$. A first-order Taylor expansion of $E[\boldsymbol{Y}|\boldsymbol{u}]$ about $\boldsymbol{X}\boldsymbol{\beta}$ is given by

$$E[\mathbf{Y}|\mathbf{u}] = g^{-1}(\boldsymbol{\eta}) \approx g^{-1}(\mathbf{X}\boldsymbol{\beta}) + \tilde{\Omega}_{MQL}(\boldsymbol{\eta} - \mathbf{X}\boldsymbol{\beta})$$
(30)

where $\tilde{\Omega}_{MQL}$ is evaluated at $X\beta$ (recall that for PQL, $\tilde{\Omega}_{PQL}$ is evaluated at $X\beta + Zu$). The unconditional expected value of Y is approximately $g^{-1}(X\beta)$ by equation (30). The variance of Y can then be derived from the relation Var(Y) = E[Var(Y|u)] + Var[E(Y|u)], which yields

$$Var[Y] = \tilde{\Omega}_{MOL} Z \Sigma Z' \tilde{\Omega}'_{MOL} + S_{n_0}. \tag{31}$$

A linearization performed at $\eta_0 = X\beta_0$ leads to

$$g^{-1}(\boldsymbol{\eta}) \approx g^{-1}(\boldsymbol{X}\boldsymbol{\beta_0}) + \tilde{\Omega}_{MQL}(\boldsymbol{\eta} - \boldsymbol{X}\boldsymbol{\beta_0}),$$
 (32)

and multiplying both sides by $\tilde{\Omega}_{MQL}^{-1}$, we can re-express equation (32) as

$$\boldsymbol{X}\boldsymbol{\beta} + \boldsymbol{Z}\boldsymbol{u} \approx \tilde{\boldsymbol{\Omega}}_{MOL}^{-1}[g^{-1}(\boldsymbol{\eta}) - g^{-1}(\boldsymbol{\eta}_0)] + \boldsymbol{X}\boldsymbol{\beta}_0. \tag{33}$$

Defining the pseudo-response \tilde{Y}_{MQL} as

$$\tilde{\mathbf{Y}}_{MQL} = \tilde{\mathbf{\Omega}}_{MQL}^{-1}[\mathbf{Y} - g^{-1}(\boldsymbol{\eta}_0)] + \mathbf{X}\boldsymbol{\beta}_0, \tag{34}$$

we next consider the linear mixed model

$$\tilde{Y}_{MQL} = X\beta + Zu + \epsilon,$$

where $Var(\boldsymbol{\epsilon})$ is given by equation (31). The estimating procedure for fixed effect parameter $\boldsymbol{\beta}$ and variance component $\boldsymbol{\Sigma}$ is the same as those in the PQL approach. Note that the pseudo-response is not a function of \boldsymbol{u} any more, so updating this quantity does not require calculating the random effects \boldsymbol{u} . Accordingly, the MQL approach is also referred to as *population-averaged* estimate approach.

Breslow and Lin [17] and Pinheiro and Chao [97] point out that the PQL approach may lead to asymptotically biased estimates and hence to inconsistency. It is not recommended to use simple PQL method in practice.

1.3.2.3 Bayes approach

As mentioned earlier, for models with higher dimensional integrals, it is not practical to evaluate the likelihood function by AGH procedure. For mixed models, a typical strategy is to treat the random effects to be missing data. Following this idea, the

problem of estimating variance components associated with random effects can be simplified. Denote the *complete data* as $\mathbf{v} = (\mathbf{y}, \mathbf{u})$, the log-likelihood of \mathbf{v} can be expressed as

$$\log \pi(\boldsymbol{\beta}, \boldsymbol{\Sigma} | \boldsymbol{v}) = \log f(\boldsymbol{y} | \boldsymbol{\beta}, \boldsymbol{u}) + \log \phi(\boldsymbol{u} | \boldsymbol{\Sigma})$$
(35)

The optimal solution for parameters in equation (35) can be obtained by *expectation-maximization* (EM) algorithm. The EM algorithm consists of two steps, readily implemented as follows:

1. **E-Step**. At (k+1)th iteration given $\boldsymbol{\beta}^{(k)}$ and $\boldsymbol{\Sigma}^{(k)}$, calculate

$$E_{\boldsymbol{\beta}^{(k)}}[\log f(\boldsymbol{\beta}, \boldsymbol{\Sigma}|\boldsymbol{v})|\boldsymbol{y}] = Q_1(\boldsymbol{\beta}, \boldsymbol{\beta}^{(k)}),$$

$$E_{\boldsymbol{\Sigma}^{(k)}}[\log \phi(\boldsymbol{\Sigma}|\boldsymbol{v})|\boldsymbol{y}] = Q_2(\boldsymbol{\Sigma}, \boldsymbol{\Sigma}^{(k)}).$$
(36)

2. **M-Step**. Maximize Q_1 and Q_2 to update $\boldsymbol{\beta}^{(k+1)}$ and $\boldsymbol{\Sigma}^{(k+1)}$.

The **E** and **M** steps are alternated until convergence is reached. Unfortunately, the expectations in equation (36) cannot be computed in closed form for GLMMs. However, they may be approximated by *Markov chain Monte Carlo* (MCMC). In light of this, McCulloch [85] developed a Monte Carlo EM (MCEM) algorithm.

The Metropolis-Hastings algorithm is used for drawing samples from difficult-tocalculate density functions. For Metropolis algorithm, a proposal distribution $g(\boldsymbol{u})$ is selected, from which an initial value of \boldsymbol{u} is drawn. The new candidate value $\boldsymbol{u}' = (u_1, u_2, \dots, u_{k-1}, u_k', u_{k+1}, \dots, u_Q)$, which has all elements the same as previous values expect the kth, is accepted (as opposed to keeping the previous value) with probability

$$A_k(\boldsymbol{u}', \boldsymbol{u}) = \min \left\{ 1, \ \frac{f(\boldsymbol{u}'|\boldsymbol{y}, \boldsymbol{\beta}, \boldsymbol{\Sigma})g(\boldsymbol{u})}{f(\boldsymbol{u}|\boldsymbol{y}, \boldsymbol{\beta}, \boldsymbol{\Sigma})g(\boldsymbol{u}')} \right\}.$$
(37)

If we choose $g(\mathbf{u}) = \phi(\mathbf{u}|\mathbf{\Sigma})$, the ratio term in equation (37) can be simplified to

$$\frac{f(\boldsymbol{u}'|\boldsymbol{y},\boldsymbol{\beta},\boldsymbol{\Sigma})g(\boldsymbol{u})}{f(\boldsymbol{u}|\boldsymbol{y},\boldsymbol{\beta},\boldsymbol{\Sigma})g(\boldsymbol{u}')} = \left[\frac{f(\boldsymbol{u}',\boldsymbol{y}|\boldsymbol{\beta},\boldsymbol{\Sigma})g(\boldsymbol{u}')}{f(\boldsymbol{y}|\boldsymbol{\beta},\boldsymbol{\Sigma})}\phi(\boldsymbol{u}|\boldsymbol{\Sigma})\right] / \left[\frac{f(\boldsymbol{u},\boldsymbol{y}|\boldsymbol{\beta},\boldsymbol{\Sigma})}{f(\boldsymbol{y}|\boldsymbol{\beta},\boldsymbol{\Sigma})}\phi(\boldsymbol{u}'|\boldsymbol{\Sigma})\right] \\
= \frac{f(\boldsymbol{y}|\boldsymbol{u}',\boldsymbol{\beta},\boldsymbol{\Sigma})\phi(\boldsymbol{u}'|\boldsymbol{\Sigma})\phi(\boldsymbol{u}|\boldsymbol{\Sigma})}{f(\boldsymbol{y}|\boldsymbol{u},\boldsymbol{\beta},\boldsymbol{\Sigma})\phi(\boldsymbol{u}|\boldsymbol{\Sigma})\phi(\boldsymbol{u}'|\boldsymbol{\Sigma})} \\
= \frac{f(\boldsymbol{y}|\boldsymbol{u}',\boldsymbol{\beta},\boldsymbol{\Sigma})\phi(\boldsymbol{u}|\boldsymbol{\Sigma})\phi(\boldsymbol{u}'|\boldsymbol{\Sigma})}{f(\boldsymbol{y}|\boldsymbol{u},\boldsymbol{\beta},\boldsymbol{\Sigma})}$$
(38)

The MCEM procedure combines the EM steps and Metropolis algorithm in estimating the fixed parameters and variance components, summarized as follows:

- 1. Choose the starting value of $\boldsymbol{\beta}^{(0)}, \boldsymbol{\Sigma}^{(0)}$. Set b=0
- 2. Generate the sequence $\boldsymbol{u}^{(1)}, \boldsymbol{u}^{(2)}, \dots, \boldsymbol{u}^{(B)}$ from the conditional distribution of \boldsymbol{u} given y with Metropolis algorithm.
- 3. Maximize $\sum_{b=1}^{B} \log f(\boldsymbol{y}|\boldsymbol{u}^{(b)},\boldsymbol{\beta})/B$ and $\sum_{b=1}^{B} \log \phi(\boldsymbol{u}^{(b)}|\boldsymbol{\Sigma})/B$ to obtain $\boldsymbol{\beta}^{(m+1)}$ and $\boldsymbol{\Sigma}^{(m+1)}$
- 4. Iterate between step 2 and 3 until convergence is reached.

This method can be easily extended to allow for multiple random effects. Yet the advantage comes at a price. A major drawback of MCEM is the computational intensity. First, the convergence of EM algorithm is usually very slow, especially at the neighborhood of maximum of marginal likelihood. Second, the chain in Metropolis algorithm has to run long enough for reliable estimation.

In the Bayes framework, there are other alternatives to estimate the parameters and variance components, for example, *Monte Carlo Newton-Raphson* (MCNR) [85] and MCMC [47].

1.3.2.4 Example of estimating parameters

We will demonstrate the estimating procedure with the Poisson log-linear mixedeffect model discussed in Section 1.3.1. The estimation procedure starts from the joint density function of $\mathbf{Y} = (Y_{jkl})'$ given $\boldsymbol{\mu} = (\mu_{jkl})'$,

$$f(\mathbf{y}|\mathbf{\mu}) = \prod_{j,k,l} f(y_{jkl}|\mu_{jkl}) = \prod_{j,k,l} \frac{[\mu_{jkl}]^{y_{jkl}} \exp(-\mu_{jkl})}{y_{jkl}!}.$$
 (39)

A re-expression of equation (14) in matrix form gives

$$\log \mu = X\beta + Z_1a + Z_2b + I_{12}\epsilon.$$

Therefore $\mu \sim \log N(\mu_0, \Sigma)$ where $\mu_0 = \xi + \log(NR)$ and $\Sigma = \sigma_1^2 Z_1 Z_1' + \sigma_2^2 Z_2 Z_2' + \sigma_0^2 I_{12}$. The density function of μ is then

$$f(\boldsymbol{\mu}|\boldsymbol{\mu}_0, \boldsymbol{\Sigma}) = \prod_{j,k,l} \mu_{jkl}^{-1} \cdot \frac{1}{\sqrt{(2\pi)^{12}|\boldsymbol{\Sigma}|}} \exp[-\frac{1}{2} (\log \boldsymbol{\mu} - \boldsymbol{\mu}_0)^T \boldsymbol{\Sigma}^{-1} (\log \boldsymbol{\mu} - \boldsymbol{\mu}_0)].$$
(40)

Since $Y_{jkl} \sim \text{Poisson}(\mu_{jkl})$, by combining equation (39) and (40), we obtain the joint distribution of Y and μ ,

$$f(\boldsymbol{y}, \boldsymbol{\mu} | \boldsymbol{\mu}_0, \boldsymbol{\Sigma}) = \frac{1}{\sqrt{(2\pi)^{12}|\boldsymbol{\Sigma}|}} \exp[-\mathbf{1}^T \boldsymbol{\mu} - \frac{1}{2} (\log \boldsymbol{\mu} - \boldsymbol{\mu}_0)^T \boldsymbol{\Sigma}^{-1} (\log \boldsymbol{\mu} - \boldsymbol{\mu}_0)] \prod_{jkl} \frac{[\mu_{jkl}]^{y_{jkl}-1}}{y_{jkl}!}.$$
(41)

Therefore we can obtain the likelihood function of or the marginal distribution of Y by integrating out the random components u,

$$L(\xi, \sigma_1^2, \sigma_2^2, \sigma_0^2 | \boldsymbol{y}) = f(\boldsymbol{y} | \boldsymbol{\xi}, \boldsymbol{\Sigma}) = \int_{\boldsymbol{a}, \boldsymbol{b}, \boldsymbol{\epsilon}} f(\boldsymbol{y}, \boldsymbol{a}, \boldsymbol{b}, \boldsymbol{\epsilon} | \boldsymbol{\mu}_0, \boldsymbol{\Sigma}) d\boldsymbol{a} d\boldsymbol{b} d\boldsymbol{\epsilon}.$$
(42)

The integral in equation (42) can be approximated by adaptive Gaussian-Hermite (AGH) quadrature or MCMC. For AGH quadrature, we first approximate the like-

lihood by equation (20) and then estimate $\boldsymbol{\theta} = (\xi, \sigma_0^2, \sigma_1^2, \sigma_2^2)'$ by maximizing the resulting likelihood. The R package lme4 [11] has an inbuilt function glmer() for this purpose. The MCMC procedure has also been implemented in several packages, such as the Rstan [116] and the MCMCPack [83].

 $\begin{tabular}{ll} \bf 2. & \bf Identifying \ stably \ expressed \ genes \ from \ multi-\\ & \bf ple \ RNA-Seq \ data \ sets \end{tabular}$

Bin Zhuo, Sarah Emerson, Jeff Chang and Yanming Di

Abstract

We examined RNA-Seq data on 211 biological samples from 24 different experiments carried out by different labs and identified genes that are stably expressed across biological samples, treatment conditions, and experiments. We fit a Poisson log-linear mixed-effect model to the read counts for each gene and decomposed the total variance into between-sample, between-treatment and between-experiment variance components. Identifying stably expressed genes is useful for count normalization and differential expression analysis. The variance component analysis that we explore here is a first step towards understanding the sources and nature of the RNA-Seq count variation. When using a numerical measure to identify stably expressed genes, the outcome depends on multiple factors: the background sample set and the reference gene set used for count normalization, the technology used for measuring gene expression, and the specific numerical stability measure used. Since DE is measured by relative frequencies, we argue that DE is a relative concept. We advocate using an explicit reference gene set for count normalization to improve interpretability of DE results, and recommend using a common reference gene set when analyzing multiple RNA-Seq experiments to avoid potential inconsistent conclusions.

2.1. Introduction

RNA sequencing (RNA-Seq) has become the technology of choice for transcriptome profiling over the last few years. The exponential growth in RNA-Seq studies have produced a large amount of *Arabidopsis thaliana* (Arabidopsis) data under a variety of experimental/environmental conditions. It is only natural to begin exploring how the large amount of existing data sets can help the analysis of future data. In this paper, we discuss identifying stably expressed genes from multiple existing RNA-Seq data sets based on a numerical measure of stability. We envision that such identified stably expressed genes could be used as a reference set or prior information for count normalization and differential expression (DE) analysis of future RNA-Seq

data sets obtained from similar or comparable experiments. We also fit a random-effect model to the read counts for each gene and decompose the total variance to into between-sample, between-treatment and between-experiment variance components. The variance component analysis is a first step towards understanding the sources and nature of the RNA-Seq count variation. To illustrate our methods, we examined RNA-Seq data on 211 Arabidopsis samples from 24 different experiments carried out by different labs and identified genes that were stably expressed across biological samples, experimental or environmental conditions, and experiments (labs).

A reference set of stably-expressed genes will be useful for count normalization. A key task of RNA-Seq analysis is to detect DE genes under various experimental or environmental conditions. Count normalization is needed to adjust for differences in sequencing depths or library sizes (total numbers of mapped reads for each biological sample) due to chance variation in sample preparation. In DE analysis, gene expression levels are often estimated from relative read frequencies. For this reason, normalization is also needed to account for the fact that non-differentially expressing genes may exhibit an apparent reduction or increase in relative read frequencies due to the respective increased or decreased relative read frequencies of truly differentially expressing genes. Many existing normalization methods, such as the trimmed mean of M-values normalization method (TMM) [106] and Anders and Huber's normalization [3], assume that the majority of the genes within an experiment are not DE, and examine the sample distribution of the fold changes between samples. If the experiment condition can affect expression levels of more than half of the genes, many of the existing normalization methods may be unreliable [78, 130]. This difficulty could be alleviated if one could identify a set of stably expressed genes whose expression levels are known or expected to not vary much under different experimental conditions. Our idea is to identify such a reference set based on a large number of existing data sets.

Our basic intuition is that a numerical quantification of expression stability—which typically measures certain aspects of RNA-Seq count variation—can be more

reliably estimated by using more data sets. There is, however, a caveat to this idea: as pointed out by Fernandes et al. [35] and Hruz et al. [52], universally stably expressed genes may not exist. Hruz et al. [52] showed that a subset of stably expressed genes from a specific biological context may have more variability than other genes if examined across a broader range of samples and conditions. Many studies have shown that stably expressed genes are subject to change from one experiment to another due to different experimental protocols, different tissue types, or other varying conditions [50, 102]. The top 100 stably expressed genes in the Arabidopsis developmental series of Czechowski et al. [24] shared only 3 genes with the top 50 stably expressed genes identified from Arabidopsis seed samples by Dekkers et al. [25]. In this study, we try to balance generality and specificity by identifying different reference gene sets for different tissue types of Arabidopsis.

We can also consider that when a normalization method is applied to a single data set, it effectively specifies an implicit reference set of stably expressed genes (those genes that have the least variation after normalization). From this perspective, we can view commonly used normalization techniques as using an internally identified reference set of genes. In contrast, what we are proposing is that one could alternatively identify a reference set externally by looking at past data sets. The internally and externally identified reference gene sets will provide different contexts for the DE analysis: in other words, one can choose to answer different scientific questions by using different reference sets. In any case, we advocate making the reference set explicit during a DE analysis and using a common reference set when analyzing multiple datasets.

We want to clarify that having stable gene expression is not equivalent to maintaining a stable biological function. Often times, we may not understand the biological functions of genes with numerically stable expression measures. From an operational point of view, however, numerical stability is more tractable. In pre-genomic era, the so-called "house-keeping genes" were often considered to be candidate reference genes for normalization [5, 19]. House-keeping genes are typically constitutive genes that maintain basic cellular function, and therefore are expected to express at relatively constant levels in non-pathological situations. However, many studies have shown that house-keeping genes are not necessarily stably expressed according to numerical measures (a review can be found in Huggett et al. [56] and reference therein). For example, in the microarray analysis of Arabidopsis, Czechowski et al. [24] showed that traditional house-keeping genes such as ACT2, TUB6, EF-1 α are not stably expressed, and thus not good reference genes for normalization. Spike-in genes have also been considered as reference genes for normalization, but Risso et al. [104] showed that spike-in genes are not necessarily stably expressed according numerical measures either.

In this paper, we identify stably expressed genes from RNA-Seq data sets based on a numerical measure—the sum of three variance components estimated from a mixedeffect model. For microarray data, there have been many efforts to numerically find stably expressed genes by quantifying the variation of measured expression levels across a large number of microarray data sets. For example, Andersen et al. [5] used a linear mixed model to estimate the between-group and within-group variances from expression profiles of microarray experiments, and then quantified expression stability by combining the two variance components using a Bayesian formulation. Czechowski et al. [24] measured the expression stability of each gene using the coefficient of variation (CV). Genes with lower CVs are considered more stably expressed. By investigating 721 arrays under 323 conditions throughout development, Czechowski et al. [24] suggested stably expressed (reference) genes under different experimental conditions for Arabidopsis. Stamova et al. [115], Dekkers et al. [25], Gur-Dedeoglu et al. [46], and Frericks and Esser [38] screened a large number of microarray data sets to identify stably expressed genes in human blood, Arabidopsis seed, breast tumor tissues, and mice respectively. Validation experiments [24, 25, 56, 115] showed that these genes are more stably expressed than traditional house-keeping genes.

Our vision is that identifying stably expressed genes is the first step towards integrative analysis of multiple RNA-Seq experiments. It will help to answer fundamental questions related to comparability, reproducibility and replicability of RNA-Seq experiments.

2.2. Methods

In Section 2.2.1, we describe the steps for collecting and processing RNA-Seq data sets from Arabidopsis experiments. In Section 2.2.2, we discuss count normalization methods and how to apply them to a subset of stably expressed genes. In Section 2.2.3, we introduce the generalized linear mixed model (GLMM, McCulloch and Neuhaus [86]) for estimating three variance components from RNA-Seq data: the between-sample, between-treatment and between-experiment variances. We define the total variance measure for expression stability as the sum of estimated variance components. In Section 2.2.4, we review the CV and M-value measures for gene expression stability.

2.2.1. RNA-Seq data collection and processing

2.2.1.1 Overview of the RNA-Seq data sets

We examined RNA-Seq data from 49 Arabidopsis experiments stored on the NCBI GEO repository (see more details below). After screening, we retained data from 211 biological samples in 24 experiments. To illustrate our methods for finding stably expressed genes, we divided the experiments into three groups: the seedling group contains 60 Arabidopsis seedling samples from 9 experiments; the leaf group contains 60 Arabidopsis leaf samples from 5 experiments; the multi-tissue group contains 91 samples from 10 experiments on multiple tissue types (shoot apical, root tip, primary root, inflorescences and siliques, hypocotyl, flower, carpels, aerial tissue, epidermis, seed). Table 2.1 summarizes the basic information about the three groups (see Appendix C.1 for more details about the 24 experiments).

Group	# experiments	# treatments	# samples	# genes
seedling	9	27	60	22207
leaf	5	28	60	20967
multi-tissue	10	39	91	23611

Table 2.1: Summary statistics for the three groups of Arabidopsis samples.

To find stably expressed genes in each group, we processed the raw sequencing data and summarized the results as count matrices of mapped RNA-Seq short reads (see details below). We removed genes with low mean numbers (less than 3) of mapped read counts for all experiments. Such genes tend to be more prone to sequencing noise, less interesting to biologists, and also cause convergence issues when fitting statistical models. Many other researchers (such as Anders et al. 4) recommend removing such genes before analyzing RNA-Seq data. The number of remaining genes in each group is also summarized in Table 2.1.

2.2.1.2 Details of the data processing steps

The Gene Expression Omnibus (GEO) repository at National Center for Biotechnology Information (NCBI, http://www.ncbi.nlm.nih.gov/) stores raw sequencing data from a large number of RNA-Seq experiments. For this study, we restrict our attention to Arabidopsis experiments satisfying the following conditions: 1. Ecotype = "Columbia" (we kept only the Columbia samples from experiments that compare Columbia samples to other ecotypes); 2. There are at least two treatments and 2 biological replicates for each treatment; 3. Library strategy= "RNA-Seq"; 4. Library source = "transcriptomic"; 5. Library selection= "cDNA"; 6. Library layout = "Single end"; 7. If there are repeated measurements over time, we choose samples from one time point.

We screened all the Arabidopsis experiments available from the NCBI GEO repository up to May 31, 2015 and downloaded raw RNA-Seq data (Sequence Read Archive files) from 49 experiments.

We assembled our own in-house pipeline to process all the raw RNA-Seq data: align the raw RNA-Seq reads to the reference genome and summarize the read counts at the gene level. In the GEO repository, the mapped read counts are unavailable for some experiments and the available ones are from different processing pipelines. Our pipeline, implemented using the software R [100], is summarized as follows:

- 1. Convert the Sequence Read Archive (SRA) files to FASTQ files using the NCBI SRA Toolkit ([67], version 2.3.5-2).
- 2. Download the reference genome

Arabidopsis_thaliana.TAIR10.22.dna.toplevel.fa

from the Ensembl plants FTP server (http://plants.ensembl.org/info/data/ftp/index.html) and build index using build() function from Subread aligner (RSubread, version 1.16.2, Liao et al. 73) in the software R [100]. The index allows fast retrieval of the sets of positions in the reference genome where the short reads are more likely to align.

- Align short reads in FASTQ files to the Arabidopsis reference genome using the align() function from Rsubread.
- 4. Summarize the read counts at the gene level using the featureCounts() function from the Subread aligner and store the read counts as data matrix. The annotation file

Arabidopsis_thaliana.TAIR10.22.gtf

is downloaded from Ensembl plants FTP server. The multi-mapping or multioverlapping reads were not counted.

Subread aligner is a recently developed sequence mapping tool that adopts a seed-andvote paradigm to map the RNA-Seq short reads to the genome. It breaks each short read into a series of overlapping segments called subreads and uses the subreads to vote on the optimal genome location of the original read. The subreads are shorter and can be mapped to the genome much faster. Compared to other aligners such as Bowite 2 [64] or BWA [69], Subread aligner is both faster and more accurate [73]. We compared results from the above pipeline to results from a pipeline described in Anders et al. [4] over several RNA-Seq experiment data, and Rsubread was more than three times faster and successfully mapped more reads to the reference genome. For researchers familiar with R, it also has the advantage that it is completely implemented in R.

We divided the experiments into three groups as summarized in Table 1. As an additional data quality control measure, we keep an experiment only when it has mapping quality (number of successfully mapped reads divided by total number of reads) $\geq 50\%$ for all samples. Then within each group, we computed an initial set of normalization factors from all samples combined using the method described in Section 2.2.2. An experiment is retained only when the normalization factors of all samples in the experiment are between 0.50 and 1.50. If the initial estimated normalization factor is too different from 1 for a sample, it often indicates that the read counts distribution in the corresponding sample is markedly different from the distributions of the rest of the samples. Such samples demand additional attention before being incorporated in the studies that we intend to do.

2.2.2. Count normalization

As explained in the introduction, count normalization is needed when analyzing RNA-Seq data to 1) adjust for differences in sequencing depths or library sizes; 2) to adjust for the apparent changes in relative read frequencies of non-DE genes that occur as a consequence of changes in relative read frequencies of truly DE genes.

For the second type of adjustment, we follow Anders and Huber's method [3] for estimating normalization factors. Let y_{ij} denote the read count for *i*th gene of the *j*th sample (m genes and n samples in total). We first create a pseudo-reference

sample where each gene's expression value is the geometric mean expression over all real samples for that gene,

$$y_{i,0} = (\prod_{j=1}^{n} y_{i,j})^{1/n}, i = 1, \dots, m.$$
 (43)

Next we calculate the median fold-change in relative frequency between each sample j and the pseudo-reference sample,

$$R'_{j} = \text{median}\left(\frac{y_{1,j}/N_{j}}{y_{1,0}/N_{0}}, \dots, \frac{y_{m,j}/N_{j}}{y_{m,0}/N_{0}}\right),$$
 (44)

where N_j is the library size for sample j (the sum of RNA-Seq counts mapped to all genes retained in each sample). Finally, the normalization factor R_j for sample j is calculated as

$$R_j = \frac{R'_j}{(\prod_{i=1}^n R'_i)^{1/n}}. (45)$$

Using the estimated normalization factors, the relative frequencies will be computed as y_{ij}/N_jR_j , which we will call the normalized relative frequency for gene i in sample j. The assumption made here is that the median fold change between normalized relative frequencies in two samples should be 1. In other words, this normalization method assumes that the majority of genes are not DE. The NBPSeq package [27] has an inbuilt function for this procedure and it will be used for count normalization in this paper. With the estimates from equation (45), we see that the median fold change in normalized relative frequencies between each sample and the pseudo-reference sample will be set to 1:

median
$$\left(\frac{y_{1,j}/N_jR_j}{y_{1,0}/N_0R_0}, \dots, \frac{y_{m,j}/N_jR_j}{y_{m,0}/N_0R_0}\right) = 1,$$
 (46)

where $R_0 = (\prod_{j=1}^n R'_j)^{-1/n}$.

We can apply equation (44) to a subset of reference genes to estimate normaliza-

tion factors. In doing so, effectively, the median fold change in equation (46) among the reference genes will be set to 1 in each sample j. Other normalization methods may make different assumptions than Anders and Huber's, but some assumptions of a similar nature seem unavoidable. For example, the TMM method of Robinson et al. [106] is based on a similar principle: assuming the majority of the genes are not DE. The TMM method can be applied to a subset of genes selected based on an initial screening of mean expression level and fold changes. In TMM method, one can also specify certain quantile (instead of the median) of the fold changes to be 1.

In this paper, we will identify stably expressed genes from multiple data sets based on numerical measure and use them as reference for estimating normalization factors (from equations (44) and (45)). However, to identify the stably expressed genes, we first need a set of initially estimated normalization factors. To tackle this circular dependence, we use a one-step iteration method to estimate the normalization factors:

- 1. First, we use all the genes to calculate the initial normalization factors;
- 2. Then, we fit a GLMM to each gene and estimate the total variance measure, incorporating the initial normalization factors as an offset term (see Section 2.2.3);
- 3. Next, we select the top 1000 stably expressed genes based on the total variance measure estimated from step 2 above, and use them as reference genes to recalculate the normalization factors.

In practice, this one-step method seems to be adequate and further iterations will only slightly change the set of 1000 stably expressed genes. For example, for the multi-tissue group of experiments, if we were to run one more iteration of steps 2 and 3, there would be 946 overlapping genes between the top 1000 genes from the first iteration and those from the second iteration.

2.2.3. Poisson log-linear mixed-effects regression model and the total variance measure of expression stability

We fit a Poisson log-linear mixed-effects regression model to the RNA-Seq counts mapped to each gene and measure gene expression stability using a total variance measure.

Let Y_{ijkl} be the number of RNA-Seq reads mapped to gene i in sample j from treatment group k in experiment l. We will fit regression models to each gene separately and suppress subscript i from the model equations. For each gene, we fit a Poisson log-linear mixed-effects regression model

$$Y_{jkl} \sim \text{Poisson}(\mu_{jkl}),$$
 (47)

$$\log(\mu_{jkl}) = \log(R_{jkl}N_{jkl}) + \xi + \alpha_l + \beta_{k(l)} + \epsilon_{jkl}, \tag{48}$$

which is a specific type of generalized linear mixed model (GLMM, McCulloch and Neuhaus [86]). In equation (48), N_{jkl} and R_{jkl} are the library size and normalization factor discussed in Section 2.2.2. We will call $R_{jkl}N_{jkl}$ the normalized library size. The parameter ξ is a fixed-effect term for the baseline log mean of the relative counts (counts divided by the normalized library sizes). The values α , β , and ϵ represent the experiment effect, the treatment effect (nested within each experiment), and the sample effect respectively. We view α , β and ϵ as random effects and assume that they are independent and follow normal distributions:

$$\alpha_l \sim N(0, \sigma_{\text{experiment}}^2), \quad \beta_{k(l)} \sim N(0, \sigma_{\text{treatment}}^2), \quad \epsilon_{jkl} \sim N(0, \sigma_{\text{sample}}^2),$$
 (49)

where $\sigma_{\text{experiment}}^2$, $\sigma_{\text{treatment}}^2$ and σ_{sample}^2 are called *variance-components*—they quantify the overall variances of the corresponding random effect terms.

The sample effect ϵ represents the extra-Poisson variation in read counts among

samples in the same treatment group and σ_{sample}^2 plays a similar role as the overdispersion parameter in a negative binomial model (Anders and Huber 3, Di et al. 26). The experiment effect, α , accounts for all sources of variation at the experiment level, including differences in lab personnel and conditions, day light hours, age of the plants, temperature, sequencing platform, and other unidentified sources. The contributions from these different experiment-level sources are often difficult to separate statistically. We treat the experiment effect α as a random effect because we view the collected experiments as a random sample from the pool of all Arabidopsis RNA-Seq experiments. We also treat the treatment effect β as a random effect. In a DE test, β is usually considered as a fixed-effect term. Here for evaluation of expression stability, we are not interested in the specific levels of the individual β 's and focus more on the overall variation of β under a range of treatment types.

We define the stability measure as the estimated total variance,

$$\hat{\sigma}^2 = \hat{\sigma}_{\text{sample}}^2 + \hat{\sigma}_{\text{treatment}}^2 + \hat{\sigma}_{\text{experiment}}^2. \tag{50}$$

The parameters $(\xi, \sigma_{\text{experiment}}^2, \sigma_{\text{treatment}}^2, \sigma_{\text{sample}}^2)$ are estimated using the glmer() function of the R package lme4 ([11], version 1.1.7), which uses a Gaussian-Hermite quadrature to approximate the likelihood function. We rank all the genes according to their values of $\hat{\sigma}^2$ in increasing order (smallest first), and consider highly ranked (e.g., top 1000) genes to be stably expressed.

Normal models (equation (49)) are commonly assumed for the random effects in the GLMM settings. The normality assumption is likely a simplification of reality, yet it is a good starting point and should be adequate for finding genes with low total variation—the stably expressed ones.

2.2.4. Other stability measures

The assessment of gene expression stability depends on the specific stability measure used. Czechowski et al. [24] and Dekkers et al. [25] used the coefficient of variation (CV) measure, computed as *standard deviation devided by mean*, to find stably expressed genes from microarray data.

The M-value in geNorm [124] is a well-cited measure. For a set of m_0 genes, the M-value measure works as follows: First, the pairwise variation between gene i_1 and gene i_2 is calculated as the standard deviation of the log fold changes between their expression levels across all the n samples:

$$V_{i_1,i_2} = st.dev \left\{ \log \left(\frac{y_{1,i_1}}{y_{1,i_2}} \right), \dots, \log \left(\frac{y_{n,i_1}}{y_{n,i_2}} \right) \right\}.$$

Next, the M-value for gene i is defined as the average pairwise variation between gene i and all other genes:

$$M_i = \frac{\sum_{k \neq i} V_{i,k}}{m_0 - 1},$$

In the Results section, we compare the *M*-value to the total variance measure on RNA-Seq data from the multi-tissue group experiments, and compare the stably expressed genes identified from these two measures to those identified from microarray data using the CV measure.

2.3. Results

In Section 2.3.1, we summarize the stably expressed genes identified from three different experiment groups and emphasize that stability is context-dependent. In Section 2.3.2, we show that traditional house-keeping genes are not necessarily stably expressed according to our numerical measure, and that microarray data and RNA-Seq data may often give different sets of stably expressed genes. In Section 2.3.3, we further demonstrate that when using a numerical measure to quantify gene expression

stability, the outcome will depend on the specific numeric measure used. These points should be intuitive, but they are not often emphasized in practice. In Section 2.3.4, we discuss results from our variance component analysis. In Section 2.3.5, we discuss how to use the identified stably expressed genes for count normalization.

2.3.1. Stably expressed genes

Using the total variance, $\hat{\sigma}^2$, from the GLMM (see equation (48) in Section 2.2.3) as a stability measure, we identified stably expressed genes from the three groups of experiments described in Section 2.2.1: the group of seedling experiments, the group of leaf experiments, and the group of experiments on different tissue types (see Table 2.1 for a summary). As we mentioned in the Introduction, absolutely stably expressed genes may not exist. Choosing different sample sets as reference allows us to identify stably expressed genes for different biological contexts.

In Tables 1–3 in the online supplementary materials, we summarize the top 1000 most stably expressed genes in each group. In Figure 2.1, we provide the histograms of the mean Count Per Million (CPM) for the 1000 most stably expressed genes identified in each group. For each gene, the CPM is computed as

$$\frac{\text{count} \times 10^6}{\text{normalized library size}}$$
 (51)

in each sample and the mean is computed over all samples.

The lists of the top 1000 genes in the three groups share 104 genes in common (see supplementary material for details). These genes are stably expressed under a wide range of experimental conditions and in different tissue types, and thus may be worth further study. This list of 104 genes has significant overlap with the top 100 stably expressed genes identified by Czechowski et al. [24] from a developmental series of microarray samples: 9 out of these 104 genes (see Table 4 in the supplementary material for details),

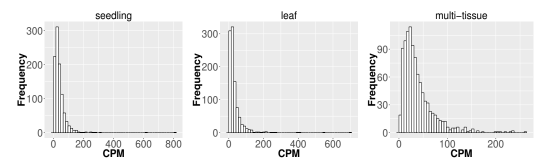


Figure 2.1: Histograms of the mean CPM (see equation (51)) for the top 1000 most stably expressed genes identified from the seedling (left), leaf (middle) and multi-tissue (right) groups using the total variance measure $\hat{\sigma}^2$. The mean CPM is computed over all samples within each respective group. Note that the x and y axis scales differ between the three plots.

AT1G13320, AT1G54080, AT2G20790, AT2G32170, AT3G10330, AT4G24550, AT5G26760, AT5G46210, AT5G46630,

appeared in the list of the top 100 stably expressed genes out of 14000 genes they examined (the probability is 4.8×10^{-9} for a list of 104 genes random selected from a set of 14000 genes to have an overlap of size 9 or more with a pre-selected list of 100 genes). In particular, one gene, AT1G13320, is in all but one of the ten lists of top 500 stably expressed genes identified by Czechowski et al. [24] for different experimental and experimental conditions (the only exception is the set of diurnal series), and is also identified by Hong et al. [50] as a stably expressed gene under all but one of the six experimental conditions they examined. This gene is ranked 159 (top 0.7%), 112 (top 0.5%), 513 (top 2.2%) in the three groups we examined, respectively, according to our stability measure. This gene is a subunit of protein phosphatase type 2A complex and is involved in regulation of phosphorylation and regulation of protein phosphatase type 2A activity. It has been used as a reference gene for normalization in many papers (e.g., Baron et al. [9], Bournier et al. [15]; these two papers cited Czechowski et al. [24] as reference).

2.3.2. Comparison to house-keeping genes and stably expressed genes identified from microarray data

Czechowski et al. [24] discussed the expression stability of house-keeping genes and showed that the house-keeping genes are not stably expressed according to their numerical measure. In particular, they compared the expression profiles of five traditional house-keeping genes (AT1G13440, AT3G18780, AT4G05320, AT5G12250, AT5G60390) and five genes (AT1G13320, AT5G59830, AT2G28390, AT4G33380 and AT4G34270) that they identified as stably expressed according to the CV measure from a developmental series of microarray experiments (see Figure 1 of that paper). In Figure 2.2, we compare the expression profiles of these 10 genes from Czechowski et al. [24] to the expression profiles of five genes (AT1G63110, AT1G79280, AT3G27530, AT4G02560, AT5G53540) that we randomly selected from the top 100 most stably expressed genes identified from the multi-tissue group RNA-Seq data according the total variance $\hat{\sigma}^2$. For each of the 15 genes, Figure 2.2 shows the expression levels measured in CPM over 91 samples in the eight experiments in the multi-tissue group, and Table 2.2 summarizes the variance components estimated from the GLMM in 2.2.3.

The five house-keeping genes show large total variation with all three variance-components relatively large as compared to the other 10 genes. This is consistent with Czechowski's observation that house-keeping genes are not necessarily stably expressed according to a numerical measure. Three of the five stably-expressed genes identified by Czechowski are among the top 1000 stably-expressed genes according to our stability measure, the total variance $\hat{\sigma}^2$. Czechowski et al. identified those five genes from microarray data and different experiments. It is not too surprising those genes might not be the most stable in RNA-Seq experiments: the two technologies differ in many aspects including coverage and sensitivity.

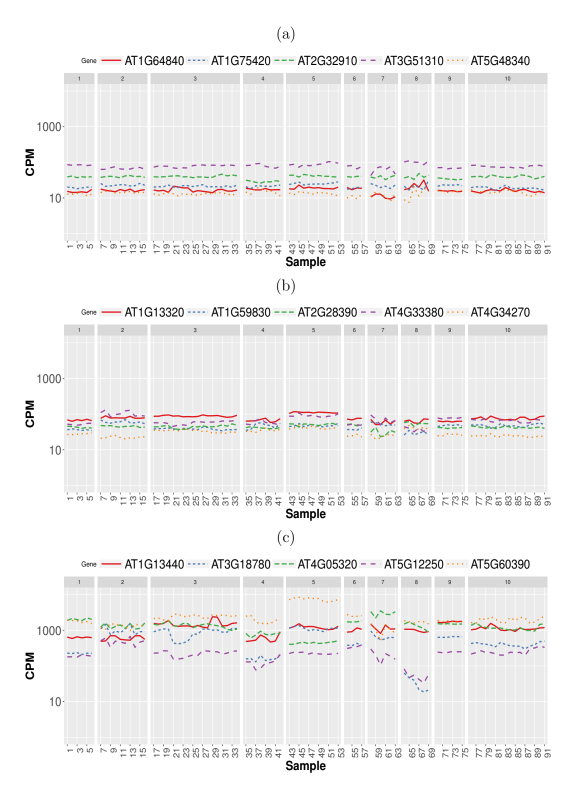


Figure 2.2: Expression profiles of 15 genes—as measured by RNA-Seq CPM—across 91 samples in the multi-tissue group. The 15 genes include (from top to bottom) a) five stably expressed genes (randomly selected out of the top 100) identified from the multi-tissue group RNA-Seq data using the total variance measure $\hat{\sigma}^2$, b) five stably expressed identified by Czechowski et al. [24] according to the CV measure from a developmental series of microarray experiments, and c) five traditional house-keeping genes (HKG) discussed in Czechowski et al. [24].

Table 2.2: Variance components estimated from the multi-tissue group RNA-Seq data for the 15 genes in Figure 2.2 (identified from different sources). Columns 3–5 are the estimated variance components. Column 6 lists the stability ranking according to the total variance $\hat{\sigma}^2$ in the multi-tissue group.

Source	Gene	betweeen-	between-	between-	Rank
		sample	${\it treatment}$	experiment	
	AT1G75420	0.0012	0.0014	0.0050	5
	AT5G48340	0.0042	0.0019	0.0074	46
RNA-Seq	AT2G32910	0.0007	0.0019	0.0113	53
	AT1G64840	0.0051	0.0008	0.0095	72
	AT3G51310	0.0028	0.0025	0.0100	73
	AT2G28390	0.0034	0.0000	0.0111	62
	AT1G13320	0.0036	0.0003	0.0258	513
Microarray	AT4G34270	0.0063	0.0000	0.0365	1074
·	AT1G59830	0.0044	0.0039	0.0370	1211
	AT4G33380	0.0103	0.0016	0.0747	3404
HKG	AT1G13440	0.0234	0.0058	0.1375	6562
	AT5G60390	0.0267	0.0068	0.2270	8867
	AT4G05320	0.0123	0.0094	0.2690	9409
	AT5G12250	0.0313	0.0128	0.3262	10589
	AT3G18780	0.0375	0.0211	1.0313	14951

2.3.3. Factors affecting stability ranking

The previous two subsections demonstrate that when using a numerical measure to quantify gene expression stability, the outcome is dependent on 1) the biological context reflected in the reference sample set used and 2) the technology used for measuring gene expression. It should also be intuitive, and we will further clarify in the second half of this subsection, that the stability ranking is also dependent on 3) the specific numerical measure used. In this section, we will first compare the lists of stably-expressed genes identified under different scenarios where one or more of the above three factors differ. We then further discuss the subtle roles played by the specific stability measure and the reference gene set by comparing the total variance $\hat{\sigma}^2$ measure from the GLMM (see equation (48)) to the M-value measure used in the geNorm method [124]. Last, we discuss the effect of an iterative elimination procedure used by geNorm.

We look at an additional five lists of stably expressed genes identified under different scenarios and examine how each of these five lists overlaps with the top stably-expressed genes identified from the multi-tissue group of RNA-Seq experiments according to the total variance measure $\hat{\sigma}^2$ (see Section 2.2.3). The five lists are:

- L_1 : 100 top stably expressed genes from the multi-tissue group according to the M-value in geNorm (applied to (count + 1)) of Vandesompele et al. [124];
- L_2 : 100 top stably expressed genes from the seedling group according to the total variance $\hat{\sigma}^2$ from the GLMM;
- L_3 : 100 top stably expressed genes from the leaf group according to the total variance $\hat{\sigma}^2$ from the GLMM;
- L_4 : 100 stably expressed genes identified from a developmental series of microarray experiments by Czechowski et al. [24] using the CV measure (see Section 2.2.4);

 L_5 : 50 stably expressed genes identified by Dekkers et al. [25] from microarray seed experiments using the CV measure.

For each list L_i above, we measure how it overlaps with the top stably expressed genes (the reference set) from the multi-tissue group using the recall percentage

$$\frac{\#\{L_i \cap \text{ reference set}\}}{\#\{L_i\}} \times 100,\tag{52}$$

where #{} denotes the number of elements in the list. In Figure 2.3, we plot the recall percentage versus the number of top stably-expressed genes we selected as reference from the multi-tissue group.

We have the following observations:

- 1. The list L_1 is identified from the same set of RNA-Seq experiments as the reference sets, but using a different stability measure (M-value in geNorm). This list has significant overlap with the top stably-expressed genes identified using the total variance measure: 29 and 98 out of the 100 genes from the list L_1 are among the top 100 and 1000 most stably-expressed genes, respectively, from the multi-tissue group identified using the total variance measure.
- 2. The lists L_2 and L_3 are identified from different sets of RNA-Seq experiments (leaf and seedling experiments) using the same stability measure as used for the reference sets. The lists L_4 and L_5 are identified from microarray experiments (a developmental series and a seed group) and using the CV measure. The overlapping (recall) percentages are still statistically significant, but much less than in the case of L_1 . This shows that differences in tissue type and in measuring technology both influence the expression stability ranking, and to comparable degrees. The lists L_3 and L_5 have the least overlapping percentages with the reference sets. These lists are identified from a leaf group and a seed group respectively. Our understanding is that the leaf group and the seed group are

more biologically homogeneous than the multi-tissue group and thus provide very different biological contexts for evaluating expression stability.

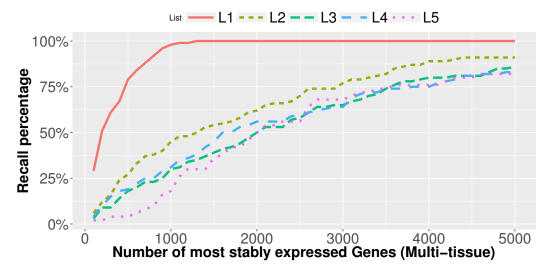


Figure 2.3: Comparison of top stably expressed genes identified under different scenarios. We choose the top 100 stably expressed genes as described in L_1 – L_4 , and the top 50 stably expressed genes in L_5 (see Section 2.3.3). and plot the recall percentages between these lists and the top most stably expressed genes identified from the multi-tissue group according to the total variance measure. The x-axis is the number of most stably expressed genes in multi-tissue group according to the total variance measure, and the y-axis shows the recall percentage (see equation (52)) for each of the five lists.

When applied to the same set of samples, the M-value and total variance measure $\hat{\sigma}^2$ give similar expression stability ranking: the rank correlation is 0.97 (see also, observation 1 above). We point out that the reason is because the M-value and normalization step needed for computing our total variance measure have similar fundamental assumptions. The basic principle behind the M-value is that the expression ratio of two stably-expressed genes should be identical in all samples. In formula, it means that the expression values of two stably-expressed genes i_1 , i_2 in

any two samples j_1 , j_2 should satisfy

$$\frac{y_{i_1,j_1}}{y_{i_2,j_1}} = \frac{y_{i_1,j_2}}{y_{i_2,j_2}}. (53)$$

Our total variance measure $\hat{\sigma}^2$ is estimated from normalized data. The basic assumption in the normalization step is that majority of genes are not DE. In formula, it means for any stably-expressed gene i_1 , its expression level as measured by the relative frequency should be stable across all samples,

$$\frac{y_{i_1,j_1}}{S_{j_1}} = \frac{y_{i_1,j_2}}{S_{j_2}},\tag{54}$$

where S_{j_1} to S_{j_2} are the normalized library sizes (i.e., $R_j N_j$ in equation (48)). This implies for any two stably-expressed genes i_1 and i_2

$$\frac{y_{i_1,j_1}}{y_{i_1,j_2}} = \frac{y_{i_2,j_1}}{y_{i_2,j_2}} = \frac{S_{j_1}}{S_{j_2}}. (55)$$

The first equation in (55) is equivalent to equation (53). (In practical application of both methods, the stability of any single gene is evaluated by comparing its expression to a set of reference genes. See the Method section 2.2.2 for more details.)

In practice, the geNorm program [124] is frequently used to rank a set of reference genes identified from other methods. An iterative elimination procedure is used along with the M-value to determine the final ranks of the expression stability: after each iteration, the gene receiving the largest M-value will be removed and a new set of M-values will be computed for the remaining genes, and the iteration will go on until there are only two genes left. We did not use such an iterative procedure in the comparisons above (i.e., we only computed one set of M-values for all genes).

This iterative elimination procedure creates an extra layer of complexity that is not well explored in literature. We use a toy example below to illustrate one subtle aspect of the iterative elimination procedure. In this example, we consider the expression values of 7 genes in two samples shown in Table 2.3. When M-value is used to rank all 7 genes, the initial ranking of expression stability is given in column 4 of the table: gene 7 is the least stable and genes 4 and 5 are considered the most stable ones. Once genes 6 and 7 are eliminated, however, the recalculated M-values will rank genes 1–3 as more stable than genes 4 and 5 (see column 5 of Table 2.3). The root cause of this reversal of ranking is that when an iterative elimination procedure is used, effectively, the reference gene set is changing after each iteration: in the initial ranking, the expression patterns genes 4 and 5 are close to the "middle of the pack" and thus considered as the most stable, and the expression patterns of genes 1–3 and genes 6 and 7 are considered relatively more extreme; once genes 6 and 7 are removed, however, the "middle of the pack" is shifted towards the expression patterns of genes 1–3, and thus genes 1–3 become the most stably expressed. With this understanding, one could and should make a conscious decision on whether such a behavior as described above is desirable or not.

The point we want to emphasize is that gene stability is a relative concept and the stability ranking depends on which set of genes we use as reference. In an iterative elimination procedure, the reference gene set will change after each iteration. The procedure can thus give surprising results and the adoption of it in practice should not be automatic.

2.3.4. Sources of variation

For each gene, the GLMM (equation (48) of section 2.2.3) allows us to decompose total count variance into between-sample, between-treatment and between-experiment variance components. The estimated variance components tell us how much each component contributes to the overall count variation. Table 2.4 summarizes the percentages—averaged over all genes—of the total variance attributable to each of the three components for three groups of RNA-Seq samples (seedling, leaf and multitissue groups in Section 2.2.1). Figure 2.4 shows the histograms of the percentages.

Table 2.3: A toy example showing the effect of iterative elimination. Columns 2 and 3 represent expression levels for seven genes in two samples, column 4 is the stability ranking of genes by M-value without iterative elimination, and column 5 is the ranking after two geNorm iterations.

	Raw Counts		Rank	
Gene	sample 1	sample 2	rank 1	rank 2
Gene1	1	1	3	1
Gene2	1	1	3	1
Gene3	1	1	3	1
Gene4	1	2	1	4
Gene5	1	2	1	4
Gene6	1	3	6	
Gene7	1	4	7	
Library Size	7	14		

Figure 2.5 shows the stacked bar plot of variance components estimated from the multi-tissue group for 20 genes randomly selected from the top 1000 stably expressed genes and 20 genes randomly selected from 23611 genes. As expected, the between-experiment variance component, on average, explains the largest proportion of the total variation. The between-experiment variation is relatively smaller among the leaf samples, indicating that the leaf samples are more homogeneous. There is more variation in the relative percentages of total variance explained by the between-sample and between-treatment variance components. In principle, the between-treatment variation will be greater when there is a higher proportion of DE genes or when the samples are more homogeneous. In practice, the between-sample variance depends greatly on what samples are used as biological replicates.

2.3.5. Reference gene set for normalization

Once we have ranked the genes according to our numerical stability measure (i.e, the total variance measure, $\hat{\sigma}^2$), one application is to use an explicit set of most stably expressed genes as reference genes for count normalization. This new approach allows

Table 2.4: Percentages—averaged over all genes—of the total variance attributable to each of the three variance components (between-sample, between-treatment, between-experiment) for the three groups of RNA-Seq samples (the seedling, the leaf and the multi-tissue groups).

Source	Seedling	Leaf	Multi-tissue
between-sample	7.2%	16.0%	7.6%
between-treatment	20.1%	28.0%	5.1%
between-experiment	72.6%	56.0%	87.3%

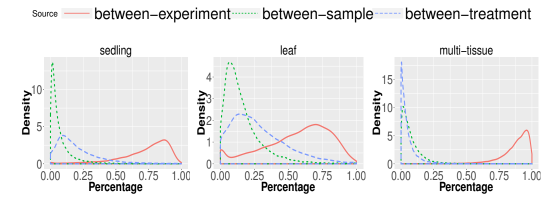


Figure 2.4: Distributions (over all genes) of the percentages of the total variance attributable to the between-sample, and between-treatment, or the between-experiment variance component, in the seedling, the leaf, and the multi-tissue groups.

investigators to prescribe a specific biological context for evaluating gene stability by choosing the most relevant reference samples and experiments when computing the stability measure. For example, the most stably expressed genes identified from the multi-tissue group and those identified from the seedling group will provide different biological contexts. In contrast, existing normalization approaches are often applied to the single data set under study, and thus provide a single, narrow context.

Even under a specific biological context, it is almost impossible to know whether the genes in any reference set are absolutely stably expressed, even though commonly used normalization methods often enforce some assumptions on the reference gene set: for example, when we use Anders and Huber's method to estimate the normalization factors based on a subset of reference genes, roughly speaking, the median fold change

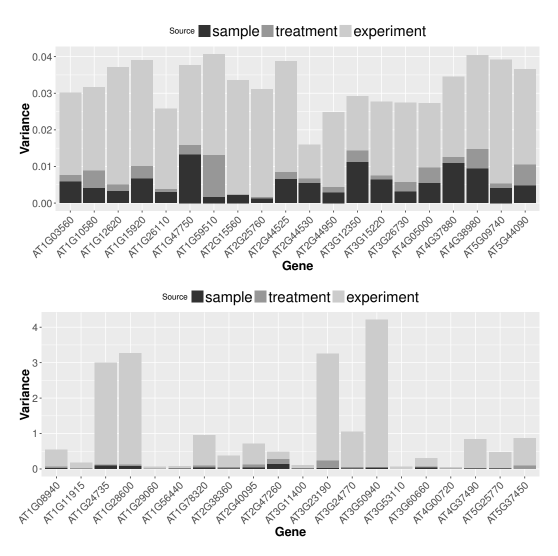


Figure 2.5: Stacked bar plots of the three variance components for selected genes in the multi-tissue group. Top: 20 genes randomly selected from top 1000 stably expressed genes; Bottom: 20 genes randomly selected from all the genes.

among the reference genes will be set to 1 (see Section 2.2.2 for more details). A subtle point we want to make is that since it is impossible to know how well such or similar assumptions on DE hold for a reference gene set, we can improve the interpretability of the DE test results by making the reference gene set explicit: we can slightly change our perspective and interpret all DE results as relative to the reference gene set. For example, a fold change of 2 inferred from the GLMM model can be interpreted as the

Table 2.5: A toy example for illustrating the importance of using a common explicit set of reference genes when comparing RNA-Seq data from multiple experiments. If a common reference gene set (e.g., genes 1–3) is used as reference for count normalization, we will notice that the DE behavior of gene 3 differs in the two experiments. If the two experiments are separately normalized using genes 1–3 as reference in experiment 1, but using genes 3–5 as reference in experiment 2, we may conclude that gene 3 is not DE in either group.

	Exp. 1		E	Exp. 2	
Gene	Control	Treatment	Control	Treatmetn	
1	10	20	10	20	
2	10	20	10	20	
3	10	20	10	10	
4	10	10	10	10	
5	10	10	10	10	

fold change of a gene is 2 times the true (but often unknowable) median fold change of the reference genes. When one estimates the normalization factors based on all genes, one is effectively specifying an implicit set of genes as a reference set. Our proposal is to make the reference set explicit and interpret DE results as relative to the reference gene set.

Interpreting the DE results as relative to an explicit reference set is especially beneficial when one wants to compare DE results from an experiment to previously published results. When the interest is in comparing different experiments, we recommend using a common reference set. For example, when two RNA-Seq data sets are separately normalized with different reference sets, a fold change of two observed in one experiment may not be directly comparable to a fold change of two observed in the other. This concern can be alleviated by using a common set of reference genes. We use a toy example to illustrate this point in Table 2.5 where we examine the mean counts for 5 genes in two two-group comparison experiments. If we use different reference gene sets for count normalization in the two experiments, for example, we use genes 1–3 as reference in experiment 1, but use genes 3–5 as reference in experiment

2, we may conclude that gene 3 is not DE in either experiment. If we use a common reference gene set—either genes 1–3 or genes 3–5—for normalization, however, we will be able to discover, in either case, that the DE behavior of gene 3 is different in the two experiments. Note that the DE conclusion in both experiments will depend on the reference genes used: if genes 1–3 are used as reference, gene 3 is not DE in experiment 1, but will be DE in experiment 2; if genes 3–5 are used as reference, gene 3 will be considered DE in experiment 1, but not DE in experiment 2. The point is, in either case, we will notice that the DE behavior of gene 3 is different between the two experiments. This information will be lost if one uses different reference sets to assess DE in the two experiments.

In practice, we recommend using the top 1000 most stably expressed genes for estimating normalization factors. The key is to avoid using too few (e.g., less than 10) or too many (e.g., using all genes) reference genes: intuitively, using too few, the estimates will be unstable; using too many, the results may be subject to influence from highly unstable genes. Our simple simulations suggest that using between 100 to 10000 genes seems to give stable results. In the first set of three examples, we used Anders and Huber's method (see equation (44)) to estimate normalization factors for samples in each of the seedling, leaf and multi-tissue groups of experiments (see Section 2.2.1). We used the top 10, 100, 1000, and 10000 stably expressed genes identified earlier (see Section 2.3.1 for details) as reference gene sets. Figure 2.6 shows the pairwise scatter plots and correlation coefficient between the normalization factors when different numbers of top stable genes are used as reference. A stronger correlation indicates the normalization factors estimated from the two settings are highly consistent. The plots and correlation coefficients suggest using between 100 and 1000 genes tend to give similar normalization factor estimates. We also used the top 10, 100, 1000, and 10000 stably expressed genes identified from the multi-tissue group as reference set for estimating normalization factors for a set of 48 root samples from a new experiment (GSE64410, [125]). The largest Pearson correlation 0.993 is between the normalization factors estimated using the top 100 and top 1000 stably expressed genes as reference. Based on the above observations, using 1000 most stably expressed genes as reference seems to be a reasonable heuristic rule.

2.4. Conclusion and discussion

In this paper, we advocate quantifying gene expression stability by applying a numerical stability measure to a large number of existing RNA-Seq data sets. Similar strategies have also been used by others to find stably expressed genes from microarray data. Since DE is measured by relative frequencies, we argue that DE is a relative concept and using an explicit reference gene set can improve interpretability of DE results, and furthermore, using a common reference gene set can avoid inconsistent conclusions when comparing multiple experiments (see Section 2.3.5).

It should be clear but worth emphasizing that when using a numerical measure to identify stably expressed genes, the outcome depends on multiple factors: the background sample set and the reference gene set used for count normalization, the technology used for measuring gene expression, and the specific numerical stability measure used. In this study, to illustrate our proposed methods, we identified three sets of stably expressed genes from three sets of Arabidopsis experiments. The major point is that stably expressed genes identified from different backgrounds will provide different biological contexts for evaluating differential expression. In practice, researchers can choose the specific context. A practical challenge in applying such a philosophy is that no two experiments will have identical settings, and researchers have to decide what experiments can be considered comparable. This is a difficult question; however, we believe it has to be asked from now on: biologists perform comparative experiments with the intent that the conclusions from a single experiment will be generalizable beyond the context of a single lab. If we do not understand comparability between different experiments, such generalization is impossible. Defining and characterizing comparability is a challenging topic that we would like to investi-

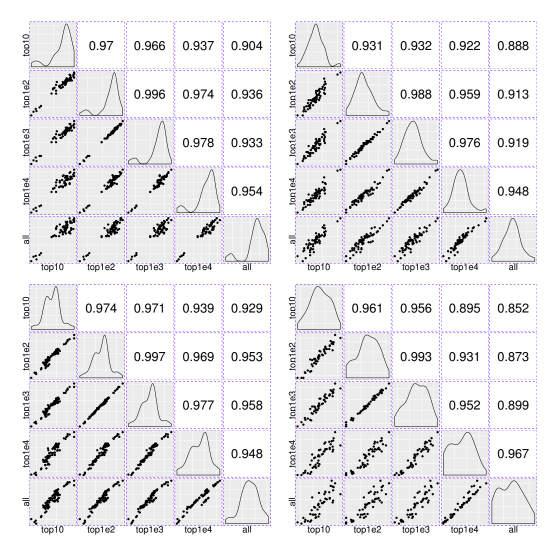


Figure 2.6: Matrices of scatter plots of normalization factors estimated using different reference gene sets. The upper-left, upper-right and lower-left plots show normalization factors estimated for the samples in the seedling, leaf, and multi-tissue groups correspondingly. In each case, the top 10, 100, 1000, and 10,000 stably expressed genes are used as reference to calculate the normalization factors. The lower right plot shows the normalization factors estimated for a new root experiment (GSE64410, with sample size 48) using the top 10–10,000 stably expressed genes identified from the multi-tissue group as reference. The normalization factors are estimated using the method described in Section 2.2.2.

gate more in the future.

To identify a set of stably expressed genes, our method still needs to estimate an

initial set of normalization factors, which requires that we must make assumptions about relative fold changes between samples. This kind of circular dependence seems unavoidable [124]. In this paper, we used a one-step iteration strategy to reduce the dependence on the initially estimated normalization factors. In future work, we intend to look at the genes through evolutionary genetics methods (e.g., 1001 genomes, [128]). For example, evolutionary genetics methods can help us test whether a gene is under negative, neutral, or positive selection and help us identify genes that are well conserved through the evolutionary history. We need to be mindful that a well conserved gene is not necessarily stably expressed, just like the house-keeping genes. However, it would be interesting to ask whether there is correlation between measures of expression stability and measures of conservativeness, and so on.

In the GLMM model we fit, the random effect terms such as the sample and treatment effects were modeled as normal random variables (Section 2.2.3). For the purpose of identifying stably expressed genes, this should be adequate, since we are mainly interested in the variances of these random effects (i.e., the variance components). In the future, it may also be of interest to model these random effects more accurately, for example, in order to build a prior distribution of the random effect terms for analyzing a new data set. A more careful examination of the individual data sets suggests that the between-sample variance varies greatly between experiments. Our observation suggests that different labs often have different understanding of what is deemed as "biological replicates".

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

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Supplemental Material

The online version of this article offers supplementary material, available to authorized users.

3. Test-statistic correlation and the correlation of underlying observed data

Bin Zhuo, Duo Jiang and Yanming Di

Abstract

We investigate the relationship between correlation among test statistics and the correlation of underlying observed data. In FDR control procedures and gene-set enrichment analysis, the sample correlation of observed data are often used to approximate the test-statistic correlation. We show, however, that such an approximation is only valid under limited settings. In particular, we derive a formula for the correlation between test statistics when they take a specific form. As a special case, we present the exact expression of test-statistic correlation for equal-variance two-sample t-test statistic under bivariate normal assumption.

3.1. Introduction

Between-gene correlations are commonly observed in gene expression data [32, 39, 55, 98, 117]. One key task of expression analysis is to detect differentially expressed (DE) genes whose expression levels are associated with experimental or environmental variables under study. In such a task, a test statistic is calculated for each gene to quantify the magnitude of DE. The test statistics may come from two-sample comparison or regression models for more complex designs. Between a pair of genes, when the observed expression levels are correlated, the test statistics calculated from the observed data will also be correlated [10, 31, 132]. This paper concerns the relationship between test-statistic correlation and the correlation of underlying observed data (e.g., expression levels).

The dependency among test statistics has brought methodological issues to multiple hypothesis testing procedures and gene-set analysis. Multiple hypothesis testing procedures determine a p-value cutoff by controlling false discovery rate (FDR) [12] or q-value [117]. Many FDR-control procedures are valid only when test statistics are independent [12] or have positive regression dependency [13]. Efron [31] showed in a simulation study that for a nominal FDR of 0.1, the actual FDR can easily vary by a

factor of 10 when correlation between test statistics exists. In a gene-set analysis, one tests for over-abundance of DE genes in a specified gene set (e.g., molecular pathways or gene ontologies) [42]. The correlation among DE test statistics, if not addressed appropriately, will undermine the validity of the gene-set test [39, 132].

A number of attempts have been made to account for test-statistic correlation in FDR control and gene-set analysis. Without replicating the experiment, we cannot directly estimate test-statistic correlation between genes since only a single test statistic value is available for each gene. For this reason, the sample correlation between observed data (after gene treatment effects accounted for) is often used as a surrogate. Efron [31] estimated a quantity called dispersion variate from the distribution of correlation among observed data, and then calculated the false discovery proportion (FDP) conditioning on this dispersion variate. Wu and Smyth [132] estimated a variance inflation factor (VIF) from the sample correlation of observed data and incorporated it into their parametric/rank-based gene-set test procedures. The same VIF was also used by Yaari et al. [134] in their gene-set test.

It is yet unclear when and to what extent the test-statistic correlations can be approximated by sample correlations of observed data. Barry et al. [10] showed by Monte Carlo simulation of gene expression data that a nearly linear relationship holds between test-statistic correlation and sample correlation of observed data for several forms of test statistics they examined. This Monte Carlo simulation results were cited by Wu and Smyth [132] and Yaari et al. [134] as a justification for estimating their VIF from observed data. Efron [31] also concluded through simulation that the distribution of z-value (the test statistic in that paper) correlation can be nearly represented by the distribution of sample correlation from observed data.

Some gene-set analysis methods based on sample-permutation implicitly assume that the joint distribution of test statistics will not change under sample permutation. Our results will reveal that such assumptions are invalid. The key issue is, as also pointed out by Efron [32], that sample permutation method has an extra assumption

(called "subset pivotality"), which states that the test statistics always follow the distribution they have under complete null that no gene is DE. We will show that that the test-statistic correlation depends on correlation between gene expression levels and also on DE status. Permutation of biological samples will change the DE status and thus alternate the correlations among (and thus the joint distribution of) the test statistics. As a consequence, in the presence of DE genes, the GSEA [119] procedure will not provide a valid null distribution for gene-set enrichment test. In a separate paper, we showed that several gene-set tests will give inflated or overly conservative type I errors in the presence of DE test correlations (see Chapter 4 for more detail).

We investigate the relationship between the correlation of test statistics and the correlation of underlying observed data. First, we present a formula for correlation between test statistics when they take a specific form and meet some assumption of independence. Then, under bivariate normal setting, we apply this formula to a special case where two-group comparison experiment is considered. We show that 1) the test-statistic correlation is equal to the correlation of underlying observed data when the test statistics are a linear combination of the observed data, and that 2) the test-statistic correlation is generally weaker than the correlation of underlying observed data when the test statistics are derived from two-sample t-test under normal assumption. We use simulation results to illustrate our findings.

3.2. General setup

Correlation is a statistical quantity used to assess a possible linear relationship between two random variables or two sets of data sets. The degree of correlation is measured by correlation coefficient, a scaler taking values on the interval [-1,1]. Correlation coefficient of +1 (-1) indicates perfect positive (negative dependence), while correlation coefficient of 0 implies no linear relationship between two random variables. Larger correlation coefficient (in absolute value) corresponds to stronger

linear correlation.

There are a number of ways to look at the correlation coefficient, many of which are special cases of *Pearson's correlation coefficient* [66]. For example, the *Kendall tau rank correlation coefficient* is computed as Pearson's correlation coefficient between the ranked variables. Throughout this paper, we will discuss Pearson's correlation under bivariate settings. We will restrict our interest to two types (following the notation of Lee Rodgers and Nicewander [66]) of Pearson's correlation coefficient. The first type of correlation, which we refer to as the correlation of underlying observed data or the population correlation, is the standardized covariance

$$\rho = \frac{\operatorname{Cov}(X, Y)}{\sqrt{\operatorname{Var}(X)\operatorname{Var}(Y)}} = \frac{E[(X - \mu_X)(Y_j - \mu_Y)]}{\sigma_X \sigma_Y}.$$
 (56)

In equation (56), μ_X and μ_Y are the expected values of random variables X and Y, and $\sigma_X < \infty$ and $\sigma_Y < \infty$ are the population standard errors. The second type of correlation, which we refer to as *sample correlation*, is a function of raw scores and means

$$r = \frac{\sum_{j} (x_{j} - \bar{x})(y_{j} - \bar{y})}{\sqrt{\sum_{j} (x_{j} - \bar{x})^{2} \sum_{i} (y_{j} - \bar{y})^{2}}},$$
(57)

where (\bar{x}, \bar{y}) is the vector of arithmetic mean of the observations. Fisher [37] proved that sample correlation r is a consistent estimator for the underlying correlation ρ .

Let (X_j, Y_j) be a bivariate random variable representing two features (genes) of sample j = 1, ..., m, and (x_j, y_j) the corresponding realization. We assume that the population mean of (X_j, Y_j) may differ across samples, but that the population covariance structure remains the same, that is,

$$E\begin{pmatrix} X_j \\ Y_j \end{pmatrix} = \begin{pmatrix} \mu_{X,j} \\ \mu_{Y,j} \end{pmatrix} \stackrel{\text{def}}{=} \boldsymbol{\mu}_j, \quad \text{for } j = 1, \dots, m$$
 (58)

and

$$\operatorname{Cov}\left(\begin{array}{c} X_j \\ Y_j \end{array}\right) = \left(\begin{array}{cc} \sigma_X^2 & \rho \sigma_X \sigma_Y \\ \rho \sigma_X \sigma_Y & \sigma_Y^2 \end{array}\right) \stackrel{\text{def}}{=} \mathbf{\Sigma}$$
 (59)

where ρ is the population correlation defined by equation (56). In addition, we assume independence across samples (note that independence implies no correlation, but not vise versa),

$$Cov(X_{j_1}, X_{j_2}) = Cov(Y_{j_1}, Y_{j_2}) = 0 \text{ for } j_1 \neq j_2$$
 (60)

In the context of gene expression study, the goal is to detect DE—whether the expression level of a gene is significantly correlated with the treatment or experimental variables. Let $\mathbf{a} := (a_1, \dots, a_m)^T$ be a vector for a contrast of interest, then DE detection for gene X can be statistically formulated as

$$H_0: \boldsymbol{a}^T \boldsymbol{\mu}_X = 0 \ Versus \ H_1: \boldsymbol{a}^T \boldsymbol{\mu}_X \neq 0, \tag{61}$$

where $\mathbf{X} = (X_1, \dots, X_m)^T$ and $\boldsymbol{\mu}_X = (\mu_{X,1}, \dots, \mu_{X,m})^T$. DE detection for gene Y can be obtained by applying the same contrast to $\mathbf{Y} = (Y_1, \dots, Y_m)$ (simply replacing the subscript X by Y in equation (61)). This hypothesis testing procedure usually results in a "t-test similar" test statistic, in which the numerator is a linear combination of \mathbf{X} and the denominator is its standard error. Without a loss of generality, we express the test statistics as follows

$$T_X = \frac{\boldsymbol{a}^T \boldsymbol{X}}{S_X}, \quad T_Y = \frac{\boldsymbol{a}^T \boldsymbol{Y}}{S_Y},$$
 (62)

where S_X and S_Y are the standard errors for $\boldsymbol{a}^T\boldsymbol{X}$ and $\boldsymbol{a}^T\boldsymbol{Y}$ respectively. Our main goal is to explore the relationship test-statistic correlation (equation (56))

$$\rho_T(m) = \operatorname{Corr}(T_X, T_Y), \tag{63}$$

and the underlying correlation of corresponding observed data

$$\rho = \operatorname{Corr}(X, Y). \tag{64}$$

We will examine under what condition and to what extent $\rho_T(m)$ converges to ρ .

3.3. Results

In this section we present the exact formula of test statistics correlation $\rho_T(m)$ by making some assumptions about T_X and T_Y , and show that the test statistics correlation ρ_T does not always equal the population correlation ρ . For the case of two-group comparison, we prove that 1) if T_X (or T_Y) is a linear transformation of X (or Y), then $\rho_T(m) = \rho$, and that 2) if T_X (or T_Y) is the two sample t-test statistic for X (or Y) under normal assumption, then $|\lim_{m\to\infty} \rho_T(m)| \leq |\rho|$. For 2), we show that the relationship between $\lim_{m\to\infty} \rho_T(m)$ and ρ depends on whether the hypothesis tests (equation 61) are true null or not. We perform simulations for the case of test statistics derived from two-sample t-test to illustrate our findings.

3.3.1. Theory

Theorem 1 Let (X_j, Y_j) , j = 1, ..., m be independent random vectors with mean and covariance structures specified in equation (58). If $(\mathbf{a}^T \mathbf{X}, \mathbf{a}^T \mathbf{Y})$ is independent of (S_X, S_Y) , then the correlation of T_X and T_Y in equation (62) can be expressed as

$$\rho_{T}(m) = \frac{\rho E(S_{X}^{-1} S_{Y}^{-1}) + \frac{\mathbf{a}^{T} \boldsymbol{\mu}_{X} \cdot \mathbf{a}^{T} \boldsymbol{\mu}_{Y}}{\sigma_{X} \sigma_{Y} \mathbf{a}^{T} \mathbf{a}} Cov(S_{X}^{-1}, S_{Y}^{-1})}{\sqrt{\left[E(S_{X}^{-2}) + \frac{(\mathbf{a}^{T} \boldsymbol{\mu}_{X})^{2}}{\sigma_{X}^{2} \mathbf{a}^{T} \mathbf{a}} Var(S_{X}^{-1})\right] \left[E(S_{Y}^{-2}) + \frac{(\mathbf{a}^{T} \boldsymbol{\mu}_{Y})^{2}}{\sigma_{Y}^{2} \mathbf{a}^{T} \mathbf{a}} Var(S_{Y}^{-1})\right]}}$$
(65)

Proof: Since samples are independent, we have

$$\operatorname{Cov}(\boldsymbol{a}^{T}\boldsymbol{X}, \boldsymbol{a}^{T}\boldsymbol{Y}) = \boldsymbol{a}^{T}\operatorname{Cov}(\boldsymbol{X}, \boldsymbol{Y})\boldsymbol{a} = \rho\sigma_{X}\sigma_{Y}\boldsymbol{a}^{T}\boldsymbol{a},$$

$$\operatorname{Var}(\boldsymbol{a}^{T}\boldsymbol{X}) = \sigma_{X}^{2}\boldsymbol{a}^{T}\boldsymbol{a},$$

$$E(\boldsymbol{a}^{T}\boldsymbol{X})^{2} = (\boldsymbol{a}^{T}\boldsymbol{\mu}_{X})^{2} + \sigma_{X}^{2}\boldsymbol{a}^{T}\boldsymbol{a},$$

$$E[(\boldsymbol{a}^{T}\boldsymbol{X})(\boldsymbol{a}^{T}\boldsymbol{Y})] = E(\boldsymbol{a}^{T}\boldsymbol{X})E(\boldsymbol{a}^{T}\boldsymbol{Y}) + \operatorname{Cov}(\boldsymbol{a}^{T}\boldsymbol{X}, \boldsymbol{a}^{T}\boldsymbol{Y})$$

$$= (\boldsymbol{a}^{T}\boldsymbol{\mu}_{X})(\boldsymbol{a}^{T}\boldsymbol{\mu}_{Y}) + \rho\sigma_{X}\sigma_{Y}\boldsymbol{a}^{T}\boldsymbol{a}$$

$$(66)$$

Note that since S_X is independent of S_X , we have

$$\operatorname{Var}(T_X) = E\left[\left(\frac{\boldsymbol{a}^T \boldsymbol{X}}{S_X}\right)^2\right] - \left[E\left(\frac{\boldsymbol{a}^T \boldsymbol{X}}{S_X}\right)\right]^2$$

$$= E[\boldsymbol{a}^T \boldsymbol{X}]^2 E[S_X^{-2}] - \left[E(\boldsymbol{a}^T \boldsymbol{X})\right]^2 \left[E(S_X^{-1})\right]^2$$

$$= \sigma_X^2 \boldsymbol{a}^T \boldsymbol{a} E(S_X^{-2}) + (\boldsymbol{a}^T \boldsymbol{\mu}_X)^2 \operatorname{Var}(S_X^{-1})$$
(67)

Similarly,

$$\operatorname{Var}(T_Y) = \sigma_Y^2 \boldsymbol{a}^T \boldsymbol{a} E(S_Y^{-2}) + (\boldsymbol{a}^T \boldsymbol{\mu}_Y)^2 \operatorname{Var}(S_Y^{-1})$$
(68)

and

$$\operatorname{Cov}(T_{X}, T_{Y}) = E\left[\frac{\boldsymbol{a}^{T}\boldsymbol{X}}{S_{X}^{-1}} \cdot \frac{\boldsymbol{a}^{T}\boldsymbol{Y}}{S_{Y}^{-1}}\right] - E\left[\frac{\boldsymbol{a}^{T}\boldsymbol{X}}{S_{X}^{-1}}\right] E\left[\frac{\boldsymbol{a}^{T}\boldsymbol{Y}}{S_{Y}^{-1}}\right]$$

$$= E[(\boldsymbol{a}^{T}\boldsymbol{X})(\boldsymbol{a}^{T}\boldsymbol{Y})] \cdot E[S_{X}^{-1}S_{Y}^{-1}] - (\boldsymbol{a}^{T}\boldsymbol{\mu}_{X})(\boldsymbol{a}^{T}\boldsymbol{\mu}_{Y})E[S_{X}^{-1}]E[S_{Y}^{-1}]$$

$$= [(\boldsymbol{a}^{T}\boldsymbol{\mu}_{X})(\boldsymbol{a}^{T}\boldsymbol{\mu}_{Y}) + \rho\sigma_{X}\sigma_{Y}\boldsymbol{a}^{T}\boldsymbol{a}]E[S_{X}^{-1}S_{Y}^{-1}] - (\boldsymbol{a}^{T}\boldsymbol{\mu}_{X})(\boldsymbol{a}^{T}\boldsymbol{\mu}_{Y})E[S_{X}^{-1}]E[S_{Y}^{-1}]$$

$$(69)$$

The result follows by plugging equations (66)—(69) into equation (56).

corollary 1 For any non zero \boldsymbol{a} , $\rho_T(m) = \rho$ if S_X and S_Y are constant with respect to $\boldsymbol{X}, \boldsymbol{Y}$.

Proof: When S_X and S_Y are constants, $Cov(S_X^{-1}, S_Y^{-1})$, $Var(S_X^{-1})$ and $Var(S_Y^{-1})$ are all 0, and equation (65) reduces to

$$\rho_T(m) = \frac{\rho E(S_X^{-1} S_Y^{-1})}{\sqrt{E(S_X^{-2})E(S_Y^{-2})}} = \rho.$$
 (70)

Corollary 1 states that test statistics correlation and expression level correlation are equal under linear transformation of X and Y.

However, if we assume that (S_X, S_Y) is a non-constant function of $(\boldsymbol{X}, \boldsymbol{Y})$, then the test statistics correlation in equation (65) can be expressed as

$$\rho_{T}(m) = \frac{\frac{E(S_{X}^{-1}S_{Y}^{-1})}{\sqrt{\text{Var}(S_{X}^{-1})\text{Var}(S_{Y}^{-1})}}\rho + \frac{(\boldsymbol{a}^{T}\boldsymbol{\mu}_{X})(\boldsymbol{a}^{T}\boldsymbol{\mu}_{Y})}{\sigma_{X}\sigma_{Y}\boldsymbol{a}^{T}\boldsymbol{a}}\rho_{s}}{\sqrt{\left[\frac{E(S_{X}^{-2})}{\text{Var}(S_{X}^{-1})} + \frac{(\boldsymbol{a}^{T}\boldsymbol{\mu}_{X})^{2}}{\sigma_{X}^{2}\boldsymbol{a}^{T}\boldsymbol{a}}\right]\left[\frac{E(S_{Y}^{-2})}{\text{Var}(S_{Y}^{-1})} + \frac{(\boldsymbol{a}^{T}\boldsymbol{\mu}_{Y})^{2}}{\sigma_{Y}^{2}\boldsymbol{a}^{T}\boldsymbol{a}}\right]}},$$
(71)

where

$$\rho_s = \frac{\text{Cov}(S_X^{-1}, S_Y^{-1})}{\sqrt{\text{Var}(S_X^{-1})\text{Var}(S_Y^{-1})}}.$$
(72)

The correlation between test statistics $\rho_T(m)$ depends on the form of test statistics, and in general, may not converge to the population correlation ρ .

3.3.2. Application of Theorem 1 under normal distribution

Many gene expression experiments are done to compare expression levels under twotreatment conditions. For the rest of this section, we discuss the relationship between ρ_T and ρ under such setting. Let $n = n_1 + n_2$ be the total number of samples, where n_1 of them are from group 1 and n_2 from group 2, and let

$$\mathbf{a} = (\underbrace{\frac{1}{n_1}, \dots, \frac{1}{n_1}}_{n_1}, \underbrace{-\frac{1}{n_2}, \dots, -\frac{1}{n_2}}_{n_2})^T$$
 (73)

be the contrast of interest. The mean expression levels are specified as

$$\boldsymbol{\mu}_{j} = (\mu_{X}, \mu_{Y})^{T}, \quad j = 1, \dots, n_{1},$$

$$\boldsymbol{\mu}_{j} = (\mu_{X}, \mu_{Y})^{T} + (\Delta_{X}, \Delta_{Y})^{T}, \quad j = n_{1} + 1, \dots, n_{1} + n_{2}.$$
(74)

If we set $S_X = 1$, then T_X corresponds to mean difference between groups 1 and 2; instead, if $S_X = \sigma_X \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}$ where σ_X is known, then T_X corresponds to the statistic for two-sample z-test. Therefore, according to Corollary 1, $\rho_T(n_1, n_2) = \rho$ if we use mean difference or z-value as test statistics.

The two-sample t-statistic is also a commonly used statistic in differential expression analysis. In the case of two sample t-test with equal variance, with the contrast a defined in equation (73), the test statistic for X is

$$T_X = \frac{\bar{X}_1 - \bar{X}_2}{S_{p,X} \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}},\tag{75}$$

where $S_{p,X}$ is the pooled variance

$$S_{p,X}^2 = \frac{(n_1 - 1)S_{X,1}^2 + (n_2 - 1)S_{X,2}^2}{n_1 + n_2 - 2}. (76)$$

Similarly, we obtain T_Y by replacing the subscript "X" with "Y" in equations (75) and (76). Under normal distribution assumption, we have the following theorem for two-sample t-test with equal variance:

Theorem 2 Let (X_i, Y_i) , i = 1, ..., n follow a bivariate normal distribution with mean specified by equations (74) and covariance Σ (see equation (58)). If T_X and T_Y are statistics for equal-variance two-sample t-test, then

$$\rho_T(n_1, n_2) = \frac{\frac{\Delta_X \Delta_Y}{\sigma_X \sigma_Y} C \rho_s + \rho B + \rho_s \rho (A - B)}{\sqrt{\left[\frac{\Delta_X^2}{\sigma_X^2} C + A\right] \left[\frac{\Delta_Y^2}{\sigma_X^2} C + A\right]}}$$
(77)

where

$$A = \frac{n_1 + n_2 - 2}{n_1 + n_2 - 4}, \quad B = \frac{\left(\frac{n_1 + n_2 - 2}{2}\right)\Gamma^2\left(\frac{n_1 + n_2 - 4}{2} + \frac{1}{2}\right)}{\Gamma^2\left(\frac{n_1 + n_2 - 2}{2}\right)},$$

$$\rho_s = Corr(S_X^{-1}, S_Y^{-1}), \quad C = \frac{(n_1 + n_2)(A - B)}{(2 + n_1 n_2^{-1} + n_1 n_2^{-1})}.$$
(78)

The proof of Theorem 2 is presented in Section 3.4. Next we present the limit of $\rho_T(n_1, n_2)$.

Theorem 3 If there exists positive constants M_1 and M_2 , such that $M_1 \leq n_1 n_2^{-1} \leq M_2$, then

$$\lim_{n_1+n_2\to\infty} \rho_T(n_1, n_2) = \frac{\rho(1 + \beta \frac{\Delta_X \Delta_Y}{\sigma_X \sigma_Y} \rho)}{\sqrt{\left[1 + \beta \frac{\Delta_X^2}{\sigma_X^2}\right] \left[1 + \beta \frac{\Delta_Y^2}{\sigma_Y^2}\right]}}$$
(79)

where
$$\beta = \lim_{n_1 + n_2 \to \infty} C = (4 + 2n_1^{-1}n_2 + 2n_1n_2^{-1})^{-1}$$
.

Theorem 3 says that as long as n_1 and n_2 grow proportionally to infinity, the quantity $\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2)$ is a function of population correlation ρ , the signal-to-noise ratios $(\delta_X,\delta_Y)=(\Delta_X/\sigma_X,\Delta_Y/\sigma_Y)$ and the sample ratio n_1/n_2 . We have the following observations:

- 1. If both tests are true null (i.e., $\Delta = 0$), then $\lim_{n_1+n_2\to\infty} \rho_T(n_1, n_2) = \rho$.
- 2. If only one test is true null, then $\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2)$ is proportional to and smaller in absolute value than ρ (i.e., $\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2) = \gamma_0 \rho$, $0 < \gamma_0 < 1$).
- 3. If both tests are true alternative (i.e., $\Delta \neq 0$), then $\rho_T \neq \rho$ in general. Specifically,
 - i) when $\Delta_X \Delta_Y > 0$ (i.e., both genes are DE towards the same direction), we have $\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2) > \rho$ for $\rho < 0$ and $0 \le \lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2) \le \rho$ for $\rho \ge 0$.
 - ii) when $\Delta_X \Delta_Y < 0$ (i.e., genes are DE towards different directions), we have $\rho < \lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2) < 0$ for $\rho < 0$ and $\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2) < \rho$ for $\rho > 0$.

Therefore in either case, we have $|\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2)| \leq |\rho|$.

We note that $|\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2)| \leq |\rho|$ when test statistics are derived from two sample t test with equal variance. In other words, T_X and T_Y are always "no more correlated" than X and Y are. It's also interesting to note that when both genes are DE, $\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2) = 0$ at $\rho = -\frac{\sigma_X\sigma_Y}{\beta\Delta_X\Delta_Y}$ provided that $\frac{\sigma_X\sigma_Y}{\beta\Delta_X\Delta_Y} \in (-1,1)$. Figure 3.1 shows the contour plots of $\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2)$ versus the signal-to-noise ratios δ_X (= Δ_X/σ_X) and δ_Y (= Δ_Y/σ_Y) for different ρ 's. The largest value of $\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2)$ (in absolute value) is always at the center, where both δ_X and δ_Y are 0 (i.e., $\Delta_X = \Delta_Y = 0$).

In addition, if $n_1/n_2 \to 0$ or ∞ , then $\beta = 0$ and we have $\lim_{n_1+n_2\to\infty} \rho_T(n_1,n_2) = \rho$. That is, when sample size of one group is not proportional to that of the other, $\operatorname{Corr}(T_X,T_Y)$ will converge to ρ regardless of whether the tests are under the null or not.

3.3.3. Simulation

We perform simulations to evaluate the correlations between test statistics and those between expression levels under two sample t-test. We simulate the expression data from normal distributions. Specifically, we let (X, Y) be the expression levels of genes X and Y, and

$$\begin{pmatrix} X_{j} \\ Y_{j} \end{pmatrix} \sim N \begin{bmatrix} \begin{pmatrix} 0 \\ 0 \end{pmatrix}, \begin{pmatrix} 1 & \rho \\ \rho & 1 \end{pmatrix} \end{bmatrix}, j = 1, \dots, n_{1}$$

$$\begin{pmatrix} X_{j} \\ Y_{j} \end{pmatrix} \sim N \begin{bmatrix} \begin{pmatrix} \Delta_{X} \\ \Delta_{Y} \end{pmatrix}, \begin{pmatrix} 1 & \rho \\ \rho & 1 \end{pmatrix} \end{bmatrix}, j = n_{1} + 1, \dots, n_{1} + n_{2}$$

$$(80)$$

For each given ρ , we consider these $n = n_1 + n_2$ pairs of (X, Y) as observations from one *simulated* experiment. Out of this experiment, we calculate $q = (T_X, T_Y)$ where T_X and T_Y are the test statistics for gene X and gene Y respectively using two-

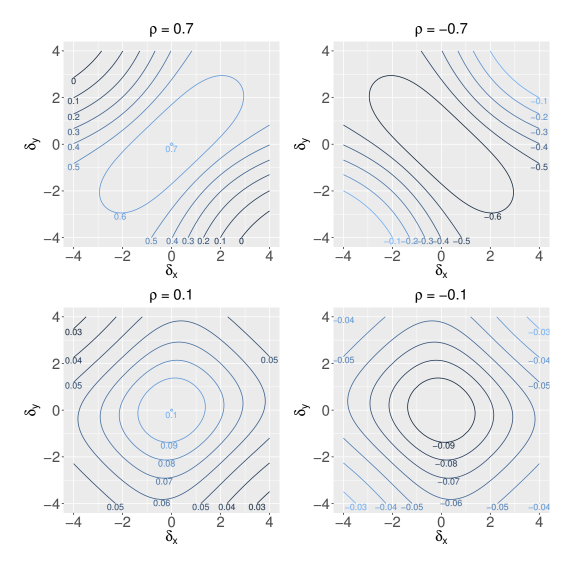


Figure 3.1: Contour plot of theoretical correlation between test statistics. For each fixed ρ and each pair of δ_X (= Δ_X/σ_X) and δ_Y (= Δ_Y/σ_Y), the theoretical correlation ρ_T is calculated according to equation (79).

sample t-test for equal variance procedure. We replicate the simulated experiment for B=1000 times, resulting in a matrix $\mathbf{Q}_{1000\times 2}$. We take the correlation between the first and the second columns of \mathbf{Q} as an estimate for test statistics correlation $r_{\text{statistics}}$. We increase ρ from -0.99 to 0.99 by fixed step size 0.01, and examine the relationship between $r_{\text{statistics}}$ and ρ under the following different cases:

a)
$$\delta_X = \delta_Y = 0;$$

- b) $\delta_X = 0, \delta_Y = 2;$
- c) $\delta_X = 0.5, \delta_Y = 2;$
- d) $\delta_X = 1, \delta_Y = 2;$
- e) $\delta_X = 3, \delta_Y = 2;$
- f) $\delta_X = -3, \delta_Y = 2.$

We conduct simulations for two different sample sizes: we set $n_1 = n_2 = 1000$ to assess asymptotic performance of $\rho_T(n_1, n_2)$ in equation (65), and set $n_1 = n_2 = 3$ to mimic small sample size scenarios which are typical in gene expression study.

In Figure 3.2, we plot $r_{\text{statistics}}$ against the underlying true population correlation ρ under both large and small sample size scenarios. In case a) where both tests are true null, $r_{\rm statistics}$ is close to the true correlation ρ when sample size is large $(n_1 = n_2 = 1000)$, but smaller (in absolute value) than ρ when sample size is small $(n_1 = n_2 = 3)$. In cases b)—f) where there is at least one true alternative, the estimate $r_{\text{statistics}}$ can be very different from ρ . In case b) where only one gene is DE, the magnitude of $r_{\text{statistics}}$ is proportional to, and smaller in absolute value than ρ . It is more interesting to note that $r_{\mathrm{statistics}}$ is not monotone with respect to ρ when both genes are DE. If genes are DE towards the same direction as in the case of e), $r_{\text{statistics}}$ first decreases until it reaches the minimum (a negative value), and then gradually increases to 1, as ρ grows from -1 to 1. When genes are DE towards opposite directions like the case f), however, the trend is reversed from that of e): $r_{\text{statistics}}$ increases from -1 to its maximum (a positive value), and then decreases. This set of simulation results is reflected in the test statistics correlation formula of equation (79). We also demonstrate the process of how ρ_T changes from being a linear function of ρ to a quadratic function, by fixing $\delta_Y = 2$ while increasing δ_X from 0 (case b) to 3 (case e).

We illustrate in Figure 3.2 the variation in $r_{\text{statistics}}$ with respect to change in sample size n. For each fixed ρ under cases a)—f), the absolute value of $r_{\text{statistics}}$ increases

when we change the sample size n from 6 to 2000. The change in $r_{\text{statistics}}$ induced by sample size could be substantial, especially when the population correlation is large (e.g., $\rho > 0.2$). This simulation shows that test statistics correlation ρ_T can be over-estimated by sample correlation, especially when sample size is small.

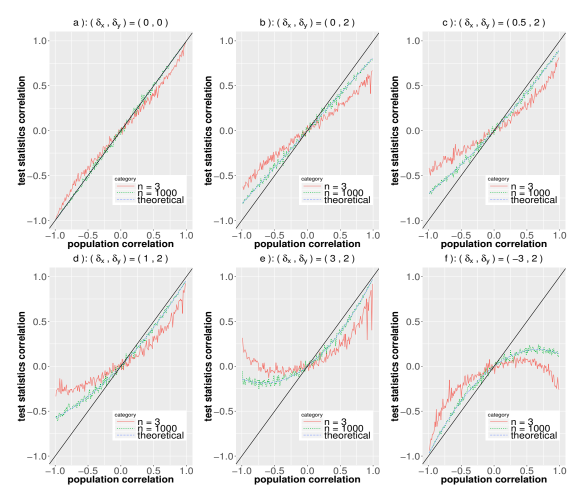


Figure 3.2: Plots of test statistics correlation against true population correlation. The test statistics are calculated using two sample t-test with equal variance, and the theoretical correlation is calculated by equation (79).

3.4. Methods

Lemma 1 The sample correlation coefficient r defined in equation (57) is a consistent estimator for the population correlation ρ ,

$$\sqrt{n}(r-\rho) \stackrel{D}{\to} N\left(0, (1-\rho^2)^2\right)$$
.

The proof of Lemma 1 can be found in Fisher [37].

To prove Theorem 2, it is useful to note that $U = (\boldsymbol{a}^T \boldsymbol{X}, \boldsymbol{a}^T \boldsymbol{Y})$ is independent of $\boldsymbol{S} = (S_X, S_Y)$, following from Lemmas 2 and 3.

Lemma 2 Let (X_j, Y_j) , j = 1..., m be independent random variables satisfying equation (60), then $\mathbf{W} = (W_X, W_Y) = (\frac{(m-1)S_X^2}{\sigma_X^2}, \frac{(n-1)S_Y^2}{\sigma_Y^2})$ follows a **bivariate chi square** distribution with density

$$f(w_x, w_y) = \frac{2^{-m} (w_x w_y)^{(n-3)/2} e^{-\frac{w_x + w_y}{2(1-\rho^2)}}}{\sqrt{\pi} \Gamma(\frac{m}{2}) (1-\rho^2)^{(m-1)/2}} \times \sum_{k=0}^{\infty} [1 + (-1)^k] \left(\frac{\rho \sqrt{w_x w_y}}{1-\rho^2}\right)^k \frac{\Gamma(\frac{k+1}{2})}{k! \Gamma(\frac{k+m}{2})}$$
(81)

for n > 3 and $-1 < \rho < 1$.

For proof of Lemma 2, interested readers are referred to Joarder [57]. It immediately follows from Lemma 2 that $\mathbf{W}_1 = (\frac{(n_1-1)S_{X,1}^2}{\sigma_X^2}, \frac{(n_1-1)S_{Y,1}^2}{\sigma_Y^2})$ follows bivariate chi-square distribution with degree of freedom $n_1 - 1$. Similarly, $\mathbf{W}_2 = (\frac{(n_2-1)S_{X,2}^2}{\sigma_X^2}, \frac{(n_2-1)S_{Y,2}^2}{\sigma_Y^2})$ follows a bivariate chi-square distribution with degree of freedom $n_2 - 1$. Note that \mathbf{W}_1 and \mathbf{W}_2 are independent since the samples are independent.

Lemma 3 $U = (U_X, U_Y)$ is independent of $S = (S_X, S_Y)$.

Proof: By Lemma 2, the density function of $W_1 + W_2$ only involves $\sigma_X^2, \sigma_Y^2, \rho$ and sample size n_1, n_2 , therefore we can denote its density by some function $g(\sigma_X^2, \sigma_Y^2, \rho, n_1 +$

 n_2). Note that $\mathbf{S}^2 = \frac{(\sigma_X^2, \, \sigma_Y^2)}{n_1 + n_2 - 2} (\mathbf{W}_1 + \mathbf{W}_2)^T$ is a linear transformation of $\mathbf{W}_1 + \mathbf{W}_2$, so its density also can be expressed in terms of $\sigma_1^2, \sigma_2^2, \rho, n_1, n_2$. Therefore $\mathbf{S} = (S_X, S_Y)$ is an ancillary statistic for Δ . On the other hand, it can be shown that $\mathbf{U} = (U_X, U_Y)$ is a complete sufficient statistic for Δ . It follows by Basu's theorem that \mathbf{U} and \mathbf{S} are independent.

Lemma 3 implies that $U_X U_Y$ is also independent of $S_X^{-1} S_Y^{-1}$, and therefore $E(\frac{U_X}{S_X} \cdot \frac{U_Y}{S_Y})$ can be expressed as $E(U_X U_Y) E(S_X^{-1} S_Y^{-1})$. We can apply Theorem 1 to calculate the correlation between T_X and T_Y under two sample t-test for equal variance.

Proof of theorem 2

First note that by Lemma 3 we have

$$Cov(T_X, T_Y) = E(T_X T_Y) - E(T_X) E(T_Y)$$

$$= \frac{1}{c_0^2} \left[E(U_X U_Y) E(S_X^{-1} S_Y^{-1}) - E(\frac{U_X}{S_X}) E(\frac{U_Y}{S_Y}) \right]$$

where $c_0 = \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}$ and $Var(T_X) = Var(\frac{U_X}{c_0 S_X}) = \frac{1}{c_0^2} Var(\frac{U_X}{S_X})$. Note that

$$\operatorname{Corr}(T_X, T_Y) = \frac{\operatorname{Cov}(T_X, T_Y)}{\sqrt{\operatorname{Var}(T_X)\operatorname{Var}(T_Y)}}$$

$$= \frac{E(U_X U_Y) E(S_X^{-1} S_Y^{-1}) - E(\frac{U_X}{S_X}) E(\frac{U_Y}{S_Y})}{\sqrt{\operatorname{Var}(\frac{U_X}{S_X})\operatorname{Var}(\frac{U_Y}{S_Y})}}$$
(82)

We need to calculate $E(U_X U_Y)$, $E(S_X^{-1} S_Y^{-1})$, $E(\frac{U_i}{S_i})$ and $Var(\frac{U_i}{S_i})$ for i = X, Y.

1. Note that $U_i \sim N\left(\Delta_i, \sigma_i^2(\frac{1}{n_1} + \frac{1}{n_2})\right), i = X, Y.$

$$E(U_X U_X) = \operatorname{Cov}(U_X, U_Y) + E(U_X) E(U_Y)$$

$$= \rho \sigma_X \sigma_Y \left(\frac{1}{n_1} + \frac{1}{n_2}\right) + \Delta_X \Delta_Y$$
(83)

2. Since $\frac{(n_1-1)S_X^2}{\sigma_X^2}$ and $\frac{(n_2-1)S_Y^2}{\sigma_Y^2}$ are independent and follow $\chi^2(n_1-1)$ and $\chi^2(n_2-1)$

respectively, , we have $W_X = \frac{(n_1+n_2-2)S_X^2}{\sigma_X^2} \sim \chi^2(n_1+n_2-2)$. It can be shown that

$$E(W_X^k) = \frac{2^k \Gamma(\frac{n_1 + n_2 - 2}{2} + k)}{\Gamma(\frac{n_1 + n_2 - 2}{2})}$$

Therefore

$$E\left(S_X^{-1}\right) = \frac{\sqrt{B}}{\sigma_X}, \quad \text{Var}\left(S_X^{-1}\right) = \frac{A - B}{\sigma_X^2}$$
 (84)

Note that $\rho_s = \operatorname{Corr}(S_X^{-1}, S_Y^{-1})$, we have

$$E(S_X^{-1}S_Y^{-1}) = E(S_X^{-1})E(S_Y^{-1}) + \rho_s \sqrt{\text{Var}(S_X^{-1})\text{Var}(S_Y^{-1})}$$

$$= \frac{B}{\sigma_X \sigma_Y} + \rho_s \frac{A - B}{\sigma_X \sigma_Y}$$
(85)

3. $U_i \sim N\left(\Delta_i, \sigma_i^2\left(\frac{1}{n_1} + \frac{1}{n_2}\right)\right)$ and $\frac{(n_1 + n_2 - 2)S_i^2}{\sigma_i^2} \sim \chi^2(n_1 + n_2 - 2)$ and by Lemma 3 U_i and $\frac{(n_1 + n_2 - 2)S_i^2}{\sigma_i^2}$ are independent for i = X, Y, we have

$$\frac{\frac{U_i - \Delta_i}{\sigma_i \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}}}{\frac{(n_1 + n_2 - 2)S_i^2}{\sigma_i^2} / (n_1 + n_2 - 2)} = \frac{U_i - \Delta_i}{S_i \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}} \sim t(n_1 + n_2 - 2)$$
(86)

It follows from

$$E\left(\frac{U_i - \Delta_i}{S_i \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}}\right) = 0, \quad \text{Var}\left(\frac{U_i - \Delta_i}{S_i \sqrt{\frac{1}{n_1} + \frac{1}{n_2}}}\right) = \frac{n_1 + n_2 - 2}{n_1 + n_2 - 4}$$
(87)

that

$$E\left(\frac{U_i}{S_i}\right) = \frac{\Delta_i}{\sigma_i} \sqrt{B}$$

$$\operatorname{Var}\left(\frac{U_i}{S_i}\right) = A\left(\frac{1}{n_1} + \frac{1}{n_2}\right) + \frac{\Delta_i^2}{\sigma_i^2} (A - B)$$
(88)

Finally, the test statistics correlation (77) is obtained by plugging equations (83–88) into equation (82).

Lemma 4 If there exists a positive number M, such that $n_1 n_2^{-1} \leq M$ and $n_1 n_2^{-1} \leq M$

M, then the following results hold:

1.
$$\lim_{n_1+n_2\to\infty} A = 1$$
.

2.
$$\lim_{n_1+n_2\to\infty} B = 1$$
.

$$3. \lim_{n_1 + n_2 \to \infty} C = \beta.$$

where A, B and C are defined in equation (78), and $\beta = (4 + n_1 n_2^{-1} + n_1^{-1} n_2)^{-1}$.

Proof: Note that

$$B = \begin{cases} \frac{(k-1)\Gamma^{2}(k-\frac{3}{2})}{\Gamma^{2}(k-1)}, & \text{if } n_{1} + n_{2} = 2k, k \geq 2\\ \\ \frac{(k-\frac{1}{2})\Gamma^{2}(k-1)}{\Gamma^{2}(k-\frac{1}{2})}, & \text{if } n_{1} + n_{2} = 2k+1, k \geq 2 \end{cases}$$

$$(89)$$

We will use second order Stirling's formula,

$$k! \approx \sqrt{2\pi k} \left(\frac{k}{e}\right)^k \left(1 + \frac{1}{12k}\right) \tag{90}$$

Using Stirling's formula (90) and $\Gamma(k+\frac{1}{2})=\frac{(2k)!}{4^kk!}\sqrt{\pi}$, it can be shown that

$$B \approx \begin{cases} \frac{(k-1)(k-2)(k-2+\frac{1}{24})^2}{(k-2+\frac{1}{12})^4}, & \text{if } n_1 + n_2 = 2k, k \ge 2\\ \\ \frac{(k-\frac{1}{2})(k-1+\frac{1}{12})^4}{(k-1+\frac{1}{24})^2(k-1)^3}, & \text{if } n_1 + n_2 = 2k+1, k \ge 2 \end{cases}$$

$$(91)$$

It can also be shown following equation (91) that

$$A - B \approx \begin{cases} \frac{\frac{1}{4}(k-1)(k-2)^3 + o((k-2)^4)}{(k-2)(k-2+\frac{1}{12})^4}, & \text{if } n_1 + n_2 = 2k, k \ge 2\\ \\ \frac{\frac{1}{4}(k-1)^3(k-\frac{1}{2})(k-3) + o((k-1)^4)}{(k-\frac{3}{2})(k-1+\frac{1}{24})^2(k-1)^3}, & \text{if } n_1 + n_2 = 2k+1, k \ge 2 \end{cases}$$

$$(92)$$

And the results immediately follow.

Lemma 5 Let (X_j, Y_j) , j = 1, ..., n be i.i.d. random variables under the two sample t-test for equal variance setting, with mean specified in equation (74) covariance structure in equation (58). Then we have $\lim_{n\to\infty} \rho_s = \rho^2$.

Proof: Let's first look at samples $j = 1, ..., n_1$. Note that

$$S_{X,1}^2 = \frac{1}{n_1} \sum_{j=1}^{n_1} (X_j - \bar{X}_1)^2$$
(93)

is the maximum likelihood estimator (MLE) for σ_X^2 . By invariance property of MLE, the pooled variance estimator

$$\begin{pmatrix} S_X^2 \\ S_Y^2 \end{pmatrix} = a_1 \begin{pmatrix} S_{X,1}^2 \\ S_{Y,1}^2 \end{pmatrix} + a_2 \begin{pmatrix} S_{X,2}^2 \\ S_{Y,2}^2 \end{pmatrix}$$
(94)

where

$$n = n_1 + n_2$$
, $a_1 = \frac{n_1 - 1}{n - 2}$, $a_2 = \frac{n_2 - 1}{n - 2}$

is also MLE for $(\sigma_X^2, \sigma_Y^2)^T$ respectively. It can be shown that

$$E[S_X^2] = \sigma_X^2, \ E[S_Y^2] = \sigma_Y^2,$$

$$Var[S_X^2] \to \frac{2\sigma_X^4}{n}, \ Var[S_Y^2] \to \frac{2\sigma_Y^4}{n}, \ Cov(S_X^2, S_Y^2) \to \frac{2\rho^2 \sigma_X^2 \sigma_Y^2}{n}$$
(95)

We have

$$\sqrt{n} \left[\begin{pmatrix} S_{X,1}^2 \\ S_{Y,1}^2 \end{pmatrix} - \begin{pmatrix} \sigma_X^2 \\ \sigma_Y^2 \end{pmatrix} \right] \xrightarrow{d} N \left[\begin{pmatrix} 0 \\ 0 \end{pmatrix}, 2 \begin{pmatrix} \sigma_X^4 & \rho^2 \sigma_X^2 \sigma_Y^2 \\ \rho^2 \sigma_X^2 \sigma_Y^2 & \sigma_Y^4 \end{pmatrix} \right]$$
(96)

If we let $g(x) = x^{-\frac{1}{2}}$, and apply δ -method to equation (96), we obtain

$$\sqrt{n} \left[\begin{pmatrix} S_X^{-1} \\ S_Y^{-1} \end{pmatrix} - \begin{pmatrix} \sigma_X^{-1} \\ \sigma_Y^{-1} \end{pmatrix} \right] \xrightarrow{d.} N \left[\begin{pmatrix} 0 \\ 0 \end{pmatrix}, \frac{1}{2} \begin{pmatrix} \sigma_X^{-2} & \rho^2 \sigma_X^{-1} \sigma_Y^{-1} \\ \rho^2 \sigma_X^{-1} \sigma_Y^{-1} & \sigma_Y^{-2} \end{pmatrix} \right] \tag{97}$$

It follows from equation (97) that $\operatorname{Corr}(S_X^{-1}, S_Y^{-1}) \to \rho^2$.

3.5. Conclusion and discussion

This article discusses the relationship between population correlation ρ and the corresponding test statistics correlation $\rho_T(m)$. We investigate $\rho_T(m)$ for test statistics of the form $(\frac{\boldsymbol{a}^T\boldsymbol{X}}{S_X}, \frac{\boldsymbol{a}^T\boldsymbol{Y}}{S_Y})$ (see equation (62)), where the denominator is the standard error of the numerator. Assuming independence between $(\boldsymbol{a}^T\boldsymbol{X}, \boldsymbol{a}^T\boldsymbol{Y})$ and (S_X, S_Y) , we derive the formula for test statistics correlation ρ_T , and show that ρ_T may not equal population correlation ρ .

In two group comparison setting, we conclude that $\rho_T(m) = \rho$ when S_X (or S_Y) is constant with respect to X (or Y). That is, $\rho_T(m) = \rho$ under linear transformation of X and Y, which is the case for two sample z-test. However, when S_X (or S_Y) is a function of X (or Y), as is the case of two sample t-test, this equality may not hold. For two sample t-test, we prove that $\lim_{m\to\infty} \rho_T(m) = \rho$ only if the null in equation (61) is true for all the tests considered, and that $\lim_{m\to\infty} \rho_T(m) \leq |\rho|$ otherwise. In the case where one test is true null and the other true alternative, $\lim_{m\to\infty} \rho_T(m)$ is directly proportional to ρ , while when both tests are true alternatives, $\lim_{m\to\infty} \rho_T(m)$ is a quadratic function of ρ .

We note that cares need to be taken when estimating correlations between test statistics. In gene expression analysis, the two-sample t-test [10, 31, 99] or moderated t-test [132] are used to calculate test statistics for DE detection, and the sample correlation (after treatment effects nullified) are used to account for correlation between those test statistics. Our study shows that, however, for DE genes, $\rho_T(m)$ may be

overestimated when two genes are positively correlated, and underestimated when they are negatively correlated. If there are true DE genes whose expression levels are correlated in either way, the VIF may not be accurately estimated in Wu and Smyth [132], resulting in biased test for their enrichment analysis as we will see in Chapter 4. Our results also indicate that the variance of $\rho_T(m)$ may also be overestimated in Efron [31], which leads to inflated variation in estimating their conditional FDP.

Theorem 1 and the subsequent results hold when the following two assumptions are met: 1) the test statistic has the of the form $\mathbf{a}^T \mathbf{X}/S_X$, and 2) $\mathbf{a}^T \mathbf{X}$ and S_X are independent. In practice, both assumptions are vulnerable. The test statistic may take different forms, depending on many factors such as the nature of the data (RNA-Seq or microarray), the experimental design structure, and the statistical hypothesis to be tested. The independence assumption between $\mathbf{a}^T \mathbf{X}$ and S_X are unlikely to hold unless the statistic is derived from two sample t-test for normally distributed random variables. Therefore, the application of Theorem 1 is somewhat limited. Yet one goal of this study is to raise awareness that the equality of $\lim_{m\to\infty} \rho_T(m)$ and ρ should not be taken for granted. In the future, we will explore the relationship between $\rho_T(m)$ and ρ for more general cases and for other types of statistics.

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

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4. Accounting for correlations in competitive geneset test for improved interpretation of genomescale data

Bin Zhuo and Duo Jiang

Abstract

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, MEACA, 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 I 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.

4.1. Introduction

Gene-set test is a statistical framework of studying the association between a test set of genes—a prior set consisting of biologically related genes—and a set of genes that are significantly correlated with experimental or environmental factors of interest. 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.

Depending on the definition of the null hypothesis, there are two types of gene set test [42]: the *self-contained* test and the *competitive* test. A self-contained test examines a set of genes by a fixed standard without reference to other genes in the genome [43, 44, 55, 123, 131]. A competitive test compares DE genes in the

test set to those not in the test set [121, 132, 134]. Many methods, regardless of the type of test, perform a three-stage analysis [60]: 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 gene-level statistic includes, among others, signal-to-noise ratio [119], ordinary t-statistic [121] or moderated t-statistic [112], log fold change [61] and Z-score [31]. 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 [119], maxmean statistic [33], and statistic derived from convoluted distribution of gene-level statistics [134], 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 [39, 42].

Many competitive gene-set tests rely on independence of gene-level statistics which implicitly requires independence among gene expression levels. For example, the PAGE [61] conducts one-sample 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 [53] for a review and references therein). The sigPathway [121] and "geneSetTest" in the limma package [112] evaluate the set-level p-values by permuting gene labels. However, tests assuming independence of genes may result in inflated false discovery rate [33, 39, 42, 132, 134], as genes within a gene set are often co-expressed and function together.

A handful of methods have been proposed to account for between-gene correlations in competitive gene-set test. One attempt is to evaluate the set-level statistic by permuting the biological sample labels [33, 119]. Permuting sample labels does not require an explicit understanding of the underlying correlation structure among genes and thus protects the test against such correlations. Since permuting sample labels is computationally inefficient, Zhou et al. [137] 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 hypotheses being tested and it is therefore difficult to characterize the null and the alternative hypotheses [42, 60, 132]. Another attempt is to use set-level statistics that directly include between-gene correlations estimated from the data. For example, CAMERA [132] calculates a variance inflation factor (VIF) from sample correlations (after the treatment effects removed) of observed data, and then incorporates it into their set-level statistics to account for between-gene correlations. QuSAGE [134], a recent extension to CAMERA, also used the same VIF in their test procedure to handle between-gene correlations. The VIF is a crucial factor and valid estimation of it relies on the assumption that correlation between any two gene-level statistics is almost the same as correlation between their corresponding expression levels. Barry et al. |10| showed by simulation that this assumption holds 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 is likely to be problematic when a fraction of genes are truly DE, in which case the correlations among gene-level statistics (e.g., t-statistics) can be badly estimated by sample correlations of observed data (as we have seen in Chapter 3).

We propose a new framework for enrichment analysis that we will call Mixedeffects Enrichment Analysis with Correlation Accounted for (MEACA). Our idea is motivated by the discrepancy between correlations among expression levels and those among gene-level statistics caused by the presence of DE genes. To tackle such discrepancy, we use differences in mean as gene-level statistics for a two group comparison experiment. We model 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 associated 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. MEACA uses a score test approach and allows for analytical assessment of p-values. Compared to existing methods including GSEA and CAMERA, MEACA enjoys robust and improved control over type I error and maintains good power in a variety of correlation structure and association settings.

The rest of the paper is organized as follows: in Section 4.2 we describe the methodology of MEACA, as well as the simulation setup for evaluating type I error rate and power, and then we summarize some existing methods; in Section 4.3 we present results from comparison of MEACA to other existing methods by simulation study, and illustrate the application of our method by two real data sets; in Section 4.4 we conclude and also specify the future work.

4.2. Methods

We consider a gene expression (e.g. RNA-Seq or microarray) experiment, in which we compare the expression levels 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" $(n_1, n_2 \geq 3)$. Suppose the expression levels 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 levels. 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 [58] or GO [6]. Our goal is to test whether this known gene set is enriched with DE signals. We will refer to the pre-defined gene set as "the test set", and genes not in this set "the background genes" which make up "the background set". We will use a

gene-level test statistic, denoted by U_i , to capture the unknown DE signal of gene i. 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 (for any given gene set G is known). As an overview,in the following sections, we will derive a model for U_i 's conditional on G, using a mixed-effect framework of the form (details to be explained later)

$$U_i = \beta_0 + \beta_1 G_i + \psi_i + \eta_i, \ i = 1, \dots, m,$$
 (98)

where β_1 is a fixed effect capturing the mean difference between the test set and the background set, and ψ_i and η_i are random effects and are mutually independent. The variance of ψ_i depends on G_i , which allows the spread of gene-level statistics to be different between the test set and the background set. The η_i 's allow for between-gene correlations.

To show why this model makes sense (??? better expression) and to elaborate the modeling assumptions on ψ_i and η_i , we will start by constructing a hierarchical model for the observed gene expression data, from which we will derive the mixed-effect model (98) for the gene-level 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 this section, our presentation of the method is conditional on G unless otherwise indicated.

4.2.1. MEACA

4.2.1.1 A hierarchical model for 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 levels 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 levels, where "baseline" refers to the average among controls. Thirdly, for any given gene, its mean expression level 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 level 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 case-control membership of the samples, with $X_i = 1$ for a case and $X_i = 0$ for a control. Let Y be an m by n matrix representing the expression data, in which each column is the expression profile for a sample and Y_{ij} $(1 \le i \le m, 1 \le j \le n)$ is the expression level of sample j at gene i. Let μ_i $(1 \le i \le m)$ be the baseline expression level for gene i. The quantities μ_i '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 i is not DE if $\Delta_i = 0$, up-regulated if $\Delta_i > 0$ and down-regulated if $\Delta_i < 0$. We model Δ as a random effect, for which we will detail our assumptions later. Given μ_i and Δ_i , the mean expression level for the control group and the treatment group are μ_i and $\mu_i + \Delta_i$, respectively. Given these means, the noise in the observed expression data for the j^{th} sample is denoted by the mean zero error vector $\epsilon_j = (\epsilon_{1j}, \dots, \epsilon_{mj})^T$, $1 \leq j \leq n$. We assume $\epsilon := (\epsilon_1, \dots, \epsilon_m)$ 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 that its empirical variance equals one before implementing our method (see Appendix (A) for more detail). For the covariance structure of ϵ , we assume

$$\epsilon_{j_1}$$
 and ϵ_{j_2} are independent, $j_1 \neq j_2$, (99)

$$Cov(\epsilon_j | \mathbf{G}) = \mathbf{C}, \quad 1 \le i \le n, \tag{100}$$

where C is an m by m between-gene correlation matrix shared by all samples and is generally unknown. Putting these elements together, we obtain the following model for the expression data Y given X and G

$$Y_{ij} = \mu_i + X_j \cdot \Delta_i + \epsilon_{ij}, \tag{101}$$

for $1 \le i \le m, 1 \le j \le n$. The term G enters this model via Δ_i and possibly μ_i .

4.2.1.2 Assumptions on the DE effects

Conditional on G, we assume that the Δ_i 's are mutually independent and come from either of the two distributions, \mathcal{D}_1 and \mathcal{D}_2 , depending on whether $G_i = 0$ 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

$$E(\Delta|G) = \beta_0 + \beta_1 G$$
, $var(\Delta|G) = \sigma_1^2 I_1 + \sigma_2^2 I_2$, (102)

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 $G_i = 1$ and those for I_2 to the genes with $G_i = 0$.

Aside from the conditions in equation (102) 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_i 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 (102)

as a special case. Suppose the m genes are independently sampled to be either DE or not. The probability for gene i to be DE is p_t if $G_i = 1$ or p_b if $G_i = 0$. 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,

$$E(\Delta_i|\mathbf{G}) = p_i \mu_\delta, \quad \text{Var}(\Delta_i|\mathbf{G}) = p_i \sigma_\delta^2 + p_i (1 - p_i) \mu_\delta^2, \tag{103}$$

where $p_i = p_t$ if $G_i = 1$ and $p_i = p_b$ if $G_i = 0$ (the derivation is provided in Appendix (B)). It can be shown that this model is a special case of equation (102).

4.2.1.3 Model for gene-level statistics

For each gene i, we consider the gene-level statistic U_i given by

$$U_i = \frac{\sum_{j:X_j=1} Y_{ij}}{n_1} - \frac{\sum_{j:X_j=0} Y_{ij}}{n_2},$$
(104)

which is sample mean difference in the expression levels between cases and controls. Given our assumption that ϵ_i has variance 1, U_i provides a DE metric for gene i. We will construct a quasi-likelihood model for $\mathbf{U} = (U_1, \dots, U_m)^T$ by deriving the mean and covariance structures of \mathbf{U} from the model for \mathbf{Y} described in Sections 4.2.1.1 and 4.2.1.2. We first observe that combining equations (104) and (101) yields

$$U_i = \Delta_i + \eta_i$$
, where $\eta_i = \frac{1}{n_1} \sum_{j:X_i=1} \epsilon_{ij} - \frac{1}{n_2} \sum_{j:X_i=0} \epsilon_{ij}$. (105)

It can be shown based on equations (99), (100) and (102) that

$$E(\boldsymbol{U}|\boldsymbol{G}) = \beta_0 + \beta_1 \boldsymbol{G},\tag{106}$$

$$\Sigma := \operatorname{Var}(\boldsymbol{U}|\boldsymbol{G}) = \sigma_0^2 \boldsymbol{C} + \sigma_1^2 \boldsymbol{I}_1 + \sigma_2^2 \boldsymbol{I}_2, \tag{107}$$

where $\sigma_0^2 = 1/n_1 + 1/n_2$ is a known parameter (the proof is provided in Appendix (B)). We note that in equation (107), 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_i 's, whereas only the C component contributes to the correlations among U_i 's.

We note that by letting $\Delta_i = \beta_0 + \beta_1 G_i + \psi_i$, equation (105) is just a re-expression of model (98) whose mean and variance are given by equations (106) and (107). The random effect ψ_i 's capture the heterogeneity of the DE effects that are conditional on whether gene i belongs to the test set $(G_i = 1)$ or not $(G_i = 0)$.

4.2.1.4 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 [10, 132]. 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 $H_0: \mathcal{D}_0 = \mathcal{D}_1$ against $H_1: \mathcal{D}_0 \neq \mathcal{D}_1$. For example, for the special scenario given by equation (103), 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 $E(\mathcal{D}_0) \neq E(\mathcal{D}_1)$, 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 4.2.1.3, the quasi-score statistic for β_1 has the form $S \propto G^T(U - \hat{\beta}_0 \mathbf{1}_m)$, 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

$$T_{\mathrm{u}} = \frac{S^2}{\widehat{\mathrm{Var}}_{0,\boldsymbol{C}=\boldsymbol{I}}(S|\boldsymbol{G})} = \frac{[\boldsymbol{G}^T(\boldsymbol{U} - \hat{\beta}_0 \mathbf{1}_m)]^2}{\boldsymbol{G}^T(\boldsymbol{I} - \boldsymbol{H})\boldsymbol{G}},$$
(108)

where $\boldsymbol{H} = \frac{1}{m} \mathbf{1}_m \mathbf{1}_m^T$. The subscript "u" stands for "uncorrelated genes". For the case of interest when between-gene correlation is present, \boldsymbol{C} is a non-trivial correlation matrix. We will again form our test statistic based on S. However, 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 (107), the variance of S is given by $\operatorname{Var}(S|\boldsymbol{G}) = \boldsymbol{G}^T(\boldsymbol{I} - \boldsymbol{H})\Sigma(\boldsymbol{I} - \boldsymbol{H})\boldsymbol{G}$. Note that $H_0: \mathscr{D}_0 = \mathscr{D}_1$ implies $\sigma_1^2 = \sigma_2^2$. Thus, under H_0 , $\Sigma := \operatorname{Var}_0(\boldsymbol{U}|\boldsymbol{G}) = \sigma_0^2 \boldsymbol{C} + \sigma_1^2 \boldsymbol{I}$, 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 $\operatorname{Var}_0(U_i) = \sigma_0^2 + \sigma_1^2$ and use $\hat{\sigma}_1^2 = \sum_{i=1}^m (U_i - \overline{U})^2/(m-1) - \sigma_0^2$. Therefore, assuming \boldsymbol{C} is known, we can obtain the MEACA test statistic given by

$$T = \frac{S^2}{\widehat{\operatorname{Var}}_0(S|\mathbf{G})} = \frac{[\mathbf{G}^T(\mathbf{U} - \hat{\beta}_0 \mathbf{1}_m)]^2}{\mathbf{G}^T(\mathbf{I} - \mathbf{H})\widehat{\mathbf{\Sigma}}(\mathbf{I} - \mathbf{H})\mathbf{G}},$$
(109)

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. Therefore we substitute C with \hat{C} , the empirical covariance matrix of the expression data after controlling for possible DE effects by centering the expression levels of cases and controls separately around zero. Formally, \hat{C} is given by $\hat{C}_{ik} = \frac{1}{n} \sum_{j=1}^{n} (Y_{ij} - \alpha_{ij})(Y_{kj} - \alpha_{kj})$ where $\alpha_{ij} = \sum_{j':X_{j'}=X_j} Y_{ij'} / \sum_{j'=1}^{n} 1\{X_{j'} = X_j\}$ is the average expression level at gene i for all samples from the same group (either treatment or control) as sample j. In real data sets, the number of genes, m, is usually much greater than the sample size m, in which case m0 is a high-dimensional parameter that cannot be efficiently estimated by m2. Interestingly, however, we find that the the test statistic m3 relies not on the accurate estimation of the entire m4, but only on three parameters involving m5, which can be much more realistically estimated by a moderate sample size. To demonstrate this, let m4 and m5 be the size of the tested set and the background set,

respectively $(m_1 + m_2 = m)$. Let ρ_1 be the average correlation between two genes in the tested set, ρ_2 be the average correlation between two background genes, and ρ_3 be the average correlation between a tested gene and a background gene. Then, ρ_1 is the mean of the off-diagonal entries in the $m_1 \times m_1$ sub-matrix of C made up of rows and columns corresponding to the test set, ρ_2 is that in the $m_2 \times m_2$ sub-matrix corresponding to the background set, and ρ_3 is the mean of the entries in the $m_1 \times m_2$ sub-matrix of C corresponding to the cross-covariance between the tested and the background sets. It can be shown that the denominator of the MEACA test statistic given in equation (109) can be written as

$$c_1(1 - \frac{1}{m_1})\rho_1 + c_1(1 - \frac{1}{m_2})\rho_2 - 2c_1\rho_3 + c_1(\frac{1}{m_1} + \frac{1}{m_2}) + c_2\hat{\sigma}_1^2, \tag{110}$$

where $c_1 = \frac{m_1^2 m_2^2}{m^2} (\frac{1}{n_1} + \frac{1}{n_2})$ and $c_2 = \frac{m_1 m_2}{m}$. Therefore, the MEACA test statistic depends on C only through ρ_1 , ρ_2 and ρ_3 .

Model (98) and equation (110) also help reveal the connections between our method and CAMERA. When the CAMERA approach is considered under our framework with $Var(\epsilon_{ij}) = 1$ and equation (104) as the gene-level statistic, its test statistic can be shown to be equivalent to equation (109) but using the following as the denominator

$$c_1(1-\frac{1}{m_1})\rho_1 + c_1(\frac{1}{m_1} + \frac{1}{m_2}).$$
 (111)

In other words, CAMERA effectively assumes that $\rho_2 = \rho_3 = 0$ (i.e. between-gene correlations is present only between genes in the test set), and that $\sigma_1^2 = \sigma_2^2 = 0$ (i.e. no gene is DE under the null hypothesis). Both assumptions are likely violated in reality. To explore how violation of them affects the performance of CAMERA, note that when m_1 and m_2 are large, the first term in equation (111) is approximately proportional to $\rho_1 + \rho_2 - 2\rho_3$. To gain insights into the type I error of CAMERA, we consider the following hypothetical scenarios where only the first assumption is violated, that is, we allow at least one of ρ_2 and ρ_3 to be nonzero: (i) if $\rho_2 > 0$ but

 $\rho_3 = 0$, CAMERA would under-estimate the denominator (because $\rho_1 + \rho_2 - 2\rho_3 > \rho_1$) and therefore would have inflated type I error; (ii) if $\rho_2 = \rho_3 > 0$, CAMERA would be too conservative in controlling type I error ($\rho_1 + \rho_2 - 2\rho_3 < \rho_1$); (iii) if $\rho_2 > 0$ but $\rho_3 < 0$, CAMERA would also produce inflated type I error ($\rho_1 + \rho_2 - 2\rho_3 > \rho_1$). These trends will be confirmed and illustrated using type I error simulations (see Section 4.3.1). On the other hand, if there are DE genes in both the test set and the background set (this corresponds to violation of the second assumption), CAMERA does not fit in the framework of MEACA.

4.2.2. Simulation methods

4.2.2.1 Simulation setup

In this section, we will specify the parameter setup for type I error and power simulations. Let Y_j be a vector denoting the expression profile of sample j and $Cov(Y_{i_1,j},Y_{i_2,j})=\rho_{i_1,i_2}$ for any two genes i_1 and i_2 . We assume that genes have the same correlation if they are from the same category (whether the test set or the background set): $Cov(Y_{i_1},Y_{i_2})=\rho_1$ if genes i_1 and i_2 are both from the test set (i.e., $G_{i_1}=G_{i_2}=1$), $Cov(Y_{i_1},Y_{i_2})=\rho_2$ if they are both from the background set (i.e., $G_{i_1}=G_{i_2}=0$). For cross-category genes, $Cov(Y_{i_1},Y_{i_2})=\rho_3$ 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): $\rho_1 = \rho_2 = \rho_3 = 0.1$; that is, all genes are correlated, with an exchangeable correlation structure.
- (c): $\rho_1 = 0.1$, $\rho_2 = \rho_3 = 0$; that is, only the genes in the test set are correlated.
- (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 consider an entire gene set containing m = 500 genes, of which $m_1 = 100$ genes are in the test set, and the remaining $m_2 = 400$ genes in the background set; second, we sample genes to be DE with probability p_t in the test set and with probability p_b in the background set, and for sampled DE genes, we simulate the DE effect Δ from a normal distribution N(2,1) (except in Table 4.3 we use N(1,0.5) to report calibrated power) and for non-DE genes we set $\Delta = 0$; 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 $MVN(\mu_1, \Sigma)$ for the control group and n_2 samples from $MVN(\mu_2, \Sigma)$ for the treatment group, where the covariance $\Sigma = [Cov(Y_{i_1}, Y_{i_2})]_{m \times m}$ may be one of the correlation structures in (a)-(e).

Further assumptions on p_t and p_b will complete our generating model used in the type I error and power simulations. We have mentioned that the test-statistic correlations among genes may not equal their sample correlations of observed data when at least one gene is truly DE (see Section 4.1). Therefore, if there are 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. For each of (a)-(e) correlation structures, we conduct two groups of simulations: genes in the background set are allowed to be DE in group A_2 but not in group A_1 . In both type I error and power simulations, we set the DE probability to be $0\%(S_0)$ in group A_1 and $10\%(S_0)$ 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 scenarios for the alternative hypothesis of the presence of enrichment: for genes in the test set, we set DE probability to be $5\%(S_1)$, $10\%(S_2)$, $15\%(S_3)$ and

 $20\%(S_4)$ in group A_1 , and $15\%(S_1)$, $20\%(S_2)$, $25\%(S_3)$ and $30\%(S_4)$ in group A_2 . Table 4.1 summarizes the simulation setup for the two groups.

Table 4.1: DE probability configurations in type I error and power simulations. S_0 is for type I error simulation. S_1 - S_4 represent the four scenarios considered in power simulations. p_b and p_t are the DE probability for genes in the background set and that in the test set, respectively.

Group	Background DE prob. (p_b)		DE prob. in test set (p_t)				
	DE prob. (p_b)	S_0	S_1	S_2	S_3	S_4	
$\overline{A_1}$	0%	0%	5%	10%	15%	20%	
A_2	10%	10%	15%	20%	25%	30%	

4.2.2.2 Other methods considered

We will compare MEACA to six previously proposed gene set tests: GSEA [119], two versions of the CAMERA procedure—CAMERA-modt and CAMERA-rank [132], SigPathway [121], MRGSE [87], and QuSAGE [134]. Except SigPathway and MRGSE, all methods incorporate features intended for between-gene correlation correction. GSEA calculates an enrichment score for the test set by examining the ranking (according to some metric, for example, the 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 [112] as gene-level statistics and estimates a VIF to account for between-gene correlations in the set-level statistic, and CAMERA-rank is the rank version of the CAMERA-modt. MRGSE is a rank-based method assuming inter-gene independence, and is recommended by Tarca et al. [120] over a class of independence-assuming methods. SigPathway is a parametric version of MRGSE, and in this simulation we use the moderated t-statistics as the gene level statistics. 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 by the convoluted PDF.

The software implementation is described as follows. The GSEA is modified from

the original R-GSEA script (http://software.broadinstitute.org/gsea/index.jsp) to accommodate single gene set test. CAMERA and MRGSE are implemented in the limma package [113] in the Bioconductor project [40]. QuSAGE is available in the Bioconductor package of the same name, and SigPathway is implemented by ourselves.

4.3. Results

According to the simulation setup in Section 4.2.2.1, 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 examine the type I error rate. As to power, we set DE probability according to each of the alternative scenarios S_1 - S_4 (see Table 4.1) and calculate the proportion of data sets for which a test would reject at a given level α . The results are based on 10,000 simulated data sets.

4.3.1. Type I error simulations

We report the type I error simulation results for group A_1 and A_2 simulations. Figure 4.1 shows the uniform quantile-quantile (QQ) plots of p-values for the seven approaches (MEACA, SigPathway, MRGSE, CAMERA-modt, CAMERA-rank, GSEA and QuSAGE) under each of the five correlation structures (each row of plots, from top to bottom, corresponds accordingly to correlation structures (a)-(e)). We also summarize the type I error rates of all the methods at a significant level of 0.05 in Table 4.2.

Table 4.2: Type I error rates for different methods (see Chapter 4). The type I error rates are summarized at a significant level of 0.05, based on 10,000 simulated data sets.

QuSAGE		0.000	0.052	0.106	0.203	0.491	0.027	0.131	0.175	0.267
GSEA	0.049	0.048	0.051	0.051	0.053	0.946	0.837	0.290	0.220	0.113
CAMERA-rank		0.000	0.044	0.104	0.197	0.050	0.000	0.021	0.049	0.116
CAMERA-modt	0.049	0.000	0.051	0.114	0.213	0.048	0.000	0.000	0.001	0.004
SigPathway	0.051	0.051	0.517	0.594	0.679	0.051	0.051	0.188	0.235	0.334
MRGSE	0.049	0.050	0.513	0.586	0.674	0.052	0.051	0.442	0.522	0.614
MEACA	0.056	0.059	0.056	0.059	0.058	0.050	0.052	0.054	0.052	0.054
Correlation	(a)	(b)		(p)	(e)	(a)	(b)		(p)	(e)
Group			A_1					A_2		

In group A_1 simulations (the left column of Figure 4.1), GSEA and MEACA 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 (c), yet they are too conservative for the case of (b) and anti-conservative for correlation structures (d) and (e). SigPathway and MRGSE procedures have well-calibrated type I error for correlation structures (a) and (b), but are anti-conservative the case of (c), (d) and (e). QuSAGE has good type I error control for only (c), and is too conservative for (a), (d) and (e), and anti-conservative for (b).

In group A_2 simulations (the right column of Figure 4.1), MEACA 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 be liberal in other situations. SigPathway and MRGSE have similar trends for p-values as they do, respectively, in group A_1 simulations. QuSAGE is conservative in (b) but anti-conservative in the remaining four correlation structures.

MEACA 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 betweengene correlation structures, and DE probability of each gene. MEACA controls the
size of type I error well because it uses difference in mean as gene-level statistic, and
the correlations between such statistics are exactly the same as correlations between
the samples (as we have seen in Chapter 3). GSEA evaluates the enrichment score of a
test set by generating its null distribution from sample permutation. When there are
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 between-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 [132]) may be over-estimated when a fraction of genes are DE (as we have seen in Chapter 3), and therefore the set-level test statistic is under-estimated. The performances of related methods—QuSAGE and two versions of CAMERA—are 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 between-gene correlations in the test set without addressing that in the background set.

Different from the five methods mentioned above, SigPathway and MRGSE rely on independence between genes. It's not surprising that such gene permutation based methods control type I error correctly when genes are "equally-correlated": in (a) genes are simulated to be independent, and in (b) genes are simulated to have an exchangeable correlation structure. However, both SigPathway and MRGSE fail to hold type I error size for the remaining three correlation structures. These simulations show that even small between-gene correlations (e.g., 0.05) will result in inflated type I error rate when the test does not account for between-gene correlations.

4.3.2. Power simulation

We compare the power of MEACA to those of the other six methods under correlation structure (a) in which genes are simulated to be independent. Since some of these tests are not well calibrated at the sample size considered (see results in Section 4.3.1), 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 4.3 summarizes the calibrated power for the two groups of simulations (i.e.,

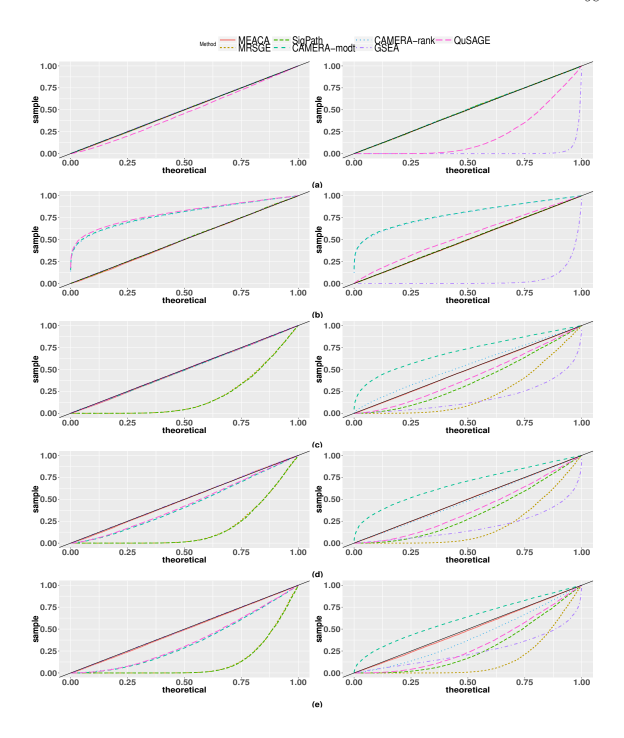


Figure 4.1: Uniform quantile-quantile plots for p-values by different methods. Each plot from top to bottom corresponds to correlation structures (a)-(e), respectively. The left column is for group A_1 simulation, and the right column for group A_2 simulation (see Table 4.1 for detail). Results are based on 10,000 simulations.

 A_1 and A_2 in Table 4.1). For A_1 simulations, GSEA has the highest, and rank based methods (MRGSE and CAMERA-rank) have the lowest, calibrated power across all four alternative scenarios. CAMERA-modt, SigPathway and MEACA have no systematic difference in the calibrated power. In group A_2 simulations, GSEA shows virtually no power. MEACA, CAMERA-modt, and SigPathway have indistinguishable calibrated power and are among the best.

Table 4.3: Recalibrated power for different methods under correlation structure (a) in which genes are simulated to be independent. The $c(\alpha)$ is the α quantile of null distribution of p-values. The powers are calculated as the proportion of $p < c(\alpha)$ for each of the four alternatives S_1 - S_4 and each of the group A_1 and A_2 simulations (see Table 4.1 for detail). Results are based on 10,000 simulations.

Group	Method	$c(\alpha)$	S_1	S_2	S_3	S_4
	MEACA	0.045	0.340	0.741	0.944	0.991
	MRGSE	0.051	0.111	0.284	0.533	0.766
	SigPathway	0.049	0.344	0.744	0.947	0.992
A_1	CAMERA-modt	0.051	0.336	0.737	0.943	0.990
	CAMERA-rank	0.053	0.108	0.280	0.519	0.758
	GSEA	0.051	0.517	0.894	0.989	0.999
	QuSAGE	0.028	0.385	0.784	0.959	0.995
	MEQLEA	0.050	0.180	0.478	0.777	0.939
	MRGSE	0.048	0.104	0.269	0.530	0.781
	SigPathway	0.049	0.175	0.473	0.773	0.936
A_2	CAMERA-modt	0.052	0.173	0.466	0.766	0.933
	CAMERA-rank	0.050	0.102	0.262	0.521	0.771
	GSEA	0.000	0.000	0.000	0.000	0.000
	QuSAGE	0.000	0.021	0.127	0.387	0.692

Figure 4.2 shows for MEACA, 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 under correlation structures (a) and (b) 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 (b) are simulated to have the same correlation 0.1. As the correlation structure becomes more complex, from (c) 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.

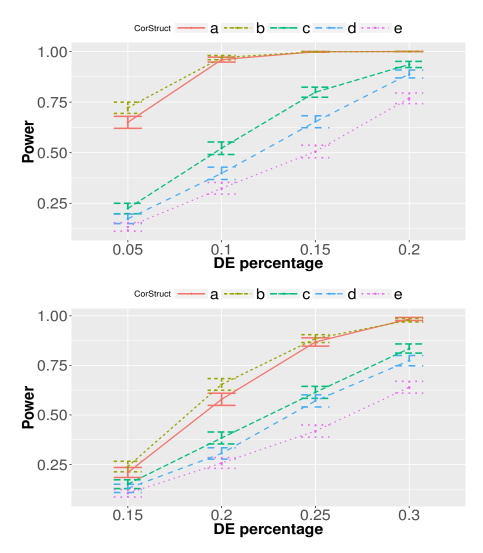


Figure 4.2: Power for MEACA under correlation structures (a)-(e) of Section 4.2.2.1. The top corresponds to group A_1 simulations, and the bottom to group A_2 simulations (see Table 4.1). The error bars are the 95% CIs based on 10,000 simulations.

4.3.3. Real data

We apply MEACA to two example data sets, and compare the lists of enriched gene sets to those obtained by other three methods (GSEA, CAMERA-modt and MRGSE). 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.

4.3.3.1 Huntington's disease data

We examine the Huntington's Disease (HD) RNA-Sequencing (RNA-Seq) data [63] to identify enriched gene sets that are potentially responsible for HD. The mRNA expression profiles in human prefrontal cortex were obtained from 20 Huntington's Disease samples and 49 neurologically normal controls. Expression values are normalized and filtered as described in the methods section of Labadorf et al. [63]. The data, containing 28,087 genes, is available as a series GSE64810 in the GEO database (http://www.ncbi.nlm.nih.gov/geo/). For each gene, we adjust for two covariates—age at death (DeathAge) and RNA Integrity Number (RIN), as also done by Labadorf et al. [63]. We follow their strategy of treating the two covariates as categorical. Briefly, DeathAge is binned into intervals 0-45, 46-60, 61-75, 76-90 and 90+, and RIN is dichotomized as > or \le 7. We regress the normalized expression levels on AgeDeath and RIN and use the resulting residuals as the covariate-adjusted expression levels.

We perform enrichment analysis on the covariate-adjusted data using the MsigDB [119] C2 Canonical Pathways (February 5, 2016, data last accessed). The C2 Canonical Pathways 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 are named by HGNC symbols in C2 and by ensembl IDs in the HD expression data set, we convert the ensembl IDs in the expression data into HGNC symbols using *BioMart* (http://uswest.ensembl.org/biomart/martview/). We retain 26,941 genes that

have corresponding HGNC symbols.

We apply four test procedures (MEACA, GSEA, CAMERA-modt and MRGSE) to run enrichment analysis for the entire C2 Canonical Pathways, and compared the four tests in terms of resulting enriched gene sets. We use the Benjamini-Hochberg [12] procedure (BH) to control the false discovery rate (FDR) for multiple hypothesis testing (unless specified otherwise, all p-values in Section 4.3.3 are adjusted by BH procedure). The BH procedure is used when the test statistics under the null have non-negative correlations [13]. We note that since many pathways have overlapped genes, the BH procedure should be appropriate in our study.

In Figure 4.3 we plot $\log 10 \, p$ -values of MEACA against those of GSEA, CAMERA-modt and MRGSE. The p-values of CAMERA-modt are overwhelmingly larger than their counterparts of GSEA or MEACA, yet smaller than those of MRGSE, even if p-values between MEACA and other three methods are highly correlated (Pearson's correlation of $\log 10 \, p$ between MEACA and GSEA, CAMERA-modt and MRGSE are 0.90, 0.96, and 0.87 respectively). This trend of p-values is consistent with our earlier simulation (see results in simulation section 4.3.1) that CAMERA-modt could produce large p values. The p-values of MRGSE are in general smaller than the corresponding p-values of MEACA, leading to more significant calls.

Using MEACA, we find 89 significant signals out of the entire 1330 gene sets at FDR level of 0.05. GSEA finds 3 enriched gene sets—2 of them were also among those 89 gene sets (the one that is not significant according to MEACA had a p-value of 0.013 and FDR 0.100). MRGSE identified 387 gene sets which include all the 89 sets MEACA identified, and CAMERA-modt identified none. Originally, Labadorf et al. [63] used the same HD data set to conduct enrichment analysis using topGo [1]. They note 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 (by nominal p-values) 18, 10 and 3 respectively in the 89

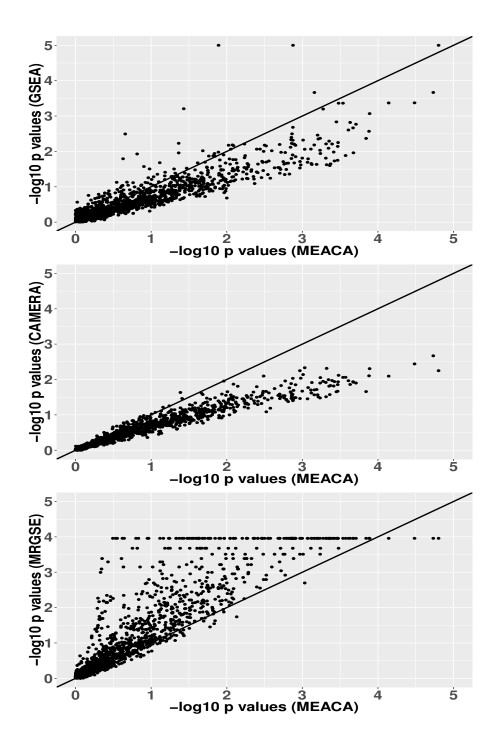


Figure 4.3: Pairwise comparisons of p-values for MEACA, GSEA, CAMERA-modtand MRGSE. The p-values are reported from enrichment test of each gene set in the C2 Canonical Pathway gene sets.

enriched gene sets.

In Table 4.4, we report the top 30 enriched gene sets (ordered by nominal p values) identified using MEACA. We also label the enriched gene sets from GSEA by "*" in the table. Many of our enriched gene sets have been shown to be closely related to HD pathogenesis. For example, the top enriched gene set by MEACA, "PID SMAD2 3NUCLEAR PATHWAY", is responsible for regulation of nuclear SMAD2/3 signaling. Katsuno et al. [59] showed that nuclear SMAD2/3 are related to polyglutamine disease, which includes HD. The third enriched gene set, "PID NFKAPPAB CANONICAL PATHWAY", is a canonical NF-kappaB pathway, and its dysregulation causes HD immune dysfunction [122]. Also, Marcora and Kennedy [81] 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 INNATE IMMUNE SYSTEM", contributes to HD pathogenesis [63, 122]. Chiang et al. [23] 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 "PID P53 DOWNSTREAM PATHWAY", Ghose et al. [41] showed the likely involvement of NFkB (RelA), p53 and miRNAs in the regulation of cell death in HD pathogenesis.

4.3.3.2 Male vs female lymphoblastoid cells data

We analyze the mRNA expression profiles from lymphoblastoid cell lines derived from 17 females and 15 males. Subramanian et al. [119] examined this data set with their GSEA method, testing the enrichment of the cytogenetic gene sets (C1). The C1 includes 24 sets, one for each of the 24 human chromosomes, and 295 sets corresponding to cytogenetic bands. For the comparison "male VS female", they expected to find gene sets on chromosome Y, not on chromosome X. We run enrichment analysis with the four tests (MEACA, GSEA, CAMERA-modtand MRGSE). In Table 4.5, we sum-

Table 4.4: Enriched gene sets (ordered by nomial p-values) identified by MEACA for HD data. The $\hat{\rho}_1$, $\hat{\rho}_2$ and $\hat{\rho}_3$, respectively, are the average estimated sample correlations of observed data between genes in the test set, between genes in the background set, and between cross-category genes. The enriched gene sets are noted by "*" for GSEA. No gene set was identified as enriched by CAMERA-modt and all the 30 gene sets are also identified as enriched by MRGSE. For all methods, a gene set is called significant when its FDR using Benjamini-Hochberg (BH) correction is < 0.05.

Gene Set	Size	$\hat{ ho}_1$	$\hat{ ho}_2$	$\hat{ ho}_3$	p-value	FDR	
PID SMAD2 3NUCLEAR PATHWAY	79	0.063	0.013	0.015	5.8E-06	5.7E-03	*
REACTOME YAP1 AND WWTR1 TAZ	23	0.121	0.013	0.014	8.5E-06	5.7E-03	
STIMULATED GENE EXPRESSION							
PID NFKAPPAB CANONICAL PATH-	22	0.127	0.013	0.019	2.3E-05	1.0E-02	
WAY							
BIOCARTA NTHI PATHWAY	23	0.130	0.013	0.023	6.2E-05	2.1E-02	
BIOCARTA TID PATHWAY	18	0.101	0.013	0.012	1.2E-04	2.2E-02	
PID HIV NEF PATHWAY KEGG PATHWAYS IN CANCER	35	0.065	0.013	0.013	1.2E-04	2.2E-02	
KEGG PATHWAYS IN CANCER	311	0.028	0.013	0.010	1.3E-04	2.2E-02	
PID MYC REPRESS PATHWAY	60	0.057	0.013	0.013	1.9E-04	2.2E-02	
PID MYC REPRESS PATHWAY BIOCARTA TOLL PATHWAY	36	0.083	0.013	0.018	2.0E-04	2.2E-02	
PID IL4 2PATHWAY	59	0.081	0.013	0.010	2.0E-04	2.2E-02	
KEGG TGF BETA SIGNALING PATH-	82	0.055	0.013	0.011	2.2E-04	2.2E-02	
WAY							
BIOCARTA DEATH PATHWAY		0.067		0.013	2.4E-04	2.2E-02	
KEGG NOD LIKE RECEPTOR SIG-	55	0.045	0.013	0.008	2.6E-04	2.2E-02	
NALING PATHWAY							
BIOCARTA CTCF PATHWAY		0.083			2.8E-04		
ST TUMOR NECROSIS FACTOR	28	0.031	0.013	0.014	3.2E-04	2.2E-02	
PATHWAY							
BIOCARTA TNFR2 PATHWAY KEGG APOPTOSIS	17	0.151	0.013	0.022		2.2E-02	
	~-	0.000				2.2E-02	
REACTOME INNATE IMMUNE SYS-	209	0.039	0.013	0.009	3.3E-04	2.2E-02	
TEM							
PID HES HEY PATHWAY		0.071				2.2E-02	
REACTOME DOWNSTREAM TCR	31	0.082	0.013	0.011	3.7E-04	2.2E-02	
SIGNALING							
PID TCPTP PATHWAY BIOCARTA 41BB PATHWAY PID FRA PATHWAY	42	0.076		0.010	3.7E-04		
BIOCARTA 41BB PATHWAY	14	0.110	0.013	0.023	3.9E-04		
PID FRA PATHWAY	34	0.154	0.013	0.008		2.2E-02	
PID P53 DOWNSTREAM PATHWAY	131	0.045	0.013	0.012		2.2E-02	
PID EPO PATHWAY BIOCARTA PPARA PATHWAY BIOCARTA EPONFKB PATHWAY BIOCARTA HIVNEF PATHWAY BIOCARTA CD40 PATHWAY	34	0.069	0.013	0.013		2.2E-02	
BIOCARTA PPARA PATHWAY	53	0.031	0.013	0.008		2.2E-02	
BIOCARTA EPONFKB PATHWAY	11	0.068	0.013	0.010		2.2E-02	
BIOCARTA HIVNEF PATHWAY	58	0.063	0.013	0.019		2.2E-02	
BIOCARTA CD40 PATHWAY	13	0.165	0.013	0.026		2.2E-02	
BIOCARTA IL7 PATHWAY	17	0.100	0.013	0.016	5.2E-04	2.3E-02	

marize all the gene sets that are called significant at FDR level 0.05. Unanimously, three gene sets—"chrY", "chrYq11" and "chrYp11"—are found to be enriched by all of the four methods. It is interesting to note that only MEACA is able to identify another Y band, "chrYp22", as enriched. In fact, these four gene sets are the only four pathways containing at least 3 genes in C1 and corresponding to chromosome Y or Y bands. MEACA does not produce small p-value (< 0.01) for the remaining three gene sets in Table 4.5, which is just as expected in that study.

Table 4.5: Enriched gene sets and their nominal p values for lymphoblastoid cells data. Reported are gene sets with FDR < 0.05 for at least one of the MEACA, GSEA, CAMERA-modt and MRGSE methods using Benjamini-Hochberg (BH) procedure.

Gene set	Size	MEACA	GSEA	CAMERA-modt	MRGSE
chrY	40	0.0E+00	0.0E+00	1.0E-05	5.9E-07
chrYq11	16	0.0E + 00	0.0E + 00	7.2E-08	8.5E-06
chrYp11	18	2.1E-15	0.0E + 00	2.8E-04	5.1E-04
chrYp22	8	3.6E-04	1.2E-02	1.0E-02	1.3E-02
chr6	614	5.6E-02	6.0E-01	6.1E-01	2.1E-04
chr1	1104	6.1E-02	5.5E-01	6.3E-01	5.3E-05
chr12	571	8.7E-02	2.6E-01	4.0E-01	5.1E-09

4.4. Conclusion and discussion

MEACA 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, MEACA controls type I error correctly and maintains good power under different correlation structures.

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 [119]. However, sample permutation method has been criticized for altering the null hypotheses being tested [42, 60]. Instead, CAMERA [132]

proposed to correct for the correlations among genes by estimating a VIF directly from the data. Incorporating VIF into set level test statistic has also been used by Yaari et al. [134] for their QuSAGE procedure. The major problem with this VIF approach is that it tries to estimate correlations among gene-level test statistics directly from sample correlations of observed data. We have argued (in Chapter 3) that the correlations among gene-level statistics may not be approximated by the sample correlation of observed data due to the presence of DE genes. The estimated VIF could be biased without taking into account such a discrepancy, which will undermine the performance of CAMERA and QuSAGE. In contrast, MEACA 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 heterogeneity of DE effects. We note that for MEACA, the estimation of covariance among gene-level statistics need not be exact: MEACA uses a score test that involves linear combinations of the entries of the covariance matrix. The denominator in the score test statistic (see equation (109)) can usually be accurately approximated given the high dimensionality of the covariance matrix. MEACA 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.

We compare the performance of MEACA to those of other existing approaches through both simulation study and real data analysis. In the simulation study, we examine the calibration of MEACA and other six method (SigPathway, MRGSE, CAMERA-modt, CAMERA-rank, GSEA and QuSAGE) in terms of type I error control and power. We demonstrate that MEACA holds correct type I error size under all correlation structures considered, whereas all other methods may fail in one or more situations. MEACA is also among the best for power performance even under independent correlation structure. In the real data analysis, we run enrichment

analysis using four methods—MEACA, CAMERA-modt, GSEA and MRGSE on two data sets. The p-values of MEACA are smaller than those produced by GSEA and CAMERA (which are intended to adjust for between-gene correlations), but larger than those of MRGSE (which assumes independence between genes). MEACA is able to identify a moderate size of enriched gene sets, some of which are insightful in the corresponding studies yet are not found by other three methods.

Currently, MEACA only supports enrichment test for two-group comparisons. In many gene expression experiments, however, researchers might use more complex design to study different factors of interest, in which case a linear model would be more appropriate. Our future work will focus on generalizing MEACA 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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5. Conclusion

5.1. Summary of the dissertation

Detecting DE genes plays a central role in gene expression analysis. The statistical questions we seek to address in this thesis help both "pre" DE analysis—finding stably expressed genes for RNA-Seq count normalization, and "post" DE analysis—investigating correlation among DE test statistics and understanding the effect of treatments on sets of genes (biological pathways or gene ontologies).

In Chapter 2 we advocate quantifying RNA-Seq gene expression stability by a numerical measure applied to a large number of data sets. We emphasize that the ranking of genes in terms of numerical stability depends on multiple factors: the background sample set and the reference gene set used for count normalization, the technology used for measuring gene expression, and the specific numerical stability measure used. Because DE is measured by relative frequencies, we argue that DE is a relative concept, and that making the reference gene set explicit can improve the interpretability of DE results. When comparing differential expression among multiple experiments, we suggest using a common set of genes as reference to avoid inconsistent conclusions. For count normalization of Arabidopsis RNA-Seq data sets, we recommend using the top 1000 stably expressed genes chosen from the appropriate reference sample (multi-tissue, leaf or seedling, etc.).

In Chapter 3 we explored the relationship between test-statistic correlation ρ_T and the correlation of underlying observed data ρ (see Section 3.2 for details about ρ and ρ_T). The test statistics investigated are of the form $T_X = \mathbf{a}^T \mathbf{X}/S_X$. Under the assumption of independence between denominator and numerator of T_X , we show that ρ_T does not necessarily equal ρ . For normally distributed random variables, we give an exact formula of ρ_T when T is the test statistic for equal-variance two-sample t-test. The formula (see equation (79)) suggests that ρ_T is a function of both

 ρ and the strength of DE (signal-to-noise ratio), and we conclude that $|\rho_T| \leq |\rho|$ for two-sample t-test of normal data. In the context of gene expression analysis, the two-sample t-test is a common choice for detecting DE genes, and the sample correlation of observed data has been used to approximate test-statistic correlation [10, 31, 132]. Our theoretical derivation shows, however, in the presence of DE, that correlation between test statistics will be over-estimated (in absolute value) by the observed-data correlation. This will have an impact on statistical inferences such as FDR control or gene-set tests that are intended to account for test-statistic correlation by using the observed-data correlation.

The MEACA, a new gene set-test framework we proposed in Chapter 4, is motivated by the observation (in Chapter 3) that correlation between test statistics may not equal the correlation of underlying observed data. MEACA uses difference in mean as gene-level statistics whose correlation can be estimated by the sample correlation of observed data, and works for gene-set test in two group comparison experiment. It has the flexibility to allow for DE genes in the background set, and does not require the data to be normally distributed. It effectively adjusts for completely unknown and unstructured correlations among genes. We demonstrated by simulation study that MEACA holds type I error size correctly, and maintains good power for all correlation structures we examined. We apply MEACA to two real data sets and find meaningful gene sets that are otherwise overlooked by methods such as GSEA or CAMERA.

5.2. Future work

To identify stably expressed genes, our model (equation (48)) still needs to estimate an initial set of normalization factors, which requires that we have to make assumptions about the relative fold changes between samples. This kind of circular dependence seems unavoidable. We use an one-step iteration to reduce the dependency of the results on the initial normalization factors. In the future, we can examine the nu-

merically stably expressed genes by evolutionary genetics methods, and ask whether there is connection between measures of expression stability and measures of genotypic/phenotypic features such as directional selection [109]. On the other hand, it would also be interesting to model the random effect terms more accurately. In Chapter 2 the random effect terms are assumed to be normally distributed, which is appropriate if the purpose is just to estimate their corresponding variance components. A more careful examination of individual data sets suggests that the between sample variances differ across experiments. Therefore, it may be beneficial to build a prior distribution of the random effect terms when analyzing a new data set.

The investigation in Chapter 3 is just a first attempt towards understanding correlation between test statistics. Our results suggest that test statistics correlation may not be approximated by observed-data correlation. In the future, we are interested in finding analytical expressions of correlation for test statistics of different hypothesis testing procedures, and as a result, improving the accuracy of FDP estimation based on the work of Efron [31].

In Chapter 4, we showed that MEACA adequately accounts for correlation among test statistics. However, everything comes at a price. It is important to choose appropriate DE measure so that correlations between test statistics can be estimated from the observed data. Our strategy is to use linear combination of the data (e.g., difference in mean under two group comparison, as we did in Chapter 4) as gene level test statistics, which may not be suitable for all cases. In many gene expression experiments, researchers might use more complicated design to study factors of interest. When the gene-level test statistics are derived from such a design based statistical analysis (e.g., generalized linear models), it is often difficult to estimate correlation among test statistics from data. In the future, we can either generalize MEACA to accommodate complicated design structures, or develop techniques that can estimate correlation between test statistics under more general statistical models.

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Appendices

A. Standardization

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

$$r_{ijk} = y_{ijk} - \sum_{j=1}^{n_k} y_{ijk}/n_k;$$
 (112)

then we calculate the pooled standard deviation from the residuals,

$$s_i = std(r_{ijk}); (113)$$

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

$$y_{ijk}^* = y_{ijk}/s_i \tag{114}$$

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

B. Covariance matrix for test statistics

Note that if we let δ_i be the DE size for gene i, then $E(\delta_i) = \mu_{\delta}$ and $Var(\delta_i) = \sigma_{\delta}^2$. To prove the equation (103), we introduce an additional random variable Z_i for DE status, where $Z_i \sim \text{Bernoulli}(1, p_t)$ if $G_i = 1$ and $Z_i \sim \text{Bernoulli}(1, p_b)$ if $G_i = 0$. It follows that $\Delta_i = Z_i \delta_i$, and we have

$$E(\Delta_i|\mathbf{G}) = E(Z_i\delta_i|\mathbf{G}) = E(\delta_i)E(Z_i|\mathbf{G}) = p_i\mu_{\delta},$$

and

$$\operatorname{Var}(\Delta_{i}|\boldsymbol{G}) = E[(Z_{i}\delta_{i})^{2}|\boldsymbol{G}] - [E(Z_{i}\delta_{i}|\boldsymbol{G})]^{2}$$

$$= \operatorname{Var}(Z_{i}|\boldsymbol{G})[E(\delta_{i})]^{2} + [(EZ_{i}|\boldsymbol{G})^{2} + \operatorname{Var}(Z_{i}|\boldsymbol{G})] \operatorname{Var}(\delta_{i})$$

$$= p_{i}\sigma_{\delta}^{2} + p_{i}(1 - p_{i})\mu_{\delta}^{2}.$$

If the gene-level test statistics U_i 's take the form of equation (104), then we have $E(U_i|\mathbf{G}) = E(\Delta_i + \eta_i|\mathbf{G}) = p_i\mu_{\delta}$. Next, note that the covariance between two genes i_1 and i_2 is given by

$$Cov(U_{i_{1}}, U_{i_{2}}|\mathbf{G}) = Cov \left[(\Delta_{i_{1}} + \eta_{i_{1}}, \Delta_{i_{2}} + \eta_{i_{2}}) | \mathbf{G} \right]$$

$$= Cov(\Delta_{i_{1}}, \Delta_{i_{2}}|\mathbf{G}) + Cov(\eta_{i_{1}}, \eta_{i_{2}}|\mathbf{G})$$

$$= Cov \left[\left(\frac{1}{n_{1}} \sum_{j:X_{j}=1} \epsilon_{i_{1},j} - \frac{1}{n_{2}} \sum_{j:X_{j}=0} \epsilon_{i_{1},j}, \frac{1}{n_{1}} \sum_{j:X_{j}=1} \epsilon_{i_{2},j} - \frac{1}{n_{2}} \sum_{j:X_{j}=0} \epsilon_{i_{2},j} \right) | \mathbf{G} \right]$$

$$= \left(\frac{1}{n_{1}} + \frac{1}{n_{2}} \right) c_{i_{1},i_{2}},$$
(115)

where c_{i_1,i_2} is the corresponding entry in C. Also

$$\operatorname{Var}(U_{i_{1}}|\boldsymbol{G}) = \operatorname{Var}(\Delta_{i_{1}} + \eta_{i_{1}}|\boldsymbol{G})$$

$$= \operatorname{Var}(\Delta_{i_{1}}|\boldsymbol{G}) + \operatorname{Var}(\frac{1}{n_{1}} \sum_{j:X_{j}=1} \epsilon_{i_{1},j} - \frac{1}{n_{2}} \sum_{j:X_{j}=0} \epsilon_{i_{1},j}|\boldsymbol{G})$$

$$= \operatorname{Var}(\Delta_{i_{1}}|\boldsymbol{G}) + \frac{1}{n_{1}} + \frac{1}{n_{2}}.$$
(116)

Equation (107) immediately follows from equations (115) and (116).

C. Supplimentary Tables

Table C.1: Summary of data sets used in the three groups: the seedling, the leaf and the multi-tissue groups.

GEO Number	Reference	Tissue	Sample Size	Mapping Quality $^1(\geq)$
GSE32202	NA	seedling	6	84.30%
GSE37159	NA	seedling	8	82.50%
GSE41766	[7]	seedling	6	85.20%
$^2\mathrm{GSE}43865$	[108]	seedling	6	92.50%
GSE51119	[136]	seedling	10	70.50%
GSE51772	[94]	seedling	8	90.60%
GSE53078	[34]	seedling	4	86.10%
$^3\mathrm{GSE}60835$	[30]	seedling	6	65.60%
GSE66666	[20]	seedling	6	91.20%
GSE36626	[129]	leaves	4	85.50%
4 GSE39463	[80]	leaves	12	74.50%
GSE48235	[76]	leaves	6	90.40%
$^{5}\mathrm{GSE}51304$	[118]	leaves	18	88.40%
⁶ GSE54677	[89]	leaves	20	85.20%
GSE35288	[92]	flower	6	76.50%
GSE35408	[8]	hypocotyl	10	77.00%
GSE52966	[22]	primary root	18	87.20%
GSE56326	NA	carpels	8	92.40%
GSE59167	[95]	root tip tissue	11	87.10%
GSE59637	[88]	inflorescences and siliques	4	71.70%
GSE60183	[62]	epidermis	6	62.30%
GSE61061	[79]	seed	6	91.60%
GSE62799	[45]	aerial tissue	6	89.30%
GSE63355	[75]	shoot apical meristem	16	87.90%

¹The number of mapped reads divided by the total number of reads in the sample

²We chose all the 6 samples at the onset of treatment out of the 42 samples.

 $^{^3\}mathrm{We}$ chose the samples of ecotype Columbia from the 12 samples.

⁴We chose all samples at 6 hours post inoculation out of the 48 samples.

⁵Out of the 48 samples, the total number of RNA-Seq samples is 18.

⁶Out of the 24 samples, the total number of RNA-Seq samples is 20.