

Matrix Ash

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Abstract

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Author Summary

Variation in gene expression is an important mechanism underlying susceptibility to complex disease. The simultaneous genome-wide assay of gene expression and genetic variation allows the mapping of the genetic factors that underpin individual differences in quantitative levels of expression (expression QTLs; eQTLs). The availability of systematically generated eQTL information could provide immediate insight into a biological basis for disease associations identified through genome-wide association (GWA) studies, and can help to identify networks of genes involved in disease pathogenesis [?]. However, most studies to date have been conducted in a single immortalized peripheral cell type, and it is unclear to what extent these findings will translate to human disease mapping across more varied cell types. Furthermore, even analyses performed on additional cell types are often performed in a single tissue framework [?,?] and fail to correlate the effect of genetics across multiple tissue types. The Genotype Tissue Expression Project, GTEx, Project will provide the data necessary to address this situation: by 2016, the resource is expected to enroll a total of approximately 900 post-mortem donors, with approximately 30 tissues collected from each donor, and the project will generate extensive genotype data and RNA-seq data on each individual. However, available methods are limited in their ability to jointly analyze data on all tissues to maximize power, while simultaneously allowing for both qualitative and quantitative differences among eQTLs present in each tissue.

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Introduction

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The availability of this information immediate insight into a biological basis for disease associations identified through genome-wide association (GWA) studies, and can help to identify networks of genes involved in disease pathogenesis

However, most studies to date have been conducted in a single immortalized peripheral cell type, and it is unclear to what extent these findings will translate to human disease mapping across more varied cell types.

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The Genotype Tissue Expression Project, GTEx, Project will provide the data necessary to address this situation: by 2016, the resource is expected to enroll a total of approximately 900 post-mortem donors, with approximately 30 tissues collected from each donor, and the project will generate extensive genotype data and RNA-seq data on each individual. However, available methods are limited in their ability to jointly analyze data on all tissues to maximize power, while simultaneously allowing for both qualitative and quantitative differences among eQTLs present in each tissue.

0.1 Aim 1

Develop methods of estimating the posterior effect size across multiple subgroups, thereby mapping eQTLs

- Combine information across tissues
- Report an effect size
- Capture distinct variation in effect sizes within and between subgroups: 'patterns of sharing'

0.2 The Setting

- How do we quantify the effect of a particular SNP on gene expression among tissues?
- Approach 1: The Isolationist Approach

Initial approaches to quantify the effect of a particular sno pon gene expression considered only one tissue at a time, and ignored the effect of the snp on gene expression in other tissues. This fails to exploit the power of shared genetic variation in effects on expression - i.e. the information that the effect of the gene snp pair in one tissue can provide about the effect in another- and limits our understanding of multiple-tissue phenotypes.

0.3 The Setting

• How do we quantify the effect of a particular SNP on gene expression among tissues?

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- Approach 2: The Joint Committee Approach the multivariate nature of this activity
- Many different patterns of sharing of effects among tissues.
- But how do we learn about the nature and frequency?

Here, we recognize the multivariate nature of the effects. Now, we aim to make an inference about the vector of true effects across tissue and thus model this effect b as an R dimensional vector composed of the true effect of the gene snp pair across multiple tissues. across tissue and thus model this effect across tissue and thus model this effect. Acknowledging the multivariate nature of this effect opens the door to illustrating many different patterns of sharing of effects among tissues. But how do we learn about the nature and freugnecy of these patterns?

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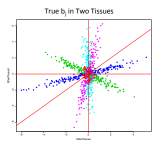
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0.4 Considering ALL the evidence!

- Each eQTL may follow a particular pattern of activity
- Within these groups, the tissues exhibit characteristic patterns of sharing of effects
- Captured by considering the covariance structure of the genetic effects among tissues.
- Natural mixture model: Each component of the mixture is defined by the prior covariance matrix U_k from which the vector of standardized effect sizes of this class is thought to be drawn.
- Learn relative frequencies from the data

Not only do we have many tissues, we also have an entire genome from which to ?learn? about these patterns of sharing. We are detectives with many tissues and many gene snp pairs at our disposal, and it is important we consider all the evidence. -For each gene snp pair, we observe a vector of standardized effect sizes and their standard error - assume that they descend from some true effect size b, - Thus as an additional level of combining information, we assume that Each eQTL may follow a particular pattern of activity characterized by its effects across tissues. - Within these groups, the tissues exhibit characteristic patterns of sharing, which can be captured by considering the covariance structure of the genetic effects among tissues. - This lends itself to a natural mixture model, in which we assume all the gene-snp pairs arise from a mixture of a finite number of Gaussian distributions with unknown parameters. - As mentioned, because of the the multivariate nature of this activity, within a particular 'pattern of sharing' some tissues may be more active than others, but not completely on or off. -Thus each component of the mixture is defined by the prior covariance matrix from which the vector of standardized effect sizes b_i is thought to be drawn. The covariance matrix of the true effects thus reflects the pattern of sharing (i.e., the diagonal elements

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of the component-specific covariance matrix then represents the variance of the effect size within and and the off-diagonal between tissues. A large prior variance in one tissue and small in another means that effects in tissue 1 tends to be large while effects in tissue 2 tend to be mall. - Because we can?t know the ?true covariance matrix for each? gene snp pair, we ?learn? the relative proportions of each pattern of sharing (i.e., the mixture weigths) from the data and model each bj as arising from a mixture that captues all the covariance patterns.

As a critical innovation on our previous method ([9, 10]), these matrices contain distinct diagonal and off-diagonal elements which reflect data-specific patterns of variation within and covariance between subgroups (tissues). This captures the variation in effect sizes within and between subgroups better than restricting effects to simply ?shared? or ?unshared? between subgroups.

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Previous work from our lab considered only the idea that the covariance between two tissues was the same across tissues thought to contain a QTL in a given pattern, or 'configuration', and thus failed to incorporate the much richer covariance structure between tissues.

The primary novelty of this proposal is to estimate this multivariate posterior distribution on the effect size in a data-sensitive way - i.e., using the mixture model to capture information about the covariance structure among subgroups (here, tissues).

0.5 So what are the Prior Covariance Matrices $U_k s$ specifying?

Suppose we have just two tissues

 \bullet Direction defined by relative ratio in effect size between tissues, specified in prior covariance of \boldsymbol{b}

Generative
$$U_k$$
 for blueSNPs $\begin{pmatrix} Var(b_1) = blue2.0 & Cov(b_1, b_2) = 0.56 \\ Cov(b_2, b_1) = 0.56 & Var(b_2) = blue0.20 \end{pmatrix}$

Additional novelty: Ratio between tissues is flexible (not simply shared or tissue-specific) and data sensitive (stay-tuned)

To illustrate the utitlity of using a variety of covariance matrices, consider that we have snps of 4 ?types? here defined, by their effects in tow tissues. Snps of the blue class tend to have large effects in tissue 1 and small in tissue 2, while snps in the purple class have very large effects in tissue 2 and small effects in tissue 2. These directions are thus specified in the prior covariance matrix which defines the direction? here simply ratio - in prior effect size between tissues,

The 45 degree angle and the lines would be simply using tissue specific or shared effects, while we can have a much richer understanding of the relationship between effect sizes using a set of covariance matrices that aims to recapitulate patterns found in the data

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0.6 Why care about the effect size?

- Comparisons among tissues in which the QTL is called active, and among gene-snp pairs with a similar degree of activity in a given tissue.
- The addition of the quantitative comparison captures the continuous nature of biological phenomenon.

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- How confident are we in the sign of the effect?
- Acknowledge the many patterns of sharing present in the data, wide array of prior covariance matrices allows our gene-snp pair to find it's true pattern of sharing

Given that a snp is called active in two tissues, we want to make more statements about our confidence in the sign and magnitude of the effect among tissues.

Similarly, if many snps are called active in a particular tissue, we can resolve differences among these gene=snp pairs

using this wide array of prior covariance matrices allows our gene snp pair to ?find it?s true pattern of sharing?

Materials and Methods

We assume the following mixture prior for the R dimensional vector of true effects, \mathbf{b}_{j} represents the genetic effect of SNP-gene pair j across R4tissues:

As mentioned above, each component of the mixture distribution is characterised by these prior covariance matrices, U_k which capture the pattern of effects across tissues.

0.7 Covariance Matrices

For a given ω_l , we specify 4 'types' of RxR prior covariance matrices $U_{k,l}$.

- 1. $U_{k=1,l} = \omega_l \, \mathbf{I}_R$
- 2. $U_{k=2,l} = \omega_l X_z$ The (naively) estimated tissue covariance matrix as estimated from the column-centered J ×R matrix of Z statistics, Z_{center} : $\frac{1}{l} Z_{center} Z_{center}$
- 3. $U_{k=3,l} = \omega_l \frac{1}{J} V_{1...p} d_{1...p}^2 V_{1..p}^t$ is the rank p eigenvector approximation of the tissue covariance matrices, i.e., the sum of the first p eigenvector approximations, where $_{1...p}$ represent the eigenvectors of the covariance matrix of tissues and $_{1...p}$ are the first p eigenvalues.
- 4. $U_{k=4:4+Q,l} = \frac{1}{J}((\Lambda \mathbf{F})^t \Lambda \mathbf{F})_q$ corresponding to the q_{th} sparse factor representation of the tissue covariance matrix
- 5. $U_{k=4+Q+1,l} = \frac{1}{J} (\Lambda \mathbf{F})^t \Lambda \mathbf{F}$ is the sparse factor representation of the tissue covariance matrix, estimated using all q factors.
- 6. $U_{k=4+Q+1,l} = \frac{1}{J} (\Lambda \mathbf{F})^t \Lambda \mathbf{F}$ is the sparse factor representation of the tissue covariance matrix, estimated using all q factors.
- 7. $U_{k=5+Q:R+4+Q,l} = \frac{1}{J} ([100..]'[100...])$
- 8. $U_{k=R+5+Q,l} = \frac{1}{7} ([111...]'[111...])$
- 9. [1000...] or [111...] represent configurations such that given membership, b_j arise from the same prior variance.

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0.8 Deconvolution

To retrieve a 'denoised' or 'deconvoluted' estimate of the non-single rank dimensional reduction matrices, I then perform deconvolution.em which initializes the EM algorithm with the matrices specified in (2), (3) and (5). The final results of this iterative procedure preserves the rank of the initialization matrix, and allows us to use the 'true' effect component as missing data in deconvoluting the prior covariance matrices. In brief, this algorithm works by treating not only the component identity but also the true effect b_j as unobserved data, and maximising the likelihood over the expectation of the complete data likelihood, considering the values b_j as extra missing data (in addition to the indicator variables

 q_{ij}). This allows us to write down the? full data? log likelihood as follows write down the? full data? log likelihood as follows:

$$\phi = \sum_{J} \sum_{K} q_{jk} ln \alpha_k N(\hat{\mathbf{b}_j} | \theta, U_k + V_j)$$

$$???? \sum_{J} \sum_{K} q_{jk} ln \alpha_k N(\mathbf{b}_j | \theta, U_k)$$
(2)

Where

 $\alpha_k represents p(\mathbf{k}) and q_{jk} is the latent identifier variable.. In brief, this is how the deconvolution. employed the property of the$

Generation of List of Covariance Matirce I then use these three non single-rank covariance matrix in place of our original choice of the empirical covariance matrix, SFA and SVD approximations. Here, I also used the Identity (K=1), 5 single-rank SFA factors (K=4-9), and the 44+1 eqtlbma.lite configurations (K=10:54 in steps (7) and (8) to a assemble a full list of covariance matrices. This is 54 matrices, and we then proceed to chooses an $^{\prime}L^{\prime}$ element grid according to the range of effect sizes present in the initial 16,069 x 44 matrix of strong Z statistics to create a KxL list of covariance matrices. In the gtex data set we choose a grid with 22 omegas for a total of 1188 covariance matrices.

0.9 Mixture Weights

We now need to compute the mixture weights pi_{kl} hierarchically - that is, using all of the data to determine the optimal mixture of covariance matrices. I use a randomly chosen set of 20,000 gene snp pairs to estimate these mixture proportions. This set does not contain the strongest gene-snp pairs, and thus will allow for substantial shrinkage, as a majority of these gene-snp pairs will have their likelihood maximized at low ω components.

To compute the likelihood at each component: For a given gene-snp pair, the Likelihood on b:

$$\hat{\boldsymbol{b}}|\boldsymbol{b} \sim N_R(\boldsymbol{b}, \hat{V})$$
 (3)

Furthermore, we take advantage of the fact that $N_R(\hat{\mathbf{b}}_i; \theta, U_k + \hat{V}_i)$

Thus we can now treat the matrices U_k as fixed and computed the JxK matrix of likelihoods to the combinations of weights which maximizes the probability of observing the data.

1 Posterior Quantities

Now that I have the estimated these prior mixture weights stored in the vector **pis**. I proceed to the inference step, where I compute the posterior weights and corresponding

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posterior quantities across all original 16069 gene snp pairs. In brief, the posterior mean, post covariance matrix and tissue specific tail probabilities are computed across all K components for each gene snp pair, and then weighted according to the posterior weights. This is performed in the **weightedquants** step.

We know that for a single multivariate *Normal* the posterior on $b|U_0$ is simply:

$$\boldsymbol{b}|\hat{\boldsymbol{b}} \sim N_R(\boldsymbol{\mu}_1, U_1)$$

where:

•
$$\mu_1 = U_1(\hat{V}^{-1}\hat{b});$$

•
$$U_1 = (U_0^{-1} + \hat{V}^{-1})^{-1}$$
.

Furthermore, a mixture-multivariate normal prior and a normal likelihood yields a mixture multivariate posterior, where the final posterior distribution is simply a weighted combination of multivariate normal distributions, each now characterized by it's posterior mean μ_{1k} and covariance $U_1 = (U_0^{-1} + \hat{V}^{-1})^{-1}$.

$$p(\mathbf{b}_{|}\hat{\mathbf{b}}, \hat{V}, \hat{\boldsymbol{\pi}}) = \sum_{k=1, l=1}^{K, L} \sim N_{R}(\boldsymbol{\mu}_{1kl}, U_{1kl}) p(z = k, l | \hat{\boldsymbol{b}}, \hat{V}, \hat{\boldsymbol{\pi}}),$$

$$= \sum_{k=1, l=1}^{K, L} \sim N_{R}(\boldsymbol{\mu}_{1kl}, U_{1kl}) \tilde{\boldsymbol{\pi}}_{k, l}$$
(4)

Furthermore, the posterior weights or responsibilites combine hierarchical and snp-specific information, as they are proportional to the product of the original mixture weights (estimated hierarchically) and the likelihood for a particular gene snp pair, which allows the gene snp pair to 'find its' true match. A large likelihood in a particular component will mean that the majority of the posterior weight is at this component, and correspondingly, the majority of the posterior mean will be comprised of the posterior mean at this component which will specify a large effect at this component. If a SNP shows a relatively 'flat likelihood' at all components, than the posterior weights will appear similar to the prior weights, and correspondingly, the posterior mean will look a lot like the null case (since the prior weights are computed from 'mostly null data' and thus the prior weights will heavily weight the components with small posterior means (as determined by small prior variance in U_k).

 $\tilde{\pi}_{k,l} = P(Component|Data) \propto P(Data|Comp.) \times P(Comp.)$ Combine hierarchical and snp-specific information

Allows pair to find its true match!

Here, the posterior weight $\tilde{\pi}_{k,l}$ is simply

$$\tilde{\pi}_{k,l} = \frac{p(\hat{\mathbf{b}}_j | \hat{V}_j, z_j = k, l) \hat{\pi}_{kl}}{\sum_{k=1,l=1}^{K,L} p(\hat{\mathbf{b}}_j | \hat{V}_j, z_j = k, l) \hat{\pi}_{kl}}$$
(5)

This will return a set of 6 files containing the JxR matrix of posterior means, marginal variances, tail probabilities, local false sign rates, and the JxK matrix of posterior weights. Checkpoint = FALSE means that the files will be created (rather than simply outputting an object array which contains the posterior quantities.

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2 Testing and Training

In order to determine the optimal number and rank of the covariance matrices, we divide our data set into a training and test data set, each containing 8000 genes.

In the training set, we proceed as above: choosing the top SNP for each of the 8000 genes, creating a list of covariance matrices through deconvolution and grid selection of these top 'training gene-snp' pairs.

Then, within the training data, we similarly choose a random set of gene-snp pairs (restricting our analysis to genes contained in the training set. Again, we choose 20,000 random-gene snp pairs and use the EM algorithm to learn the mixture proportions π from this data set.

We then use the KxL vector of π from the training set to estimate the log likelihood of each data point in the test data set. If our model is 'overfit' to the training data set, than a larger number of covariance matrices may actually decrease the test log-likelihood.

I found that the K=1188 set of covariance matrices containing the Identity, the denoised empirical covariance matrix, rank 5 SFA approximation and rank 3 SVD approximation as well as 5 single-rank SFA factors and the 45 eqtl.bma.lite configurations maximized this likelihood.

3 Training and Testing Procedure: Estimating Hierarchical Weights

We wish to choose the model which best maximizes the probability of observing the data set.

Incomplete Data likelihood:

$$L(\boldsymbol{\pi}; \hat{\boldsymbol{b}}) = \prod_{j=1}^{J} \sum_{k}^{K} \pi_k P(\hat{\boldsymbol{b}_j} | z_j = k)$$

$$(6)$$

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- To estimate the hierarchical prior weights π_k : compute the likelihood at each each gene snp pair j by evaluating the probability of observing $\hat{\boldsymbol{b}}_j$ given that we know the true \boldsymbol{b}_j arises from component k
- Use the EM algorithm to estimate the optimal combination of weights: How often does this particular covariance matrix occur in the data?

We then use these weights to estimate the test set log likelihood.

Results

Discussion

Supporting Information

Acknowledgments

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