An atlas of the genetic architecture of gene expression traits across the entire human body

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# Abstract

For most complex traits, gene regulation is known to play a crucial mechanistic role as demonstrated by the consistent enrichment of expression quantitative trait loci (eQTLs) among trait-associated variants. For this reason, the GTEx project has generated RNA-Seq data on hundreds of individuals across more than 40 tissues providing a comprehensive atlas of gene expression traits. Here, we systematically examined the local versus distant heritability as well as the sparsity versus polygenicity of protein coding gene expression traits in tissues across the entire human body. To determine tissue context specificity, we decomposed the expression levels into cross-tissue and tissue-specific components via orthogonal tissue decomposition (OTD). Regardless of tissue type, we found that local heritability can be well characterized with current sample sizes. Unless strong functional priors are used, the heritability due to distant variants cannot be estimated. We also find that the distribution of effect sizes is more consistent with a sparse architecture across all tissues. We also show that the cross-tissue and tissue-specific expression phenotypes constructed with our OTD model recapitulate complex Bayesian multi-tissue analysis results demonstrating that they reflect the expected biology. Finally, this knowledge was applied to develop prediction models of gene expression traits for all tissues. The prediction models, heritability for each tissue, and prediction performance R2 are made publicly available (<https://github.com/hakyimlab/PrediXcan>).

# Introduction

Regulatory variation plays a key role in the genetics of complex traits [1–3]. While many common diseases have are likely polygenic [4–6], it is unclear whether gene expression levels are also polygenic or instead have simpler genetic architectures. It is also unclear how much these expression architectures vary across genes [7]. Most human expression quantitative trait loci (eQTL) studies have focused on how local genetic variation affects gene expression in order to reduce the multiple testing burden that would be required for a global analysis [7,8]. Furthermore, when both local and distal eQTLs are reported [9–11], effect sizes and replicability are much higher for local eQTLs. Indeed, while the heritability of gene expression attributable to local genetic variation has been estimated accurately, large standard errors have prevented accurate estimation of the contribution of distal genetic variation to gene expression variation [11,12].

We assessed the ability of various models, with different underlying assumptions, to predict gene expression in order to both understand the underlying genetic architecture of gene expression and to further optimize predictors for our complex trait prediciton method, PrediXcan [13]. In our PrediXcan paper, we showed that a polygenic score model was suboptimal to more sparse models like the elastic net model with . However, a top eQTL only model did not do as well as the elastic net either [13], suggesting that for many genes, the genetic architecture is sparse, but not regulated by one SNP. Gene expression traits with sparse architecture should be better predicted with models such as LASSO (Least Absolute Shrinkage and Selection Operator), which prefers solutions with fewer parameters, each of large effect [14]. Conversely, highly polygenic traits should be better predicted with ridge regression or similarly polygenic models that prefer solutions with many parameters, each of small effect [15–17]. To obtain a more thorough understanding of gene expression achitecture, we used the hybrid approaches of the elastic net and BSLMM (Bayesian Sparse Linear Mixed Model) [18] to quantify sparse and polygenic effects.

Most previous human eQTL studies were performed in whole blood or lymphoblastoid cell lines due to ease of access or culturabilty [9,19,20]. The Genotype-Tissue Expression (GTEx) Project aims to examine the genetics of gene expression more comprehensively and recently published a pilot analysis of eQTL data from 1641 samples across 43 tissues from 175 individuals confirming that eQTLs are highly shared across tissues [21]. We also showed that gene expression predictors perform well across tissues [13]. In order to harness this cross-tissue effect for prediction and to better understand the genetic architecture of tissue-specific and cross-tissue gene regulation, we developed a mixed effects model called orthogonal tissue decomposition (OTD) to determine the cross-tissue and tissue-specific components of gene expression in the rich GTEx dataset. We modeled the underlying genetic architecture of the cross-tissue and tissue-specific gene expression components and developed predictors for use in PrediXcan [13].

# Results

## Local genetic variation can be well characterized for all tissues

We estimated the heritability of gene expression in whole blood from the Depression Genes and Networks (DGN) cohort (n=922) [20] using a mixed-effects model (see Methods) and calculated variances using restricted maximum likelihood as implemented in GCTA [22]. We fit two joint models, each with the same local genetic relationship matrix (GRM) and a two different distal GRMs. The local GRM was derived from SNPs within 1 Mb of each gene in both models. In model 1, the distal GRM was derived from all SNPs on non-gene chromosomes. In model 2, the distal GRM was derived from SNPs that are located on non-gene chromosomes and are eQTLs in the Framingham Heart Study (FHS) cohort (n=5257, FDR < 0.05) [23]. The mean local h2 was similar in both models, 0.123 in model 1 and 0.130 in model 2 (Fig 1). Of expressed genes, 52.6% and 54.6% had a positive 95% confidence interval (CI) for the local estimate in models 1 and 2, respectively. The mean distal h2 was 0.123 with a standard error (SE) of 0.284 in model 1, while the mean distal h2 was 0.076 with a SE of 0.142 in model 2. Using functional priors (known eQTLs) to define distal h2 in model 2 increased the percentage of genes with a positive CI from 3.2% to 4.2% (Fig 1). In model 2, the maximum local h2 was 0.93 with a standard error (SE) of 0.009 while the maximum distal h2 was 0.91 with a SE of 0.16.

We also estimated the heritability of gene expression in the 9 tissues from the GTEx Project with the largest sample sizes, which ranged from 190-361 samples (see Methods). Mean local h2 estimates ranged from 0.028-0.044 with 4.4-8.6% of genes having a positive CI (Fig 2). While the reduced sample sizes reduced the overall mean, maximum local h2 estimates in each tissue ranged from 0.79-0.91, similar to the maximum seen in DGN. As expected, distal h2 (eQTLs on non-gene chromosomes) could not be reliably estimated in these smaller sample sizes as less than 1% of genes in each tissue had a positive CI in the joint model (Fig 3-**[SUP]**)). Thus, we focus on the local architecture in subsequent sections.

## The effect of local genetic variation on gene expression is sparse rather than polygenic

We performed 10-fold cross-validation using the elastic net [24] to test the predictive performance of local SNPs for gene expression across a range of mixing parameters, . The that gives the largest cross-validation R2 informs the sparsity of each gene expression trait. That is, at one extreme, if the optimal (equivalent to ridge regression), the gene expression trait is highly polygenic, whereas if the optimal (equivalent to LASSO), the trait is highly sparse. We found that for most gene expression traits, the cross-validated R2 was suboptimal for and , but nearly identically optimal for through in the DGN cohort (Fig 4). An was also clearly suboptimal for gene expression prediction in the nine GTEx tissues, while models with or 1 had similar predictive power (Fig 5-**[SUP]**). This suggests that for most genes, the effect of local genetic variation on gene expression is sparse rather than polygenic.

To further examine sparsity and polygenicity, we used BSLMM [18] to define the total proportion of variance in expression explained by sparse and polygenic effects together (PVE) and the proportion of this genetic variance explained by sparse effects (PGE) for local SNPs in each gene in the DGN cohort. The PVE can be thought of as a Bayesian estimate of chip heritability and, indeed, there is a strong correlation between BSLMM-estimated PVE and GCTA-estimated h2 (Fig 6A, R=0.96). For genes with large PVE, the PGE also was large, indicative of a sparse genetic architecture. For example, all genes with PVE > 0.50 had PGE > 0.82 and their median PGE was 0.989 (Fig 6B). The median PGE for genes with PVE > 0.1 was 0.949. Fittingly, for most (96.3%) of the genes with PVE estimates > 0.10, the median number of SNPs included in the model was no more than 10 (Fig 6C).

Interestingly, when we applied BSLMM to the GTEx data, we found that many genes had strikingly larger BSLMM-estimated PVE than GCTA-estimated h2 (Fig 7). This likely reflects the increased power of the BSLMM method at the lower sample sizes () present in GTEx to estimate variance explained when the trait is more sparse than polygenic. GCTA assumes an underlying polygenic model, but as we saw in DGN, BSLMM-estimated PVE and GCTA-estimated h2 are more correlated when the sample size is larger (n=922, Fig 6A). As we observed in DGN, genes with larger PVE estimates were more likely to have a PGE estimate approaching 1 with a lower credible set greater than 0.01 in each of the nine GTEx tissues (Fig 8).

## Cross-tissue expression phenotype has increased predictive power and recapitulates known multi-tissue eQTL target genes

In order to better understand the context specificity of gene expression regulation, we developed a method called orthogonal tissue decomposition (OTD), which uses a mixed effects model to generate cross-tissue and tissue-specific gene expression levels (see Methods). Using a marginal model with just the local GRM, we estimated the local h2 of cross-tissue gene expression and tissue-specific gene expression in the nine tissues with the most samples. The cross-tissue heritabilities were larger and the standard errors were smaller than the tissue-specific estimates (Fig 9-**[SUP]**). The percentage of GCTA h2 estimates with positive CIs was much larger for cross-tissue expression (17.3%) than the tissue-specific expressions (all less than 3%, Fig 10). Similarly, the percentage of BSLMM PVE estimates with a lower credible set greater than 0.01 was 49% for cross-tissue expression, but ranged from 24-27% for tissue-specific expression (Fig 11).

We also compared the cross-tissue h2 from the OTD to h2 estimates from the pre-OTD measures of gene expression in each of the nine tissues, which we term whole tissue expression. Again, the cross-tissue heritabilities were larger and the standard errors were smaller than the whole tissue estimates (Fig 12-**[SUP]**), though less striking than the tissue-specific comparison. The percentage of whole tissue h2 estimates with positive CIs ranged from 4.4-8.6% and thus were all larger than the tissue-specific postive CI percentages, but smaller than the cross-tissue percentage (Fig 2). Cross-tissue BSLMM PVE estimates had lower error than whole tissue PVE (Fig 8, Fig 11). Like whole tissue expression, cross-tissue and tissue-specific expression showed better predictive performance using the elastic-net when than when (Fig 13-**[SUP]**). Cross-tissue predictive performance exceeded that of both tissue-specific and whole tissue expression as indicated by higher cross-validated R2 (Fig 5-**[SUP]**, Fig 13-**[SUP]**).

We compared our OTD results to those from a joint multi-tissue eQTL analysis method [25], which was previously performed on a subset of the GTEx data [21]. The results of this analysis include eQTL posterior probabilities for each of the nine tissues, which can be interpreted as the probability a SNP is an eQTL in tissue *x* given the data. Using the top eQTL for each gene, we defined an entropy statistic (see Methods) that combines the nine posterior probabilities into one value such that higher entropy values mean the gene is more likely to be regulated by the same eQTL across all nine tissues, rather than in just a subset of the nine. We observed a strong correlation between entropy and both the cross-tissue expression heritability (R = 0.082, Fig 14A) and PVE (R = 0.12, Fig 14B) estimates, using the cross-tissue expression derived from the OTD. Thus, genes with high cross-tissue heritability are more likely to have cross-tissue eQTLs, confirming that OTD is capturing the cross-tissue component of gene expression. Also, the correlation between tissue-specific OTD gene expression PVE and the posterior probability that the gene has an eQTL in that tissue is strongest in each respective tissue, confirming that OTD also captures tissue-specific components of gene expression (Fig 15).

# Discussion

Because regulatory variation plays a key mechanistic role in the genetics of complex traits [1–3], we sought to comprehensively characterize the genetic architecture of gene expression across tissues. We accurately quantify the local heritability of gene expression in DGN whole blood and nine GTEx tissues. In DGN, the mean local h2 was 0.13, similar to that found in family studies of blood expression, where mean h2 ranged from 0.07-0.11 [11,12]. While we found that functional priors (known trans-eQTLs) can reduce the error of the estimate by reducing the number of genetic markers included in the genetic relationship matrix, larger sample sizes (n > 1000) are needed to accurately estimate distal heritability.

Using the hybrid polygenic-sparse approach of BSLMM (Bayesian Sparse Linear Mixed Model) [18], we show that the local architecture of gene expression is sparse (high PGE) for most heritable genes in both DGN and GTEx. Using the elastic net [24], we observed improved cross-validated expression prediction for across tissues, confirming the sparsity result. This result demonstrates that sparse effects can be identified with sample sizes in the hundreds rather than the thousands and is supported by many prior studies with sample sizes near 100 that identified replicable eQTLs near the transcription start sites of genes[9]. Conversely, for traits that are highly polygenic, e.g. height, BMI, schizophrenia, and bipolar disorder, thousands to tens of thousands of samples are needed to identify significant genetic signals [26]. Therefore, the distal contributions to expression h2 are likely to be more polygenic because they could not be accurately estimated here with sample sizes in the hundreds.

Our BSLMM analysis quantified the optimal number of SNPs to include for each gene. For example, the median number of SNPs for *ERAP2* was 1. When we previously plotted out-of-sample observed vs. predicted expression using elastic net () generated predictors for this gene, we saw three clusters, corresponding to each of the three genotypes for the causal variant [13]. Similarly, BSLMM estimated 2, 3, and 5 SNPs for *PEX6*, *NUDT2*, and *ERAP1*, respectively, consistent with the out-of-sample observed vs. predicted expression plots in our PrediXcan paper (see Figure 5 in [13]). Potentially due to variations in imputation quality of the input SNPs for expression prediction, it is useful to include more than the likely causal variants (i.e. elastic net rather than BSLMM) in the prediction for robustness. In addition, the elastic net is amenable cross-validation, while genome-wide cross-validation with BSLMM is impractical at current runtimes.

We developed a mixed effects model called orthogonal tissue decomposition (OTD) to determine the cross-tissue and tissue-specific components of gene expression in the GTEx dataset. Previous studys have shown that many eQTLs are shared across tissues [21,25]. In addition, because expression data from multiple tissues were available from the same individuals in GTEx, we could effectively use the multiple tissue samples as subject replicates in our OTD model. However, the tissue availability is unbalanced across individuals because of the difficulties of sample collection and the uneven quality of the tissues. By combining all available expression data in our OTD model, we found that estimates of the local heritability of cross-tissue gene expression have larger magnitude and improved standard errors compared to single tissue estimates due to the borrowing of information across all samples. Thus, OTD effectively increases power to estimate heritability. Comparing our OTD results to a previously performed joint multi-tissue eQTL analysis method [25], we show that genes with high cross-tissue heritability are more likely to have cross-tissue eQTLs, confirming that OTD is capturing the cross-tissue component of gene expression.

We confirmed that OTD also captures tissue-specific components of gene expression by showing the correlation between tissue-specific OTD gene expression PVE and the posterior probability that the gene has an eQTL in that tissue is strongest in each respective tissue. Interestingly, whole blood and thyroid appear to be outliers in that they each have a negative correlation with all the other tissues (Fig 15). In the GTEx pilot analysis, whole blood had the lowest levels of eQTL sharing with other tissues and thyroid had the largest number of cis-eQTL genes, which implies a higher number of tissue-specific eQTLs [21].

In this paper, we quantitate the genetic architecture of gene expression and develop predictors across tissues. We show that local heritability can be accurately estimated across tissues, but distal heritability cannot be reliably estimated at current sample sizes. Using two different approaches, the elastic net and BSLMM, we show that for local gene regulation, the genetic architecture is mostly sparse rather than polygenic. Using new expression phenotypes generated in our OTD model, we show that cross-tissue predictive performance exceeded that of both tissue-specific and whole tissue expression as indicated by higher elastic net cross-validated R2. Predictors generated in this study of gene expression architecture have been added to our PrediXcan database (<https://github.com/hakyimlab/PrediXcan>) [13] for use in future studies of complex trait genetics.

# Methods

## Genomic and Transcriptomic Data

### DGN Dataset

We obtained whole blood RNA-Seq and genome-wide genotype data for 922 individuals from the Depression Genes and Networks (DGN) cohort [20], all of European ancestry. For our analyses, we used the HCP (hidden covariates with prior) normalized gene-level expression data used for the *trans*-eQTL analysis in Battle et al. [20] and downloaded from the NIMH repository. The 922 individuals were unrelated (all pairwise < 0.05) and thus all included in downstream analyses. Imputation of approximately 650K input SNPs (minor allele frequency [MAF] > 0.05, Hardy-Weinberg Equilibrium [P > 0.05], non-ambiguous strand [no A/T or C/G SNPs]) was performed on the University of Michigan Imputation-Server (<https://imputationserver.sph.umich.edu/start.html>) [27,28] with the following parameters: 1000G Phase 1 v3 ShapeIt2 (no singletons) reference panel, SHAPEIT phasing, and EUR population. Approximately 1.9M non-ambiguous strand SNPs with MAF > 0.05, imputation R2 > 0.8 and, to reduce computational burden, inclusion in HapMap Phase II were retained for subsequent analyses.

### GTEx Dataset

We obtained RNA-Seq gene expression levels from 8555 tissue samples (53 unique tissue types) from 544 unique subjects in the GTEx Project [21] data release on 2014-06-13. Of the individuals with gene expression data, genome-wide genotypes (imputed with 1000 Genomes) were available for 450 individuals. While all 8555 tissue samples were used in the OTD model (described below) to generate cross-tissue and tissue-specific components of gene expression, we used the nine tissues with the largest sample sizes when quantifying tissue-specific effects. Tissues and sample sizes (both RNA-seq and genotypes available) included cross-tissue (), skeletal muscle (), whole blood (), skin from the sun-exposed portion of the lower leg (), subcutaneous adipose (), tibial artery (), lung (), thyroid (), tibial nerve () and left ventricle heart (). Approximately 2.6M non-ambiguous strand SNPs included in HapMap Phase II were retained for subsequent analyses.

## Partitioning local and distal heritability of gene expression

To investigate the proximity of gene expression regulation to each gene, we partitioned the proportion of gene expression variance explained by SNPs in the DGN cohort into two components: local (SNPs within 1Mb of the gene) and distal (eQTLs on non-gene chromosomes) as defined by the GENCODE [29] version 12 gene annotation. We calculated the proportion of the variance (narrow-sense heritability) explained by each component using the following mixed-effects model:

Assuming a random effects for and , where is the identity matrix, we calculated the total variability explained by local and distal components by estimating with restricted maximum likelihood (REML) using GCTA software [22]. For heritability analyses in the GTEx cohort, we removed the term from the model and only estimated marginal h2 due to the smaller sample sizes of both cross-tissue and tissue-specific expression levels compared to DGN.

## Determining polygenicity versus sparsity using the elastic net

We applied the elastic net [24] to model the effect of local genetic variation (SNPs within 1 Mb of gene) on the genetic architecture of gene expression. We used the cv.glmnet function in the R package glmnet [30,31] to perform 10-fold cross-validation of the elastic net across a range of mixing paramaters () to find the that maximized predictive performance, measured by Pearson's R2. Specifically, glmnet solves the following problem:

**[[Haky, should/can we simplify this equation? I just took it from** [**http://web.stanford.edu/~hastie/glmnet/glmnet\_alpha.html**](http://web.stanford.edu/~hastie/glmnet/glmnet_alpha.html)**, but they don't explain all the terms]]**

over a grid of values of covering the entire range [30,31]. This tuning parameter controls the overall strength of the penalty.

The elastic net penalty is controlled by mixing parameter , which spans LASSO (, the default) [14] at one extreme and ridge regression () [15] at the other. The ridge penalty shrinks the coefficients of correlated SNPs towards each other, while the LASSO tends to pick one of the correlated SNPs and discard the others. Thus, an optimal prediction R2 for means the gene expression trait is highly polygenic, while an optimal prediction R2 for means the trait is highly sparse. An optimal prediction R2 in between (e.g. ) means the trait has a mixed genetic architecture.

In the DGN cohort, we tested 21 values of the mixing parameter () for optimal prediction of gene expression of the 341 genes on chromosome 22. For the rest of the autosomes in DGN and for whole tissue, cross-tissue, and tissue-specific expression in the GTEx cohort, we tested .

## Quantifying sparsity with Bayesian Sparse Linear Mixed Models (BSLMM)

We used BSLMM [18] to model the effect of local genetic variation (SNPs within 1 Mb of gene) on the genetic architecture of gene expression. The BSLMM consists of a standard linear mixed model, with one random effect term, and with sparsity inducing priors on the regression coefficients [18]. We used the software GEMMA [32] to implement BSLMM for each gene using the following parameters:

gemma -g [localGenoFile] -p [geneExpFile] -a [snpAnnotFile] -bslmm 1 -s 100000 -o [outFile]

The -bslmm 1 option specifies a linear BSLMM and the -s 100000 option specifies the number of sampling steps per gene. The BSLMM estimates the PVE (the total proportion of variance in phenotype explained by the sparse effects and random effects terms together) and PGE (the proportion of genetic variance explained by the sparse effects terms). From the second half of the sampling iterations for each gene, we report the median and the 95% credible sets of the PVE, PGE, and the || parameter (the number of SNPs with non-zero coefficients).

## Orthogonal tissue decomposition

To better understand the context specificity of gene expression regulation, we developed a method called orthogonal tissue decomposition (OTD). This approach is an extension of our method to develop an intrinsic growth phenotype [33]. We applied OTD to GTEx Project [21] data and decomposed the expression of each gene into cross-tissue and tissue-specific components. The tissue availability is unbalanced across individuals because of the difficulties of sample collection and the uneven quality of the tissues. OTD decomposes the expression traits into orthogonal components as represented by the following model:

Specifically, to generate cross-tissue and tissue-specific expression levels, we used the lmer function in the R [34] package lme4 [35] to fit the following mixed-effects model:

fit <- lme4::lmer(expression ~ (1|SUBJID) + TISSUE + GENDER + PEERs)

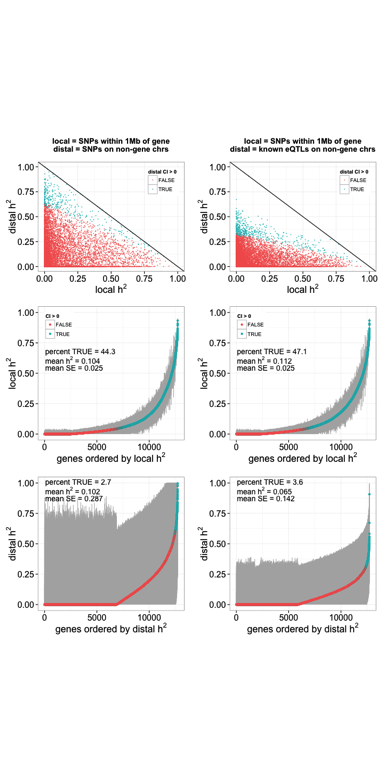
The model included whole tissue gene expression levels in 8555 GTEx tissue samples from 544 unique subjects. A total of 17,647 Protein-coding genes (defined by GENCODE [29] version 18) with a mean gene expression level across tissues greater than 0.1 RPKM (reads per kilobase of transcript per million reads mapped) were included in the model. SUBJID was a random effect and the covariates TISSUE, GENDER, and PEERs were fixed effects used to predict whole tissue expression levels (expression in the model). PEERs included the top 15 PEER factors estimated across all tissues using the R package PEER [36] to control for batch effects and experimental confounders. Cross-tissue expression was defined as the random effects from the model (ranef(fit)) and tissue-specific expression as the residuals (resid(fit)).

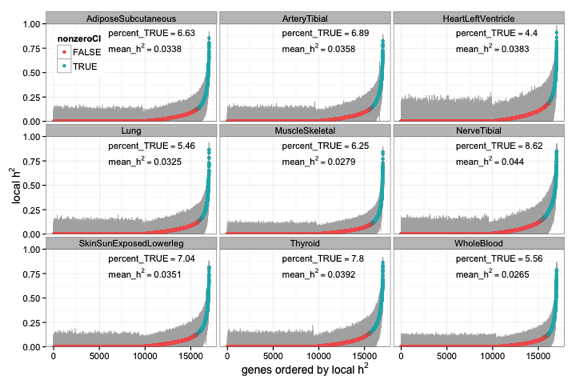
## Comparison of OTD PVE to multi-tissue eQTL results

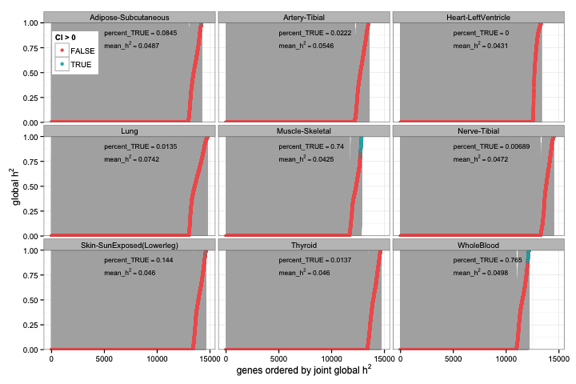
Using results from a joint multi-tissue eQTL analysis method [25] performed with a subset of the GTEx data (maximum n=175 in the nine tissues of the pilot phase, see [21]), we defined an entropy statistic to compare these results to those from our OTD method. The results of the multi-tissue analysis include eQTL posterior probabilities for each of the nine tissues, which can be interpreted as the probability a SNP is an eQTL in tissue given the data. Using the top eQTL for each gene , we defined the entropy as:

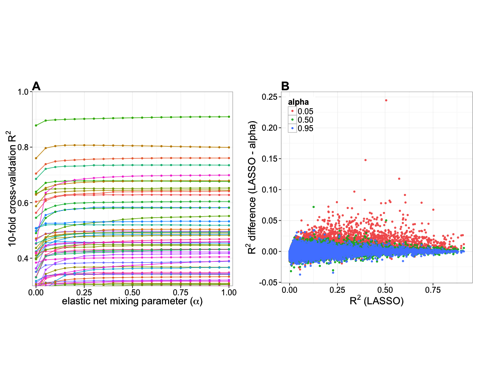
where is the eQTL probability in tissue normalized to 1 for each gene . Thus, eQTLs with higher entropy statistics are more likely to be cross-tissue eQTLs, rather than only regulating gene expression in one or a few tissues. We calculated the Pearson correlation between and the cross-tissue expression heritability and PVE for each gene to verify that our OTD method captures cross-tissue effects. We also calculated a Pearson correlation matrix between the posterior probabilities in each tissue from the multi-tissue eQTL method and the tissue-specific gene expression PVE from the OTD method.

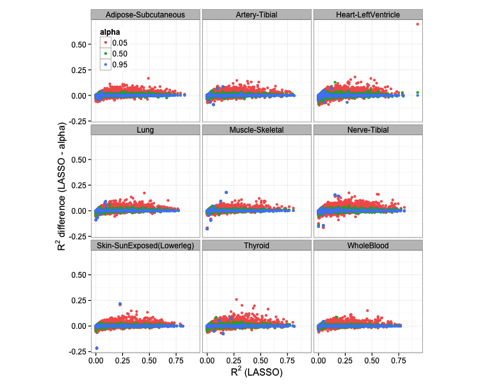
# Figures

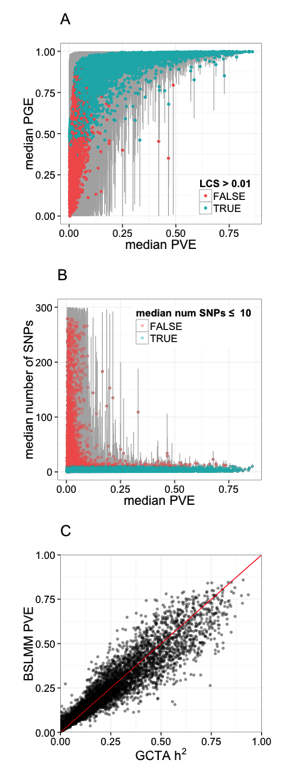


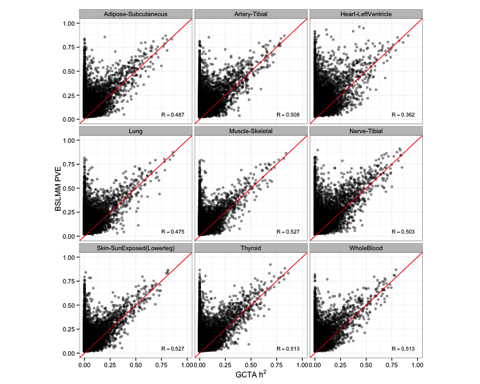


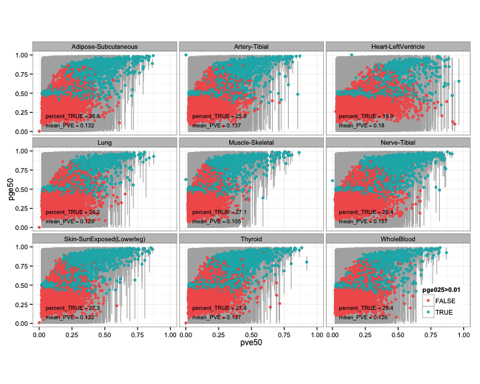


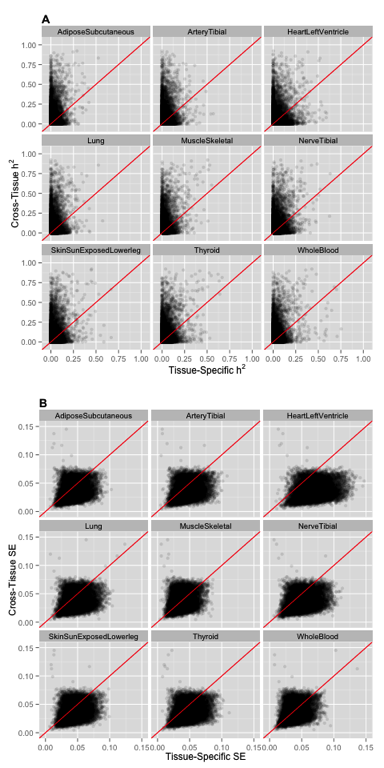


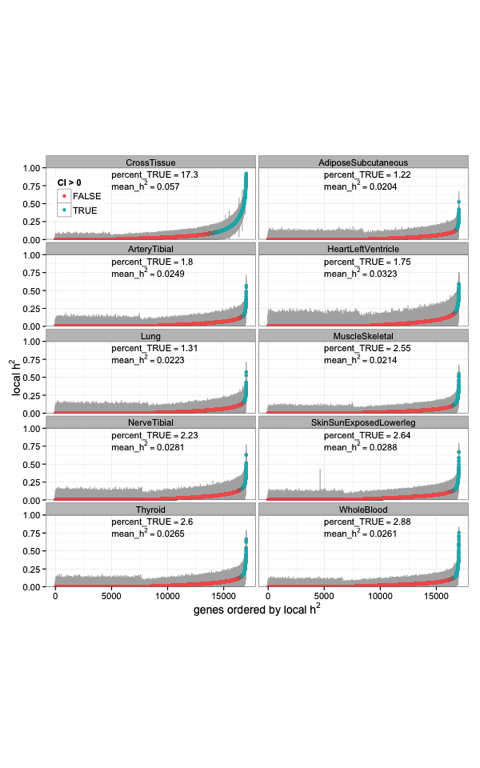


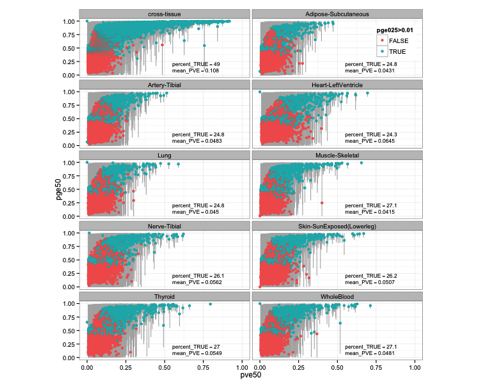


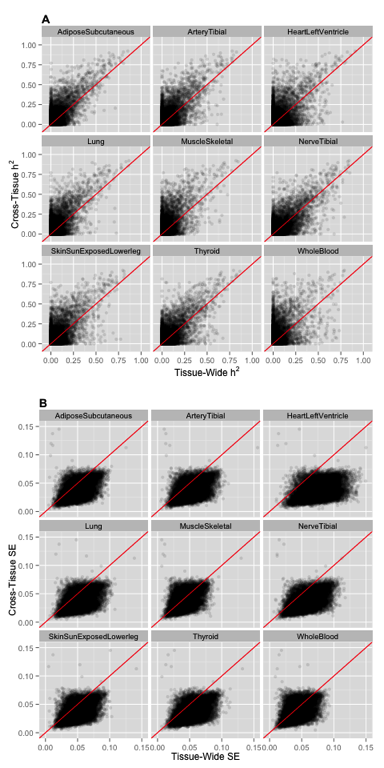


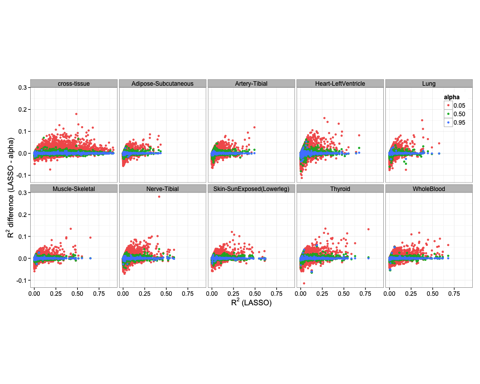


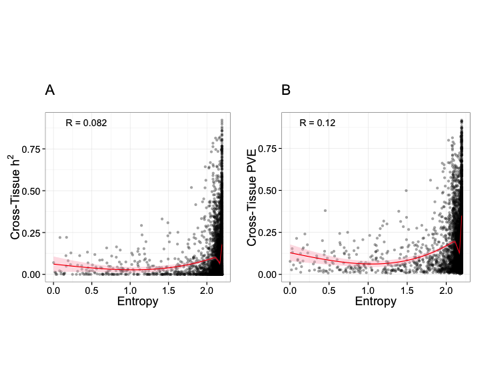


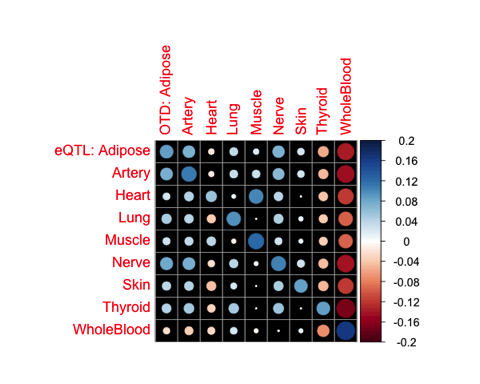












# Supplemental Figures

# Acknowledegments

We thank Nicholas Knoblauch and Jason Torres for initial pipeline development and planning.

# Grants

We acknowledge the following US National Institutes of Health grants: K12 CA139160 (H.K.I.), T32 MH020065 (K.P.S.), R01 MH101820 and R01 MH090937 (GTEx), P30 DK20595 and P60 DK20595 (Diabetes Research and Training Center), P50 DA037844 (Rat Genomics), UO1 GM61393 (Pharmacogenomics of Anticancer Agents Research), P50 MH094267 (Conte), and U19 HL065962 (PGRN Statistical Analysis Resource). H.E.W. was supported in part by start-up funds from Loyola University Chicago.

## GTEx data

The Genotype-Tissue Expression (GTEx) Project was supported by the Common Fund of the Office of the Director of the National Institutes of Health (commonfund.nih.gov/GTEx). Additional funds were provided by the NCI, NHGRI, NHLBI, NIDA, NIMH, and NINDS. Donors were enrolled at Biospecimen Source Sites funded by NCI Leidos Biomedical Research, Inc. subcontracts to the National Disease Research Interchange (10XS170), Roswell Park Cancer Institute (10XS171), and Science Care, Inc. (X10S172). The Laboratory, Data Analysis, and Coordinating Center (LDACC) was funded through a contract (HHSN268201000029C) to the The Broad Institute, Inc. Biorepository operations were funded through a Leidos Biomedical Research, Inc. subcontract to Van Andel Research Institute (10ST1035). Additional data repository and project management were provided by Leidos Biomedical Research, Inc.(HHSN261200800001E). The Brain Bank was supported supplements to University of Miami grant DA006227. Statistical Methods development grants were made to the University of Geneva (MH090941 & MH101814), the University of Chicago (MH090951,MH090937, MH101825, & MH101820), the University of North Carolina - Chapel Hill (MH090936), North Carolina State University (MH101819),Harvard University (MH090948), Stanford University (MH101782), Washington University (MH101810), and to the University of Pennsylvania (MH101822). The datasets used for the analyses described in this manuscript were obtained from dbGaP at <http://www.ncbi.nlm.nih.gov/gap> through dbGaP accession number phs000424.v3.p1.

## DGN data

NIMH Study 7 (GenRED I) - Data and biomaterials were collected in six projects that participated in the National Institute of Mental Health (NIMH) Genetics of Recurrent Early-Onset Depression (GenRED) project. From 1999-2003, the Principal Investigators and Co-Investigators were: New York State Psychiatric Institute, New York, NY, R01 MH060912, Myrna M. Weissman, Ph.D. and James K. Knowles, M.D., Ph.D.; University of Pittsburgh, Pittsburgh, PA, R01 MH060866, George S. Zubenko, M.D., Ph.D. and Wendy N. Zubenko, Ed.D., R.N., C.S.; Johns Hopkins University, Baltimore, MD, R01 MH059552, J. Raymond DePaulo, M.D., Melvin G. McInnis, M.D. and Dean MacKinnon, M.D.; University of Pennsylvania, Philadelphia, PA, RO1 MH61686, Douglas F. Levinson, M.D. (GenRED coordinator), Madeleine M. Gladis, Ph.D., Kathleen MurphyEberenz, Ph.D. and Peter Holmans, Ph.D. (University of Wales College of Medicine); this preprint is the author/funder. It is made available under a CC-BY 4.0 International license. bioRxiv preprint first posted online June 17, 2015; doi: <http://dx.doi.org/10.1101/020164>; The copyright holder for University of Iowa, Iowa City, IW, R01 MH059542, Raymond R. Crowe, M.D. and William H. Coryell, M.D.; Rush University Medical Center, Chicago, IL, R01 MH059541- 05, William A. Scheftner, M.D., Rush-Presbyterian. NIMH Study 18 - Data and biomaterials were obtained from the limited access datasets distributed from the NIH-supported “Sequenced Treatment Alternatives to Relieve Depression” (STAR\*D). STAR\*D focused on non-psychotic major depressive disorder in adults seen in outpatient settings. The primary purpose of this research study was to determine which treatments work best if the first treatment with medication does not produce an acceptable response. The study was supported by NIMH Contract # N01MH90003 to the University of Texas Southwestern Medical Center. The ClinicalTrials.gov identifier is NCT00021528. NIMH Study 52 (GenRED II) – Data and biomaterials in this release were collected in six projects that participated in the National Institute of Mental Health (NIMH) Genetics of Recurrent Early-Onset Depression (GenRED) project (1999-2009). The Principal Investigators and Co-Investigators were: New York State Psychiatric Institute, New York, NY, R01 MH 060912, Myrna M. Weissman, Ph.D.; Johns Hopkins University, Baltimore, MD, R01 MH059552, J. Raymond DePaulo, M.D., and James B. Potash, M.D., M.P.H.; University of Pennsylvania, Philadelphia, PA (1999-2005), and Stanford University (2006-2009), R01 MH61686, Douglas F. Levinson, M.D. (GenRED coordinator); University of Iowa, Iowa City, IW, R01 MH059542e, Raymond R. Crowe, M.D., and William H. Coryell, M.D.; Rush University Medical Center, Chicago, IL, R01 MH059541-05, William A. Scheftner, M.D.; and University of Pittsburgh, Pittsburgh, PA (1999-2003), R01 MH060866, George S. Zubenko, M.D., Ph.D., and Wendy N. Zubenko, Ed.D., R.N., C.S. NIMH Study 88 -- Data was provided by Dr. Douglas F. Levinson. We gratefully acknowledge the resources were supported by National Institutes of Health/National Institute of Mental Health grants 5RC2MH089916 (PI: Douglas F. Levinson, M.D.; Coinvestigators: Myrna M. Weissman, Ph.D., James B. Potash, M.D., MPH, Daphne Koller, Ph.D., and Alexander E. Urban, Ph.D.) and 3R01MH090941 (Co-investigator: Daphne Koller, Ph.D.).

## Computing resources

This work made use of the Open Science Data Cloud (OSDC) which is an Open Cloud Consortium (OCC)-sponsored project. This work was supported in part by grants from Gordon and Betty Moore Foundation and the National Science Foundation and major contributions from OCC members like the University of Chicago. this preprint is the author/funder. It is made available under a CC-BY 4.0 International license. bioRxiv preprint first posted online June 17, 2015; doi: <http://dx.doi.org/10.1101/020164>; The copyright holder for <https://www.opensciencedatacloud.org/> Grossman RL, Greenway M, Heath AP, Powell R, Suarez R, Wells W, White KP, Atkinson M, Klampanos I, Alvarez H, Harvey C and Mambretti J, The Design of a Community Science Cloud: The Open Science Data Cloud Perspective. (2012) <doi:10.1109/SC.Companion.2012.127> This work made use of the Bionimbus Protected Data Cloud (PDC), which is a collaboration between the Open Science Data Cloud (OSDC) and the IGSB (IGSB), the Center for Research Informatics (CRI), the Institute for Translational Medicine (ITM), and the University of Chicago Comprehensive Cancer Center (UCCCC). The Bionimbus PDC is part of the OSDC ecosystem and is funded as a pilot project by the NIH. <https://www.bionimbus-pdc.opensciencedatacloud.org/> Heath AP, Greenway M, Powell R, Spring J, Suarez R, Hanley D, Bandlamudi C, McNerney ME, White KP and Grossman RL, Bionimbus: A Cloud for Managing, Analyzing and Sharing Large Genomics Datasets. J Am Med Inform Assoc (2014) <doi:10.1136/amiajnl-2013-002155>

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