

# Survey of the Heritability and Sparsity of Gene Expression Traits Across Human Tissues

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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. Thus, understanding the genetic architecture of gene expression traits is key to elucidating the underlying mechanisms of complex traits. However, a systematic survey of the heritability and the distribution of effect sizes across all representative tissues in the human body has not been reported.

Here we fill this gap through analyses of the RNA-seq data from a comprehensive set of tissue samples generated by the GTEx Project and the DGN whole blood cohort. We find that local  $h^2$  can be relatively well characterized with 49% of expressed genes

showing significant  $h^2$  in DGN and 2-10% in GTEx. However, the current sample sizes ( $n < 362$  in GTEx) only allow us to compute distal  $h^2$  for a handful of genes (3% in DGN and  $<0.8\%$  in GTEx). Thus, we focus on local regulation. Bayesian Sparse Linear Mixed Model (BSLMM) analysis and the sparsity of optimal performing predictors provide compelling evidence that local architecture of gene expression traits is sparse rather than polygenic across DGN and all 40 GTEx tissues examined.

To further delve into the tissue context specificity, we decompose the expression traits into cross-tissue and tissue-specific components. Heritability and sparsity estimates of these derived expression phenotypes show similar characteristics to the original traits. Consistent properties relative to prior GTEx multi-tissue analysis results suggest that these traits reflect the expected biology.

Finally, we apply this knowledge to develop prediction models of gene expression traits for all tissues. The prediction models, heritability, and prediction performance  $R^2$  for original and decomposed expression phenotypes are made publicly available (<https://github.com/hakyimlab/PrediXcan>).

## Author Summary

Gene regulation is known to contribute to the underlying mechanisms of complex traits. 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. Regardless of tissue type, we found that local heritability can be well characterized with current sample sizes. Unless strong functional priors and large sample sizes 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 orthogonal tissue decomposition model recapitulate complex Bayesian multi-tissue analysis results. This knowledge was applied to develop prediction models of gene expression traits for

all tissues, which we make publicly available.

## Introduction

Regulatory variation plays a key role in the genetics of complex traits [1–3]. Methods that partition the contribution of environment and genetic components are useful tools to understand the biology underlying complex traits. Partitioning heritability into different functional classes has been successful in quantifying the contribution of different mechanisms that drive the etiology of diseases [3–5].

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 [6, 7]. Furthermore, when both local and distal eQTLs are reported [8–10], 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 [10, 11].

While many common diseases are likely polygenic [12–14], 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 [6].

The relative prediction performance of sparse and polygenic models can provide useful information about the underlying distribution of effect sizes. For example, if the true model of a trait is polygenic, it is natural to expect that polygenic models will predict better than sparse ones. 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 gene-level association method, PrediXcan [15]. When we calibrated the prediction model that was used in the PrediXcan paper, we showed that sparse models such as LASSO performed better than a polygenic score model. We also showed that a model that uses the top eQTL variant outperformed the polygenic score but did not do as well as LASSO or elastic net [15], suggesting that for many genes, the genetic architecture is sparse, but not regulated by a single SNP.

Thus, 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 [16]. 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 [17–19]. To obtain a more thorough understanding of gene expression architecture, we used the hybrid approaches of the elastic net and BSLMM (Bayesian Sparse Linear Mixed Model) [20] 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 culturability [8, 21, 22]. Although studies with a few other tissues have been published, comprehensive coverage of human tissues was not available until the launching of the Genotype-Tissue Expression (GTEx) Project. GTEx aims to examine the genetics of gene expression more comprehensively and has recently published a pilot analysis of eQTL data from 1641 samples across 43 tissues from 175 individuals [23]. Here we use a much larger set of 8555 samples across 53 tissues corresponding to 544 individuals. One of the findings of this comprehensive analysis was that a large portion of the local regulation of expression traits is shared across multiple tissues. Corroborating this finding, our prediction model based on whole blood showed robust prediction across the 9 core GTEx tissues chosen by initial sample sizes [15].

This shared regulation implies that there is much to be learned from large sample studies of easily accessible tissues. Yet, a portion of gene regulation seems to be tissue dependent [23]. 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 use a mixed effects model called orthogonal tissue decomposition (OTD) to decouple the cross-tissue and tissue-specific mechanisms 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 [15].

## Results

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

We estimated the local and distal heritability of gene expression levels in 40 tissues from the GTEx consortium and whole blood from the Depression Genes and Networks (DGN) cohort. The sample size in GTEx varied from 72 to 361 depending on the tissue, while 922 samples were available in DGN [22]. We used mixed-effects models (see Methods) and calculated variances using restricted maximum likelihood as implemented in GCTA [24].

For the local heritability component, we used variants within 1Mb of the transcription start and end of each protein coding gene, whereas for the distal component, we used variants outside of the chromosome where the gene was located. Different approaches to pick the set of distal variants were explored, but results were robust to different selections. See more details in Methods.

Table 1 summarizes the unconstrained heritability estimate results across all tissues. In order to obtain unbiased estimates of mean  $h^2$ , we allow the values to be negative when fitting the REML, as done previously [10,11]. This approach reduces the standard error of the estimated mean of heritability, especially important for the distal component. Even though each individual gene's distal heritability is noisy, averaging across all genes reduces the error substantially. For the DGN dataset, we were able to estimate the mean distal  $h^2$ . However for the GTEx samples, the sample size was too small and the REML algorithm became unstable when allowing for negative values. This numeric instability would cause a small number of genes with large positive (and noisy) heritability values to converge biasing the mean value. For this reason we do not show mean distal heritability estimates for GTEx tissues.

The left column of Fig. 1 shows the estimated local and distal  $h^2$  from DGN, this time using REML constrained between 0 and 1 (GCTA default) [24]. Even though many genes show relatively large point estimates of distal  $h^2$ , only the ones colored in blue are significantly different from zero. The local component of  $h^2$  is relatively well estimated in DGN with 48.6% of genes (6180 out of 12719) showing  $h^2$  values significantly different from zero. In contrast, the distal heritability is significantly different from zero for only 2.7% (343 out of 12719) of the genes.

**Table 1. Estimates of local  $h^2$  across whole tissues.**

tissue	n	mean $h^2$	mean se	prop CI > 0	num CI > 0	num expressed
DGN-WholeBlood	922	0.143	0.0292	0.486	6180	12719
Adipose-Subcutaneous	298	0.0385	0.0381	0.0735	1040	14205
AdrenalGland	126	0.0432	0.075	0.0425	601	14150
Artery-Aorta	198	0.0421	0.0561	0.0649	898	13844
Artery-Coronary	119	0.0371	0.0773	0.0337	476	14127
Artery-Tibial	285	0.0417	0.0402	0.0798	1080	13504
Brain-Anteriorcingulatecortex(BA24)	72	0.0275	0.133	0.031	450	14515
Brain-Caudate(basalganglia)	100	0.0367	0.091	0.0343	502	14632
Brain-CerebellarHemisphere	89	0.0492	0.11	0.0509	728	14295
Brain-Cerebellum	103	0.0504	0.094	0.0544	788	14491
Brain-Cortex	96	0.0451	0.0937	0.0393	578	14689
Brain-FrontalCortex(BA9)	92	0.0379	0.101	0.0341	496	14554
Brain-Hippocampus	81	0.0368	0.114	0.0285	414	14513
Brain-Hypothalamus	81	0.017	0.115	0.0235	347	14759
Brain-Nucleusaccumbens(basalganglia)	93	0.0293	0.0965	0.0292	426	14601
Brain-Putamen(basalganglia)	82	0.0324	0.108	0.0291	419	14404
Breast-MammaryTissue	183	0.0289	0.053	0.0365	537	14700
Cells-EBV-transformedlymphocytes	115	0.0578	0.0814	0.0497	619	12454
Cells-Transformedfibroblasts	272	0.0508	0.0424	0.0925	1180	12756
Colon-Sigmoid	124	0.0327	0.0807	0.0389	557	14321
Colon-Transverse	170	0.0358	0.059	0.0436	640	14676
Esophagus-GastroesophagealJunction	127	0.0318	0.0761	0.0332	469	14125
Esophagus-Mucosa	242	0.0416	0.0479	0.0764	1090	14239
Esophagus-Muscularis	219	0.0393	0.0505	0.069	969	14047
Heart-AtrialAppendage	159	0.0418	0.0638	0.0475	660	13892
Heart-LeftVentricle	190	0.0342	0.0558	0.0503	670	13321
Liver	98	0.033	0.0935	0.0299	405	13553
Lung	279	0.0315	0.0401	0.0577	853	14775
Muscle-Skeletal	361	0.0327	0.0324	0.0732	939	12833
Nerve-Tibial	256	0.0523	0.0445	0.095	1380	14510
Ovary	85	0.0369	0.0988	0.0258	364	14094
Pancreas	150	0.0469	0.0674	0.0557	777	13941
Pituitary	87	0.0379	0.107	0.0358	544	15183
Skin-NotSunExposed(Suprapubic)	196	0.0407	0.0532	0.0502	735	14642
Skin-SunExposed(Lowerleg)	303	0.0385	0.0385	0.0765	1120	14625
SmallIntestine-TerminalIleum	77	0.0365	0.112	0.0281	418	14860
Spleen	89	0.059	0.0998	0.0417	603	14449
Stomach	171	0.0315	0.0579	0.0367	533	14531
Testis	157	0.0539	0.0679	0.0727	1230	16936
Thyroid	279	0.0436	0.0413	0.0861	1260	14642
WholeBlood	339	0.0326	0.0328	0.0677	823	12160

Except for DGN-WholeBlood, all tissues are from the GTEx Project.

It has been shown that local-eQTLs are more likely to be distal-eQTLs of target

genes [25]. Thus, we tested whether restricting the distal genetic similarity computation

to QTLs (as determined in the Framingham mRNA dataset of over 5000 individuals [26]

independent of the DGN and GTEx cohorts) for other genes could improve distal

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heritability precision by prioritizing functional variants. We exclude eQTLs on the same chromosome as the tested gene to avoid contaminating distal  $h^2$  with cis associations.

Using functional priors (known eQTLs) to define distal  $h^2$  increased the percentage of genes with a positive CI from 2.7% (343 genes) to 3.6% (458) in whole blood (Fig. 1). A total of 125 genes have significant distal  $h^2$  by both approaches, i.e. all variants on other chromosomes or only known eQTL variants on other chromosomes.

However, using the subset of known eQTLs (from an independent source) in other chromosomes for computing distal heritability reduced the mean value from 3.4% to 1.6%. Therefore, while we gain some power to detect significant distal heritability by using cis eQTL priors, a good portion of the distal regulation is lost when using only the smaller subset of known cis-eQTL variants. We used functional priors to estimate distal  $h^2$  in the GTEx cohort, but less than 1% of genes had a positive CI (S1 Fig).

Given the limited sample size we will focus on local regulation for the remainder of the paper.

## Sparse local architecture implied by sparsity of best prediction models

Next, we sought to determine whether the local genetic contribution to gene expression is polygenic or sparse. In other words, whether many variants with small effects or a small number of large effects were contributing to expression trait variability. For this, we first looked at the prediction performance of a range of models with different degrees of polygenicity, such as the elastic net model with mixing parameter values ranging from 0 (fully polygenic, ridge regression) to 1 (sparse, LASSO).

More specifically, we performed 10-fold cross-validation using the elastic net [27] to test the predictive performance of local SNPs for gene expression across a range of mixing parameters ( $\alpha$ ). The mixing parameter that yields the largest cross-validation  $R^2$  informs the degree of sparsity of each gene expression trait. That is, at one extreme, if the optimal  $\alpha = 0$  (equivalent to ridge regression), the gene expression trait is highly polygenic, whereas if the optimal  $\alpha = 1$  (equivalent to LASSO), the trait is highly sparse. We found that for most gene expression traits, the cross-validated  $R^2$  was smaller for  $\alpha = 0$  and  $\alpha = 0.05$ , but nearly identically for  $\alpha = 0.5$  through  $\alpha = 1$  in the

DGN cohort (Fig. 2). An  $\alpha = 0.05$  was also clearly suboptimal for gene expression prediction in the GTEx tissues, while models with  $\alpha = 0.5, 0.95$ , or 1 had similar predictive power (S2 Fig). This suggests that for most genes, the effect of local genetic variation on gene expression is sparse rather than polygenic.

## Direct estimation of sparsity using BSLMM also points to sparse local architecture

To further confirm the local sparsity of gene expression traits, we turned to the BSLMM [20] approach, which models the genetic contribution as the sum of a sparse and a polygenic component. The parameter PGE in this model represents the proportion of genetic variance explained by sparse effects. Another parameter, the total variance explained (PVE) by additive genetic variants, is a more flexible Bayesian equivalent of the chip heritability we have estimated using a linear mixed model (LMM) as implemented in GCTA.

As anticipated, we find that for highly heritable genes, the sparse component is large. For example, all genes with  $PVE > 0.50$  had  $PGE > 0.82$  and their median PGE was 0.989 (Fig. 3A). 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. 3B).

## BSLMM outperforms LMM in estimating $h^2$ for small samples

Also as expected, we find that when the sample size is large enough, such as in DGN, there is a strong correlation between BSLMM-estimated PVE and GCTA-estimated  $h^2$  (Fig. 3C,  $R=0.96$ ). In contrast, when we applied BSLMM to the GTEx data, we found that many genes had measurably larger BSLMM-estimated PVE than GCTA-estimated  $h^2$  (Fig. 4). This is further confirmation of the local sparse architecture of gene expression traits: the underlying assumption in the GCTA (LMM) approach to estimate heritability is that the genetic effect sizes are normally distributed, i.e. most variants have small effect sizes. LMM is quite robust to departure from this assumption, but only when the sample size is rather large. For the relatively small sample sizes in GTEx ( $n \leq 361$ ), we found that a model that directly addresses the sparse component such as



BSLMM outperforms GCTA for estimating  $h^2$ .

## Orthogonal decomposition of cross-tissue and tissue-specific expression traits

Since a substantial portion of local regulation was shown to be common across multiple tissues [23], we sought to decompose the expression levels into a component that is common across all tissues and tissue-specific components. For this we use a linear mixed effects model with a person-level random effect. See details in Methods. We use the posterior mean of this random effect as an estimate of the cross tissue component. We consider the residual component of this model as the tissue specific component. Below we describe the properties of these derived phenotypes.

We call this approach orthogonal tissue decomposition (OTD) because the cross-tissue and tissue-specific components are assumed to be independent in the model. The decomposition is applied at the expression trait level so that the downstream genetic regulation analysis is performed separately for each derived trait, cross-tissue and tissue-specific expression, which greatly reduces computational burden. For all the derived phenotypes, one cross-tissue and 40 tissue-specific ones, we computed the local heritability and generated prediction models.

## Cross-tissue expression phenotype is less noisy and more predictable

Our estimates of  $h^2$  for cross tissue expression traits are larger than the corresponding estimates for each whole tissue expression traits (S3 Fig). This means that our OTD approach increases the ratio of genetically regulated component to noise by averaging across multiple tissues. In addition to the increased  $h^2$  we observe reduction in standard errors of the estimated  $h^2$ . This is partly due to the increased  $h^2$  – higher  $h^2$  are better estimated – but also due to the larger effective sample size for cross tissue phenotypes. There were 450 samples for which cross tissue traits were available whereas the maximum sample size for whole tissue phenotypes was 362. As consequence of this increase  $h^2$  and decreased standard errors, the percentage of cross  $h^2$  estimates with positive CIs was 19.8% whereas for whole tissue expression traits they ranged from

2.4-9.5%. Similarly, cross-tissue BSLMM PVE estimates had lower error than whole tissue PVE (S4 Fig, S5 Fig).

As for the tissue specific components, the cross-tissue heritability estimates were also larger and the standard errors were smaller reflecting the fact that a substantial portion of regulation is common across tissues (S6 Fig). The percentage of GCTA  $h^2$  estimates with positive CIs was much larger for cross-tissue expression (19.8%) than the tissue-specific expressions (all less than 4%, S1 Table). 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 (S5 Fig).

Cross-tissue predictive performance exceeded that of both tissue-specific and whole tissue expression as indicated by higher cross-validated  $R^2$  (S2 Fig, S7 Fig). Like whole tissue expression, cross-tissue and tissue-specific expression showed better predictive performance when using more sparse models. In other words elastic-net models with  $\alpha \geq 0.5$  predicted better than the ones with  $\alpha = 0.05$  (S7 Fig).

## Cross Tissue expresion phenotype recapitulates published multi-tissue eQTL results

We compared our OTD results to those from a joint multi-tissue eQTL analysis method [28], which was previously performed on a subset of the GTEx data [23] covering 9 tissues. In particular, we used the posterior probability of a gene being actively regulated (PPA) in a tissue. These analysis results are available on the GTEx portal.

First, we reasoned that genes with high cross tissue  $h^2$  would be actively regulated in most tissues so that the PPA of a gene would be roughly uniform across tissues. By contrast, a gene with tissue specific regulation would have concentrated posterior probability in one or a few tissues. Thus we decided to define a measure of uniformity of the posterior probability vector across the 9 tissues using the concept of entropy. More specifically, for each gene we normalized the vector of posterior probabilities so that the sum equaled 1. Then we applied the usual entropy definition (negative of the sum of the log of the posterior probabilities weighted by the same probabilities, see Methods). In other words, we defined an entropy statistic that combines the nine posterior

probabilities into one value such that higher entropy values mean the gene regulation is more uniform across all nine tissues, rather than in just a small subset of the nine.

Reassuringly, we found that the cross tissue phenotype heritability estimates were well correlated with the entropy statistics indicating that our derived cross tissue phenotype is capturing the common regulatory component. Figure 5 shows the comparison between this entropy statistics and cross-tissue expression heritability ( $R=0.082$ ) and PVE ( $R = 0.12$ ).

For the original whole tissue, we expected the whole tissue expression heritability to correlate with the posterior probability of a gene being actively regulated in a tissue as shown in Figure 6A. Strikingly, the tissue specific phenotype heritability showed highest correlation with the PPA of the matching tissue and lower correlations with other tissues, confirming that OTD also captures tissue-specific components of gene expression (Fig. 6B). However, for whole blood there is a consistently negative correlation with other tissues. This issue needs to be further investigated.

## Discussion

Motivated by the key role that regulatory variation plays in the genetic control of complex traits [1–3], we performed a survey of the heritability and patterns of effect sizes of gene expression traits across a comprehensive set of human tissues. We quantified the local and distal heritability of gene expression in DGN and 40 different tissues from the GTEx consortium. For the DGN dataset, we estimate the relative proportion of mean local and distal genetic contribution to gene expression traits. For GTEx samples it was not possible to estimate the mean distal heritability because of the limited sample size. As the number of GTEx samples grows to near 1000 individuals, we expect to be able to estimate these values.

In DGN (whole blood), the mean local  $h^2$  was 14.3% and the mean distal  $h^2$  was 3.4% such that the local variation contribution is estimated as  $14.3/(3.4+14.3) = 81\%$ . This is much higher than the 37% reported by Price et al. [11] based on blood expression data from a cohort of Icelandic individuals. This potentially underestimation of the distal component could be due to over-correction of confounders used in the preprocessing of the expression trait data we used. Indeed, PEER [29], SVA [30], and

other types of hidden confounder corrections have been shown to increase local eQTL replicability, but their consequences on distal regulation is not well understood. As larger sample sizes become available, we will test this hypothesis in GTEx data by computing the distal  $h^2$  without PEER factor correction.

We showed that restricting distal variants to known functional variants such as eQTL data from independent studies improves the precision of distal heritability estimates, but also reduces mean distal heritability estimates by half. For GTEx tissues, we computed the distal heritability using only the subset of functional (known eQTLs) variants to reduce the errors in the estimates at the expense of missing variants that contribute to the traits.

Using results implied by the improved predictive performance of sparse models and by directly estimating sparsity using BSLMM (Bayesian Sparse Linear Mixed Model), we show evidence that for highly heritable genes, local regulation is sparse across all the tissues analyzed here. For genes with moderate and low heritability the evidence is not as strong, but results are consistent with a sparse local architecture. Better methods to correct for hidden confounders that do not dilute distal signals and larger sample sizes will be needed to determine the properties of distal regulation.

Given that a substantial portion of local regulation is shared across tissues, we propose here to decompose the expression traits into cross-tissue and tissue-specific components. This approach, called orthogonal tissue decomposition, aims to decouple the shared regulation from the tissue-specific regulation. We examined the genetic architecture of these derived traits and find that they follow similar patterns to the original whole tissue expression traits. The cross-tissue component benefits from an effectively larger sample size than any individual tissue trait, which is reflected in more accurate heritability estimates and consistently better prediction performance. Encouragingly, heritability estimates of cross-tissue traits correlate well with a measure of uniformity of regulation across tissues defined as the entropy of the vector of probability for a gene to be regulated in a given tissue. Higher entropy genes will show more uniform regulation across tissues. As for the tissue-specific expression traits, we found that they recapitulate correlation with the vector of probability of tissue-specific regulation. The main application for which these traits were devised is to be used as prediction models in PrediXcan [15]. We expect results from the cross-tissue models to

relate to mechanisms that are shared across multiple tissues whereas results from the tissue-specific models will inform us about the context specific mechanisms.

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  $R^2$ . Predictors generated in this study of gene expression architecture have been added to our PredictDB database (<https://github.com/hakyimlab/PrediXcan>) for use in future studies of complex trait genetics.

## Materials and 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 [22], 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. [22] and downloaded from the NIMH repository. The 922 individuals were unrelated (all pairwise  $\hat{\pi} < 0.05$ ) 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 Michigan Imputation Server (<https://imputationserver.sph.umich.edu/start.html>) [31,32] 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  $R^2 > 0.8$  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 [23] 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 40 tissues with the largest sample sizes when quantifying tissue-specific effects (see Table 1). Approximately 2.6M non-ambiguous strand SNPs included in HapMap Phase II were retained for subsequent analyses.

**Framingham Expression Dataset.** We obtained exon array expression and genotype array data from [Tzumi, what is your final n?] 5257 individuals from the Framingham Heart Study [26].

We used the Affymetrix power tools (APT) suite to perform the preprocessing and normalization steps. First the robust multi-array analysis (RMA) protocol was applied which consists of three steps: background correction, quantile normalization, and summarization [33]. The background correction step uses antigenomic probes that do not match known genome sequences to adjust the baseline for detection, and is applied separately to each array. Next, the normalization step utilizes a "sketch" quantile normalization technique instead of a memory-intensive full quantile normalization. The benefit is a much lower memory requirement with little accuracy trade-off for large sample sets such as this one. Finally, the adjusted probe values were summarized (by the median polish method) into log-transformed expression values such that one value is derived per exon or gene. Additionally an analysis of the detection of probes above the background noise (DABG) was carried out. It provides further diagnostic information which can be used to filter out poorly performing probes and weakly expressed genes. The summarized expression values were then annotated more fully using the annotation databases contained in the huex10stprobeset.db (exon-level annotations) and huex10sttranscriptcluster.db (gene-level annotations) R packages available from Bioconductor [34,35]. In both cases gene annotations were provided for each feature.

Plink [36] was used to carry out several data wrangling and cleaning steps. The data wrangling steps included updating probe IDs, unifying data to "+" strand, and

updating locations to GRCh37. The data cleaning steps included a step to filter for variant and subject missingness and minor alleles, one to filter variants with Hardy-Weinberg exact test, and a step to remove unusual heterozygosity. Additionally we used the HRC-check-bin tool in order to carry out data wrangling steps required to make our data compatible with the Haplotype Reference Consortium (HRC) panel (<http://www.well.ox.ac.uk/wrayner/tools/>). Having been prepared thusly, the data were split by chromosome and pre-phased with SHAPEIT [37] using the 1000 Genomes phase 3 panel and converted to vcf format. These files were then submitted to the Michigan Imputation Server (<https://imputationserver.sph.umich.edu/start.html>) [31,32] for imputation with the HRC version 1 panel [38].

\*cite Shabalín AA: Matrix eQTL: ultra fast eQTL analysis via large matrix operations. *Bioinformatics* 2012, 28:1353?1358. [39]

## Partitioning local and distal heritability of gene expression

Motivated by the observed differences in regulatory effect sizes of variants located in the vicinity of the genes and distal to the 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 [40] version 12 gene annotation. We calculated the proportion of the variance (narrow-sense heritability) explained by each component using the following mixed-effects model:

$$Y_g = \sum_{k \in local} w_{k,g} X_k + \sum_{k \in distal} w_{k,g} X_k + \epsilon$$

Assuming a random effects for  $w_{k,g} \sim N(0, \sigma_w^2)$  and  $\epsilon \sim N(0, \sigma_\epsilon^2 I_n)$ , where  $I_n$  is the identity matrix, we calculated the total variability explained by local and distal components by estimating  $\sigma_w^2$  with restricted maximum likelihood (REML) using GCTA software [24]. For heritability analyses in the GTEx cohort, we removed the *distal* term from the model and only estimated marginal *local*  $h^2$  due to the smaller sample sizes of both cross-tissue and tissue-specific expression levels compared to DGN.

Approximate confidence intervals were computed as the point estimate  $\pm 2$  times the

estimated standard error. The intervals were also forced to be  $\geq 0$  or  $\leq 1$ . Genes were considered to have heritability significantly different from 0 if the confidence interval did not include 0.

By default we restricted the heritability estimates to be in the 0 to 1 interval. However, for the purpose of estimating the mean heritability (see Table 1 and S1 Table), we performed separate runs allowing the heritability estimates to take negative values with the `--reml-no-constrain` option in GCTA. Despite the lack of obvious biological interpretation of a negative heritability, it is an accepted procedure used in order to avoid bias in the estimated mean [10,11].

## Determining polygenicity versus sparsity using the elastic net

We used the glmnet package to fit an elastic net model where the tuning parameter is chosen via 10 fold cross validation to maximize prediction performance measure by Pearson's  $R^2$  [41,42].

The elastic net penalty is controlled by mixing parameter  $\alpha$ , which spans LASSO ( $\alpha = 1$ , the default) [16] at one extreme and ridge regression ( $\alpha = 0$ ) [17] 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  $R^2$  for  $\alpha = 0$  means the gene expression trait is highly polygenic, while an optimal prediction  $R^2$  for  $\alpha = 1$  means the trait is highly sparse.

In the DGN cohort, we tested 21 values of the mixing parameter ( $\alpha = 0, 0.05, 0.1, \dots, 0.90, 0.95, 1$ ) 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  $\alpha = 0.05, 0.5, 0.95, 1$ .

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

We used BSLMM [20] to model the effect of local genetic variation (SNPs within 1 Mb of gene) on the genetic architecture of gene expression. The BSLMM is a linear model with a polygenic component (small effects) and a sparse component (large effects)



enforced by sparsity inducing priors on the regression coefficients [20]. We used the software GEMMA [43] to implement BSLMM for each gene using the following parameters:

```
gemma -g [genoFile] -p [expFile] -a [snpFile] -bslmm 1 -s 100000 -o [out]
```

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  $|\gamma|$  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 [44]. We applied OTD to GTEx Project [23] 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:

$$Y_i = T_{i,cross} + T_{i,tissue}$$

Specifically, to generate cross-tissue and tissue-specific expression levels, we used the `lmer` function in the R [45] package `lme4` [46] 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 [40] version 18) with a mean gene expression level across tissues greater than 0.1 RPKM (reads per kilobase of transcript per million reads mapped) and RPKM > 0 in at least 3 individuals 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** [29] 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 [28] performed with a subset of the GTEx data (maximum n=175 in the nine tissues of the pilot phase, see [23]), 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  $t$  given the data. Using the top eQTL for each gene  $g$ , we defined the entropy  $S_g$  as:

$$S_g = - \sum_t p_{t,g} \log p_{t,g}$$

where  $p_{t,g}$  is the eQTL probability in tissue  $t$  normalized to 1 for each gene  $g$ . 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  $S_g$  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.

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## Supporting Information

### S1 Fig

**GTE<sub>x</sub> whole tissue distal heritability ( $h^2$ ) estimation.** Distal (SNPs that are eQTLs in the Framingham Heart Study on other chromosomes [FDR < 0.05]) gene expression  $h^2$  estimates from a joint model in the nine GTE<sub>x</sub> tissues with the largest sample sizes are ordered by increasing  $h^2$ . The 95% confidence interval (CI) of each  $h^2$  estimate is in gray and genes with a lower bound greater than zero are in blue.

## S2 Fig

**GTEx whole tissue cross-validated predictive performance across the elastic net.** Predictive  $R^2$  difference between LASSO ( $\alpha = 1$ ) and several other values of  $\alpha$  compared to LASSO predictive  $R^2$  for all autosomal genes per tissue.

## S3 Fig

**Cross-tissue and whole tissue comparison of heritability ( $h^2$ , A) and standard error (SE, B).** Cross-tissue local  $h^2$  is estimated using the cross-tissue component (random effects) of the mixed effects model for gene expression and SNPs within 1 Mb of each gene. Whole tissue local  $h^2$  is estimated using the measured gene expression for each respective tissue and SNPs within 1 Mb of each gene.

## S4 Fig

**GTEx whole tissue expression Bayesian Sparse Linear Mixed Model.** Comparison of median PGE (proportion of PVE explained by sparse effects) to median PVE (total proportion of variance explained) for expression of each gene. The 95% credible set of each PGE estimate is in gray and genes with a lower credible set (LCS) greater than 0.01 are in blue.

## S5 Fig

**GTEx orthogonal tissue decomposition cross-tissue and tissue-specific expression Bayesian Sparse Linear Mixed Model.** Comparison of median PGE (proportion of PVE explained by sparse effects) to median PVE (total proportion of variance explained) for expression of each gene. The 95% credible set of each PGE estimate is in gray and genes with a lower credible set (LCS) greater than 0.01 are in blue.

## S6 Fig

**Cross-tissue and tissue-specific comparison of heritability ( $h^2$ , A) and standard error (SE, B) estimation.** Cross-tissue local  $h^2$  is estimated using the cross-tissue component (random effects) of the mixed effects model for gene expression

and SNPs within 1 Mb of each gene. Tissue-specific local  $h^2$  is estimated using the tissue-specific component (residuals) of the mixed effects model for gene expression for each respective tissue and SNPs within 1 Mb of each gene.

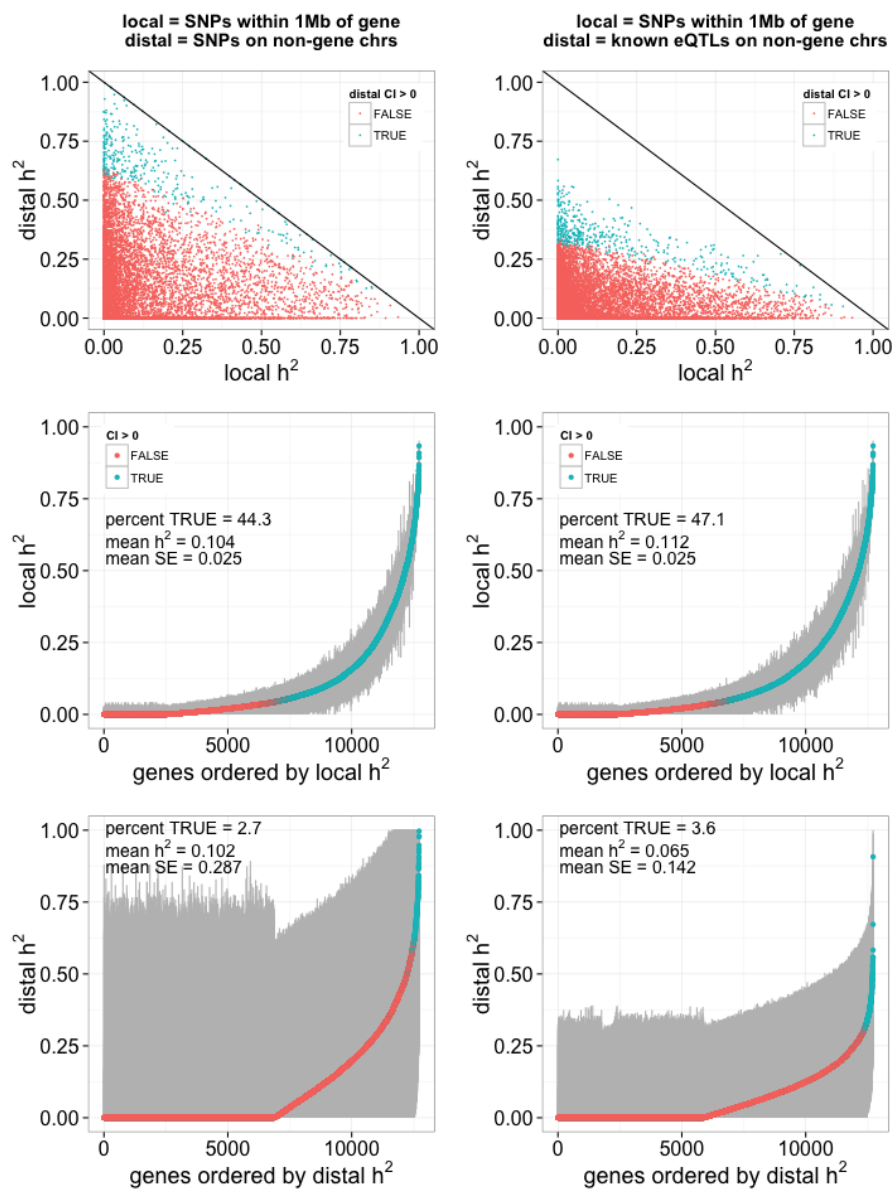
## S7 Fig

**GTEx orthogonal tissue decomposition cross-tissue and tissue-specific expression cross-validated predictive performance across the elastic net.**

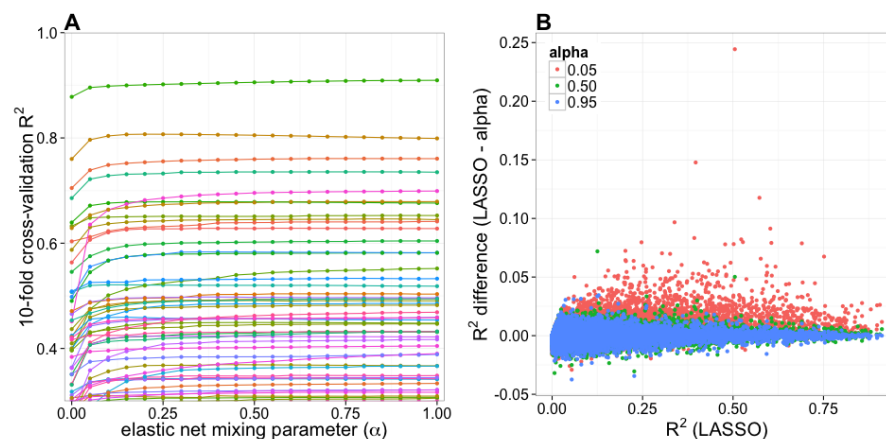
Predictive  $R^2$  difference between LASSO ( $\alpha = 1$ ) and several other values of  $\alpha$  compared to LASSO predictive  $R^2$  for all autosomal genes per tissue.

## S1 Table

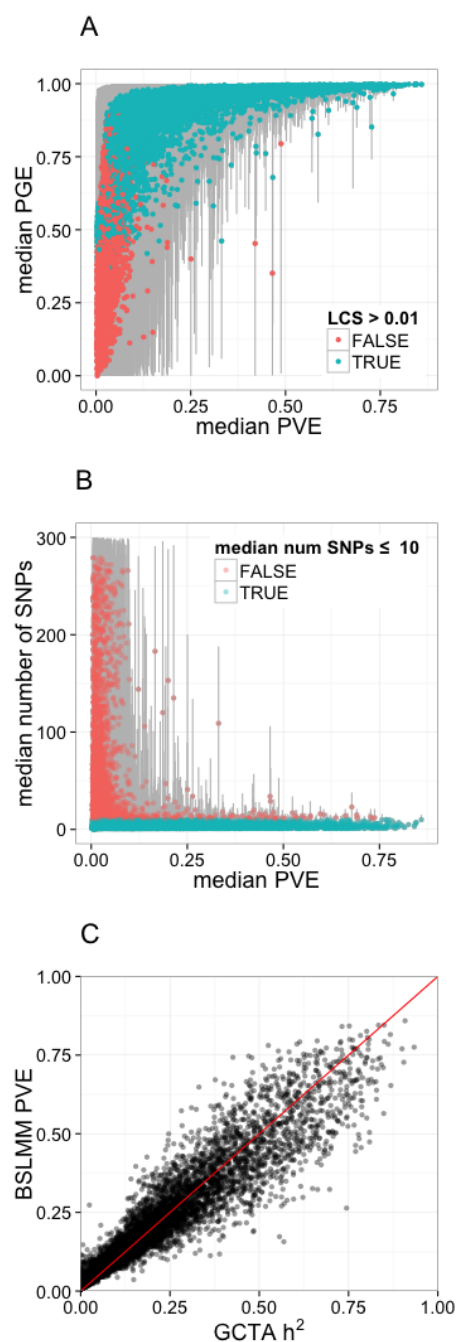
**Estimates of cross-tissue and tissue-specific local  $h^2$ .** Expression levels derived by Orthogonal Tissue Decomposition and  $h^2$  estimated using the `--reml-no-constrain` method.



**Figure 1. DGN whole blood expression joint heritability ( $h^2$ ).** Local (SNPs within 1 Mb of each gene) and distal (Left: SNPs on non-gene chromosomes. Right: SNPs that are eQTLs in the Framingham Heart Study on other chromosomes [FDR < 0.05])  $h^2$  for gene expression were jointly estimated. (**Top**) Distal  $h^2$  compared to local  $h^2$  per gene in each model. (**Middle**) Local and (**Bottom**) distal gene expression  $h^2$  estimates ordered by increasing  $h^2$ . The 95% confidence interval (CI) of each  $h^2$  estimate is in gray and genes with a lower bound greater than zero are in blue.

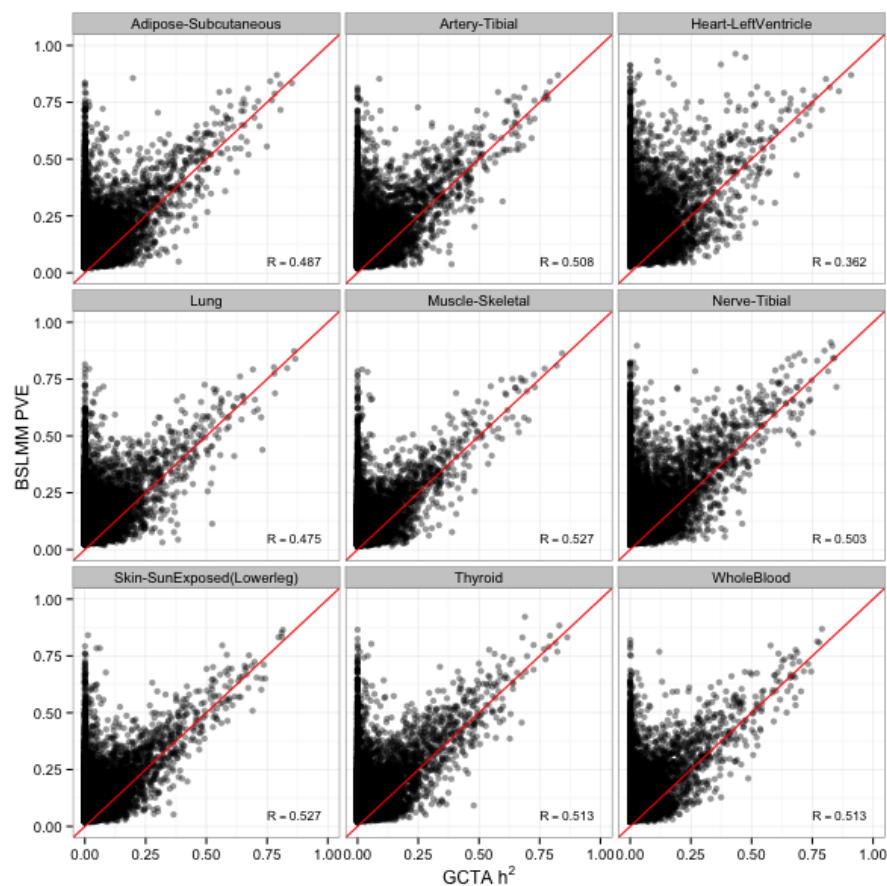


**Figure 2. DGN cross-validated predictive performance across the elastic net.** (A) 10-fold cross-validated  $R^2$  of predicted vs. observed expression in DGN whole blood compared to a range of elastic net mixing parameters ( $\alpha$ ) for genes on chromosome 22 with  $R^2 > 0.3$ . (B) Predictive  $R^2$  difference between LASSO ( $\alpha = 1$ ) and several other values of  $\alpha$  compared to LASSO predictive  $R^2$  for all autosomal genes.

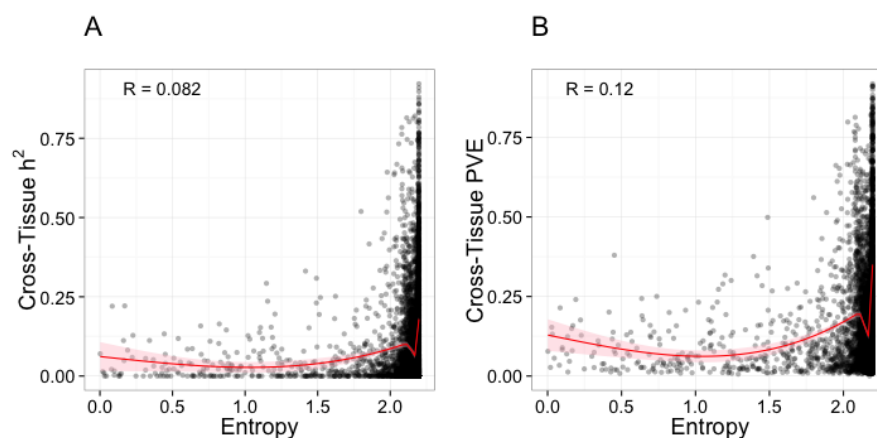


**Figure 3. Bayesian Sparse Linear Mixed Models reveal the sparsity of gene expression architecture.** (A) Comparison of median PGE (proportion of PVE explained by sparse effects) to median PVE (total proportion of variance explained) for expression of each gene. The 95% credible set of each PGE estimate is in gray and genes with a lower credible set (LCS) greater than 0.01 are in blue. (B) Comparison of the median number of SNPs included in the model of each gene to median PVE. The 95% credible set of each SNP-number estimate is in gray and genes with a median of 10 or fewer SNPs are in blue. (C) BSLMM-estimated PVE compared to GCTA-estimated heritability per gene ( $R=0.96$ ).

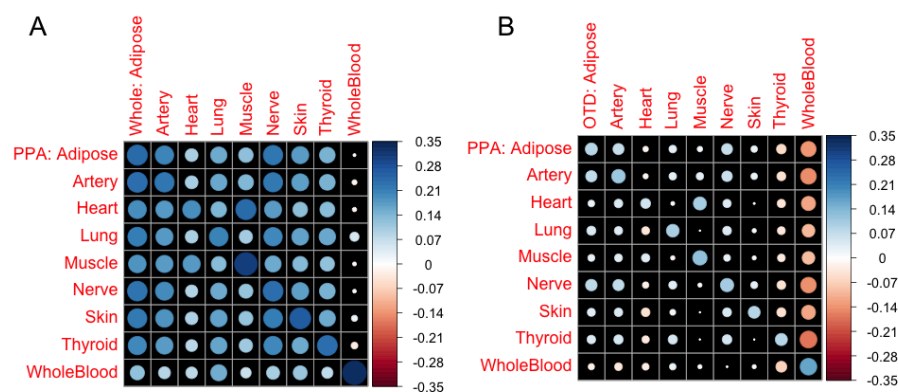




**Figure 4. GTEx PVE vs.  $h^2$ .** GTEx whole tissue expression BSLMM-estimated PVE (total proportion of variance explained) compared to GCTA-estimated heritability per gene. R = Pearson correlation.



**Figure 5. Cross-tissue expression from the orthogonal tissue decomposition (OTD) compared to multi-tissue eQTL results.** Pearson correlation (R) between the entropy of the posterior probabilities from the Flutre et al. multi-tissue eQTL method and the estimates of (A) heritability and (B) PVE of cross-tissue gene expression derived from the OTD. The generalized additive model smoothing line is in red.



**Figure 6. Comparison of whole tissue or tissue-specific expression PVE to the posterior probability of a gene being actively regulated in a tissue.** Pearson correlation (R) between the posterior probability the top multi-tissue eQTL regulates its gene in a given tissue (PPA, Flutre et al. method) and the PVE of (A) whole tissue gene expression or (B) tissue-specific gene expression from the orthogonal tissue decomposition (OTD). Area of each circle is proportional to the absolute value of R.